



Statistical organelle dissection of *Arabidopsis* guard cells using image database LIPS

SUBJECT AREAS:

IMAGING

PLANT CELL BIOLOGY

PLANT SCIENCES

BIOINFORMATICS

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Received
6 March 2012

Accepted
30 April 2012

Published
11 May 2012

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To comprehensively grasp cell biological events in plant stomatal movement, we have captured microscopic images of guard cells with various organelles markers. The 28,530 serial optical sections of 930 pairs of *Arabidopsis* guard cells have been released as a new image database, named Live Images of Plant Stomata (LIPS). We visualized the average organellar distributions in guard cells using probabilistic mapping and image clustering techniques. The results indicated that actin microfilaments and endoplasmic reticulum (ER) are mainly localized to the dorsal side and connection regions of guard cells. Subtractive images of open and closed stomata showed distribution changes in intracellular structures, including the ER, during stomatal movement. Time-lapse imaging showed that similar ER distribution changes occurred during stomatal opening induced by light irradiation or femtosecond laser shots on neighboring epidermal cells, indicating that our image analysis approach has identified a novel ER relocation in stomatal opening.

Quantitative and statistical analysis of intracellular structures or dynamics from a large scale image set has become trends in cell biological study. This is because that these approaches help reducing biased information from a single or a small number of representative microscopic images. Recent improvements of high throughput microscopy encourage the trends by simplifying the acquisition of a large-scale image set. The establishment and subsequent web-release of image databases are essential steps to public use and efficient data mining from such large-scale image data sets. In the plant sciences, the 10 or so image databases that have so far been released¹ can be roughly classified into two categories; a database for “protein localizome”, and a database for “cell biological dynamics”. The former group, derived from a so-called post-genomic approach, aims to determine the intracellular localization of all proteins and includes the typical FTFLP Database (<http://gfp.stanford.edu/>)²; GFP localizome database (<http://www.psb.ugent.be/papers/cellbiol/>)³; GFP database (<http://data.jic.bbsrc.ac.uk/cgi-bin/gfp/>)⁴; and AtNoPDB (<http://bioinf.scri.sari.ac.uk/cgi-bin/atnopdb/home/>)⁵. The latter group was established by researchers specializing in cell biology, and includes the typical databases of Plant Cell Imaging (<http://deepgreen.stanford.edu/>)⁶; the Illuminated Plant Cell (<http://www.illuminatedcell.com/>)⁷; Plant Organelles Database (<http://podb.nibb.ac.jp/Organelleme/>)⁸; and the Plant Organelles World (<http://podb.nibb.ac.jp/Organelleme/PODBworld/en/index.html>)⁹. These databases have highlighted the dynamics of intracellular structures in plants, and their high-resolution images have promoted a deeper understanding of cellular processes in the plant sciences, while their attractive pictures and movies have helped develop curiosity in the non-scientific community. The database is thought to have particular potential in addressing academic issues, however, there is no actual case in which a cell biological hypothesis has been formulated using an image database as far as we are aware.

Stomatal movement is an essential phenomenon for gas exchange and transpiration in higher plants, and is accomplished by significant changes in guard cell volume¹⁰. The key molecules identified in stomatal movement include photoreceptors, proton ATPase, protein kinases and ion channels, and recent studies have shown these molecules to be deeply involved in intracellular structure dynamics. For example, the potassium channel, KAT1,



Table 1 | List of the Arabidopsis transgenic lines used in this study

Fluorescent probe name	Target intracellular structures	Fluorescently tagged protein	References
HistoneH2B-RFP	Cell nuclei	<i>Nicotiana tabacum</i> Histone H2B2	44
GFP-tubulin	Microtubules	<i>Arabidopsis thaliana</i> β -Tubulin 6	30
GFP-EB1	Microtubule plus-ends	<i>Arabidopsis thaliana</i> End binding 1	31
GFP-ABD2	Actin microfilaments	Actin binding domain 2 of <i>Arabidopsis thaliana</i> Fimbrin 1	15
GFP-mTn	Actin microfilaments, cell nuclei	Actin binding domain of <i>Mus musculus</i> Talin	32
Mt-GFP	Mitochondria	Mitochondrial targeting domain of <i>Saccharomyces cerevisiae</i> Cytochrome c oxidase IV	33
CT-GFP	Chloroplasts	Chloroplast stroma targeting domain of <i>Arabidopsis thaliana</i> RecA	34
Autofluorescence in Wild type (Col-0)	Chloroplasts	-	-
GFP-PIP2a	Plasma membrane	<i>Arabidopsis thaliana</i> Plasma membrane intrinsic protein (PIP) 2a	6
GFP-ER	Endoplasmic reticulum	C-terminal HDEL (Endoplasmic reticulum retention signal) sequence	35
YFP-inserted MFP2	Peroxisomes	<i>Arabidopsis thaliana</i> Multifunctional protein (MFP) 2	39
ARA6-GFP	Endosomes	<i>Arabidopsis thaliana</i> Rab isoform (ARA) 6	36
GFP-VAMP727	Endosomes	<i>Arabidopsis thaliana</i> Vesicle-associated membrane protein (VAMP) 727	17
ERD2-GFP	<i>cis</i> -Golgi, ER	<i>Arabidopsis thaliana</i> ER retention defective (ERD) 2	37, 41
XYLT-YFP	<i>medial</i> -Golgi	N-terminus of <i>Arabidopsis thaliana</i> β (1,2)-xylosyltransferase (XYLT)	45
ST-mRFP	<i>trans</i> -Golgi, vacuoles, apoplast	N-terminus of <i>Rattus norvegicus</i> α (2,6)-sialyltransferase (ST)	40
GFP-VAM3	Vacuolar membrane	<i>Arabidopsis thaliana</i> Syntaxin of plant (SYP) 22/Vacuolar morphology (VAM) 3	38
GFP- δ TIP	Vacuolar membrane, ER	<i>Arabidopsis thaliana</i> δ -Tonoplast intrinsic protein (TIP)	6

