Cross-membranes orchestrate compartmentalization and morphogenesis in Streptomyces

Katherine Celler1,*, Roman I. Koning2,*, Joost Willemse1, Abraham J. Koster2 & Gilles P. van Wezel1

Far from being simple unicellular entities, bacteria have complex social behaviour and organization, living in large populations, and some even as coherent, multicellular entities. The filamentous streptomycetes epitomize such multicellularity, growing as a syncytial mycelium with physiologically distinct hyphal compartments separated by infrequent cross-walls. The viability of mutants devoid of cell division, which can be propagated from fragments, suggests the presence of a different form of compartmentalization in the mycelium. Here we show that complex membranes, visualized by cryo-correlative light microscopy and electron tomography, fulfil this role. Membranes form small assemblies between the cell wall and cytoplasmic membrane, or, as evidenced by FRAP experiments, large protein-impermeable cross-membrane structures, which compartmentalize the multinucleoid mycelium. All areas containing cross-membrane structures are nucleoid-restricted zones, suggesting that the membrane assemblies may also act to protect nucleoids from cell-wall restructuring events. Our work reveals a novel mechanism of controlling compartmentalization and development in multicellular bacteria.

1 Department of Molecular Biotechnology, Institute of Biology Leiden, Leiden University, PO Box 9505, 2300 AB Leiden, The Netherlands. 2 Department of Molecular Cell Biology, Leiden University Medical Centre, PO Box 9600, 2300 RC Leiden, The Netherlands. * These authors contributed equally to this work. Correspondence and requests for materials should be addressed to G.P.v.W. (email: g.wezel@biology.leidenuniv.nl).
Bacteria have a complex organization on both a unicellular and multicellular level, with many species demonstrating multicellular traits or behaviours analogous to those seen in eukaryotic systems. This multicellularity can manifest itself in different ways, from intercellular communication between individual cells in a population, to a lifecycle spent as a coherent group of cells, such as in biofilms, fruiting bodies, filaments or mycelia. A case in point are the streptomycetes, filamentous bacteria, which grow forming highly structured multicellular colonies composed of physiologically distinct hyphae.

The streptomycetes are soil-dwelling saprophytes, well-known for their complex secondary metabolism and ability to produce an abundance of natural products including antibiotics, anticancer agents and immunosuppressants. After a spore has germinated, a complex network is formed as the hyphae grow by tip extension and branching into a syncytial vegetative mycelium separated by occasional peptidoglycan-based septa, with a spacing of some 5–10 μm. Upon nutrient depletion, the streptomycetes undergo extensive chemical and physiological differentiation, with the coordinated production of aerial hyphae and antibiotics. At this stage in the life cycle, programmed cell death (PCD) occurs in certain parts of the vegetative mycelium and not others, which may be interpreted as a strategy to provide nutrients to the developing aerial mycelium. The aerial hyphae eventually metamorphose into chains of unigenomic spores.

Though cell division is the hallmark of life, in Streptomyces it is non-essential and the gene encoding the cell division scaffold protein FtsZ can be deleted. Surprisingly, although it no longer undergoes reproductive cell division and fails to form the cross-walls that are needed to compartmentalize vegetative hyphae, the ftsZ-deletion mutant can be propagated from fragments obtained by mechanical maceration of a large syncytial mycelium. It is a mystery why in the absence of any physical barrier the hyphae do not ‘bleed’ to death from open hyphal ends. This strongly suggests that a yet unknown form of compartmentalization of the hyphae exists. Compartmentalization is also implied by the infrequent nature of cross-wall division in the vegetative hyphae, yet prevalence of short live and dead portions of hyphae undergoing differentiation and PCD.

The streptomycetes grow by hyphal extension at apical sites, with highly dynamic cell wall construction and remodelling. In these multicellular bacteria pose an interesting problem: how can these processes take place without damaging the many chromosomes in the multinucleoid hyphae? In addition, during sporulation-specific cell division up to a hundred septa are formed in the sporogenic aerial hyphae, visualized as spectacular Z ladders over nucleoids that have not yet segregated. The canonical DNA-damage control systems such as the SOS response or nucleoid occlusion, which mediate division control in cells with a planktonic lifestyle, have not been identified in Streptomyces. An alternative system must therefore exist to prevent lethal DNA damage by the cell-wall synthetic machineries.

