

Title	Stomagen positively regulates stomatal density in Arabidopsis.
Author(s)	Sugano, Shigeo S; Shimada, Tomoo; Imai, Yu; Okawa, Katsuya; Tamai, Atsushi; Mori, Masashi; Hara-Nishimura, Ikuko
Citation	Nature (2009), 463: 241-244
Issue Date	2009-12-09
URL	<a href="http://hdl.handle.net/2433/91250">http://hdl.handle.net/2433/91250</a>
Right	c 2009 Nature Publishing Group All Rights Reserved. 許諾条件により本文は2010-06-10に公開.
Type	Journal Article
Textversion	author

# Stomagen positively regulates stomatal density in *Arabidopsis*

Shigeo S. Sugano<sup>1\*</sup>, Tomoo Shimada<sup>1\*</sup>, Yu Imai<sup>1</sup>, Katsuya Okawa<sup>2</sup>, Atsushi Tamai<sup>3</sup>, Masashi Mori<sup>3</sup> & Ikuko Hara-Nishimura<sup>1</sup>

<sup>1</sup>Graduate School of Science, Kyoto University, Sakyo-ku, Kyoto 606-8502, Japan. <sup>2</sup>Innovative Drug Research Laboratories, Kyowa Hakko Kirin Co., Ltd., Taksasaki 370-1295, Japan. <sup>3</sup>Research Institute for Bioresources and Biotechnology, Ishikawa Prefectural University, Nonoichi-machi, Ishikawa 921-8836, Japan.

\*These authors contributed equally to this work.

**Stomata in the epidermal tissues of leaves are valves through which passes CO<sub>2</sub>, and as such they influence the global carbon cycle<sup>1</sup>. The two-dimensional pattern and density of stomata in the leaf epidermis are genetically and environmentally regulated to optimize gas exchange<sup>2</sup>. Two putative intercellular signalling factors, EPF1 and EPF2, function as negative regulators of stomatal development in *Arabidopsis*, possibly by interacting with the receptor-like protein TMM<sup>3-6</sup>. One or more positive intercellular signalling factors are assumed to be involved in stomatal development, but their identities are unknown<sup>7</sup>. Here we show that a novel secretory peptide, which we designate as stomagen, is a positive intercellular signalling factor that is conserved among vascular plants. Stomagen is a 45-amino-acid, cysteine-rich peptide that is generated from a 102-amino-acid precursor protein designated as STOMAGEN. Both an *in planta* analysis and a semi-*in-vitro* analysis with recombinant and chemically synthesized stomagen peptides showed that stomagen has stomata-inducing activity in a dose-dependent manner. A genetic analysis showed that TMM is epistatic to STOMAGEN (At4g12970), suggesting that stomatal development is finely regulated by competitive binding of positive and negative regulators to the same receptor. Notably, STOMAGEN is expressed in inner tissues (the mesophyll) of immature leaves but not in the epidermal tissues where stomata develop. This study provides evidence of a mesophyll-derived positive regulator of stomatal density. Our findings provide a conceptual advancement in understanding stomatal development: inner photosynthetic tissues optimize their function by regulating stomatal density in the epidermis for efficient uptake of CO<sub>2</sub>.**

Stomatal development in leaves is negatively regulated by putative cell-surface receptors TOO MANY MOUTHS (TMM) and ERECTA family receptor-like kinases (ER, ERL1 and ERL2)<sup>3,8</sup>. Two possible ligands of these receptors, EPIDERMAL PATTERNING FACTOR (EPF) 1 and EPF2, act as negative signalling factors at distinct steps during

stomatal development<sup>4-6</sup>. Another negative regulator, STOMATAL DENSITY AND DISTRIBUTION 1 (SDD1), which is a subtilisin-type proteinase, is reported to be dependent on TMM<sup>9</sup>. Although positive signalling factors are thought to exist<sup>6,7,10</sup>, no signalling factors are known to positively regulate stomatal development in leaves.

To identify novel positive signalling factors, we first selected genes that are co-expressed with a set of TMM, SDD1 and EPF1 by using *in silico* screening with a transcriptome database of *Arabidopsis thaliana*, ATTED-II<sup>11</sup> (Supplementary Table 1). This analysis identified a gene (At4g12970) that encodes EPFL9, a member of the EPF family, whose function was unknown<sup>6</sup>. Interestingly, overexpression of the At4g12970 gene resulted in an increase of stomatal density of various organs that naturally produce stomata (leaves, cotyledons, stems, fruits and anthers) in each of 13 overexpression lines that we established ( $n = 7$  for each T2 line, Fig. 1a and Supplementary Fig. 1). This phenotype contrasts with a decreased stomatal density phenotype in transgenic plants overexpressing EPF1 or EPF2<sup>4-6</sup>. We named the At4g12970 gene STOMAGEN after its stoma-generating activity.

Stomata in most dicotyledonous plants are separated by at least one non-stomatal cell<sup>12</sup>, which is thought to improve the efficiency of gas exchange<sup>2</sup>. The overexpression lines of STOMAGEN (STOMAGEN-OX) exhibited many clustered stomata in mature leaves (Fig. 1a) and formed satellite meristemoids (stomatal precursor cells) adjacent to guard cells or their precursors in immature cotyledons (Supplementary Fig. 2), indicating that the orientation of cell division in the stomatal lineage is defective in these lines. The abnormal clustering is probably due to the 35S promoter-driven expression of STOMAGEN in the epidermis (see Fig. 2c, d).

We next examined the loss-of-function phenotype by gene silencing using an artificial microRNA that targeted the 3' side of the coding region of STOMAGEN (Supplementary Fig. 3a). Fourteen independent T2 lines in which STOMAGEN was silenced by RNA interference (STOMAGEN-RNAi lines) had reduced stomatal densities in various organs (Fig. 1a and Supplementary Fig. 1). Other STOMAGEN-silenced lines made with another artificial microRNA (12 independent T2 lines) and

double-stranded RNA (dsRNA; 4 independent T2 lines) exhibited the same phenotype (Supplementary Fig. 3), indicating that *STOMAGEN* is required for proper stomatal development. A quantitative analysis revealed a positive correlation between the expression level of *STOMAGEN* messenger RNA and either stomatal density or the stomatal index (defined as the ratio of stomata to stomata plus other epidermal cells) (Fig. 1b, c). Both the gain-of-function and loss-of-function phenotypes indicate that *STOMAGEN* is a positive regulator of stomatal development.

*STOMAGEN* was predominantly expressed in immature organs including leaves, stems and flower buds (Fig. 2a).  $\beta$ -Glucuronidase (GUS) activity in p*STOMAGEN*::GUS lines was observed in immature cotyledons and developing leaves, but not in fully expanded cotyledons or leaves (Fig. 2b). These expression patterns are consistent with the fact that stomatal development starts in immature organs<sup>2</sup>. Notably, GUS activity was detected in the mesophyll tissues but not in the epidermal tissues where stomata develop (Fig. 2c). *In situ* RNA hybridization gave the same result (Fig. 2d). These results indicate that *STOMAGEN* is produced in mesophyll cells and that inter-tissue signalling regulates stomatal development in the epidermis. This provides a counterexample to previous reports that most regulatory genes for stomatal development are expressed in epidermal tissues<sup>10,13,14</sup>. To determine whether *STOMAGEN* itself migrates from the mesophyll to the epidermis, we generated transgenic lines (12 independent T2 lines) that expressed a *STOMAGEN*-Venus fusion protein, which exhibited the same phenotype as *STOMAGEN*-OX (see Fig. 3d). Venus fluorescence was observed outside the plasmolysed cells (Fig. 2e and Supplementary Fig. 4), indicating that *STOMAGEN* is secreted from the cells to the apoplast. In previous studies, mesophyll-epidermis interactions have been limited to coordinating the airspace and stomata<sup>12,15</sup>. The discovery of *STOMAGEN* will provide a valuable insight into mesophyll-epidermis interaction for stomatal density.

The *STOMAGEN* gene encodes a small protein of 102 amino acids with a putative signal peptide at its amino terminus (Fig. 3a). *STOMAGEN*-Venus, which was immunoprecipitated from the transgenic leaves with anti-GFP antibody, gave a single band on an immunoblot with anti-stomagen antibody (Fig. 3b). Edman degradation of the immunoprecipitate gave an N-terminal sequence of IGSTAPTXY (where X indicates undetermined; Supplementary Fig. 5), which is identical to the sequence starting from Ile 58 of *STOMAGEN*. The N-terminal sequence is consistent with the N-terminal sequences obtained by matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF/MS) analyses of lysyl endopeptidase and trypsin digests of the immunoprecipitate,

IGSTAPTCTYNECRGCRYK and IGSTAPTCTYNECR, respectively (Fig. 3c). These results indicate that *STOMAGEN* is processed *in vivo* to produce the 45-amino-acid peptide that contains six cysteine residues. We designated the cysteine-rich peptide as stomagen. Expression of stomagen with a signal peptide at the N terminus, which lacked the propeptide domain, increased the stomatal density in the transgenic plants (10 independent T2 lines; Fig. 3d), indicating that the 45-amino-acid peptide is sufficient to induce stomatal formation. Genes for stomagen-like peptides were found in various plant species (Supplementary Fig. 6). Interestingly, a similar peptide was found in *Selaginella moellendorffii*, a primitive vascular plant. Together, these results indicate that stomagen is widely distributed in vascular plants.

To obtain pure stomagen, tobacco-cultured BY-2 cells were transformed with the *STOMAGEN* gene. The culture medium was found to be immunopositive for stomagen and was fractionated with reverse-phase high-performance liquid chromatography (HPLC; Supplementary Fig. 7a, b). Several fractions were found to contain a single ~5-kDa band (Fig. 3e and Supplementary Fig. 7c), the N-terminal sequence of which matched that of stomagen (Supplementary Fig. 7d). Treatment with the purified stomagen for 3 days induced stomatal formation of *Arabidopsis* plants (Fig. 3f). We also used chemically synthesized stomagen (45 amino acids residues). Both the synthetic and recombinant stomagens increased stomatal density dose-dependently at concentrations from 300 nM to 10  $\mu$ M (Fig. 3g). The induction of stomatal formation was suppressed by adding anti-stomagen antibody (Fig. 3g). These results demonstrate that stomagen is a stomata-inducing factor.

The stomatal lineage is triggered by *SPEECHLESS* (*SPCH*)<sup>13,14</sup>, which is a basic helix-loop-helix (bHLH) type transcriptional factor. Both overexpression of *STOMAGEN* in a *spch* mutant (three lines of T1 plants) and application of purified stomagen to a *spch* mutant ( $n = 20$ ) failed to induce stomatal formation (Fig. 4a). These results indicate that the stomata-inducing activity of stomagen is dependent on the *SPCH* pathway. *SPCH* regulates the expressions of both the receptor-like protein TMM and its putative ligands EPF1 and EPF2 (refs 6, 13, 14).

