Pulses of Ca$^{2+}$ coordinate actin assembly and exocytosis for stepwise cell extension

Norio Takeshita$^{ab,1}$, Minoas Evangelinos$^{a,c}$, Lu Zhou$^{d,e}$, Tomoko Serizawa$^b$, Rosa A. Somera-Fajardo$^a$, Ling Lu$^f$, Naoki Takayab$^a$, G. Ulrich Nienhaus$^{d,g,h}$, and Reinhard Fischer$^a$

*Department of Microbiology, Institute for Applied Bioscience, Karlsruhe Institute of Technology, 76131 Karlsruhe, Germany; $^d$Faculty of Life and Environmental Sciences, University of Tsukuba, Tsukuba 305-8572, Japan; $^g$Faculty of Biology, University of Athens, Athens 15784, Greece; $^h$Institute of Applied Physics, Karlsruhe Institute of Technology, 76131 Karlsruhe, Germany; $^i$Institute of Nanotechnology, Karlsruhe Institute of Technology, 76344 Eggenstein-Leopoldshafen, Germany; $^j$College of Life Sciences, Nanjing Normal University, 210023 Nanjing, China; $^k$Department of Physics, University of Illinois at Urbana–Champaign, Urbana, IL 61801; and $^l$Institute of Toxicology and Genetics, Karlsruhe Institute of Technology, 76344 Eggenstein-Leopoldshafen, Germany

Many eukaryotic cells grow by extending their cell periphery in pulses. The molecular mechanisms underlying this process are not yet fully understood. Here we present a comprehensive model of stepwise cell extension by using the unique tip growth system of filamentous fungi. Live-cell imaging analysis, including superresolution microscopy, revealed that the fungus *Aspergillus nidulans* extends the hyphal tip in an oscillatory manner. The amount of F-actin and secretory vesicles (SV) accumulating at the hyphal tip oscillated with a positive temporal correlation, whereas vesicle amounts were negatively correlated to the growth rate. The intracellular Ca$^{2+}$ level also pulsed with a positive temporal correlation to the amount of F-actin and SV at the hyphal tip. Two Ca$^{2+}$ channels, MidA and CchA, were needed for proper tip growth and the oscillations of actin polymerization, exocytosis, and the growth rate. The data indicate a model in which transient Ca$^{2+}$ pluses cause depolymerization of F-actin at the cortex and promote SV fusion with the plasma membrane, thereby extending the cell tip. Over time, Ca$^{2+}$ diffuses away and F-actin and SV accumulate again at the hyphal tip. Our data provide evidence that temporally controlled actin polymerization and exocytosis are coordinated by pulsed Ca$^{2+}$ influx, resulting in stepwise cell extension.

Significance

The day–night rhythm in living organisms controls many processes, which then oscillate in a circadian manner. In addition, there are many processes that appear continuous but the underlying mechanisms oscillate with distinct periods. For example, eukaryotic cells grow by extending their cell periphery in pulses rather than by continuous extension. Here we investigate the molecular basis for such oscillatory growth by using the unique tip growth system of filamentous fungi, where actin assembly, exocytosis, and growth rate were observed simultaneously. We provide evidence that temporally controlled actin assembly and exocytosis are coordinated by pulsed Ca$^{2+}$ influxes, resulting in stepwise cell extension. This mechanism allows the cells to respond more quickly to both internal and environmental cues, including chemical and mechanical stimuli.

Edited by Jay C. Dunlap, Geisel School of Medicine at Dartmouth, Hanover, NH, and approved April 13, 2017 (received for review January 6, 2017)


The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

1To whom correspondence should be addressed. Email: takeshita.norio.gf@u.tsukuba.ac.jp

This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1700204114/-/DCSupplemental.

www.pnas.org/cgi/doi/10.1073/pnas.1700204114

PNAS | May 30, 2017 | vol. 114 | no. 22 | 5701–5706

C

ells extend their periphery before division and thereby maintain their original size and shape. Spatial and temporal observations have shown that cell extension does not occur continuously but stepwise. This phenomenon is shared by fission yeast (1), filamentous fungi (2), mammalian polarized neurons (3), nonpolarized cells (4), plant root hairs (5), and pollen tubes (6). Stepwise cell extension is driven by oscillations in the supply of proteins and lipids mediated by vesicle trafficking and cytoskeleton dynamics. Intracellular Ca$^{2+}$ levels regulate actin assembly and vesicle fusion (7, 8). The Ca$^{2+}$ level also oscillates and, consequently, could be closely related with stepwise cell extension (4–6). The molecular mechanism underlying discontinuous cell extension, however, has remained elusive.