In this work, we demonstrate the presence of a membrane system in the vegetative hyphae of Streptomyces, existing apart from the peptidoglycan-based cell compartmentalization mediated by the FtsZ-guided cell division machinery. These so-called ‘cross-membranes’ form protein impermeable barriers between hyphal segments and effectively compartmentalize the multinucleoid hyphae.

Results

Intracellular membranes exist in Streptomyces vegetative hyphae.

To investigate membrane remodelling in the Streptomyces vegetative mycelium, anionic phospholipids including phosphatidylglycerol (PGL), were stained with the membrane dye FM5–95, cross-walls stained with FITC-WGA (FITC-wheat germ agglutinin, or alternatively, BODIPY-vancomycin, which binds to the terminal d-Ala within the wall peptide of peptidoglycan) and their localization studied with fluorescence light microscopy (fLM). This not only revealed membrane structures associated with peptidoglycan cross-walls, but also in surprisingly large assemblies, several micrometres in length, spanning the width of vegetative hyphae (Fig. 1b). Lipid assemblies were present within hyphae in differing amounts, with some hyphae being devoid of membranes and others containing multiple formations, from small blebs to large structures. Quantification experiments indicated that of 218 membrane localizations counted, 75.3% localized in large or small apparently non-septal assemblies. The remaining 24.7% were at septal locations—either clearly co-localizing with BODIPY-vancomycin or in the absence of BODIPY-vancomycin—still appearing in thin, cross-wall-like form (Supplementary Table 1 and Supplementary Figs 5–8).

We applied cryo-correlative light and electron microscopy (cryo-CLEM), combining cryo-fLM with cryo-electron microscopy (cryo-CLEM). Positions with extended lipids were observed (Fig. 1d), consisted of extended tubular membrane structures. A light region, devoid of ribosomes and other macromolecular complexes, can be seen forming a faint ribbon behind a membrane-filled tip (arrows, f, for tomogram see Supplementary Movie 1). Scale bars, 5 μm (a–c), 500 nm (d–f).

Figure 1 | Membrane assemblies in vegetative hyphae. (a) In vegetative hyphae of S. coelicolor M145 stained with the membrane dye FM5–95 (red) and the cell wall dye FITC-WGA (green), cross-walls are evident (arrowheads). (b) Large membrane structures/agglomerates are also found within hyphae, which do not co-localize with WGA-stained cross-walls (arrowheads). (c–f) To investigate their ultrastructure, membranes in S. albus were fluorescently labelled with FM5–95 and imaged with cryo-CLEM. Positions with extended lipids were observed (d–f). Cross-membrane assemblies within hyphae (d,e) and at tips (e,f) consisted of extended tubular membrane structures. A light region, devoid of ribosomes and other macromolecular complexes, can be seen forming a faint ribbon behind a membrane-filled tip (arrows, f, for tomogram see Supplementary Movie 1). Scale bars, 5 μm (a–c), 500 nm (d–f).
tomography (cryo-ET), to study the fine structural detail of the membrane assemblies in three dimensions at nanometre resolution. Cryo-CLEM enables direct fluorescent labelling and targeting of molecules or molecular assemblies (such as intracellular membranes, DNA or cyto-structural elements) in the same cryo-immobilized samples that are then visualized by cryo-ET, avoiding the possible artefacts of chemical fixation, plastic embedding and/or metal staining. In this manner, sites of interest can be quickly located without imposing cryogenic samples to extended electron beam damage. We used *Streptomyces albus* in our cryo-CLEM experiments as this strain is morphologically very similar to model organism *S. coelicolor* on agar plates, with abundant sporulation, yet has thinner hyphae in submerged cultures and therefore allows for high-resolution three-dimensional imaging by whole-cell cryo-ET. By performing cryo-CLEM on *S. albus* hyphae stained with FM5–95, we were able to readily target membrane assemblies within hyphae (Fig. 1c). In young vegetative hyphae (12h after inoculation), membranes could be seen forming bundles of tube-like structures varying from small assemblies along the cell wall to large structures completely delimiting the hyphae (Fig. 1d and Supplementary Fig. 1). The membranes were also identified at apical sites (Fig. 1e,f and Supplementary Movie 1), as seen in fLM of *S. coelicolor*24. Similar membrane assemblies were also found in *S. coelicolor* when examined by cryo-CLEM (Supplementary Fig. 2).