TMM acts negatively in leaves but positively in stems<sup>16</sup>, whereas EPF1 and EPF2 act negatively in both organs<sup>4,5</sup>. Surprisingly, 44.8  $\pm$  16.7-fold overexpression of *STOMAGEN* in a *tmm* mutant background failed to increase stomatal density in either stems or leaves (ten independent T2 lines; Fig. 4b and Supplementary Fig. 8). Gene silencing of *STOMAGEN* in *tmm* did not affect the stomatal density of *tmm* leaves (ten independent T2 lines; Fig. 4b and Supplementary Fig. 8). These results indicate that *STOMAGEN* requires *TMM* for positive

regulation of stomatal development. EPF1 and EPF2 are thought to negatively regulate stomatal development by interacting with TMM as ligands. It is possible that stomagen competes with either EPF1 or EPF2 for the same receptor, TMM.

We silenced *STOMAGEN* in *epf* mutants and determined two indices: stomatal density and non-stomatal cell density (density of epidermal cells other than stomata). Stomatal density was higher in each *epf* mutant than in the wild type and the elevation of stomatal density was reduced in *STOMAGEN* gene silencing lines (Fig. 4c and Supplementary Fig. 9). This result suggests that *STOMAGEN* regulates the stomatal density in an EPF-independent manner. On the other hand, non-stomatal cell density was much higher in *epf2* mutants than in the wild type (Fig. 4d), consistent with the role of *EPF2* as a negative regulator of the production of stomatal lineage cells<sup>6,17</sup>. Gene silencing of *STOMAGEN* did not reduce the high density of non-stomatal cells in *epf2* (11 independent T2 lines), although it reduced non-stomatal cell density in the wild type (14 independent T2 lines) (Fig. 4d and Supplementary Fig. 9). This result suggests that *STOMAGEN* regulates the production of stomatal lineage cells in an *EPF2*-dependent manner. It seems that *STOMAGEN* has two functions: it regulates the production of stomatal lineage cells *EPF2*-dependently, and the differentiation of stomatal lineage cells to stomata independent of *EPF1* and *EPF2*.

Taken together, three putative signalling factors—stomagen, EPF1 and EPF2—regulate stomatal development through the common receptor-like protein TMM. In addition to these signalling factors, *SDD1* is known to act upstream of *TMM*<sup>9</sup>. *SDD1* has been thought to negatively regulate stomatal development by proteolytically processing some negative precursor factors<sup>9,18</sup>. However, *SDD1* is genetically independent of *EPF1* and *EPF2*<sup>4-6</sup>. Another possibility is that *SDD1* exerts its negative effect on stomatal development by degrading stomagen, a positive regulator.

Our findings show that stomatal development is regulated by two classes of signals which have opposing effects: a positive signal, stomagen, and the negative signals, EPF1 and EPF2. Such a bi-directional signalling system would enable plants to control stomatal density more precisely. Our results also show that mesophyll-derived stomagen positively regulates the differentiation of stomatal lineage cells into stomata in the epidermis. This inter-tissue signalling would provide a conceptual advancement: stomatal density is optimized by photosynthetic tissues for efficient uptake of CO<sub>2</sub>. Stomagen could be a useful tool, either through genetic engineering or by spraying, for developing crops and trees with high stomatal densities that have a high absorption capacity for CO<sub>2</sub>.

## METHODS SUMMARY

### Plant materials and phenotypic analysis

*Arabidopsis thaliana* accession Columbia-0 (CS60000) was used as the wild type. Mutants used are *tmm* (SALK\_011958), *epf1* (SALK\_137549), *epf2* (SALK\_047918) and *spch* (SAIL\_36\_B6). Stomata stained with Safranin-O (1  $\mu\text{g ml}^{-1}$ ) were inspected with a microscope (Axioskop 2, Zeiss).

### Plasmid constructions

Plant binary vectors based on Gateway cloning technology (Invitrogen) were used. Two artificial microRNAs for *STOMAGEN* were designed as described<sup>19</sup>. The promoter (2 kb) and coding region of genomic DNA of *STOMAGEN* were cloned into pENTR/D-TOPO (Invitrogen). The DNA fragments were transferred into pBGWFS7 for promoter analysis and into either pB2GW7 or pH2GW7 for overexpression.

### Immunoprecipitation

Extracts from transgenic plants expressing *STOMAGEN*-Venus (3 g fresh weight) were subjected to immunoprecipitate at 4 °C with 200  $\mu\text{l}$  of anti-GFP microbeads ( $\mu\text{MACS GFP tag protein isolation kit}$ , Milteny Biotech).

### Recombinant stomagen

Stomagen was produced in tobacco-cultured BY-2 cells by using the Tomato mosaic virus based expression vector and the inducible virus infection system<sup>20</sup>. The BY-2 medium was ultra-centrifuged at 1,000,000g for 1 h and filtered through a 0.22- $\mu\text{m}$  filter (Millex). Filtrate was charged to a reverse-phase HPLC column ( $\mu\text{RPC C2/C18 SP4/6}$ ) and washed with 5% methanol and 0.5% TFA. Elution was conducted with 90% methanol and 0.5% TFA. Each fraction (1 ml) was concentrated up to approximately 100  $\mu\text{l}$ .

### Stomata induction assay

Stomagen was applied to *Arabidopsis* seedlings at 2 days after germination that germinated in B5 sterilized liquid medium. After a 3-day incubation at 22 °C, abaxial epidermis of cotyledons was inspected with a confocal microscope to determine stomatal densities.

Received 9 June; accepted 17 November 2009; Published online 9 December 2009.

1. Hetherington, A. M. & Woodward, F. I. The role of stomata in sensing and driving environmental change. *Nature* **424**, 901–908 (2003).

2. Nadeau, J. A. & Sack, F. D. Stomatal development in *Arabidopsis*. In *The Arabidopsis Book* (eds Someville, C. & Meyerowitz, E) doi:10.1199/tab.0066 (American Society of Plant Biologists, 2002).
3. Nadeau, J. A. & Sack, F. D. Control of stomatal distribution on the *Arabidopsis* leaf surface. *Science* **296**, 1697–1700 (2002).
4. Hara, K., Kajita, R., Torii, K. U., Bergmann, D. C. & Kakimoto, T. The secretory peptide gene *EPF1* enforces the stomatal one-cell-spacing rules. *Genes Dev.* **21**, 1720–1725 (2007).
5. Hunt, L. & Gray, J. E. The signaling peptide EPF2 controls asymmetric cell divisions during stomatal development. *Curr. Biol.* **19**, 864–869 (2009).
6. Hara, K. *et al.* Epidermal cell density is autoregulated via a secretory peptide, EPIDERMAL PATTERNING FACTOR 2 in *Arabidopsis* leaves. *Plant Cell Physiol.* **50**, 1019–1031 (2009).
7. Bergmann, D. C. Integrating signals in stomatal development. *Curr. Opin. Plant Biol.* **7**, 26–32 (2004).
8. Shpak, E. D., McAbee, J. M., Pillitteri, L. J. & Torii, K. U. Stomatal patterning and differentiation by synergistic interactions of receptor kinases. *Science* **309**, 290–293 (2005).
9. Berger, D. & Altmann, T. A subtilisin-like serine protease involved in the regulation of stomatal density and distribution in *Arabidopsis thaliana*. *Genes Dev.* **14**, 1119–1131 (2000).
10. Nadeau, J. A. Stomatal development: new signals and fate determinants. *Curr. Opin. Plant Biol.* **12**, 29–35 (2009).
11. Obayashi, T., Hayashi, S., Saeki, M., Ohta, H. & Kinoshita, K. ATTED-II provides coexpressed gene networks for *Arabidopsis*. *Nucleic Acids Res.* **37**, D987–D991 (2009).
12. Sachs, T. The development of spacing patterns in the leaf epidermis. In *Clonal Basis of Development* 161–182 (Academic Press, 1987).
13. MacAlister, C. A., Ohashi-Ito, K. & Bergmann, D. C. Transcription factor control of asymmetric cell divisions that establish the stomatal lineage. *Nature* **445**, 537–540 (2007).
14. Pillitteri, L. J., Sloan, D. B., Bogenschutz, N. L. & Torii, K. U. Termination of asymmetric cell division and differentiation of stomata. *Nature* **445**, 501–505 (2007).
15. Serna, L., Torres-Contreras, J. & Fenoll, C. Specification of stomatal fate in *Arabidopsis*: evidences for cellular interactions. *New Phytol.* **153**, 399–404 (2002).
16. Geisler, M., Yang, M. & Sack, F. D. Divergent regulation of stomatal initiation and patterning in organ and suborgan regions of the *Arabidopsis* mutants *too many mouths* and *four lips*. *Planta* **205**, 522–530 (1998).
17. Geisler, M., Nadeau, J. & Sack, F. D. Oriented asymmetric divisions that generate the stomatal spacing pattern in *Arabidopsis* are disrupted by the *too many mouths* mutation. *Plant Cell* **12**, 2075–2086 (2000).
18. von Grolla, U., Bergera, D., & Altmann, T. The subtilisin-like serine protease SDD1 mediates cell-to-cell signaling during *Arabidopsis* stomatal development. *Plant Cell.* **14**, 1527–39 (2002).
19. Schwab, R., Ossowski, S., Riester, M., Warthmann, N. & Weigel, D. Highly specific gene silencing by artificial microRNAs in *Arabidopsis*. *Plant Cell* **18**, 1121–1133 (2006).
20. Dohi, K., Tamai, A. & Mori, M. Insertion in the coding region of the movement protein improves stability of the plasmid encoding a tomato mosaic virus-based expression vector. *Arch. Virol.* **153**, 1667–1675 (2008).

**Supplementary Information** is linked to the online version of the paper at [www.nature.com/nature](http://www.nature.com/nature).

**Acknowledgements** We thank T. Nakagawa for donating the TMM::GFP seeds; K. Yamada for his technical support of quantitative PCR; and Y. Nakata for her technical support of *in situ* hybridization. We thank J. Raymond for reading and editing the manuscript. We thank the ABRC for providing seeds of *Arabidopsis* T-DNA insertion lines. This work was supported by Grants-in-Aid for Scientific Research (no. 16085203 and 17107002) from the Ministry of Education, Culture, Sports, Science and Technology (MEXT) of Japan, by the Global Center of Excellence Program ‘Formation of a Strategic Base for Biodiversity and Evolutionary Research: from Genome to Ecosystem’ of MEXT, and by Japan Science and Technology Agency (JST).