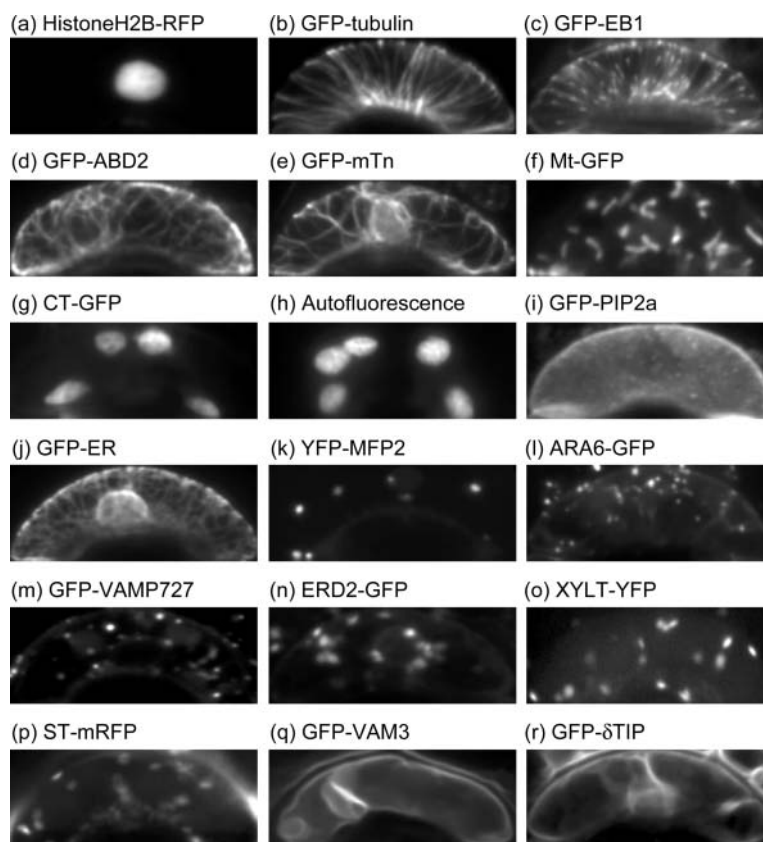


Figure 1 | Shape-normalized cell images in Arabidopsis guard cells. (a) Cell nuclei labeled by HistoneH2B-RFP. (b) Microtubules labeled by GFP-tubulin. (c) Microtubule plus-ends labeled by GFP-EB1. (d, e) Actin microfilaments labeled by GFP-ABD2 (d) and GFP-mTn (e). (f) Mitochondria labeled by Mt-GFP. (g, h) Chloroplasts labeled by CT-GFP (g) and autofluorescence (h). (i) Plasma membrane labeled by GFP-PIP2a. (j) Endoplasmic reticulum labeled by GFP-ER. (k) Peroxisomes labeled by YFP-MFP2. (l, m) Endosomes labeled by ARA6-GFP (l) and GFP-VAMP727 (m). (n) *cis*-Golgi labeled by ERD2-GFP. (o) *medial*-Golgi labeled by XYLT-YFP. (p) *trans*-Golgi labeled by ST-mRFP. (q, r) Vacuolar membranes labeled by GFP-VAM3 (q) and GFP- δ TIP (r). All of the image sizes were normalized into a mean size of 304 \times 119 pixels (19.5 \times 7.6 μ m).

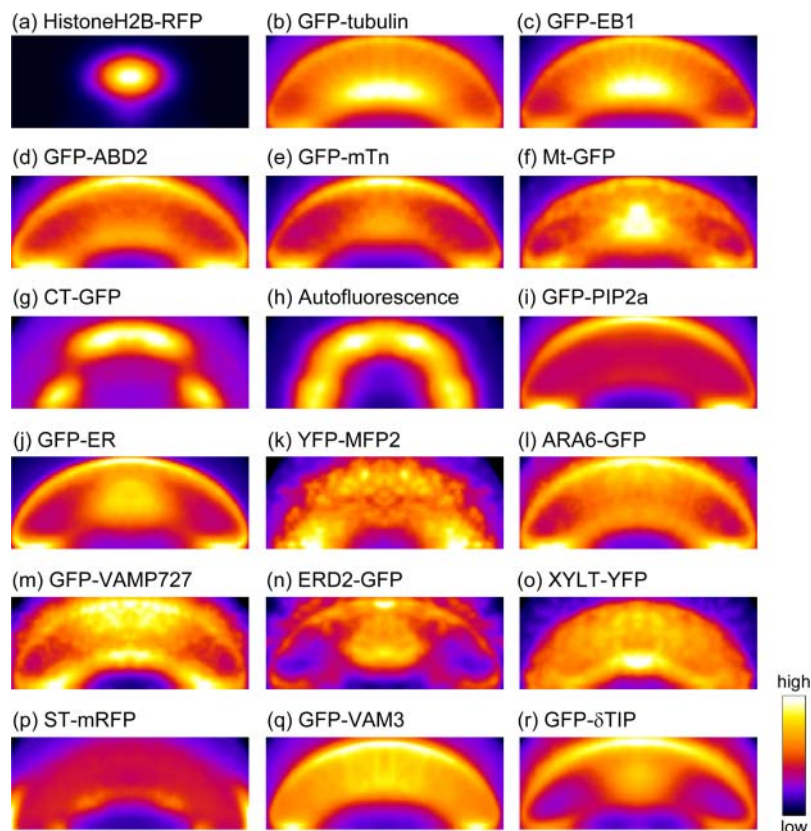


Figure 2 | Probability maps of the intracellular structures in guard cells. (a) Cell nuclei labeled by HistoneH2B-RFP. (b) Microtubules labeled by GFP-tubulin. (c) Microtubule plus-end labeled by GFP-EB1. (d, e) Actin microfilaments labeled by GFP-ABD2 (d) and GFP-mTn (e). (f) Mitochondria labeled by Mt-GFP. (g, h) Chloroplasts labeled by CT-GFP (g) and autofluorescence (h). (i) Plasma membrane labeled by GFP-PIP2a. (j) Endoplasmic reticulum labeled by GFP-ER. (k) Peroxisomes labeled by YFP-MFP2. (l, m) Endosomes labeled by ARA6-GFP (l) and GFP-VAMP727 (m). (n) *cis*-Golgi labeled by ERD2-GFP. (o) *medial*-Golgi labeled by XYLT-YFP. (p) *trans*-Golgi labeled by ST-mRFP. (q, r) Vacuolar membranes labeled by GFP-VAM3 (q) and GFP- δ TIP (r). These maps were obtained from group-averaged images from 100–120 guard cells, and are shown in pseudocolor. All of the image sizes were normalized into a mean size of 304×119 pixels ($19.5 \times 7.6 \mu\text{m}$).

is selectively internalized by abscisic acid from the plasma membrane with endosomes¹¹. On the other hand, the microtubule-binding protein, VfPIP, that was isolated as a protein interacting with the photoreceptor phototropin¹², supports previous observations that cortical microtubules undergo dynamic changes to their orientations with diurnal cycles¹³ and in response to light conditions¹⁴. Although intracellular dynamics are thought to be important for key signal molecule functions, comprehensive cell biological studies in guard cells have not yet been conducted.