Cryo-ET and data segmentation enabled visualization and analysis of the membrane formations in three dimensions, and demonstrated that the assemblies form in the space between the cytoplasmic membrane and cell wall (Fig. 2). Membranes were observed to vary from several small tubes to larger structures, which appeared to be constricting, decreasing the connection between hyphal compartments, likely in an intermediate stage of membrane closure (Fig. 2c). Evidence of septum synthesis could also be seen associated with cross-membranes, and mostly with full cross-membrane structures, which spanned the entire hyphae (Fig. 2d). Tomograms and full 3D surface renderings can be found in Supplementary Movies 2–5.

**Cross-membranes create a chromosome-free area.** Excited by the novel visualization of cross-membranes that cryo-ET afforded and to specifically determine the spatial localization of the membranes relative to chromosomes, we combined membrane staining by FM5–95 with chromosome visualization using 4′,6-diamidino-2-phenylindole (DAPI; Fig. 3a–c). It was previously demonstrated that nucleoids are absent from areas enriched in anionic phospholipids24. We observed that all areas that showed a distinct gap in DAPI fluorescence in cryo-fLM, and therefore were devoid of DNA, correlated to large hyphae-spanning membrane assemblies (red arrows, Fig. 3a,d) or vesicle structures (blue arrows, Fig. 3b,d). Again, we observed evidence of septum synthesis within full cross-membrane structures, which spanned the entire hyphae (Fig. 3e–h). This correlation, as well as the frequent presence of membranes at hyphal tips, suggested that in *Streptomyces* vegetative hyphae membrane assemblies may act to occlude chromosomal DNA from sites of active cell wall restructuring, either for septum synthesis (Fig. 3f–h), tip growth or branching.

**Membranes form independent of cell wall remodelling.** To investigate the relationship between membrane formation and cell wall remodelling, we stained samples for cryo-CLEM with BODIPY-vancomycin and looked for membrane assemblies at locations of BODIPY-vancomycin staining. Experiments revealed numerous examples of BODIPY-vancomycin staining present at septa (Fig. 4a), and to a lesser extent at growing tips (Fig. 4b), and new branch points (Fig. 4c). High-resolution transmission electron micrographs confirmed that some regions stained with BODIPY-vancomycin also contained cross-membranes (Fig. 4d). A statistical analysis on fLM images co-stained with BODIPY-vancomycin and FM5–95 indicated that of a total of 335 locations measured (containing FM5–95 staining, BODIPY-vancomycin staining or both), 117 (34.9%) demonstrated only BODIPY-vancomycin staining (and no FM5–95 staining) at septa or points of cell wall remodelling. Of the remaining locations, 153 (45.6%) demonstrated only FM5–95 staining, and 65 (19.4%) demonstrated both (Supplementary Table 1 and Supplementary Figs 5–8). This result suggests that membranes are either not essential for new cell wall formation, or that if membranes do co-localize with newly forming cell wall, the co-localization is transient. To obtain insight into the dynamics of co-localization during peptidoglycan synthesis, we therefore performed live imaging experiments. These experiments, however, repeatedly failed due to the toxicity of the immunofluorescence stains to young vegetative hyphae.

*Streptomyces* grow by hyphal tip extension, with polar growth and branching depending on DivIVA25. As we observed many membrane-filled tips during cryo-CLEM experiments, we were
interested to see if these membranes mark the location of DivIVA. We performed fLM studies to quantify the co-localization of this morphoprotein with membranes and found that 27.0% of DivIVA foci co-localize with membrane assemblies (Supplementary Table 2 and Supplementary Fig. 9).