**Author Contributions** T.S. performed *in silico* screening to identify *STOMAGEN*. S.S.S. established the *STOMAGEN*-RNAi lines, performed genetic interaction experiments and other experiments including antibody production and *in situ* hybridization. S.S.S. also purified recombinant stomagen, refolded chemically synthesized stomagen, and evaluated their stomata inducing activities. Y.I. developed the *STOMAGEN*-OX and p*STOMAGEN*::GUS lines. A.T. and M.M. produced stomagen in tobacco-cultured cells by using their ToMV-expression system. K.O. conducted MS analysis of *STOMAGEN*-Venus. I.H.-N. and T.S. designed this project and supervised. S.S.S., T.S. and I.H.-N. wrote the paper with input from co-authors.

**Author Information** Reprints and permissions information is available at [www.nature.com/reprints](http://www.nature.com/reprints). The authors declare no competing financial interests. Correspondence and requests for materials should be addressed to I.H.-N. ([ihnishi@gr.bot.kyoto-u.ac.jp](mailto:ihnishi@gr.bot.kyoto-u.ac.jp)).

## METHODS

### *In silico* screening

The genes co-expressed with a set of *TMM*, *SDD1* and *EPF1* were searched by using ATTED-II (<http://atted.jp/>). The program used was NetworkDrawer in the tool section of ATTED-II. The procedure was composed of three steps: first was to select 'Execute' and 'MR-rank method'<sup>11</sup> and 'top 300 genes in 22,263 genes' in Function 3; second was to input three AGI codes of At1g80080 (*TMM*), At1g04110 (*SDD1*) and At2g20875 (*EPF1*); third was to submit the codes.

### Plant materials and growth conditions

*Arabidopsis* accession Columbia-0 (CS60000) was used as the wild type. Plants were initially grown on GM plates (MS salts, 1% sucrose, 0.5% MES-KOH, pH 5.7 and 0.5% Gellan gum) under continuous light at 22 °C for 2–3 weeks before being transferred to soil. The mutants used were as follows: *tmm* (SALK\_011958), *epf1* (SALK\_137549), *epf2* (SALK\_047918) and *spch* (SAIL\_36\_B6). Their T-DNA inserts were confirmed using primers designed by iSECT tools (Supplementary Table 2). The pTMM::GFP line was provided by T. Nakagawa<sup>21</sup>. Three types of *STOMAGEN*-silenced lines (*STOMAGEN*-RNAi) were generated (see below). *STOMAGEN*-RNAi generated with ami-A was used except for Supplementary Fig. 3.

### Plasmid construction

Plant binary vectors based on Gateway cloning technology (Invitrogen) were used for most manipulations. DNA fragments for *STOMAGEN* overexpression and silencing by dsRNA were PCR-amplified from genomic DNA. An upstream region (2 kb) from the start codon for *STOMAGEN* was used as a promoter for *STOMAGEN* expression analysis. Translational fusions for *STOMAGEN* were constructed by joining three independently amplified PCR fragments, namely Venus with 4 glycines at its C terminus (Venus), the signal peptide for pumpkin 2S albumin<sup>22</sup> (SP), and the *STOMAGEN* open reading frame without its stop codon (*STOMAGEN*). The two independent artificial microRNAs for *STOMAGEN* (ami-A and ami-B) were designed according to ref. 19. The PCR products were cloned into pENTR/D-TOPO (Invitrogen) before being transferred to pBGWFS7 for promoter analysis, to either pB2GW7 or pH2GW7 for overexpression, or to pB2GWI(2)WG7 for dsRNA.

Plants were stably transformed by using standard protocols for *Agrobacterium* (strain GV3101)-mediated transformation. Transgenic lines were selected on GM plates containing 10 mg l<sup>-1</sup> glufosinate-ammonium (Sigma) or 10 mg l<sup>-1</sup> hygromycin (Sigma). For expression of stomagen in *Escherichia coli*, a DNA fragment coding stomagen was PCR-amplified and then cloned into pET32-derived vector<sup>23</sup> (Novagen).

### Phenotype analysis

Phenotypic analysis was performed using at least seven individual plants of each of ten independent T2 lines ( $n > 70$ ) except for *sph* *STOMAGEN*-OX (three lines of T1 plant). Plant organs, including 23 DAG cotyledons, 23 DAG first leaves, fully expanded second internodes, mature fruits and anther of flowering buds in the same size were collected. The organs were fixed in ethanol:acetic acid (9:1) for over 7 h and cleared in chloral hydrate solution (chloral hydrate:water:glycerol (8:2:1, w/v/w)). After removal of chloral hydrate, stomata were stained with 1 µg ml<sup>-1</sup> Safranin-O. Leaf epidermis was examined under a microscope at ×20 (Axioskop 2 plus system; Zeiss) with a CCD camera (model VB-7010; Keyence). Four square-areas of 0.22 mm<sup>2</sup> were analysed for quantification of stomatal density and non-stomatal cell density. Statistical analysis was conducted using R statistical computing software (<http://www.R-project.org>). We conducted Kruskal-Wallis test for Fig. 4b, and two-tailed Student's *t*-test for Figs 3d, g and 4c, d.

### RNA isolation and quantitative RT-PCR

Total RNA was extracted from 12th leaves of wild type, mutants and transgenic lines (23 DAG) as described previously<sup>24</sup>. Real-time PCR was performed using the 7500 Fast Real-Time PCR system (Applied Biosystems) and the TaqMan gene expression assay kit (Applied Biosystems) according to the manufacturer's instructions. The assay identifiers were At02219575\_g1 for *STOMAGEN* and At02270958\_gH for *ACTIN8*. We used the threshold cycle method for relative quantification<sup>25</sup>.

### GUS staining

GUS staining was performed as described previously<sup>26</sup>, except that the samples were fixed with 90% acetone at 4 °C for 7–10 h.

### *In situ* hybridization

*In situ* hybridization was performed as described previously<sup>24</sup>. Labelled probes were prepared from the coding region of the *STOMAGEN* cDNA (306 bp).

### Confocal fluorescent microscopy

The fluorescent images were inspected under a confocal laser scanning microscope (model LSM510

META; Zeiss) by using the 488-nm and 544-nm lines of a 40-mW Ar/Kr laser. We used filter sets for Venus, GFP and FM1-43 fluorescences (an excitation filter of BP450-490, a dichroic mirror of FT510, and a barrier filter of BP515-565; Zeiss) and for FM4-64 fluorescence (an excitation filter of BP515-560, a dichroic mirror of FT510, and a barrier filter of BP515-565; Zeiss).

### Expression of recombinant proteins by using tobacco-cultured BY-2 cells

The recombinant proteins including stomagen, STOMAGEN-GFP and EPF1-GFP were produced in cultured BY-2 cells using the tomato mosaic virus (ToMV)-based expression vector and the inducible virus infection system (IVIS)<sup>20</sup>. The ToMV vector for expression of each recombinant protein was constructed using pBICER8-ToMV-C0.3-HuIFN $\gamma$ -SRz22 by replacing the IFN $\gamma$  gene with the cDNA encoding each of stomagen, STOMAGEN-GFP and EPF1-GFP. GFP cDNA was PCR-amplified and inserted into the 3' end of *STOMAGEN* or *EPF1* gene of ToMV-STOMAGEN or ToMV-EPF1 vectors to produce the constructs. These constructs were introduced to the transgenic BY-2 cells expressing the estrogen-activatable transcription factor XVE. Induction by  $\beta$ -estradiol was performed essentially according to ref. 20. The cell cultures were centrifuged for 3 min at 800 r.p.m. The supernatant containing the secreted recombinant stomagen was subjected to purification (see below). The pellets containing STOMAGEN-GFP or EPF1-GFP were subjected to immunoblot analysis.

### Purification of recombinant stomagen

Culture medium of the BY-2 cells expressing stomagen was ultra-centrifuged at 1,000,000g for 1 h and the supernatant was filtered through a 0.22- $\mu$ m filter (Millex). The filtrate was applied to a reverse-phase HPLC column ( $\mu$ RPC C2/C18 SP4/6). After washing the column with a solution of 5% methanol and 0.5% TFA, elution was conducted with a linear gradient of methanol (5% to 90%). Each fraction (1 ml) was collected and then concentrated to approximately 100  $\mu$ l by evaporation.

### Specific antibody

Stomagen was expressed in *E. coli* strain BL21. The recombinant protein was purified by affinity chromatography using a Hi-Trap chelating column (GE Healthcare) followed by cation-exchange chromatography using a H-SP column (GE healthcare). The purified preparation was injected into a rabbit as described previously<sup>26</sup>. To check this, we did immunoblots of STOMAGEN-GFP and EPF1-GFP with either the antibody raised against stomagen or anti-GFP antibody. We confirmed that

the antibody against stomagen recognized STOMAGEN-GFP but not EPF1-GFP, whereas anti-GFP antibody recognized both (Supplementary Fig. 10).

### Immunoprecipitation

Immunoprecipitation was conducted by using the GFP tag protein isolation kit ( $\mu$ MACS; Milteny Biotech). Extract from wild type or STOMAGEN-Venus plants (7 g fresh weight) in 5 ml of 50 mM Tris-HCl, pH 7.5, 50 mM NaCl, and 1% CHAPS was incubated with 200  $\mu$ l of anti-GFP microbeads at 4 °C. The beads were trapped with magnetic columns and the bound proteins were eluted with a loading buffer of the kit.

### Immunoblot

Immunoblot was basically performed as described previously<sup>24</sup>. Anti-stomagen antibody was diluted 1:1,000 (v/v). Horseradish peroxidase-conjugated goat antibody against rabbit IgG (diluted 5,000-fold, NA934, GE Healthcare) was used as the secondary antibody.

### N-terminal sequencing

The immunoprecipitated samples were subjected to SDS-PAGE followed by blotting on PVDF membranes (Immobilon-P, Millipore) and then subjected to either Coomassie blue (CBB) staining or immunostaining. The CBB-stained band that corresponded to the positive signal on the immunoblot was subjected to N-terminal sequencing analysis with an automatic peptide sequencer (Procise 492cLC, Applied Biosystems), as described previously<sup>22</sup>.

### Refolding of synthetic stomagen

Chemically synthesized peptides (Invitrogen, 0.5 mg ml<sup>-1</sup>) that were dissolved in 20 mM Tris-HCl, pH 8.8, and 50 mM NaCl were dialysed (Spectra/Por3 MWCO, 3,500; Spectrum Laboratories) for 1 day at 4 °C against 0.5 mM glutathione (reduced and oxidized forms, Wako) and 200 mM L-arginine (Sigma) at pH 8.0, and were further dialysed three times against 20 mM Tris-HCl, pH 8.8, and 50 mM NaCl for 1.5 days to remove glutathione.