Filamentous fungi grow as highly polarized tubular cells that indefinitely extend the cell body at one end in a process called “tip growth.” During tip growth the cell-extension site is maintained at the hyphal tip, where accumulation of F-actin, active exocytosis, and growth rate can be monitored simultaneously. Thus, filamentous fungi are excellent systems for the analysis of stepwise cell extension and the understanding of cell-shape determination (9–11). The extension of hyphal tips requires constant enlargement of the cell membrane. The supply of proteins and lipids is achieved by vesicle trafficking via the actin and microtubule cytoskeletons with their corresponding motor proteins (12–16). Microtubules serve as tracks of secretory vesicles (SV) for long-distance transport to the hyphal tip and are important for rapid hyphal growth (17). Actin cables, formed from the hyphal tip in retrograde direction, are involved in exocytosis and SV accumulation before exocytosis (18, 19). Vesicle accumulation before secretion results in the formation of a fungal-specific structure, called the Spitzkörper (20). Besides these fungal-specific microstructures, most constituents of the fungal-tip growth apparatus are shared with other eukaryotic cells that extend in a stepwise fashion. Hence, filamentous fungi are great models for studying the mechanism of oscillatory growth.

Stepwise cell extension at hyphal tips was reported in several filamentous fungi 20 y ago (2); still, only a few details on the molecular mechanism have been revealed since. Our recent study using superresolution fluorescence microscopy revealed that the membrane-associated polarity marker, TeaR (21), transiently assembles at the hyphal tip membrane of *Aspergillus nidulans* and disperses along the membrane after exocytosis, which inserts a new membrane and results in local membrane extension (22). This observation suggested a “transient polarity assembly model,” which explains that fungal-tip cells extend by a repetition of coordinated steps, assembly/disassembly of TeaR, actin polymerization, and exocytosis, rather than by constant elongation (22, 23). Oscillations of the Ca$^{2+}$ level have been observed at hyphal tips in filamentous fungi (24). These observations strongly suggest that stepwise extension of hyphal tips requires the concerted action of Ca$^{2+}$, actin, and exocytosis.
Here we demonstrate critical correlations between the intracellular Ca\(^{2+}\) level, actin polymerization, exocytosis, and cell extension at fungal tips, and provide evidence that temporally controlled actin polymerization and exocytosis drive stepwise cell extension, which are coordinated by pulsed Ca\(^{2+}\) influx. Because the switch from yeast to the hyphal growth form is often a pathogenicity factor in animal and plant pathogens, and the hyphal secretion machinery is also required for the production of commercial enzymes and reagents, understanding the tip-growth machinery may impact medical and agricultural antifungal agent development and the improvement of fungi as valuable biotechnological work horses (25–27).

**Results**

**Oscillation of the Fungal-Tip Cell-Extension Rate.** We investigated the cell-extension rate of hyphae in *A. nidulans* for 30 min in 10-s intervals (Fig. 1A and Movie S1). The average cell-extension rate of 1.8 \(\mu\)m/min (0.03 \(\mu\)m/s) was derived from a kymograph along the growth axis (Fig. 1B). When the extension distance of the tip position was measured every 20 s (Fig. 1C), a line graph showed unevenness and fluctuation of the distance from 0.4 to 1 \(\mu\)m in 20 s, indicating pulsed growth of the hyphae (Fig. S1 A). Because the dynamics was not reliably measurable as a result of the limited spatial resolution of conventional light microscopy, we applied superresolution photoactivation localization microscopy (PALM) (28, 29). We imaged hyphae producing a fusion protein of chitin synthase class III (ChsB), which localizes at the plasma membrane and is transported by the SV (30), and mEosFP\(_{\text{thermo}}\), a thermostable, monomeric variant of the green-to-red photoconverting fluorescent protein, EosFP (32, 33), is widely used for conventional fluorescence imaging. The PALM images showed ChsB clusters at the apical membrane. We applied a “moving-window binning” technique to generate sequences of PALM images in 20-s intervals for a total duration of 140 s (Fig. 1D) (22). The temporal changes of the apical cell edges were compared in 20-s intervals (Fig. 1E). Larger extensions of the apical membrane were occasionally observed from 40 to 60 s, from 100 to 120 s, and from 120 to 140 s (Fig. 1E and Fig. S1B). These results revealed that the cell-extension rate of hyphae is not constant but varies in an oscillatory manner.