Given the incomplete spatio-temporal co-localization of membranes with newly forming cell wall at cross-walls or hyphal tips, we wondered whether cross-membranes form independent of septum synthesis. We therefore quantified membrane and cell wall formation in the *S. coelicolor* ftsZ deletion strain, which grows without forming septa. fLM experiments staining the hyphae with FM5–95 for membranes and WGA for newly forming cell wall demonstrated that in 77 hyphae, with a total length of roughly 2,635 μm, 368 cross-membranes were present and no cross-walls (Fig. 5). This amounts to one cross-membrane per 7.16 ± 0.66 μm (313 membranes counted), whereas the number of cross-walls formed is one per 9.03 ± 0.46 μm (99 cross-walls counted). Cryo-electron microscopy (EM) experiments confirmed that the cross-membranes in the *ftsZ* deletion strain appear indistinguishable to those in wild-type *S. coelicolor* (Supplementary Fig. 3). The complete absence of cross-walls and abundance of cross-membranes in the *ftsZ* deletion strain indicates that septum synthesis is not required for the formation of—and that septa are not necessarily present in—cross-membranes.

**Cross-membranes compartmentalize hyphae.** Staining of *Streptomyces* mycelium with propidium iodide and SYTO 9 demonstrates alternating live and dead compartments in very young vegetative mycelium, suggesting that the PCD linked to development already occurs in very young hyphae. We wondered if
the large cross-membrane structures we observe (and not septa) form the cytosol impermeable barrier that enables this type of  

differentiation. We therefore applied fluorescence recovery after photobleaching (FRAP) on a S. coelicolor strain constitutively expressing enhanced green fluorescent protein (eGFP) and stained with FM5–95 for membranes, bleaching an entire compartment on one side of a cross-membrane to establish whether green fluorescent protein (GFP) molecules from the adjacent compartment can restore fluorescence. FRAP experiments (N = 91) were performed on regions in close proximity to membranes. These experiments demonstrated that in 71% of cases cross-membranes are cytosol permeable (Fig. 6a and Supplementary Movie 6) and in 29% of cases impermeable (Fig. 6b and Supplementary Movie 7), preventing fluorescent GFP migration into the adjacent compartment. Additional FRAP experiments were performed to distinguish cross-membranes containing cross-walls from those that do not contain any cell wall material. In these experiments, the S. coelicolor strain constitutively expressing eGFP was stained with both FM5–95 for membranes and far-red dye WGA-Alexa 633 to stain cross-walls. Membranes with no cell wall present were targeted to start. In these cases, cross-membranes were found to be permeable in 23% of cases (green arrows, Supplementary Fig. 4, Supplementary Movie 8) or impermeable in 77% of cases (red arrows, Supplementary Fig. 4, Supplementary Movie 8), revealing that cross-membranes alone can block diffusion within hyphae. Next, membranes with cell-wall material present were targeted. For these cross-membranes, the permeable fraction was 71%, whereas large membrane assemblies near a cross-wall completely blocked diffusion. This increased permeability of cross-membranes with cell-wall material present can be explained by the fact that cross-walls are likely permeable, once cross-membranes disassemble, transport can occur over cross-walls. Taken together, our experiments show that in the absence of cross-walls, cross-membranes can act as impermeable physical barriers between adjacent compartments.

**Discussion**

In this work, we discovered the presence of a vast internal membrane system in Streptomyces species that can be found within and throughout vegetative hyphae, forming bundles of tubular structures between the cytoplasmic membrane and cell wall. Using cryo-CLEM, we observed membrane formations in small blebs along the cell wall or at hyphal tips, or as large assemblies and hyphae-delimiting structures we termed cross-membranes. Similar structures were observed in the 1960s (refs 27,28), although their function was unknown. Our present investigation of these membranes resolves several intriguing questions surrounding their role in the Streptomyces lifecycle. Although the production of antibiotics and other clinically important secondary metabolites by the streptomycetes has been linked to PCD in the vegetative mycelium, to date, little was known about how the hyphae separate into live and ‘dead’ segments.

Peptidoglycan-based septa, or cross-walls, are infrequent in the vegetative hyphae. As beautifully illustrated by live imaging experiments, during hyphal growth of S. coelicolor cross-walls (that is, vegetative septa) are produced seemingly randomly at sites distant from the growing tips, delimiting the hyphae into compartments of 5–10 μm (ref. 29). However, the compartments can be up to 40 μm for some other streptomycetes30 and these compartments contain approximately one chromosome per micrometre, or tens of copies of the genome per compartment. Cross-walls do not form in the ftsZ deletion strain, which raises the question as to how a colony can ensure its survival during PCD or following physical damage to the mycelium. It is now clear that membranes play an important role, offering a less energy-intensive compartmentalization solution than cross-wall formation.