Therefore, in order to provide new insights into the mechanisms involved in stomatal movement, we have conducted a comprehensive imaging analysis, focusing on the distribution and dynamics of intracellular structures in guard cells. Our microscopic image set has now been released as the Live Images of Plant Stomata (LIPS) database and is freely available at <http://hasezawa.ib.k.u-tokyo.ac.jp/lips/>. To provide a new understanding of the cell biological events during stomatal movements from this database, we developed an image processing framework to quantify the behavior of the various organelles at once. Application of this framework has identified a novel ER localization changes in stomatal opening.

Results

Establishment of the LIPS database. The wild type Col-0 and 17 transgenic Arabidopsis plants were prepared as a marker set of representative plant organelles (Table 1) and their guard cells were captured by $0.5\text{-}\mu\text{m}$ -interval serial optical sections with two channels

(fluorescent and bright field images) at random times during the diurnal cycle. Some fluorescent markers showed multiple localization patterns (Supplementary Fig. 1a–c, Table 1). A total of 28,530 sectional images have been designed as LIPS (Live Images of Plant Stomata) database and freely available at <http://hasezawa.ib.k.u-tokyo.ac.jp/lips/>. The LIPS database contains images of 50–60 pairs of guard cells in each probe, thus providing a sufficient number of images for statistical analyses. It can therefore be expected that visitors to the LIPS database will be able to obtain an integrative understanding of organellar form and distribution without being limited to only one representative image.

Probabilistic mapping to visualize average distributions of the organelles in guard cells. Guard cells have the dorsal-ventral polarity that deeply involves in stomatal movement. Therefore it is valuable to reveal the organelle locations for understanding its function in stomatal movement. However, a single microscopic snapshot is not enough to grasp an overall trend of the organelle localizations (Fig. 1). To overcome this problem, we visualized localization trends of each intracellular structure with probabilistic mapping techniques to average the localization information from the LIPS database (Fig. 2). The following is the image processing scheme developed in this study. First, the guard cell region was manually segmented with a bounding box (Supplementary Fig. 2a, b). All of the segmented image sizes were then normalized into a mean size of 304×119 pixels ($19.5 \times 7.6 \mu\text{m}$), with their fluorescent intensities also normalized to an average intensity of 0 with a standard