### Mass spectrometry

Mass spectrometric identification was performed as described<sup>27</sup>. Proteins were separated by SDS-PAGE followed by silver staining. The respective bands were excised separately and then subjected to the in-gel digestions with either trypsin or lysyl endopeptidase (Promega) in a buffer containing 50 mM ammonium bicarbonate, pH 8.0, and 2% acetonitrile at 37 °C overnight. The digests were subjected to matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF/MS)

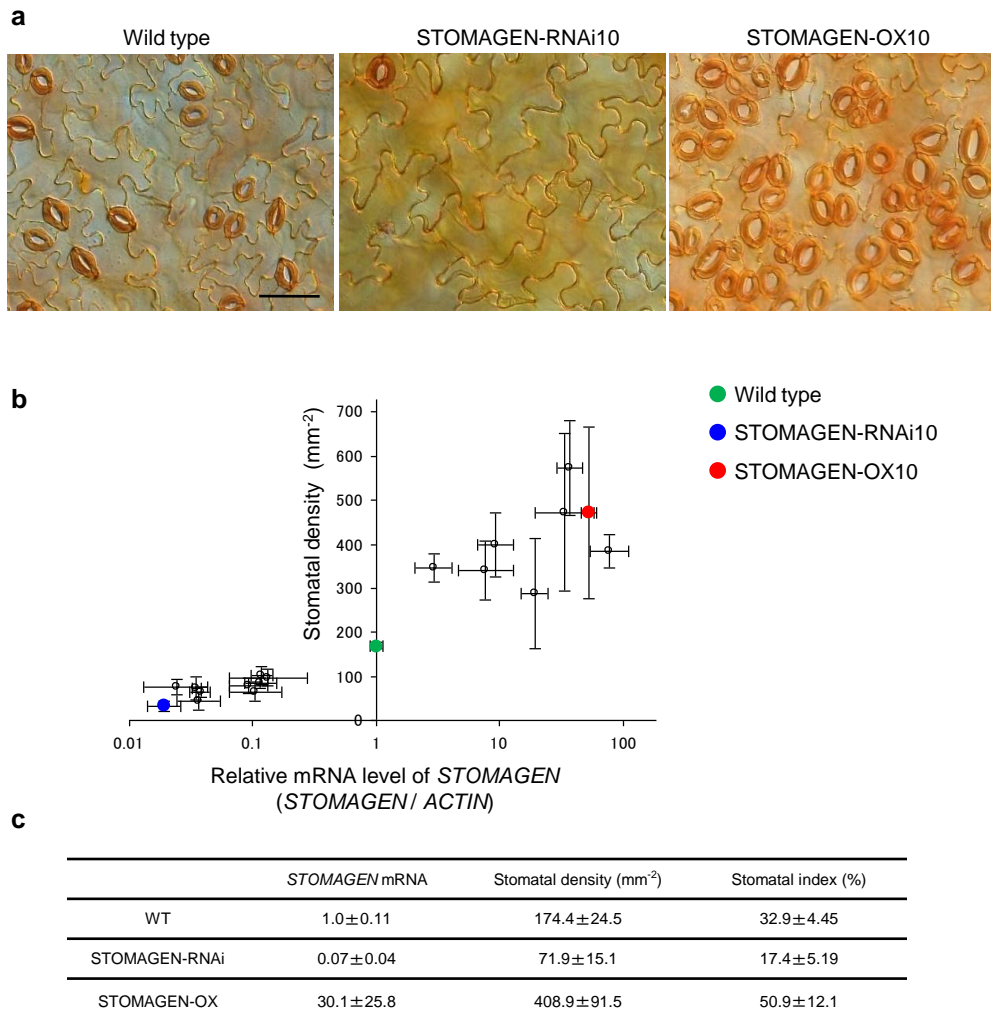
with an ultraflex TOF/TOF (MALDI-TOF/MS, Bruker Daltonics).

### **Stomata induction assay**

Stomagen was applied to 2 DAG *Arabidopsis* plants that had germinated in B5 sterilized liquid medium. After further 3 days of incubation at 22 °C, stomatal densities of abaxial cotyledon were determined by inspecting with a confocal microscope. An effect of ethylene on stomatal development in liquid medium with or without stomagen was evaluated by using an ethylene action inhibitor, silver thiosulphate. We confirmed that stomata-inducing activity of stomagen was independent on ethylene.

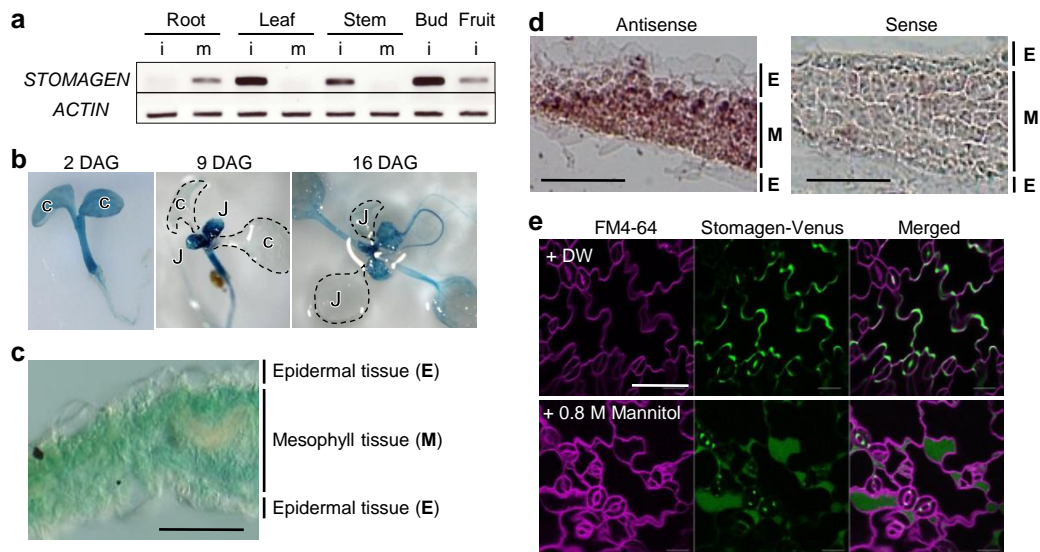
21. Kono, A. *et al.* The *Arabidopsis* D-type cyclin CYCD4 controls cell division in the stomatal lineage of the hypocotyl epidermis. *Plant Cell* **19**, 1265–1277 (2007).
22. Hara-Nishimura, I., Takeuchi, Y., Inoue, K. & Nishimura, M. Vesicle transport and processing of the precursor to 2S albumin in pumpkin. *Plant J.* **4**, 793–800 (1993).
23. Shimada, T. *et al.* AtVPS29, a putative component of a retromer complex, is required for the efficient sorting of seed storage proteins. *Plant Cell Physiol.* **47**, 1187–1194 (2006).
24. Kunieda, T. *et al.* NAC family proteins NARS1/NAC2 and NARS2/NAM in the outer integument regulate embryogenesis in *Arabidopsis*. *Plant Cell* **20**, 2631–2642 (2008).
25. Livak, K. J. & Schmittgen, T. D. Analysis of relative gene expression data using real-time quantitative PCR and the  $2^{-\Delta\Delta C(T)}$  method. *Methods* **25**, 402–408 (2001).
26. Kinoshita, T. *et al.* Vacuolar processing enzyme is up-regulated in the lytic vacuoles of vegetative tissues during senescence and under various stressed conditions. *Plant J.* **19**, 43–53 (1999).
27. Jensen, O. N., Podtelejnikov, A. & Mann, M. Delayed extraction improves specificity in database searches by matrix-assisted laser desorption/ionization peptide maps. *Rapid Commun. Mass Spectrom.* **10**, 1371–1388 (1996).





**Figure 1. *STOMAGEN* positively regulates stomatal development.**

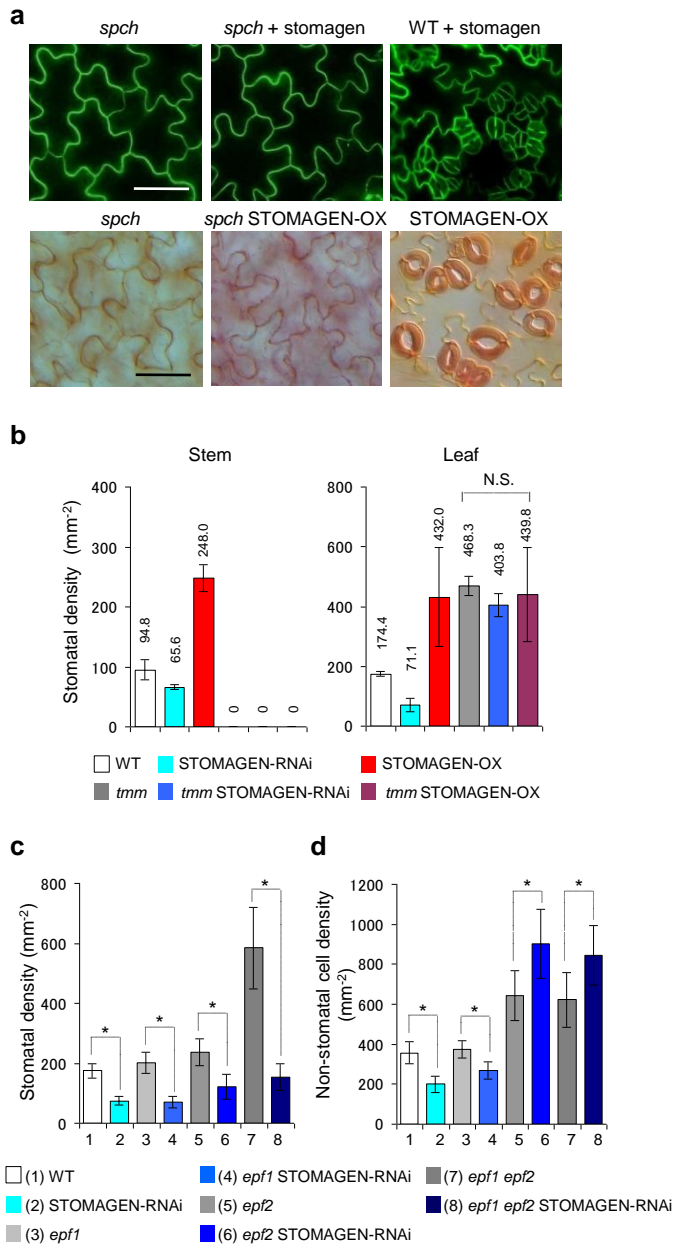
**a**, Differential interference contrast (DIC) images of the abaxial epidermis of mature first leaves of wild type, a *STOMAGEN*-silenced line (STOMAGEN-RNAi10) and a *STOMAGEN*-overexpressing line (STOMAGEN-OX10). Stomata were stained with Safranin-O. Scale bar, 50  $\mu$ m. **b**, Correlation between stomatal density in mature first leaves and the relative mRNA level of *STOMAGEN* measured by real-time PCR. Green, blue and red dots correspond to wild type, STOMAGEN-RNAi10 and STOMAGEN-OX10. Error bars indicate s.d. **c**, *STOMAGEN* mRNA level, stomatal density and stomatal index in wild type, STOMAGEN-RNAi and STOMAGEN-OX.