Interestingly, although compartmentalization is rare in bacteria, it does exist in a variety of species31–33. Recently, a protein-mediated diffusion barrier was also discovered in the polar stalk of Caulobacter crescentus, acting to prevent exchange of membrane and soluble proteins between the stalk and cell body34. The cross-membranes in Streptomyces similarly delimit and prevent diffusion of soluble proteins between diverse hyphal segments, as evidenced by FRAP experiments on the diffusion of molecules of GFP.

Our FRAP experiments revealed that 29% of cross-membranes are impermeable. Although unlikely, it is conceivable that laser-mediated damage to the cross-membranes may have contributed to leakage of GFP from adjacent compartments, increasing the number of permeable membranes we counted. The percentage of impermeable cross-membrane barriers may therefore be higher than what we observed. Molecules smaller than GFP may also be able to pass the cross-membranes. The high percentage of permeable membrane assemblies we observed are likely membrane structures that are not yet fully closed, that is, in an earlier stage of cross-membrane formation, or perhaps these membranes serve a different purpose during the Streptomyces lifecycle.

Indeed, the abundance of lipid structures localizing in small blebs adjacent to the cell wall, in addition to the full cross-membranes delimiting hyphae, could indicate that smaller membrane assemblies may be involved in cellular functions other than compartmentalization. At locations along the hyphae, small blebs are likely mainly precursors for cross-membranes. However, membrane assemblies are also frequently observed at hyphal tips, which suggests that they may be involved in tip growth. It follows that some of the membrane blebs along the hyphae may not be precursors for cross-membranes, but rather the future sites of branching. At growing tips and new branches, it is likely that the assemblies observed have a function in creating a nucleoid-free zone in a region of high activity.

In the Streptomyces growing apex, a highly dynamic tip-organizing complex (the so-called TIPOC) exists, consisting of many components—including various cytoskeletal elements—

---

**Figure 5 | Membrane and cell wall localization in an S. coelicolor ftsZ-deletion strain.** Membrane assemblies are still abundant in S. coelicolor ftsZ-deletion strain (arrows, a, stained with FM5–95), but no cross-walls form (arrows, b, note N-acetylglucosaminyl residues present in cell wall stained with WGA-Oregon green). An overlay image is provided (c), as well as the light image (d). Scale bar, 5 μm. DIC, differential interference contrast.
and ensuring that DNA replication takes place at some distance behind the tip\textsuperscript{16,35,36}. Extensive statistics using cryo-ET are not feasible considering the high-resolution nature of these experiments, so instead we performed static fLM experiments on vegetative hyphae stained with FM5–95 and peptidoglycan marker BODIPY-vancomycin and tip growth protein DivIVA to determine the degree of membrane co-localization with new cell wall synthesis. These experiments (see Supplementary Tables 1 and 2 for details) did not indicate complete spatio-temporal co-localization. It should be noted, however, that studying dynamic events with static experiments is notoriously difficult. Further experiments using other cell division or morphoproteins can perhaps provide more insight into the role of the membranes in providing a suitable environment for protein localization. In addition, use of an integrated system for live light microscopy and cryo-fixation (for example, the Rapid Transfer System\textsuperscript{37} or the MAVIS\textsuperscript{38}) could enable the targeting of actively growing tips or cross-walls at different stages of formation, and elucidating the role of membranes therein.