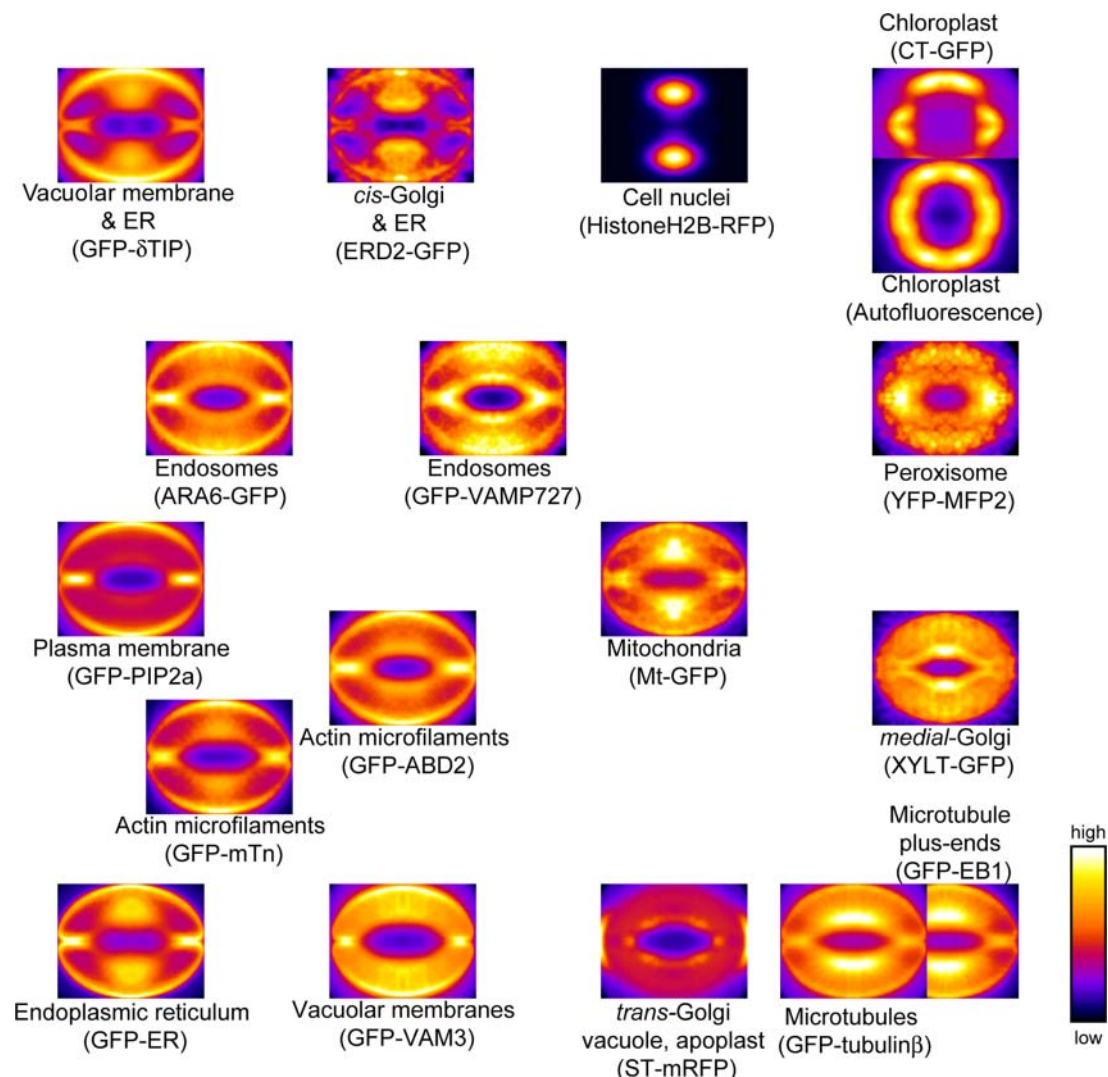


Figure 3 | Clustering of probability maps for each probe. Self-organizing map (SOM) of the 18 probability maps. The SOM was produced based on raster-scanned intensity profiles (**Supplementary Fig. 4**). Note that mitochondria, *medial*-Golgi and microtubules are mainly localized to the ventral side (lower right portion of the SOM), whereas endosomes, actin microfilaments, and the endoplasmic reticulum (ER) are mainly localized to the dorsal side (left portion of the SOM). The images are shown in pseudocolor. All of the image sizes were normalized into a mean size of 304×238 pixels ($19.5 \times 15.2 \mu\text{m}$).

derivation of 1 (**Fig. 1**, **Supplementary Fig. 2c**). Thereafter, group-averaged images were obtained for 100–120 guard cells (**Supplementary Fig. 2d**). For the 18 probes used in this study, the intracellular structure distributions were found to be symmetrical with respect to the guard cell major axis (**Supplementary Fig. 3**), and so we were able to divide the group-averaged guard cells in the middle and then again average the half-separated cell images (**Supplementary Fig. 2e, f**). Finally, we were able to derive the probabilistic maps in which higher intensities represented higher intracellular localization probabilities (**Fig. 2**).

Clustering analysis of the organelle probabilistic maps. Whereas the probabilistic maps of the actin microfilaments markers GFP-ABD2 and GFP-mTn were similar in appearance (**Fig. 2d, e**), the chloroplast autofluorescence and CT-GFP maps showed some differences in their patterns (**Fig. 2g, h**). We consider that this was because the numbers of chloroplasts in guard cells were slightly reduced in the CT-GFP expression line compared with wild type (**Supplementary Fig. 1d**). The probabilistic map of chloroplast autofluorescence in CT-GFP expression line showed similar patterns to that of CT-GFP (**Supplementary Fig. 1e**).

To objectively estimate such distribution differences of intracellular structures within the overall framework, we subsequently performed microscopic image clustering analysis¹⁵. As metrics to evaluate the intracellular localizations, we used raster-scanned intensity profiles in the probability maps (**Supplementary Fig. 4**). The three different clustering methods used, namely principle component analysis (PCA) (**Supplementary Fig. 5a**), hierarchical clustering (**Supplementary Fig. 5b**) and self-organizing map (SOM) (**Fig. 3**), all showed similar distributions. Nearby residents of the different probes, but similar targeting structures, including microtubules (GFP-tubulin β vs. GFP-EB1), chloroplasts (Autofluorescence vs. CT-GFP), actin microfilaments (GFP-ABD2 vs. GFP-mTn), and endosomes (ARA6-GFP vs. GFP-VAMP727) verified the validity of the metric patterns used in this study (**Fig. 3**). In the dendrogram, actin microfilaments markers GFP-ABD2 and GFP-mTn appeared in different clades (**Supplementary Fig. 5b**). It might be because GFP-mTn fluorescence leaked into cell nuclei (**Supplementary Fig. 1c**). The SOM also showed near location of the multiple organelle markers GFP- δ TIP and ERD2-GFP, in which the ER was commonly labeled (**Fig. 3**). This clustering analysis thus successfully provided an overall picture of structural locations within guard cells, with *medial*-Golgi and microtubules being located to the

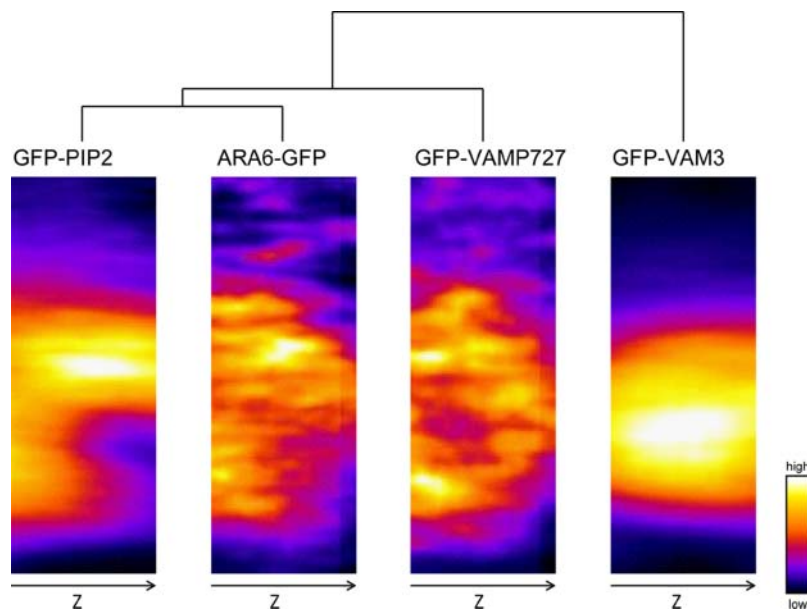


Figure 4 | Clustering of the probability maps of YZ sections of GFP-PIP2a, ARA6-GFP, GFP-VAMP727 and GFP-VAM3. We performed Z-axis registration using the cross-point in the bright field YZ sections (see **Supplementary Fig. 6a, b**). To facilitate GFP visualization, images are presented in pseudo-color. These maps were obtained from group-averaged images from 100–120 guard cells. All of the image sizes were normalized into a mean size of 73×200 pixels ($4.67 \times 12.8 \mu\text{m}$). The dendrogram was produced based on raster-scanned intensity profiles (**Supplementary Fig. 4**) with Euclidean distance and Ward's algorithm. Note that the image of ARA6-GFP is more similar to GFP-PIP2a than to GFP-VAMP727, which appears to be located around the vacuolar membrane as shown by GFP-VAM3.

ventral side (**Fig. 3**, lower right portion of the SOM) and actin microfilaments and ER located to the dorsal side and connecting regions (**Fig. 3**, lower left portion of the SOM).