**Figure 2. Stomagen is expressed in inner tissues of immature organs and secreted out of the cells to mediate inter-tissue signalling.**

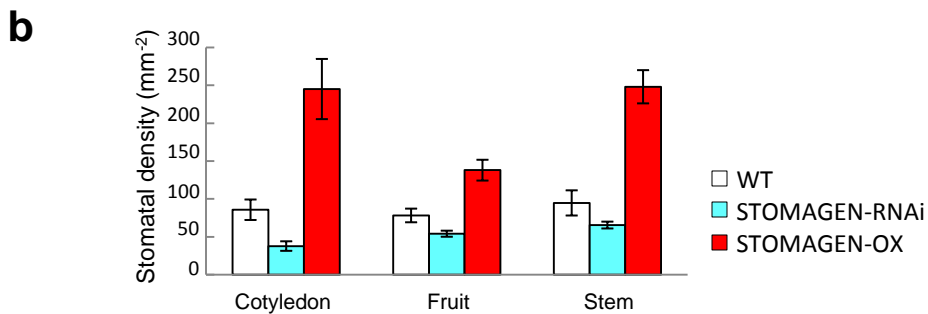
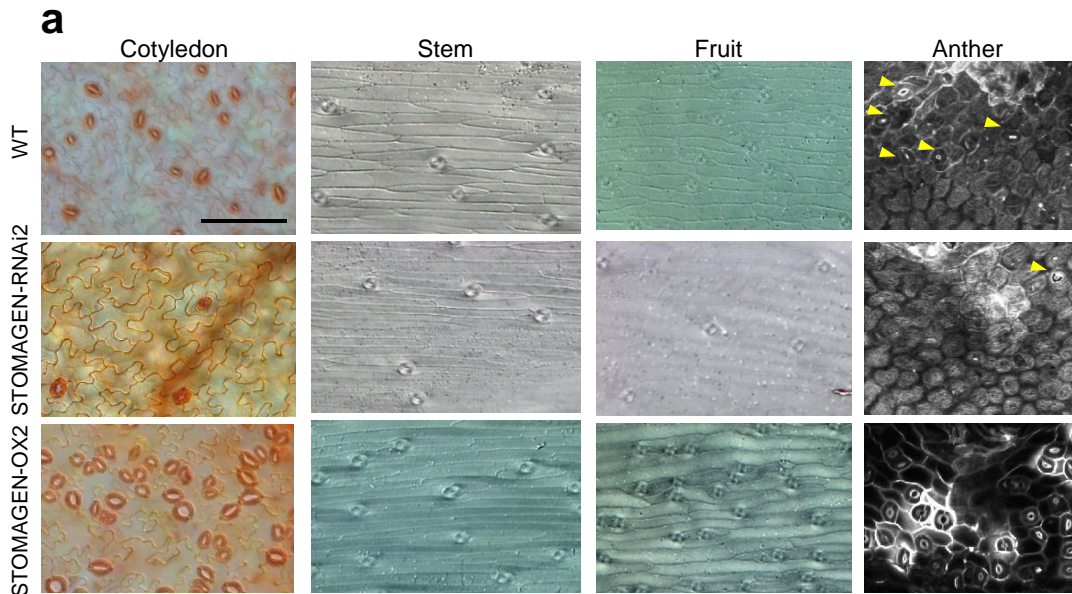
**a**, RT-PCR showing *STOMAGEN* mRNA levels in mature (m) and immature (i) organs including roots, leaves, stems, buds and fruits. *ACTIN* was used as an internal control. **b**, GUS staining of a p*STOMAGEN*::GUS line at 2, 9 and 16 days after germination (DAG). C, cotyledons; J, juvenile leaves. **c**, A cross-section of the 9th leaf of an 18-DAG p*STOMAGEN*::GUS plant showing GUS staining in mesophyll. **d**, *In situ* hybridization in leaf primordia with both antisense and sense *STOMAGEN* probes. **e**, Confocal images of abaxial sides of 3-DAG cotyledons stained with FM4-64 of a *STOMAGEN*-Venus line. Untreated cells in distilled water (DW) (upper) and plasmolysed cells in 0.8 M mannitol (lower) are shown. All scale bars, 50  $\mu$ m.





**Figure 4. *STOMAGEN* regulates stomatal density *TMM*-dependently in combination with negative regulators *EPF1* and *EPF2***

**a**, *SPCH*-dependent stomatal inducing activity of stomagen. Confocal images of abaxial epidermis of cotyledons of wild type (WT) and *spch* with or without 2  $\mu$ M stomagen treatment are shown in the top panel. DIC images of abaxial epidermis of cotyledons of *spch*, STOMAGEN-OX in *spch* and STOMAGEN-OX are shown in the bottom panel. Scale bars, 50  $\mu$ m. **b**, A genetic interaction between *TMM* and *STOMAGEN*. Stomatal density of stems (left) and an abaxial epidermis of the first leaves (right) of the indicated plants. NS, no significant difference among the three genotypes (*tmm*, *tmm* STOMAGEN-RNAi, *tmm* STOMAGEN-OX). **c**, **d**, A genetic interaction of *STOMAGEN* with each of *EPF1* and *EPF2*. Stomatal density (**c**) and non-stomatal cell density (**d**) of abaxial epidermis of the first leaves of the indicated plants are shown. Asterisk,  $P < 0.01$ .

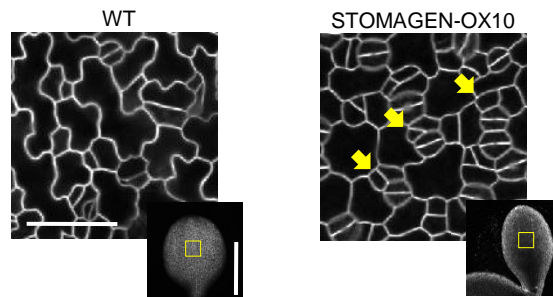


**Supplementary Figure 1. Stomatal phenotype of STOMAGEN-OX and STOMAGEN-RNAi lines.**

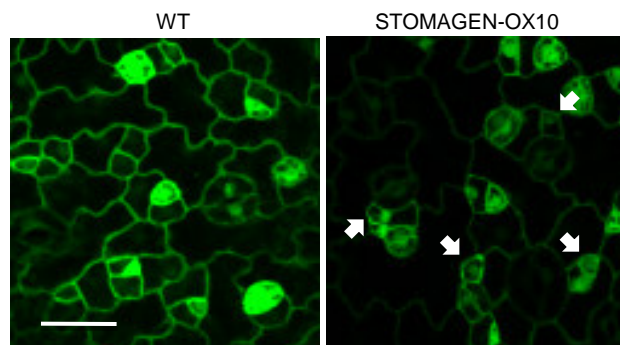
**a**, Epidermis of each of cotyledons, stems and fruits (DIC images) and of anthers (confocal images) of the wild type (WT), STOMAGEN-RNAi2 and STOMAGEN-OX2. Cell outlines in anthers were visualized with FM1-43. Yellow arrowheads indicate stomata of anthers. Bar, 100  $\mu$ m.

**b**, Quantification of stomatal density in each organ with significant differences in ANOVA ( $P < 0.01$ ). Error bars indicate S.D.

**a**



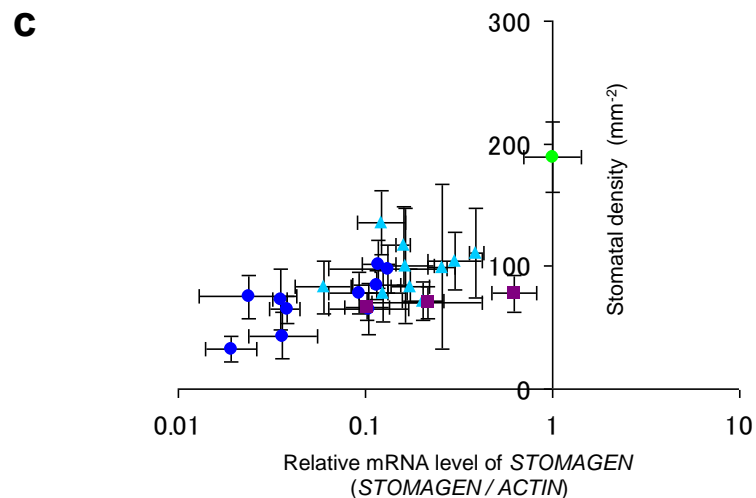
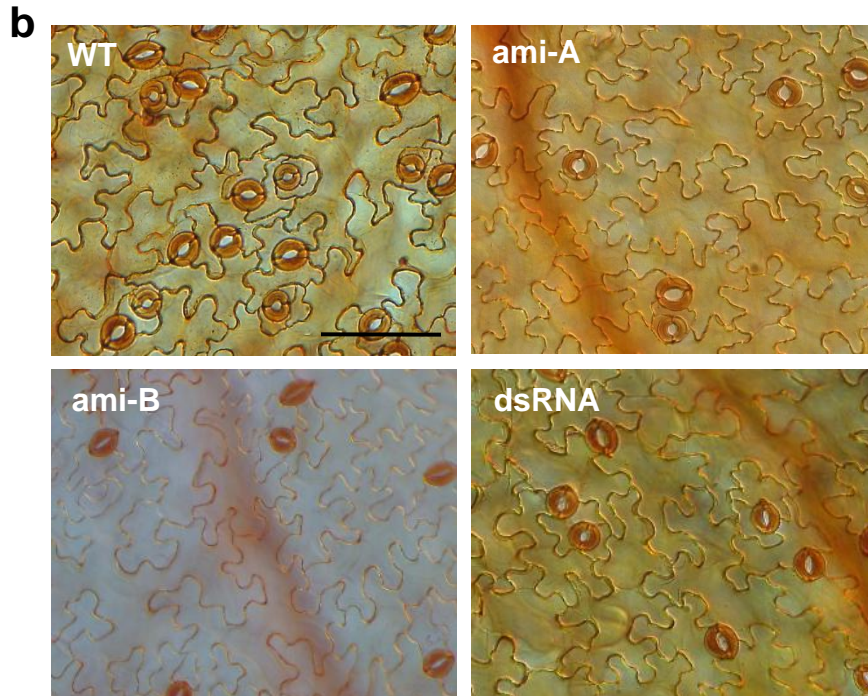
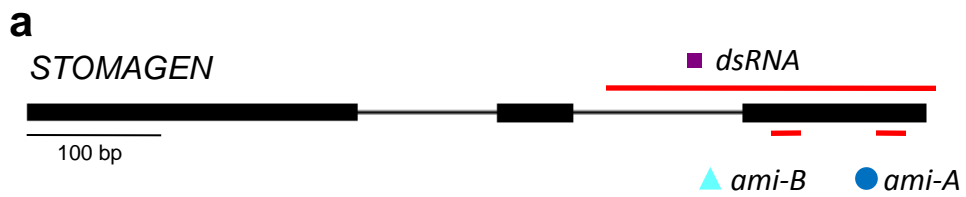
**b**