Next, we asked whether polymerization of actin and the amount of SV at the hyphal tips correlate with the oscillation of the cell-extension rate. F-actin and SV were visualized by fluorescence of GFP-tagged tropomyosin (TpmA) and mCherry-tagged ChsB, respectively (18, 30). Prominent signals were visible at the hyphal

![Fig. 1. Stepwise growth of hyphae.](image-url)
tip, and time-resolved recording and frame analysis by kymographs revealed that the signal intensity oscillated (Fig. 1 F and G and Movie S2). The line profiles corresponding to the signal intensity of F-actin (green) and SV (red) indicated oscillations with peaks at similar times (Fig. 1H). The mean interval of the intensity of F-actin peaks was 29 \pm 8 \text{s} (n = 50 peaks in 20 hyphae), whereas the mean interval of peaks of the SV was comparable to the one of F-actin, 30 \pm 7 \text{s} (n = 50 peaks in 20 hyphae). To determine the temporal relationship between the presence of F-actin and the SV, we calculated the normalized cross-correlation of their signal intensities (Fig. U and Fig. S1C). The value at the central peak is 0.50, indicative of a positive correlation between the signals of F-actin (Fig. 1H, green) and the SV (Fig. 1H, red) (1 and –1 represent perfect positive and negative correlation, respectively). There were a few second delays in the signals of the SV in comparison with the signals of F-actin. These results indicate that SV accumulate during actin polymerization phases and SV are depleted because of exocytosis during actin depolymerization. The distance of tip extension (measured every 20 s) and the SV signal showed a negative correlation (Fig. S1D).

**Negative Correlation Between the SV Amount at Hyphal Tips and the Cell-Extension Rate.** Another model filamentous fungus, *Neurospora crassa*, is advantageous for cell imaging because of its larger cell size, faster cell-extension rate, and more distinct Spitzenkörper (20). The cell-extension rate of *N. crassa* was \( \sim 10 \text{ times faster} \) (0.2 \( \mu \text{m/s} \)) than the one of *A. nidulans* (2.4 \( \mu \text{m/s} \)) and Movie S3) and oscillated between 0.15 and 0.45 \( \mu \text{m/s} \) (see below). GFP-tagged chitin synthase class III (CHS-1) continuously localized to the Spitzenkörper at the hyphal tip (Fig. 2A) (34). The signal intensity of CHS-1-GFP at the Spitzenkörper was displayed in a kymograph along the growth axis (Fig. 2B). The line graph indicates oscillations of the GFP intensity (Fig. 2C). The mean period between peaks was 17 \pm 6 \text{s} (n = 30 peaks in 4 hyphae) (Fig. 2D).

The signal intensity of CHS-1-GFP at the Spitzenkörper and the hyphal cell-extension rate were measured every 2 s (Fig. 2E). When the GFP intensity increased, the cell-extension rate decreased, whereas a decrease in the GFP intensity was accompanied by an increase in the cell-extension rate. Normalized cross-correlation analysis yielded a clear negative correlation between CHS-1-GFP intensity and cell-extension rate, with a value of –0.54 at the central peak (Fig. 2F). These results again reveal that SV accumulate at the hyphal tip during slow growth phases and that the SV amount decreases by active exocytosis during fast growth phases. Sometimes, the Spitzenkörper moved from the center of the hyphal apex to the left or to the right; up and down movement was not resolved in our experiment (Fig. S1E). Whereas the cell-extension rate in *N. crassa* was \( \sim 10 \text{ times faster} \) than that of *A. nidulans*, the oscillation period of SV accumulation of *N. crassa* was half that of *A. nidulans*. It appears that faster growing hyphae exhibit shorter oscillation periods of their cell extension (2).

**Intracellular Ca\textsuperscript{2+} Oscillation Synchronizes Actin Polymerization and SV Accumulation.** Intracellular Ca\textsuperscript{2+} levels regulate actin assembly and vesicle fusion (7, 8). The red-fluorescent Ca\textsuperscript{2+} biosensor R-GECO was produced in *A. nidulans*. No obvious morphological changes of the cells because of the expression of the foreign protein were observed under the tested conditions. R-GECO emission varies with the concentration of Ca\textsuperscript{2+} (Fig. S2A) (35). When grown in media supplemented with 1 \( \mu \text{M} \) CaCl\textsubscript{2}, pulses of the R-GECO signal were observed (Fig. 3A and Movie S4). The R-GECO pulses in Fig. 3A were visualized in a kymograph as sharp lines and quantified in a line graph (Fig. 2B and C). The mean interval between peaks was 26 \pm 7 \text{s} (n = 52 peaks in 15 hyphae) (Fig. 3D), comparable to the one of F-actin and the SV (Fig. 1F). The R-GECO signal appeared as a tip-high gradient and diffused backward (Fig. S2B and C). Each hypha showed the signal with different timing (Fig. S2D). Such R-GECO pulses continued for multiple times before they disappeared (maximum \( n = 8 \) in 180 s), probably because of a limited turnover of R-GECO. The fluorescence of R-GECO could not be detected in media without CaCl\textsubscript{2} or with 1 \( \mu \text{M} \) CaCl\textsubscript{2} + 10 mM EGTA (Fig. S3A and B), indicating that the increase of the intracellular Ca\textsuperscript{2+} level is induced by the influx of Ca\textsuperscript{2+} to the cells at hyphal tips. R-GECO pulses were observed with this method in rapidly growing hyphae but not in germlings.