Detection of detailed localization differences between ARA6-GFP and GFP-VAMP727. The endosomal markers ARA6-GFP and GFP-VAMP727 labeled dot-like endosomal components, and recent studies have shown that ARA6 is mainly involved in endosome-plasma membrane trafficking¹⁶, whereas VAMP727 acts in both endosome-plasma membrane and endosome-vacuolar membrane trafficking^{16,17}. Image clustering in this study showed that ARA6-GFP was more closely located to the plasma membrane marker GFP-PIP2a than to GFP-VAMP727 (**Fig. 3**, **Supplementary Fig. 5a**). The similar patterns of ARA6-GFP and GFP-PIP2a were more clearly demonstrated not only by clustering of XY sections but also by YZ sections (**Fig. 4**, **Supplementary Fig. 6c, d**), which were generated via Z-axis registration using the cross-point of bright field images (**Supplementary Fig. 6a, b**).

Changes in organelle distributions during stomatal movements.

We subsequently visualized intracellular localization changes during stomatal movements using subtractive images of the probability map in open and closed stomata. Although some transgenic plants, including lines expressing GFP-mTn, showed slightly decreased mean stomatal apertures as previously reported^{15,18}, all the transgenic plants generally showed differences in stomatal aperture, suggesting some degree of variation in their ability for stomatal movement (**Supplementary Fig. 7**). We first manually determined the apertures of all 930 stomata, which are also available in the LIPS database, and collected the top 30% (defined as opened stomata) and bottom 30% (defined as closed stomata) images. We then obtained the subtractive images of the probability map of open and closed stomata with registration by affine transformation (**Supplementary Fig. 8, 9, 10**), and finally visualized the comprehensive intracellular structural changes at their localizations during stomatal movement (**Fig. 5**). Noteworthy changes included highly condensed microtubules in the ventral pore edge side of opened stomata (**Fig. 5b, c**). This localization of microtubules was

suggestive of their polymerization from the pore edge regions when stomata opened. More interestingly, we found that the ER was located to the connecting region when stomata closed and in the dorsal side when stomata opened (**Fig. 5j**, arrow).

Direct observations of ER distribution changes during stomatal opening. Compared with cytoskeletons in guard cells^{13–15}, little is known about ER distribution changes during stomatal movement. To verify the ER dynamics that we found through image database analysis (**Fig. 5j**, arrow), we performed time-lapse imaging on GFP-ER during light-induced stomatal opening. We first captured the three-dimensional images of the ER in guard cells of closed stomata after 72 hours of dark treatment (**Fig. 6a**, before light treatment), before the guard cells were treated with $200 \mu\text{mol m}^{-2}\text{s}^{-1}$ white light treatment under microscopy. After 2 hours of light treatment, the stomata opened (**Fig. 6a, b**) and the three-dimensional images of the ER were recaptured (**Fig. 6a**, after light treatment). During light-induced stomatal opening, the GFP-ER fluorescence intensity increased on the dorsal side (**Fig. 6a**, arrowheads). Measurement of the dorsal/ventral fluorescence intensity ratio indicated that GFP-ER relocated to the dorsal side (**Fig. 6c**, **Supplementary Fig. 11**), suggestive of ER accumulation in the dorsal side during stomatal opening.

To further understand ER dynamics during stomatal movement, we performed cell manipulation experiments by micro-perforation using a focused femtosecond laser. As the pushing force from neighboring epidermal cells on guard cells is a key factor regulating stomatal movement, we attempted to perforate the neighboring epidermal cells and reduce the pushing force by focusing an intense femtosecond laser pulse train through the objective lens. The ultrashort infrared laser pulse (150 fs, 800 nm) induces laser ablation and results in multiphoton absorption at the laser focal point¹⁹. When the laser pulse was focused on the surface of epidermal cells, the nearest stoma opened with shrinkage of the targeted epidermal cells (**Fig. 7a, b**). Immediately after the laser-induced stomatal opening, the fluorescence intensity of GFP-ER increased at the dorsal side (**Fig. 7c**), similar to light-induced stomatal opening (**Fig. 6**). The very quick