Finally, although we demonstrated that cross-membranes form independent of septum synthesis, during our experiments, we frequently observed cross-membranes containing emerging

**Figure 6 | FRAP analysis of GFP distribution over cross-membranes.** Fluorescence recovery after photobleaching (FRAP) on a S. coelicolor strain constitutively expressing eGFP and stained with FM5–95 for membranes. An entire compartment was bleached on one side of a cross-membrane to establish whether GFP molecules from the adjacent compartment would restore fluorescence. Membranes can be both cytosol permeable (a and Supplementary Movie 6), which can be seen when bleached GFP molecules spread through the membrane as indicated by the white arrows, or impermeable (b and Supplementary Movie 7), which can be seen when the membrane prevents the flow of bleached GFP molecules. Recovery curves demonstrate the recovery in fluorescence intensity in the case of the permeable membrane, together with a decrease in fluorescence in the adjacent compartment. In the case of the impermeable membrane, a slight increase in fluorescence is seen in the bleached compartment due to diffusion of GFP molecules from within the compartment, whereas there is no decrease in fluorescence in the adjacent compartment. Further imaging results in bleaching and a decrease in fluorescence in both the bleached and adjacent compartments.
cross-walls. (Fig. 3f–h). In the multicellular streptomycetes, hyphal compartments are multigenomic; it is likely that some form of subcellular organization exists that serves to separate the chromosomes. One obvious reason for such compartmentalization would be to allow cross-walls to form in a chromosome-free area.

In conclusion, we show here that cross-membranes exist in the hyphae of the streptomycetes, acting to compartmentalize the hyphae. These membranes can be cytosol permeable or impermeable, and provide a plausible answer to the question why hyphae of the ftsZ null mutant do not ‘bleed’ to death but rather can be fragmented and propagated. Further, the cross-membranes create DNA-free zones and offer a solution to the long-standing problem of how cell wall remodelling can take place without DNA damage in the multicellular hyphae. Future technological advances in correlative microscopy, in which fluorescently tagged proteins can be superimposed on high-resolution images of cytostructural elements in tomograms, should provide further insights into the exciting cell biology of the streptomycetes.

**Methods**

**Bacterial strains and constructs.** *S. coelicolor* and *S. albus* strains were grown on R2YE or soya flour mannitol agar. Liquid cultures grown in 2YT medium supplemented with 10% sucrose. All of the growth medium recipes used are those mentioned in the ref. 39. For fluorescence microscopy, samples from liquid cultures were collected from samples that had been inoculated at the acute-angle junction of coverslips inserted at a 45° angle in soya flour mannitol agar plates. Strains used in this study include *S. coelicolor* M145 (ref. 39), obtained from the John Innes Centre strain collection, its derivative K202 containing plasmid pKF41 that expresses FtsZ-GFP17, *S. albus* subspecies albus G. ATCC 25426 and, for FRAP, a *S. coelicolor* strain constitutively expressing GFP.

This strain was created by performing *S. coelicolor* M145 with pCG4, a pBluescript derivative harbouring the constitutive FtsZ1 promoter of *S. coelicolor* A32) M145 cloned upstream of the eGFP gene (B. Zacchetti and D. Claessen, unpublished observations).

**Microscopy.** Fluorescence microscopy. Fluorescence and corresponding light micrographs were obtained with a Zeiss Axioscope A1 upright fluorescence microscope (with an AxioCam MrC5 camera at a resolution of 37.5 nm per pixel), with, for the green channel, 470–490 nm excitation and 515 long-pass detection; and for the red channel, 530–550 nm excitation and 590 long-pass detection. For cryo-stage imaging, the green fluorescent images were created using 470/40 nm band-pass detection and 525/50 nm long-pass detection; for the red channel, 550/25 nm band-pass detection and 605/70 nm long-pass detection were used. For staining of the cell wall (peptidoglycan), we used FITC-WGA; for membrane staining, we used FM5 (Molecular Probes). FM5 fluorescence experiments were carried out on a Zeiss Imager system, using 488 nm excitation for eGFP with 505–530 detection and 543 nm excitation for FM5 (refs 27, 28) 5–95 with LP 560 detection. Bleaching was performed with 25% laser intensity for 8 s. Images were recorded every 17 s and post-bleaching was monitored for 5 min. All images were corrected with contrast. Images outside the hyphae were set to 0 to obtain a sufficiently dark background. These corrections were made using Adobe Photoshop CS5. Triple coloured FRAP experiments were performed with 488 nm excitation, which does not excite WGA-Alexa 483 and therefore the same band-passes were used for signal detection: for example, for eGFP, band-pass filter 505–530; for FM5–95, long-pass 560. WGA-Alexa 633 was imaged with 633 excitation and long-pass 650 detection.