We compared the signals of R-GECO and GFP-TpmA (F-actin) or BglA-GFP (β-glycosidase, secreted protein as a marker for the SV) (22). The fungal cells producing R-GECO and GFP-TpmA, cultured in medium supplemented with 1 \( \mu \text{M} \) CaCl\textsubscript{2}, showed oscillating signals of GFP and R-GECO (Fig. 3E and Movie S5). Kymographs along the growth axis indicated temporal changes of signal intensities (Fig. 3E). Line profiles corresponding to the signal intensities of F-actin (Fig. 3E, green) and the Ca\textsuperscript{2+} concentration (Fig. 3E, red) at the tip indicated that oscillations of peak intensities had similar periods (29 \pm 8 and 26 \pm 7 \text{s}, respectively) and were synchronized (Fig. 3F). Normalized cross-correlation analysis yielded a positive correlation between the concentrations of F-actin (Fig. 3F, green) and Ca\textsuperscript{2+} (Fig. 3F, red), with the central peak at 0.40 (Fig. 3G) and indicated that the peaks of Ca\textsuperscript{2+} appeared a few seconds earlier than those of F-actin. These results are in agreement with the notion that Ca\textsuperscript{2+} influx at the growing hypha induces actin depolymerization.

Hyphae producing R-GECO and BglA-GFP (SV) showed oscillations of the GFP intensity at the tip and R-GECO pulses (Fig. 3H and Movie S6), with similar periods (30 \pm 7 and 26 \pm 7 \text{s}, respectively), which were synchronized (Fig. 3J). Normalized cross-correlation analysis yielded a positive correlation with a
central peak value of 0.43 (Fig. 3J) and a few seconds delay of BglA-GFP. These results indicate that Ca^{2+} influx affords exocytosis mediated by fusion of the SV with the plasma membrane, as well as actin depolymerization.

**Ca^{2+} Channels Coordinate Ca^{2+} Influx, Actin Polymerization, and Cell-Extension Rate.** Several Ca^{2+} channels, pumps, and transporters function in different organelles, such as the plasma membrane, endoplasmic reticulum, Golgi, mitochondria, and vacuole in fungi (36). The Ca^{2+} channels at the plasma membrane, Mid1 and Cch1p, of *Saccharomyces cerevisiae* share the single pathway that responds to environmental stresses and ensures cellular Ca^{2+} homeostasis (37–39). Deletion of these orthologs in *A. nidulans*, *midA* and *cchA*, caused a defect in polarized growth and cell wall synthesis (40). Deletion of *midA* or *cchA* often caused abnormal branching and de-polarized hyphae (Fig. 4A and Fig. S3C), and addition of 10 mM CaCl_{2} partially restored normal morphology.

Fluorescence imaging showed no pulse of R-GECO fluorescence in the *midA*- or *cchA*-deletion strain grown in the media supplemented with 1 μM CaCl_{2}, and no oscillation of actin polymerization determined by GFP-TpmA fluorescence (Fig. 4B and Fig. S3D), in contrast to the wild-type (Fig. 3E). No oscillation of the cell-extension rate was apparent (Fig. 4C), and cell extension in the deletion strain was too slow for microscopy to show the oscillations of the hyphal tip extension. Addition of 10 mM CaCl_{2} occasionally caused Ca^{2+} entry into the deletion strains, but restored neither the oscillation of Ca^{2+} influx nor the oscillation of actin polymerization (Fig. 4D). Ca^{2+} influx and actin polymerization in the wild-type strain remained oscillatory in the presence of 10 mM CaCl_{2} (Fig. S3E). These results indicate that pulsed Ca^{2+} influx effected by Ca^{2+} channels is required for the oscillation of actin polymerization and the cell-extension rate, resulting in proper polarized growth of hyphae, which is also supported by the result that the treatment of calcium ionophore A23187 disrupted the oscillation of Ca^{2+} and actin polymerization and caused swollen hyphal tips by depolarized growth (Fig. S3F).