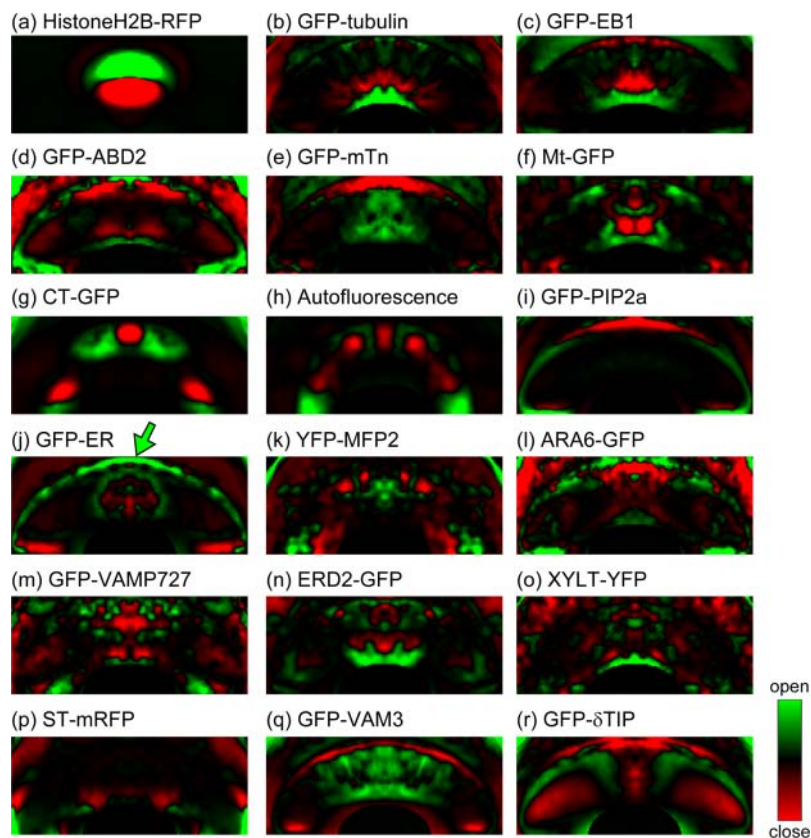


Figure 5 | Subtractive images of probability maps of the top 30% and bottom 30% of stomatal apertures. Green and red colors show the most prominent localizations in opened and closed stomata, respectively. (a) Cell nuclei labeled by HistoneH2B-RFP. (b) Microtubules labeled by GFP-tubulin. (c) Microtubule plus-end labeled by GFP-EB1. (d, e) Actin microfilaments labeled by GFP-ABD2 (d) and GFP-mTn (e). (f) Mitochondria labeled by Mt-GFP. (g, h) Chloroplasts labeled by CT-GFP (g) and autofluorescence (h). (i) Plasma membrane labeled by GFP-PIP2a. (j) Endoplasmic reticulum labeled by GFP-ER. (k) Peroxisomes labeled by YFP-MFP2. (l, m) Endosomes labeled by ARA6-GFP (l) and GFP-VAMP727 (m). (n) *cis*-Golgi labeled by ERD2-GFP. (o) *medial*-Golgi labeled by XYLT-YFP. (p) *trans*-Golgi labeled by ST-mRFP. (q, r) Vacuolar membranes labeled by GFP-VAM3 (q) and GFP- δ TIP (r). These subtractive images were obtained from 60–72 guard cell images, and are shown in pseudocolor. These maps were obtained from group-averaged images from 60–72 guard cells. All of the image sizes were normalized into a mean size of 303×128 pixels ($19.4 \times 8.2 \mu\text{m}$). Note that localization of GFP-ER increased in dorsal side of guard cells of open stomata (arrow).

localization changes suggested responding to mechanical stress associated with stomatal opening.

Discussion

In this study, we have established a new microscopic image database of *Arabidopsis* guard cells, which we refer to as LIPS. Although the LIPS database may be considered a type of cell biological image database, it can be distinguished from existing plant image databases by the following features. First, we focused on only one specific cell type, namely guard cells and, as far as we are aware, no image database focusing on a particular cell type exists in the plant sciences. Guard cells are attractive targets not only because of their importance for gas exchange and transpiration in higher plants, but also because of the ease by which their proteins can be localized. Second, we standardized the kinds and settings of microscopy used. Although image acquisition conditions are frequently standardized in image databases for protein localizomes, conditions are generally not standardized for cell biological image databases because the emphasis is on organellar shapes and movements, and observation with different magnifications is to be expected. Furthermore, in the case of a collective image database from numerous researchers, such as the Plant Organelle Database⁸, it is almost impossible to use the same types of microscopy. Finally, the LIPS database contains images of 50–60 pairs of guard cells in each probe, thus providing a sufficient number of images for statistical analyses. It can therefore be expected

that visitors to the LIPS database will be able to obtain an integrative understanding of organellar form and distribution without being limited to only one representative image. This series of features allowed us to perform data mining of organellar localization through simple probabilistic mapping techniques.

To visualize average distributions of the organelles, we introduced an image analysis pipeline combined with probabilistic mapping and image clustering. In the analysis of a magnetic resonance image (MRI), probabilistic mapping has proved useful for measuring specific brain region volume²⁰, visualizing neural activation²¹, and determining the orientation of white matter fibers²². In principle, probabilistic maps provide a powerful means of visualizing cell biological phenomena, such as organellar distribution. However, normalization of cell shapes and size are often a bottle-neck for practical use. In cultured animal cells, fibronectin micropatterns, which enforce cells to take a certain shape and prevent their migration, were developed to normalize cell shapes^{23–25}. The resulting probabilistic map, with shape-normalized cells, has been useful for quantitatively understanding the distribution of actin stress fiber²³ and endomembrane organization²⁴. Although controlling the shape and size of plant cells is technically difficult, mature guard cells fortunately show less individual size and shape variations, and we were able to successfully exploit this characteristic to perform spatial normalization by simple image processing.

Clustering of probabilistic maps allowed visualization and quantification of organelle distribution similarities (Fig. 3, 4,