**Supplementary Figure 2. Abnormal meristemoids in STOMAGEN-OX.**

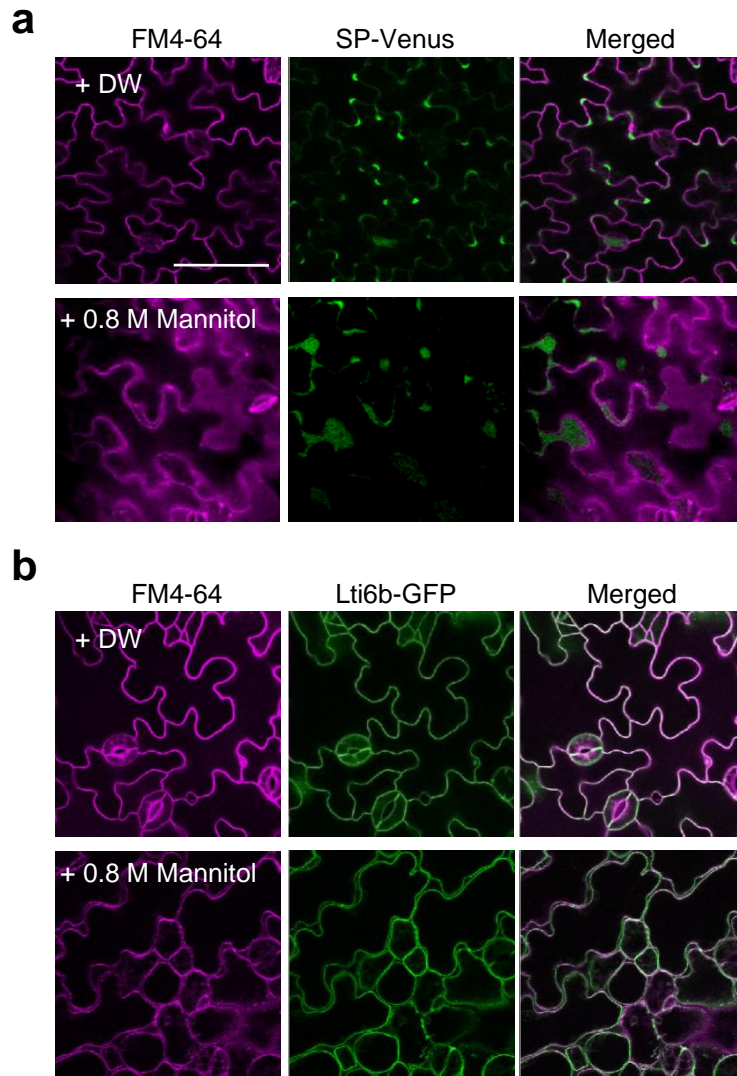
**a**, Abaxial sides of 2.5-DAG cotyledons of the wild type (WT) and STOMAGEN-OX plants that were stained with FM4-64 were inspected with a confocal microscope. The yellow-boxed areas of the cotyledons are enlarged to obtain the confocal images. Arrows indicate abnormal asymmetric divisions. Horizontal bar, 50  $\mu\text{m}$ ; vertical bar, 500  $\mu\text{m}$ .

**b**, Confocal images of *pTMM::GFP* in the epidermis of developing first leaves (5 DAG) of the wild type (WT) and STOMAGEN-OX plants. Abnormal meristemoids adjacent to guard cells are shown by arrows. Bar, 50  $\mu\text{m}$



**Supplementary Figure 3. Generation of three types of *STOMAGEN*-RNAi lines and correlation between stomatal density and the relative reduction level of *STOMAGEN* mRNA.**

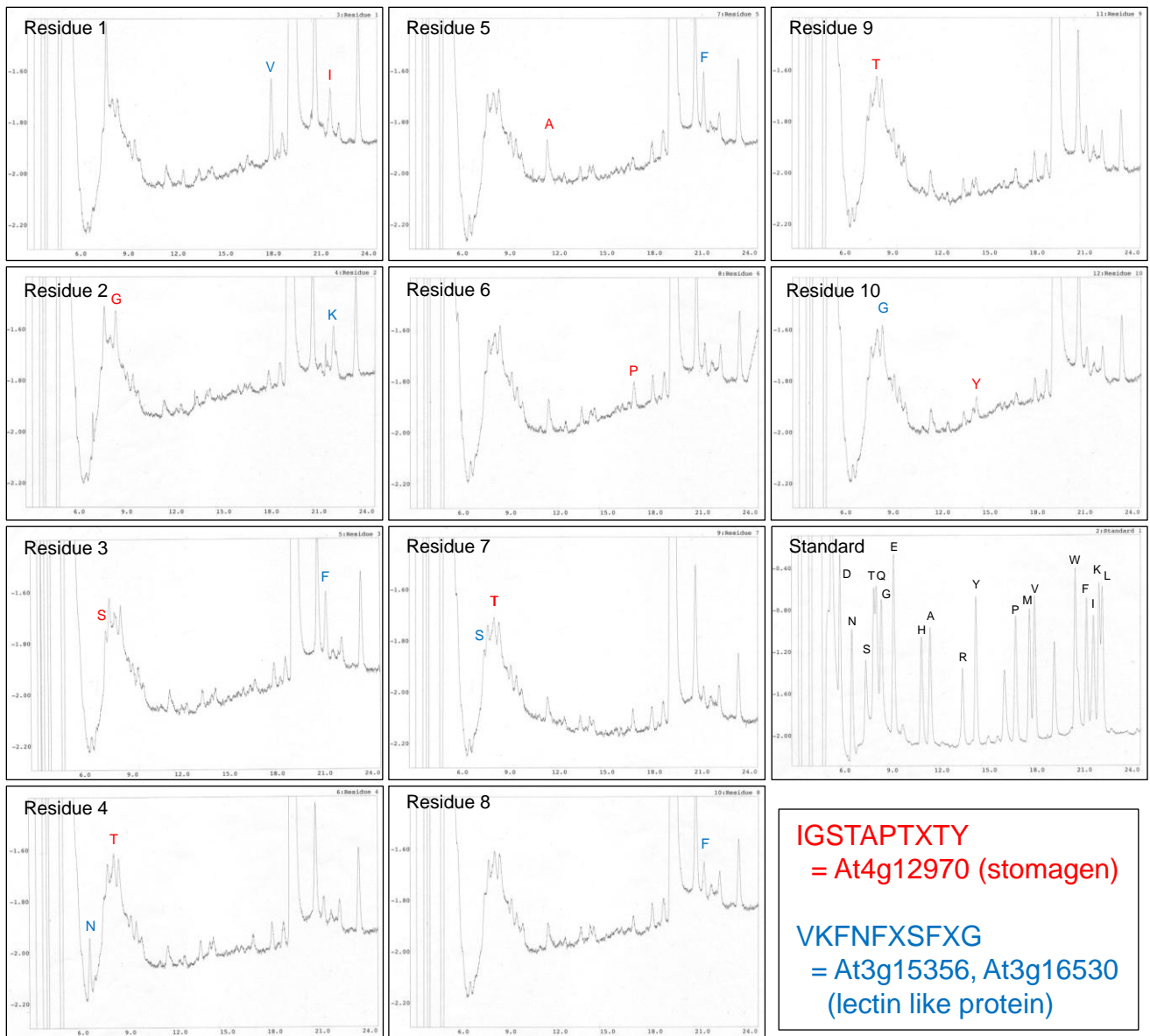
**a**, A schematic representation of three target sites of artificial microRNA-A (*ami-A*), *ami-B* and *dsRNA* for RNAi. The *STOMAGEN* gene is drawn with black boxes (exons) and lines (introns).  
**b**, DIC images of the abaxial sides of mature first leaves of *STOMAGEN*-gene silenced lines (*STOMAGEN*-RNAi) by using *ami-A*, *ami-B*, and *dsRNA*. Bar, 50  $\mu\text{m}$ .  
**c**, Correlation between stomatal density in mature first leaves in the *STOMAGEN*-RNAi lines and the relative reduction level of *STOMAGEN* mRNA measured by real-time PCR. Green circle, the wild type; blue circle, *STOMAGEN*-RNAi (*ami-A*); blue triangle, *STOMAGEN*-RNAi (*ami-B*); red square, *STOMAGEN*-RNAi (*dsRNA*). Error bars indicate S.D.



**Supplementary Figure 4. Control experiment of Figure 2e showing secretion of stomagen outside the cells.**

Confocal images of abaxial sides of 3-DAG cotyledons of transgenic plants that overexpressed a secreted form of GFP (SP-Venus, **a**) or a plasma membrane marker (Lti6b-GFP, **b**). Both untreated cells (+DW) and plasmolyzed cells in 0.8 M mannitol (+Mannitol) were inspected. Plasma membrane was counterstained with FM4-64. Bars, 50  $\mu$ m.