**Discussion**

The hyphal tip of filamentous fungi comprises an excellent model to investigate polarity determination and maintenance, as well as polar cell growth, which occurs in a pulsed fashion. We were able to simultaneously analyze actin polymerization, SV fusion, and the intracellular Ca^{2+} concentration. Our findings

---

**Fig. 3.** Oscillation of Ca^{2+} influx correlated with F-actin and the SV. (A) Fluorescence image sequence of R-GECO. The elapsed time is given in seconds. (Scale bar, 5 μm.) (B) Kymograph along the hypha shown in A. Total 120 s. (Scale bar, 5 μm.) (C) Fluorescence intensity around the tip of hypha in B is plotted. Arrows and numbers indicate the time of Ca^{2+} peaks. (D) Distribution of intervals between two peaks. (Scale bar, 2 μm.) (E) Fluorescence intensity of F-actin (green) and Ca^{2+} (red) along the apex of the growing hypha between dotted lines in E. (G) Normalized cross-correlation of F-actin signal (GFP-TpmA) and Ca^{2+} signal (R-GECO). (H) Fluorescence images and kymographs along the growth axis of SV (BglA-GFP; green) and Ca^{2+} (R-GECO; red). Total 180 s. (Scale bar, 1 μm.) (I) Fluorescence intensity of SV (green) and Ca^{2+} (red) along the apex of the growing hypha between dotted lines in H. (J) Normalized cross-correlation of SV signal (BglA-GFP) and Ca^{2+} signal (R-GECO).

**Fig. 4.** Deletion of Ca^{2+} channel. (A) Hyphal morphology of the *midA* deletion strain. (Scale bar, 20 μm.) (B) Fluorescence images of GFP-TpmA and R-GECO in the *midA*-deletion hyphae growing in the media with 1 μM CaCl_{2}. (Scale bar, 10 μm.) (C) The length traveled by the hyphal apex over a 20-s interval in the wild-type and *midA*-deletion strain. Total period of 600 s. (D) Fluorescence images of GFP-TpmA and R-GECO in the *midA*-deletion hyphae growing in the media with 10 mM CaCl_{2}. (Upper) and time course of R-GECO (Left). (Scale bars, *Upper and Right*, 5 μm.) Kymographs along the hyphae in B and D. Total 240 s, every 2 s. (Scale bar, *Bottom*, 1 μm.)
can be summarized in the following model (shown in Fig. 5). During slow growth phases (i.e., low exocytosis activity), polymerized actin filaments accumulate at the hyphal tips, followed by SV accumulation before exocytosis. The transient increase of the intracellular Ca^{2+} level as a result of Ca^{2+} influx causes depolymerization of actin and fusion of the SV with the plasma membrane, resulting in more exocytosis and fast cell-extension phases. Reports that Ca^{2+} levels induce actin depolymerization and vesicle fusion in vitro support our model (7, 8). Similar oscillation periods of Ca^{2+} influx, actin polymerization, and SV accumulation at the tips of A. nidulans (∼30 ± 8 s) (Figs. 1H and 3 F and I) indicate that the cells coordinate these processes during stepwise cell extension. A few seconds of delay between these processes indicate that Ca^{2+} influx, actin depolymerization, and SV fusion are sequential mechanisms that proceed in this order to extend the cell periphery at the hyphal tips. Negative correlation between the oscillation peaks of the SV amounts and the cell-extension rate without a time gap (Fig. 2E) is consistent with the notion that exocytosis activity limits cell extension. This model is in agreement with our recent transient polarity assembly model of fungal tip growth, which describes the assembly/disassembly of polarity markers, polymerization of actin, and local exocytosis resulting in local membrane extension (18, 22, 23).

Here we demonstrate that normal oscillation of actin polymerization and stepwise growth require pulses of Ca^{2+} influx through the Ca^{2+} channels, CChA and MidA, in A. nidulans. The key event hence appears to be the activation of Ca^{2+} channels. One attractive possibility is that the Ca^{2+} channels could be stretch-activated. During slow growth phases cells gradually build up turgor pressure against the membrane and the cell wall. When the membrane tension exceeds a threshold, the Ca^{2+} channels could be activated. Following the entry of Ca^{2+}, exocytosis is promoted and leads to cell extension, which in turn decreases the turgor pressure and inactivates the channels. Indeed, stretch activation of the S. cerevisiae ortholog Mid1 has been reported (41). Another simple possibility is that the calcium channel firing is refractory to further activation until the cytoplasmic Ca^{2+} level is restored to resting levels by diffusion or efflux. The oscillation of the cytoplasmic Ca^{2+} level might also play a role in other physiological processes as well and not only for hyphal tip growth.

Our oscillatory model of cell growth implies a transient waiting time of SV before exocytosis. This raises the question about an advantage of pulsed versus continuous growth. Massive exocytosis at hyphal tips, which is the basis for rapid hyphal growth, imposes a huge problem for polarity maintenance. The newly inserted membrane dilutes locally polarity markers and the exocytosis machinery (i.e., the exocyst complex and SNARE proteins), which serve to direct vesicles to specific locations on the plasma membrane and to mediate their tethering to the membrane immediately before fusion (22, 42). If the number of fused vesicles exceeds the capacity of the cell to assemble focused polarity sites, the polar marker would diffuse on the plasma membrane. This has been nicely shown in S. cerevisiae (42). The waiting time of the SV till the next exocytosis event may allow the polarity components to rearrange at newly forming polarity sites and thus assemble enough actin cables that serve as tracks for directed vesicle transport (16, 19). In contrast, cortical actin meshes anchor the vesicles and, at the same time, function as barriers for vesicle fusion with the plasma membrane in mammalian cells (4). The transient increase of the intracellular Ca^{2+} level by Ca^{2+} influx is likely to be an important signal to coordinate depolymerization of actin meshes and thereby allow the anchored SV to fuse with the plasma membrane for exocytosis.