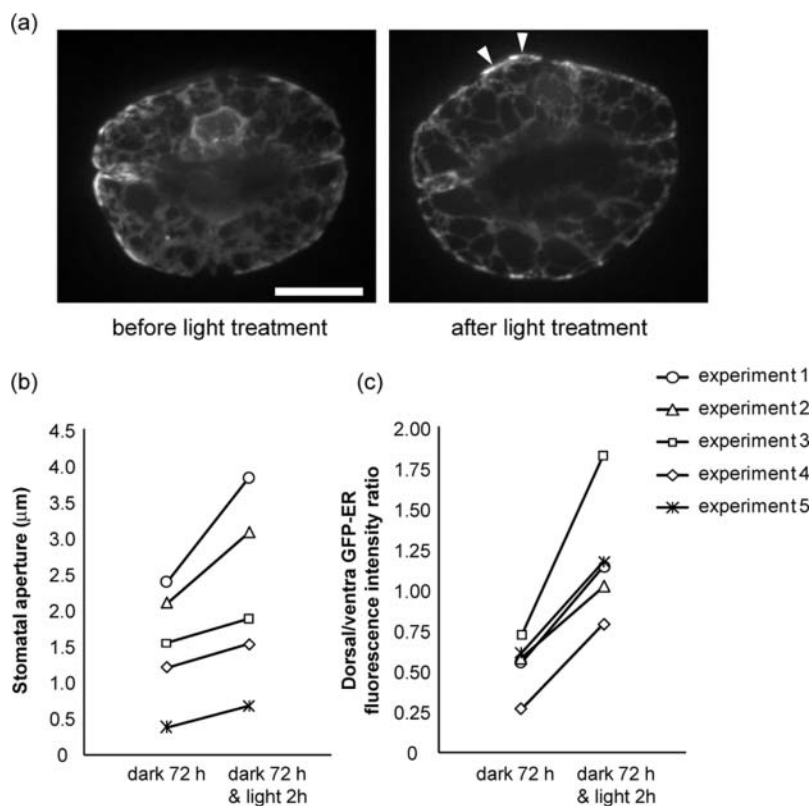


Figure 6 | Changes in ER distribution during light-induced stomatal opening. (a) Time-lapse images of GFP-ER in guard cells. The guard cells were dark-treated for 72 hours (before light treatment) and then irradiated with $200 \mu\text{mol m}^{-2}\text{s}^{-1}$ white light for 2 hours (after light treatment). Note that GFP-ER became much in the dorsal side after light treatment (arrowheads). Scale bar indicates $10 \mu\text{m}$. (b) Changes in stomatal aperture before and after the 2 hour white light treatment. (c) Changes in the dorsal/ventral GFP-fluorescence intensity ratios before and after light treatment. We measured the mean intensities of the line along the dorsal and ventral side (**Supplementary Fig. 11**). Results from five pairs of guard cells are shown.

Supplementary Fig. 5, 6c, d). We previously introduced image metrics specialized for cytoskeletal organization and image clustering methods using GFP-ABD2-labelled guard cell actin microfilaments¹⁵. In this study, we introduced new intensity profiles that have a broad utility for various intracellular structures as metrics for distribution patterns (**Supplementary Fig. 4**). We were thus able to detect detailed differences between ARA6-GFP and GFP-VAMP727 localization, with ARA6-GFP localization being more similar to the plasma membrane marker, GFP-PIP2a (**Fig. 4**). ARA6 is a plant-unique RAB5 member that acts in endosome-to-plasma membrane trafficking while VAMP727 executes membrane fusion at both vacuoles and the plasma membrane¹⁶. Thus our results clearly demonstrate the potential of our image analysis approach to detect the differences in molecular properties of specific proteins.

Our analysis using image subtraction favors the hypothesis that microtubules appear from the ventral side and reorganize into a radial orientation during stomatal opening as previously reported^{13,14}. A component of the microtubule organization center (MTOC), γ -tubulin, is localized adjacent to the ventral cell wall²⁶, and microtubule plus-ends grow outward from the ventral side²⁷, supporting the view that cortical microtubules become radially reorganized by growing from the MTOC beneath the ventral cell wall in stomatal opening. In addition, our analysis also demonstrated possible changes in ER distribution during stomatal movement. Time-lapse imaging directly visualized the hypothetical ER accumulation during stomatal opening (**Fig. 6, 7**). Although, the significance of the ER distributions on stomatal movement are unclear at present, it is likely that these structures are intimately involved in the turnover of guard cell membrane proteins that regulate stomatal movement via membrane trafficking. For example, ER-localized phospholipase A₂ β -mediated hydrolysis products have been suggested to move to

the plasma membrane and activate proton ATPase in light-induced stomatal opening²⁸. Alternatively, such ER accumulation may be some form of cellular mechanical response. It has been reported that pressure from a microneedle on the epidermal cell surface induces rapid ER accumulation²⁹. In the case of our femtosecond laser micro-perforation experiments, the suction force of the neighboring epidermal cells would suddenly be eliminated, and the guard cells would then be forced to swell up and to increase their tension forces, especially on the dorsal side. Future studies will clarify the significance of the organellar distribution changes during stomatal movement.

Methods

Plant materials and growth conditions. The transgenic lines used in this study were listed in **Table 1**. Seeds of transgenic Arabidopsis were kindly provided by Drs Minami Matsui (HistoneH2B-RFP); Takashi Hashimoto (GFP-tubulin³⁰); Jaideep Mathur (GFP-AtEB1³¹); Miyo Terao-Morita and Masao Tasaka (GFP-mTn³²); Andreas Nebenfuhr (Mt-GFP³³ (CS16263)); Maureen Hanson (CT-GFP³⁴); David W. Ehrhardt (GFP-PIP2a⁶ (CS84725); GFP- δ TIP⁶ (CS84727)); Scott Poethig (GFP-ER³⁵, E1728 (CS70123)); Takashi Ueda (ARA6-GFP³⁶, GFP-VAMP727¹⁷, ERD2-GFP³⁷, XYLT-YFP, ST-mRFP); Masa H Sato (GFP-VAM3³⁸); and the FTFLP Consortium (YFP-inserted MFP2³⁹ (CS37004)). ST-mRFP labels not only Golgi bodies but the excess is also localized to vacuoles and the apoplast⁴⁰. Furthermore, ERD2-GFP and GFP- δ TIP not only label Golgi bodies and the vacuolar membrane, respectively, but their excess is localized to the ER^{41,42}. Arabidopsis plants were grown on soil-vermiculite, as described by Naito et al. (1994)⁴³, in growth chambers set at 23.5°C , a 60% relative humidity, and a 12/12-hour light/dark cycle using $100 \mu\text{mol m}^{-2}\text{s}^{-1}$ white lights. Fully-expanded rosette leaves of 4- to 5-week-old plants were used for image acquisition.

Microscopic image acquisition and analysis. To capture stomatal images, The abaxial epidermis was peeled off and floated, with its inner surface facing down, on a glass slide covered with basal buffer [50 mM KCl, 5 mM 2-(N-morpholine)ethanesulphonic acid (MES)-Tris, pH 6.5, and 10 mM CaCl_2], and then covered by a cover glass (Matsunami, Osaka, Japan). The slide glass was placed onto the inverted