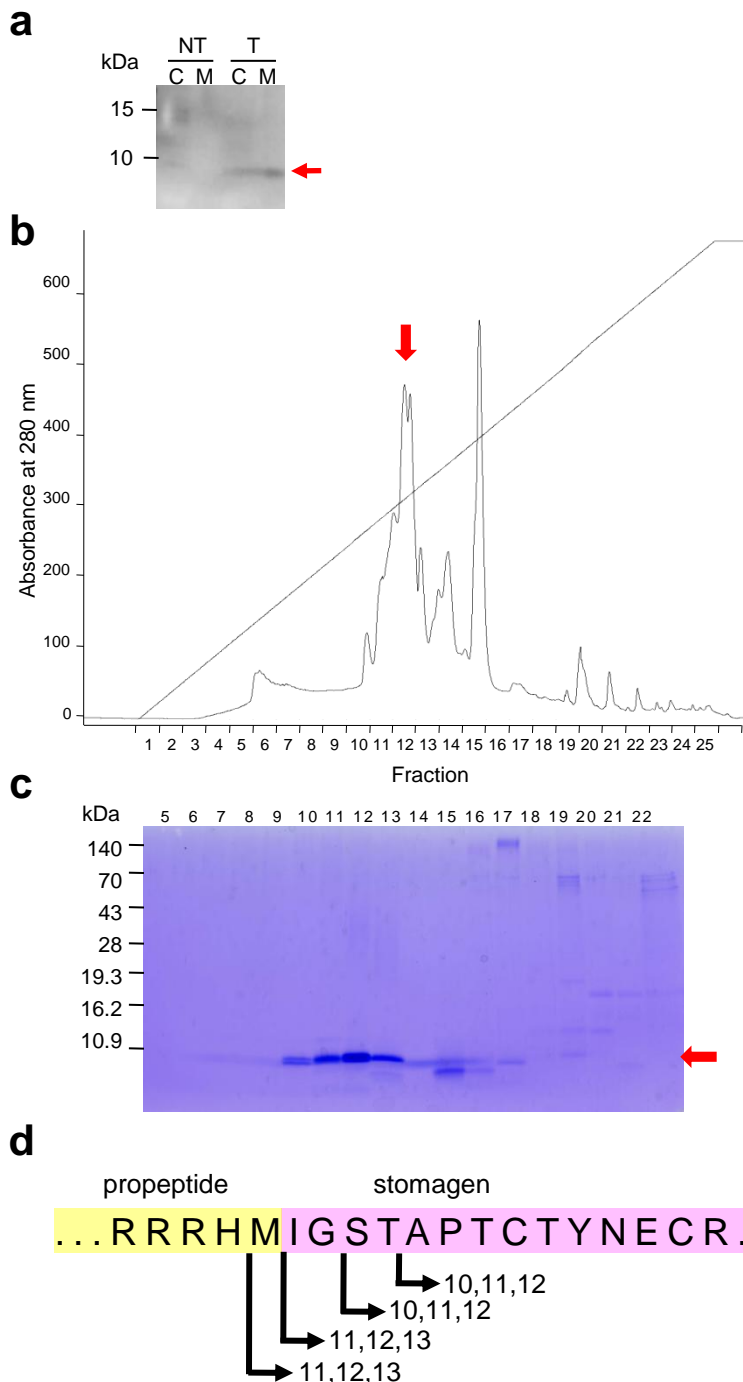
### Supplementary Figure 5. Determination of the N-terminus of stomagen.

STOMAGEN-Venus that was expressed in *Arabidopsis* was pulled down with anti-GFP beads. The pulled down protein was subjected to automatic peptide sequencer to determine the N-terminal sequence. Each panel shows the chromatogram of each cycle of the Edman degradation. The N-terminal sequence of stomagen was determined to be IGSTAPTXTY (X is not determined). Another sequence (VKFNFXSFXG) detected in this analysis was derived from contaminated lectin-like protein of *Arabidopsis* (At3g15356 or At3g16530).

<i>Arabidopsis thaliana</i>	-----MKHEMNIKPRCITIFLLFALLLGNVYVQASRPRSI-----ENTVSLLP	45
<i>Olyza sativa</i>	MANACPTSTTSSLPLFFLFCFLFSHARCNGHHGSISGTDYGEQYPHQTLPPEEHIHLQE	60
<i>Vitis vinifera</i>	-----MLISKQPNL-FLVLLFTLLFAAYFTQGSKSQVVPYPYHQREIVHLQS	45
<i>Populus trichocarpa</i>	-----MANTRLCY-LLSLLFTFILAAFVIQGSRNQELLPYHQSISTPSQE	44
<i>Selaginella moellendorffii</i>	-----	0
	▼	
<i>Arabidopsis thaliana</i>	QVHLL-----NSRRRHMIGSTAPTCTYNECRGCRYKCRAEQVPVEGNDPINSAYHYRCV	99
<i>Olyza sativa</i>	NIKVLNKERLPKYARRMLIGSTAPICTYNECRGCRFKCTAEQVPVDANDPMNSAYHYKCV	120
<i>Vitis vinifera</i>	SGEQW---MNRNSRRLMIGSTAPICTYNECRGCKYKCRAEQVPVEGNDPINSAYHYRCV	101
<i>Populus trichocarpa</i>	DSQALGGNEEQMSSKRLMIGSTAPICTYNECRGCKYKCRAEQVPVEGNDPIHSAYHYKCI	104
<i>Selaginella moellendorffii</i>	---MINEWQYPSDRRAMVVGSTAPSCTYNECRDCKTECRAEQVPLNGRNPKESAYRYLCV	56
<i>Arabidopsis thaliana</i>	CHR--	102
<i>Olyza sativa</i>	CHR--	123
<i>Vitis vinifera</i>	CHR--	104
<i>Populus trichocarpa</i>	CHR--	107
<i>Selaginella moellendorffii</i>	CRKRH	61

### Supplementary Figure 6. Stomagen orthologous proteins in vascular plants.

Multiple alignment of stomagen orthologous proteins conducted by Clustal W. Stomagen-like proteins are found in various plants, including *Oryza sativa* (Os01g0914400), *Vitis vinifera* (AM444732), and *Populus trichocarpa* (Pt02g2557). A similar sequence was also found in *Selaginella moellendorffii* (Sm084711), a primitive vascular plant. Conserved amino acids among these species are shown in reverse. The N-terminus of stomagen in *Arabidopsis* is shown by a red arrowhead.



**Supplementary Figure 7. Purification of the recombinant stomagen that was expressed in tobacco-cultured BY-2 cells.**

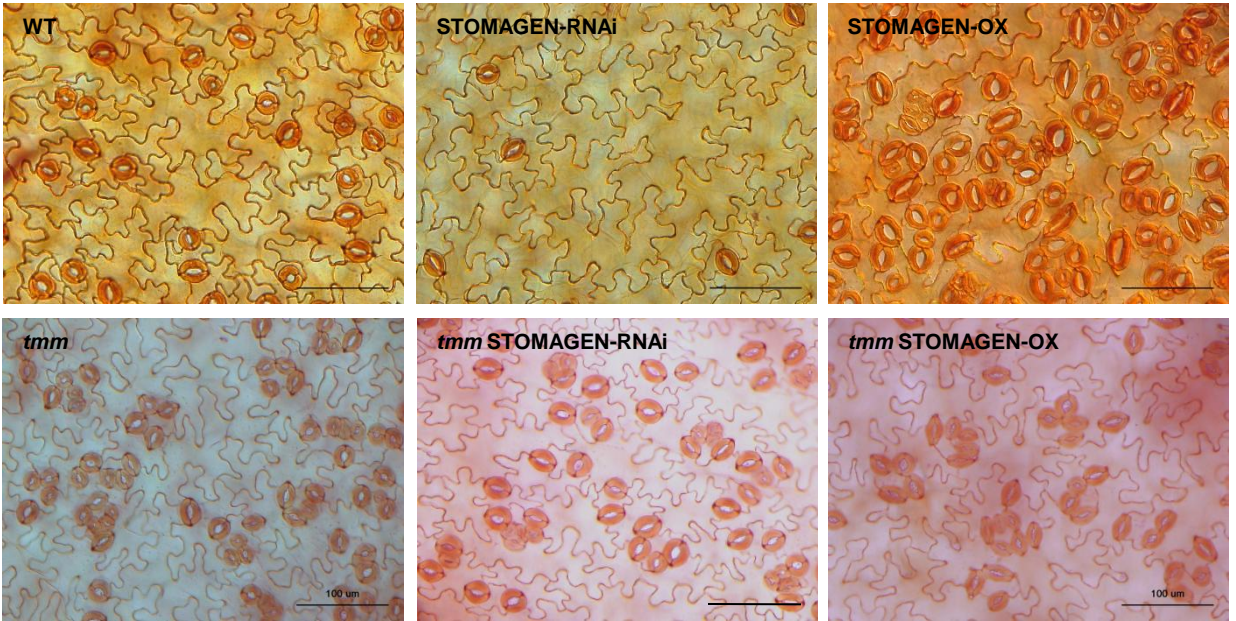
**a**, An immunoblot with anti-stomagen antibody showing that stomagen was detected in the culture medium (M) but not in transformed cells (C). T indicates transformed BY-2 cells and NT indicates non-transformed BY-2 cells.

**b**, A chromatogram of purification of recombinant stomagen using a reverse-phase C-18 column. A red arrow indicates the elution position of the recombinant stomagen.

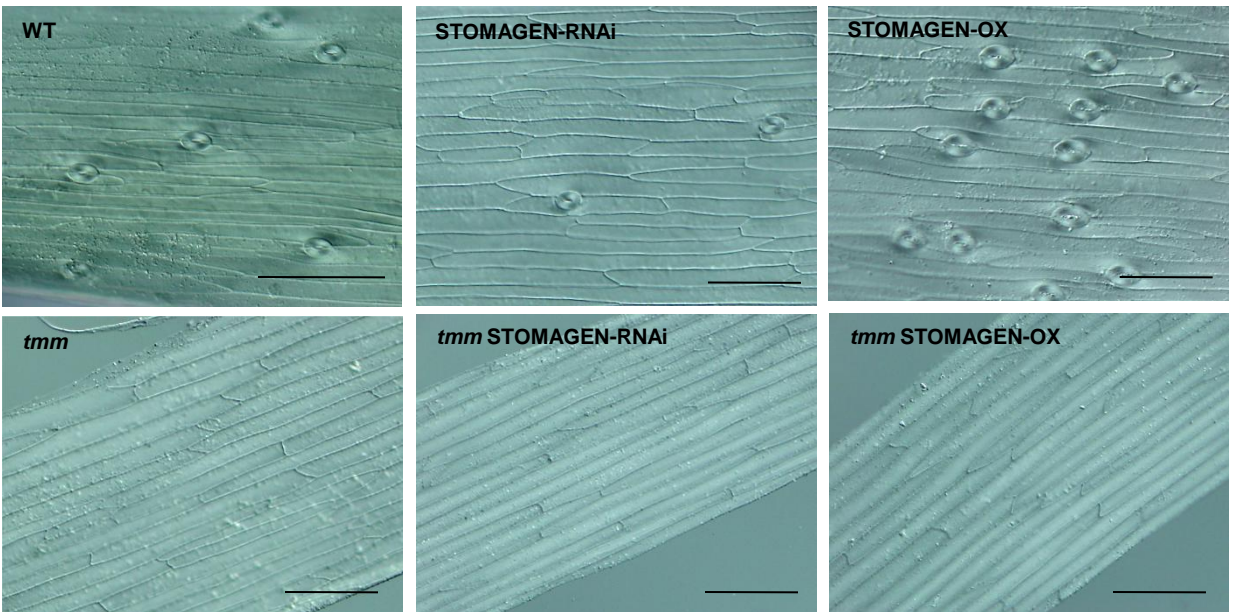
**c**, SDS-PAGE with CBB staining of each eluted fraction of the chromatography. The stomagen was recovered in fraction numbers from 10 to 13. A red arrow indicates the position of purified recombinant stomagen in the chromatography in **b**.