Our data reveal correlations among oscillations of four cell peripheral machineries of cell extension, F-actin, exocytosis, and Ca^{2+} levels, and provide comprehensive understanding of pulsed cell extension at the molecular level. Coordinated cycles or oscillations of these machineries are expected to be ubiquitous in all eukaryotes. Indeed, oscillations of intracellular Ca^{2+} levels and actin polymerization have been revealed in mammalian and plant cells (4–6). Although the time needed for assembly of polarity markers may be one reason for oscillatory growth, it may also be a mechanism for the cells to respond more quickly to both internal and environmental cues, including chemical and mechanical stimuli. Indeed, Ca^{2+} influx by the Ca^{2+} channel Cch1 in Candida albicans is involved in the control of directional growth of hyphae (43). The stepwise growth coordinated by transient Ca^{2+} influx could link growth with chemotropism and chemotaxis. Oscillatory recruitment of MAP kinase to cell tips during cell fusion in N. crassa is also involved in the process (44).

Because filamentous fungi maintain the growth site at their hyphal tips, this system is ideal for elucidating the mechanism of pulsed growth. Further analysis of the contribution of other processes, such as endocytosis, cell signaling, cell wall synthesis, and turgor pressure, will be necessary to complete the picture.

Materials and Methods

Strains, Plasmids, and Culture Conditions. Supplemented minimal medium for A. nidulans was prepared as described, and standard strain construction procedures were used (45). Two percent of glucose was used as carbon source; 70 mM sodium nitrate and 0.9 μM ammonium molybdate were used as nitrogen source for solid media, if not stated otherwise. For liquid media, the carbon sources were 2% glucose, 2% threonine, or 2% glycerol. A. nidulans strains used in this study are listed in Table S1. Standard laboratory Escherichia coli strains (Top 10 F) were used. N. crassa (CHS-1–GFP) strain was grown on Vogel's minimal medium agar at 32 °C for 3 days and observed using the “inverted agar block method” (46).

Plasmid and Strain Construction. The sequence of R-GECO was amplified from plasmid pPD60 REGEO.1 (35), using primers R-GECO-KpnI-fw and R-GECO-XbaI-r, generating KpnI and XbaI restriction enzyme sites, respectively, and subsequently cloned into plasmid pET12. The constructed plasmid was further digested with KpnI and XbaI and the corresponding to R-GECO ORF band was gel-extracted, purified, and cloned in the KpnI-XbaI-digested pCM817apx (for proteins of interest expressed under the regulatable alcA promoter; contains N. crassa pyr-4) (21) plasmid downstream of the alcA promoter sequence, yielding pNT76. The DNA fragment for GFP tagging at the C terminus of BglA was amplified by fusion-PCR with a GFP-pyrG cassette (47) using primers BglA_P1, BglA_P2, BglA_P3, BglA_P5, and BglA_P6, and was subsequently cloned into plasmid pET12, yielding plasmid pET12. The sequence of chsB was amplified from genomic DNA by using primers chsB_AscI_fwd and chsB_PacI_rev, was digested with Ascl and PacI, and cloned into the Ascl-PacI-digested pSH44 (for mCherry tagging at N terminus of proteins of
Fluorescence Microscopy. Cells were grown in eight-well glass-bottom slides (b)idi) with minimal medium at 28 °C overnight. Images were captured using an Axiophot microscope using a Plan-Apochromatic 63x 1.4 Oil objective lens, the ZEISS AxioCam MRM camera (Zeiss), and the HBO 103 mercury arc lamp (Osram) or HXP 120 (Zeiss), featuring faster wavelength switching. Images were collected and analyzed by using the Zen system (Zeiss) and Image software. Photobleaching corrections were not applied to the data.

Super-Resolution Microscopy. PALM images were acquired at room temperature on a modified inverted microscope (Axiovert 200, Zeiss) as described previously (42). The fluorescent proteins were converted from their green to red emitting forms using a 405-nm laser (Stradux 405-250, Vortran Laser Technology) with an intensity of 0–50 W/cm². Fluorescence was excited by a 561-nm laser (Gem 561, Laser Quantum) with simultaneous 405-nm illumination (200–400 W/cm²). PALM images were analyzed with custom-written analysis software, a-livePALM (50) running under the MATLAB R2015b (The Mathworks) environment.