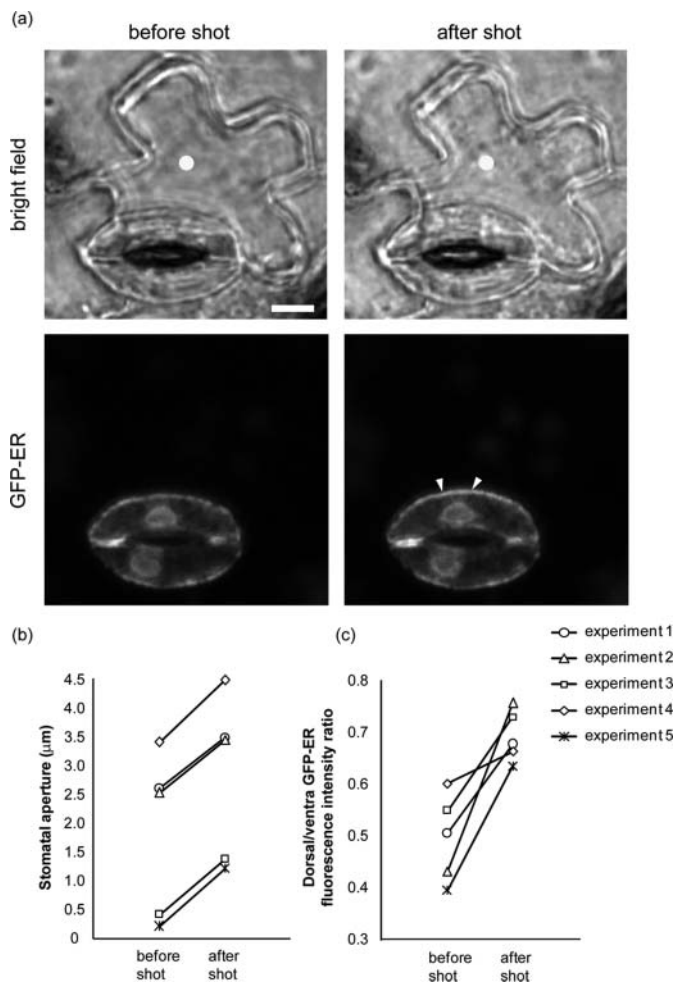


Figure 7 | Stomatal opening and ER dynamics induced by femtosecond laser micro-perforation. (a) The bright field and GFP-ER fluorescence images in guard cells before and after laser perforation. Laser focal points are indicated as yellow dots. Note that GFP-ER localization increased in stomatal opening (arrowheads). Scale bar indicates 10 µm. (b) Changes in stomatal aperture before and after the laser perforation. (c) Changes in the dorsal/ventral GFP-fluorescence intensity ratios before and after a femtosecond laser shot. We measured the mean intensities of the line along the dorsal and ventral side (**Supplementary Fig. 11**). Results from five pairs of guard cells are shown.

platform of a fluorescence microscope (IX70, Olympus) equipped with an UplanApo 100x/1.35 oil iris objective lens and a CSU10 scanning head (Yokogawa, Tokyo, Japan) together with a cooled CCD camera head system (CoolSNAP HQ, PhotoMetrics). At image acquisition, pixel width (0.064 µm) and step size for three-dimensional imaging (0.5 µm) were fixed. Image processing was performed using ImageJ (<http://rsbweb.nih.gov/ij/>) with our original JAVA plug-in package (<http://hasezawa.ib.ku-tokyo.ac.jp/zp/Kbi/>) and ImageJ macro (<http://hasezawa.ib.ku-tokyo.ac.jp/zp/Kbi/HigPDMClustering>). Intensity normalization was performed by the “avg0sd1” mode of KbiStkFilter plug-in. The probability map images were generated by the ImageJ macro `hig_Pdmapping.ijm`. The raster-scanned intensity profiles were measured by the ImageJ macro `hig_Lineprofile.ijm`. SOM clustering was performed by the “som” mode of KbiClustering plug-in. The affine transformation was performed with Adobe Photoshop CS4. PCA analysis and hierarchical clustering were performed using MINITAB (Minitab Inc., <http://www.minitab.com>).

Time-lapse observations of light-induced stomatal opening. The seven-day seedlings were transferred onto a cover glass (Matsunami) and placed onto the inverted platform of a microscope with a CSU10 scanning head (Yokogawa). To prevent drying, 3 mm-thick 1% agarose gel with basal buffer was placed onto the seedling. For light treatment, the seedlings were irradiated with 200 µmol m⁻²s⁻¹ white light for 2 hours.

Experimental setup of the femtosecond laser irradiation. Femtosecond laser pulses from a regeneratively amplified Ti:sapphire femtosecond laser system (Spectra-

Physics, Hurricane, 800 nm, 150 femtoseconds, 1 kHz) were introduced to an inverted microscope (IX71, Olympus) with a FV300 laser scanning confocal system (Olympus), and focused onto the peeled epidermal cells through a 40× objective lens (UMPlanFI, 0.75 NA, Olympus). The 125 pulse repetitions were detected with a mechanical shutter (gate time 1/8 sec.) and delivered to a fixed point on the sample. The laser pulse energy was tuned by a half-wavelength (λ/2) plate and dual polarizers. Laser pulses were collimated by dual convex lenses before the microscope, and the laser focal point was tuned to the plane of the image on the microscope. The laser pulse energy through the objective was measured by attaching a laser power meter (ORIEL, model AN/2) to the objective and was tuned to be 70 nJ/pulse.

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Acknowledgements

The authors thank the ABRC for providing seeds of transgenic Arabidopsis lines. We also thank Dr. Sachihiko Matsunaga of Tokyo University of Science for advice on the cell manipulation experiments. This work was financially supported in part by a Grant-in-Aid for Scientific Research on Priority Areas to S.H. (No. 23012009), by a Grant-in-Aid for Scientific Research on Innovative Areas to S.H. (No. 22114505), by a Grant-in-Aid for Young Scientists (B) to N.K. (No. 24770038), by a Grant-in-Aid for Research Activity start-up to T.H. (No. 22870004) from the Japanese Ministry of Education, Science, Culture, Sports and Technology, and an Advanced Measurement and Analysis grant from the Japan Science and Technology Agency (JST) to S.H.

Author contributions

T.H. designed and performed experiments, analyzed and wrote the paper. N.K. developed the ImageJ plug-ins. Y.H. developed femtosecond laser irradiation system. K.A. observed Golgi markers. K.E. and T.U. established endosomes and Golgi markers, and contributed through discussion and revision of the paper. N.K. and S.H. supervised this project. All authors reviewed the manuscript.

Additional information

Supplementary information accompanies this paper at <http://www.nature.com/scientificreports>

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

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How to cite this article: Higaki, T. *et al.* Statistical organelle dissection of Arabidopsis guard cells using image database LIPS. *Sci. Rep.* **2**, 405; DOI:10.1038/srep00405 (2012).