**d**, N-terminal sequences of the recombinant stomagen in each fraction from 10 to 13 as determined by Edman degradation.

**a**



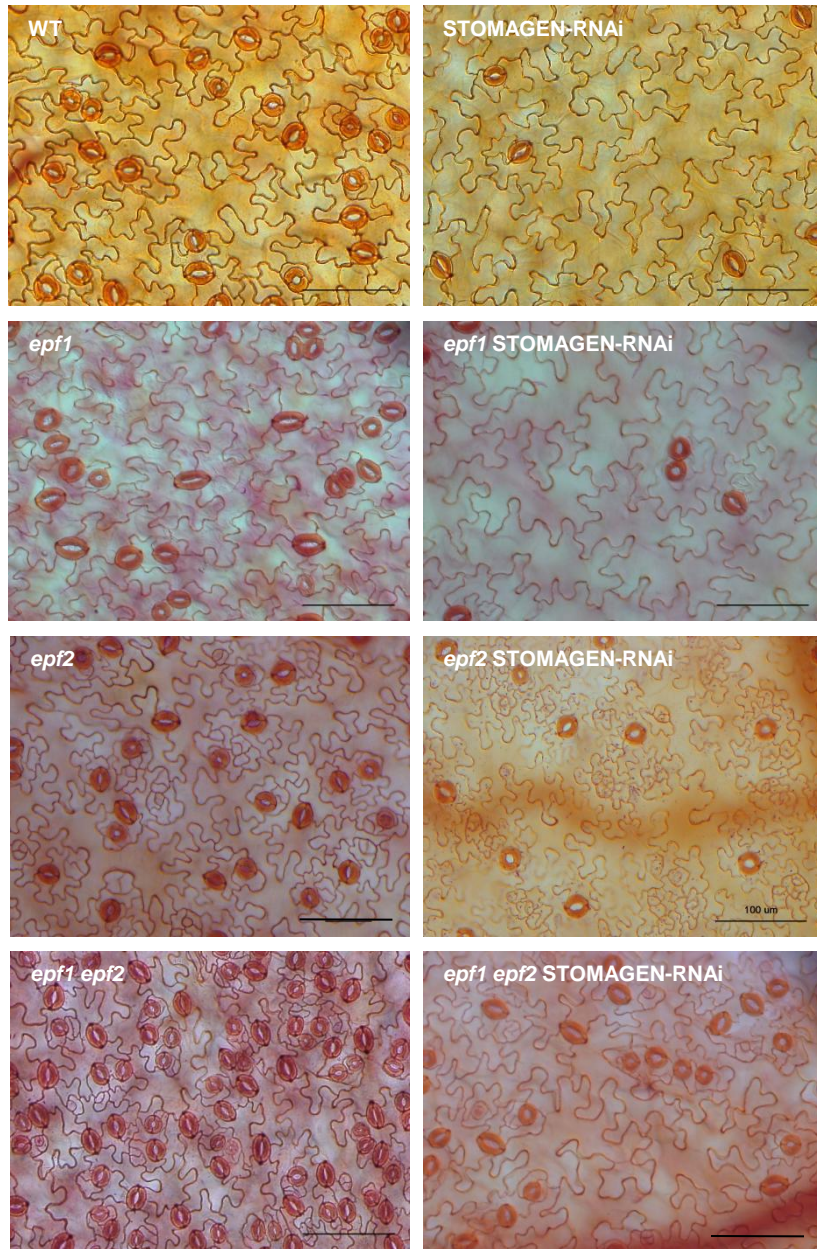
**b**



**Supplementary Figure 8. Representative images of STOMAGEN-OX or STOMAGEN-RNAi in *tmm* background.**

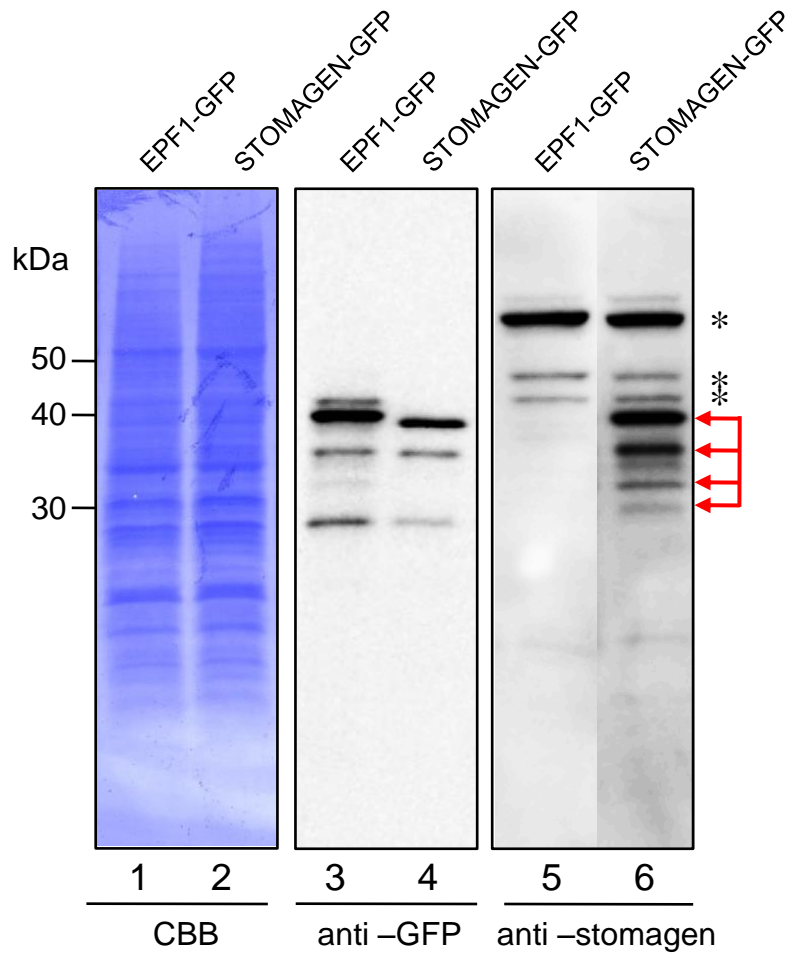
**a**, DIC images of the abaxial epidermis of mature first leaf of the wild type (WT), STOMAGEN-RNAi10, STOMAGEN-OX10, *tmm*, *tmm* STOMAGEN-RNAi8, *tmm* STOMAGEN-OX8. Quantitative data of stomatal density in each line are presented in Fig. 4b. Bars, 100  $\mu$ m.

**b**, DIC images of the stems of the wild type (WT), STOMAGEN-RNAi10, STOMAGEN-OX10, *tmm*, *tmm* STOMAGEN-RNAi8, *tmm* STOMAGEN-OX8. Quantitative data of stomatal density in each line are presented in Fig. 4b. Bars, 100  $\mu$ m.



**Supplementary Figure 9. Representative images of STOMAGEN-RNAi in *epf* backgrounds.**

DIC images of the abaxial epidermis of mature first leaf of each of the wild type (WT), STOMAGEN-RNAi10, *epf1*, *epf1* STOMAGEN-RNAi8, *epf2*, *epf2* STOMAGEN-RNAi8, *epf1 epf2*, and *epf1 epf2* STOMAGEN-RNAi8. Quantitative data of stomatal density in these lines are presented in Fig. 4 (c, d). Bars, 100 μm.



**Supplementary Figure 10. Specificity of anti-stomagen antibody used in this study.**

Both EPF1-GFP (lanes 1, 3 and 5) and STOMAGEN-GFP (lanes 2, 4 and 6) were expressed in tobacco-cultured BY-2 cells. Whole homogenate of the BY-2 cells were subjected to SDS-PAGE followed by either Coomassie blue (CBB) staining or by immunoblot with anti-GFP antibody or anti-stomagen antibody. Red arrows indicate the specific bands of STOMAGEN-GFP. Asterisks indicate non-specific signals.

**Supplementary Table 1. Co-expressed genes with *TMM*, *SDD1*, and *EPF1*.** *STOMAGEN* gene is colored in red. Known genes regulating stomatal development are colored in blue. This table was made using ATTED-II (<http://atted.jp/>). Details are in Full Method.

Rank	AGI code	Gene annotations
1	At1g04110	SDD1 (STOMATAL DENSITY AND DISTRIBUTION); serine-type endopeptidase
2	At1g33930	avirulence-responsive family protein / avirulence induced gene (AIG1) family protein
3	At1g80080	TMM (TOO MANY MOUTHS); protein binding / receptor
4	At5g60880	BASL (BREAKING OF ASYMMETRY IN THE STOMATAL LINEAGE )
5	At2g20875	EPF1 (EPIDERMAL PATTERNING FACTOR 1)
6	At5g19170	unknown protein
7	At1g34245	EPF2 (EPIDERMAL PATTERNING FACTOR 2)
8	At2g39690	unknown protein
9	At3g49260	iqd21 (IQ-domain 21); calmodulin binding
10	At5g53210	SPCH (SPEECHLESS); DNA binding / transcription factor
11	At1g26600	CLE9 (CLAVATA3/ESR-RELATED 9); protein binding / receptor binding
12	At1g06100	fatty acid desaturase family protein
13	At5g63050	EMB2759 (embryo defective 2759)
14	256752_at	
15	At3g06120	MUTE (MUTE); DNA binding / transcription factor
16	At3g27920	ATMYB0 (MYB DOMAIN PROTEIN 0); DNA binding / protein binding / transcription activator/ transcription factor
17	At4g12970	stomagen
18	At1g12860	SCRM2 (SCREAM 2); DNA binding / transcription factor
19	At3g57600	AP2 domain-containing transcription factor, putative
20	At1g17850	
21	At3g21950	methyltransferase
22	At3g61840	unknown protein
23	At1g24290	AAA-type ATPase family protein
24	At3g57180	GTP binding
25	At4g29020	glycine-rich protein
26	At2g40690	GLY1; glycerol-3-phosphate dehydrogenase (NAD+)
27	At3g61970	NGA2 (NGATHA2); transcription factor
28	At2g41340	RPB5D (RNA POLYMERASE II FIFTH LARGEST SUBUNIT, D); DNA binding / DNA-directed RNA polymerase
29	At3g63510	FAD binding / catalytic/ tRNA dihydrouridine synthase
30	At5g47380	unknown protein