Acknowledgments. We thank our laboratory members for valuable discussion and critical reading of the manuscript, and B. Schreckenberger and E. Wohlmann for excellent technical assistance. This work was supported by the Deutsche Forschungsgemeinschaft Grants (TA1912/1, FOR1334, NI291/12-1, and GRK2039), the Helmholtz Association Program STN, the Japan Society for the Promotion of Science KAKENHI Grant 15K18663, and the Japan Science and Technology Agency Exploratory Research for Advanced Technology (ERATO) Grant JPMJER1502.

Fig. S1. Stepwise growth of hyphae. (A) Kymograph of the growing hypha along the growth axis (Left) and enhanced image (Right). Total 60 min. (Scale bar, Left, 10 μm.) (B) The length of the interval traveled by the hyphal apex along the line in overlay of cell profiles from a series of six PALM images expressing hypha (Fig. 1E). Larger extensions of the apical membrane were indicated by arrows. (Scale bar, 300 nm.) (C) Cross-correlation between GFP-TpmA and mCherry-ChsB. To account for the effect of photobleaching during imaging (3 min), the baseline of each signal was defined as a line with negative slope, based on the average value of first and second half of the signal. The cross-correlation was then calculated based on the corrected signal. (D) Fluorescence intensity of secretory vesicles (red) along the apex of the growing hypha between dotted lines in (Fig. 1G). The length of the interval traveled by the hyphal apex was measured every 20 s. (E) The position of Spitzenkörper was aligned every 3 s from Movie S3.
Fig. S2. Intracellular Ca\(^{2+}\) level by R-GECO. (A) Scheme of the Ca\(^{2+}\) biosensor, R-GECO. (B) Fluorescence image sequence of R-GECO. The elapsed time is given in seconds. (Scale bar, 5 μm.) (C) The time-lapse signal intensity of R-GECO at different points from the tip shown in B. (D) Fluorescence image sequence of R-GECO in the three hyphae. Arrows indicate hypha showing the R-GECO signal with different timing. The elapsed time is given in seconds. (Scale bar, 10 μm.)
**Fig. S3.** Effect of CaCl$_2$ on the oscillation of R-GECO. (A and B) Fluorescence image of R-GECO in the hypha growing in the media without CaCl$_2$ (A) and 1 μM CaCl$_2$ + 10 mM EGTA (B). (Scale bars, 10 μm.) Kymographs (Lower) along the hyphae. Total 240 s, every 2 s. (Scale bars, 1 μm.) (C, Upper) Hyphal morphology of the cchA-deletion strain. (Scale bar, 20 μm.) (Lower) Colonies of wild-type, midA-deletion and cchA-deletion strain grown on the minimal media plate for 3 d. (D and E) Fluorescence image of R-GECO and GFP-TpmA, in the cchA-deletion hyphae growing in the media with 1 μM CaCl$_2$ (D), and in the wild-type hypha growing in the media with 10 mM CaCl$_2$ (E). (Scale bars, 5 μm.) (Lower) Kymographs along the hypha. Total 240 s, every 2 s. (Scale bars, 1 μm.) (F) Fluorescence image of R-GECO and GFP-TpmA in the wild-type hypha growing in the media with 1 μM CaCl$_2$ after the treatment of calcium ionophore A23187 (Sigma) 5 μg/mL for 30 min. (Scale bars, 5 μm.)