Sample preparation for electron microscopy. A small drop (3 μl) of Streptomyces liquid culture was applied to EM grids and vitrified by plunging into a liquid ethane/propane mixture using a vitrobot Mark IV (FEI Company) operated at 22°C and 100% humidity using 1–2 s blotting. Plunge-frozen grids were stored in liquid nitrogen until further use. For cryo-electron tomography, 15 nm colloidal gold particles coupled to protein A (CMMC) were added to samples as fiducial markers. For correlative light and electron microscopy work, holey carbon grids were used. Plunge-frozen grids were stored in liquid nitrogen until use for EM grids.

**Image analysis and visualization.** Tomographic tilt series were processed using IMOD version 4.5 (ref. 40). Projection images were pre-processed by hot pixel removal and rough alignment by cross-correlation. Final alignment was done using fiducial gold markers. The tomograms were obtained using a weighted back-projection or a simultaneous iterative reconstruction technique. Cryo-electron tomograms were Fourier filtered and denoized with a nonlinear anisotropic diffusion to enhance the visibility of structures. For 3D surface rendering, the tomographic volumes were imported into AMIRA (FEI) for further processing and representation. Cryo-CLEM overlays were produced by manual overlay (Adobe Photoshop) of different magnification TEM and FLM images using grid bars, streptomycetes, holey carbon support film and fiducial gold markers for alignment.

**Data availability.** The data that support the findings of this study are available within the article or from the corresponding author upon request.

**References**

1. Shapiro, J. A. Thinking about bacterial populations as multicellular organisms. *Annu. Rev. Microbiol.* 52, 81–104 (1998).
2. Claessen, D., Rozen, D. E., Kuipers, O. P., Sogaard-Andersen, L. & van Wezel, G. P. Bacterial solutions to multicellularity: a tale of biofilms, filaments and fruiting bodies. *Nat. Rev. Microbiol.* 12, 115–124 (2014).
3. Lyons, N. A. & Kolter, R. On the evolution of bacterial multicellularity. *Curr. Opin. Microbiol.* 24, 21–28 (2015).
4. Miguez, E. M., Hardisson, C. & Manzanal, M. B. Hyphal death during colony development in *Streptomyces* antibiotics: morphological evidence for the existence of a process of cell deletion in a multicellular prokaryote. *J. Cell Biol.* 121, 515–525 (1999).
5. Miguez, E. M., Hardisson, C. & Manzanal, M. B. Streptomycetes: a new model to study cell death. *Int. Microbiol.* 3, 153–158 (2000).
6. Hopwood, D. A. *Streptomyces in Nature and Medicine: the Antibiotic Makers* (Oxford Univ. Press, 2007).
7. Chater, K. F. & Losick, R. in *Bacteria as Multicellular Organisms* (eds Shapiro, J. A. & Dworkin, M.) 149–182 (Oxford Univ. Press, 1997).
8. Barz, E. A. et al. Taxonomy, physiology, and natural products of the *Actinobacteria*. *Microbiol. Mol. Biol. Rev.* 80, 1–43 (2016).
9. Flardh, K. & Buttner, M. J. *Streptomyces* morphogenetics: dissecting differentiation in a filamentous bacterium. *Nat. Rev. Microbiol.* 7, 36–49 (2009).
10. Chater, K. F. in *Streptomyces: Molecular Biology and Biotechnology* (ed. Dyson, P.) 43–86 (Caister Academic, 2011).
11. McCormick, J. R., Su, E. P., Driks, A. & Losick, R. Growth and viability of *Streptomyces coelicolor* mutant for the cell division gene ftsZ. *Mol. Microbiol.* 14, 243–254 (1994).
12. McCormick, J. R. Cell division is dispensable but not irrelevant in *Streptomyces*. *Curr. Opin. Biotechnol.* 12, 689–698 (2001).
13. Jablonskiwicz, M. & van Wezel, G. P. Cell division and DNA segregation in *Streptomyces*: how to build a septum in the middle of nowhere? *Mol. Microbiol.* 59, 393–404 (2012).
14. Manteca, A., Alvarez, R., Salazar, N., Yague, P. & Sanchez, J. Mycelium differentiation and antibiotic production in submerged cultures of *Streptomyces coelicolor*. *Appl. Environ. Microbiol.* 74, 3877–3886 (2008).
15. Flardh, K. Cell polarity and the control of apical growth in *Streptomyces*. *Curr. Opin. Microbiol.* 13, 758–765 (2010).
16. Holmes, N. A. et al. Coiled-coil protein Sce is a key component of a multiprotein assembly controlling polarized growth in *Streptomyces*. *Proc. Natl Acad. Sci. USA* 110, E397–E406 (2013).
17. Granettchavara, N., Lustig, U. & Flardh, K. Dynamics of FtsZ assembly during sporulation in *Streptomyces coelicolor* A32). *J. Bacteriol.* 187, 3227–3237 (2005).
18. Schwedock, J., McCormick, J. R., Angert, E. R., Nodwell, J. R. & Losick, R. Assembly of the cell division protein FtsZ into ladder like structures in the aerial hyphae of *Streptomyces coelicolor*. *Mol. Microbiol.* 25, 847–858 (1997).
19. Willensme, J. & van Wezel, G. P. Imaging of Streptomyces coelicolor A32) with reduced auto-fluorescence reveals a novel stage of FtsZ localization. *PLoS ONE* 4, e4242 (2009).
20. de Boer, P. A. Advances in understanding E. coli cell fission. *Curr. Opin. Microbiol.* 13, 730–737 (2010).
21. Schumann, J., Mollering, H. & Jaenicke, R. Intrinsic stability and extrinsic stabilization of creatin from *Pseudomonas putida*. *Biochem. Hoppe-Seyler’s Z. Physiol. Chem.* 374, 427–434 (1993).
22. Wu, L. J. & Errington, J. Nucleoid occlusion and bacterial cell division. *Nat. Rev. Microbiol.* 10, 8–12 (2011).
23. Reynolds, P. E. Structure, biochemistry and mechanism of action of glycopeptide antibiotics. *Eur. J. Clin. Microbiol. Infect. Dis.* **8**, 943–950 (1989).