**Table S1.** *Aspergillus nidulans* strains used in this study

<table>
<thead>
<tr>
<th>Strain</th>
<th>Genotype</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>TN02A3</td>
<td>pyrG89; argB2; ΔnkuA::argB; pyroA4</td>
<td>Nayak et al. (49)</td>
</tr>
<tr>
<td>SNT147</td>
<td>pyrG89; argB2; ΔnkuA::argB; pyroA4; [alcA(p)-gfp-tpmA::pyr-4]</td>
<td>Bergs et al. (18)</td>
</tr>
<tr>
<td>CIA08</td>
<td>pyrG89; riboB2; argB2; ΔnkuA::argB; pyroA4; [ΔmidA::pyrG]</td>
<td>Wang et al. (40)</td>
</tr>
<tr>
<td>W5A05</td>
<td>pyrG89; riboB2; argB2; ΔnkuA::argB; pyroA4; [ΔcchA::pyrG]</td>
<td>Wang et al. (40)</td>
</tr>
<tr>
<td>SNG10</td>
<td>pyrG89; argB2; ΔnkuA::argB; pyroA4; [bgA-gfp::pyr-4]</td>
<td>Present study</td>
</tr>
<tr>
<td>SNT167</td>
<td>pyrG89; argB2; ΔnkuA::argB; pyroA4; [alcA(p)-mEeosthermofp-chsB::pyr-4]</td>
<td>Present study</td>
</tr>
<tr>
<td>SNT161</td>
<td>pyrG89; argB2; ΔnkuA::argB; pyroA4; [alcA(p)-gfp-tpmA::pyr-4]; [alcA(p)-mcherry-chsB::pyrA]</td>
<td>Present study</td>
</tr>
<tr>
<td>SNT162</td>
<td>pyrG89; argB2; ΔnkuA::argB; pyroA4; [alcA(p)-R-GECO::pyr-4]</td>
<td>Present study</td>
</tr>
<tr>
<td>SNT163</td>
<td>pyrG89; argB2; ΔnkuA::argB; pyroA4; [pyroA] [alcA(p)-gfp-tpmA::pyr-4]; [alcA(p)-R-GECO::pyr-4]</td>
<td>Present study</td>
</tr>
<tr>
<td>SNT164</td>
<td>pyrG89; argB2; ΔnkuA::argB; pyroA4; [pyroA] [bgA-gfp::pyr-4]; [alcA(p)-R-GECO::pyr-4]</td>
<td>Present study</td>
</tr>
<tr>
<td>SNT165</td>
<td>pyrG89; riboB2; argB2; ΔnkuA::argB; pyroA4; [ΔmidA::pyrG]; [pyroA]; [alcA(p)-gfp-tpmA::pyr-4]; [alcA(p)-R-GECO::pyr-4]</td>
<td>Present study</td>
</tr>
<tr>
<td>SNT166</td>
<td>pyrG89; riboB2; argB2; ΔnkuA::argB; pyroA4; [ΔcchA::pyrG]; [pyroA]; [alcA(p)-gfp-tpmA::pyr-4]; [alcA(p)-R-GECO::pyr-4]</td>
<td>Present study</td>
</tr>
</tbody>
</table>

**Table S2.** Plasmids used in this study

<table>
<thead>
<tr>
<th>Plasmid</th>
<th>Description</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>pPD60</td>
<td>RGEKO.1</td>
<td>Zhao et al. (35)</td>
</tr>
<tr>
<td>pNT77</td>
<td>alcA(p)-mcherry-chsB::pyrA</td>
<td>Present study</td>
</tr>
<tr>
<td>pNG3</td>
<td>bgA-gfp::pyr-4</td>
<td>Present study</td>
</tr>
<tr>
<td>pNT76</td>
<td>alcA(p)-R-GECO::pyr-4</td>
<td>Present study</td>
</tr>
</tbody>
</table>
Table S3. Primers used in this study

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence (5′ → 3′)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RGECO-KpnI-fw</td>
<td>gcggtaccATGGTCGACTCATCACGTCG</td>
</tr>
<tr>
<td>RGECO-XbaI-r</td>
<td>gctctagaCTACTTCGCTGTCATCATTG</td>
</tr>
<tr>
<td>BglA_P1</td>
<td>GCGGTGACGGACAACTG</td>
</tr>
<tr>
<td>BglA_P2</td>
<td>CAAGACGCGGTCTGAGTCTAC</td>
</tr>
<tr>
<td>BglA_P3</td>
<td>tggcgcggttgcccAGAGGAAGCTTGCTGAGG</td>
</tr>
<tr>
<td>BglA_P5</td>
<td>GACGGACATGCATGCATGCTAGCAC</td>
</tr>
<tr>
<td>BglA_P6</td>
<td>GCATAACAGGAAACACGCT</td>
</tr>
<tr>
<td>chsB_AscI_fwd</td>
<td>cggccgctATGGCCTACCACGCTC</td>
</tr>
<tr>
<td>chsB_PacI_rev</td>
<td>cttacatattaGGCAACACATGACATATCC</td>
</tr>
</tbody>
</table>

Movie S1. Image sequences of the DIC A. nidulans wild-type hyphae growing in the minimal media at 28 °C; every 10 s, total 1,800 s. (Scale bar, 10 μm.) See Fig. 1A.

Movie S1

Movie S2. Image sequences of GFP-TpmA and mCherry-ChsB; every 2 s, total 180 s. (Scale bar, 2 μm.) See Fig. 1F.

Movie S2
Movie S3. Image sequences of DIC and CHS-1- GFP in *Neurospora crassa*; every 2 s, total 600 s. (Scale bar, 10 μm.) See Fig. 2A.

Movie S4. Image sequences of R-GECO; every 1 s, total 120 s. (Scale bar, 5 μm.) See Fig. 2B.

Movie S5. Image sequences of GFP-TpmA and R-GECO; every 2 s, total 180 s. (Scale bar, 2 μm.) See Fig. 3E.
Movie S6. Image sequences of BglA-GFP and R-GECO; every 2 s, total 180 s. (Scale bar, 2 μm.) See Fig. 3H.