24. Jyothikumar, V. et al. Cardiolipin synthase is required for *Streptomyces coelicolor* morphogenesis. *Mol. Microbiol.* **84**, 181–197 (2012).

25. Flärdh, K., Richards, D. M., Hempel, A. M., Howard, M. & Buttner, M. J. Regulation of apical growth and hyphal branching in *Streptomyces*. *Curr. Opin. Microbiol.* **15**, 737–743 (2012).

26. Manteca, A., Fernandez, M. & Sanchez, J. A death round affecting a young *NATURE COMMUNICATIONS*| DOI: 10.1038/ncomms11836 | www.nature.com/naturecommunications

24. Jyothikumar, V.

27. Glauert, A. M. & Hopwood, D. A. A membranous component of the cytoplasm in *Streptomyces coelicolor*. *J. Biophys. Biochem. Cytol.* **6**, 515–516 (1959).

28. Glauert, A. M. & Hopwood, D. A. The fine structure of *Streptomyces coelicolor*. I. The cytoplasmic membrane system. *J. Biophys. Biochem. Cytol.* **7**, 479–488 (1960).

29. Jyothikumar, V., Tilley, E. J., Wali, R. & Herron, P. R. Time-lapse microscopy of *Streptomyces coelicolor* growth and sporulation. *Appl. Environ. Microbiol.* **74**, 6774–6781 (2008).

30. Kretschmer, S. Location of branches within the apical hyphal region of *Streptomyces granaticolor* mycelia. *J. Basic Microbiol.* **32**, 35–42 (1992).

31. Junglas, B. et al. *Ignicoccus hospitalis* and *Nanoarchaeum equitans*: ultrastructure, cell-cell interaction, and 3D reconstruction from serial sections of freeze-substituted cells and by electron cryotomography. *Arch. Microbiol.* **190**, 395–408 (2008).

32. Santarella-Mellwig, R., Pruggnaller, S., Roos, N., Mattaj, I. W. & Devos, D. P. Three-dimensional reconstruction of bacteria with a complex endomembrane system. *PLoS Biol.* **11**, e1001565 (2013).

33. Boekema, E. J., Scheffers, D. J., van Bezouwen, L. S., Bolhuis, H. & Folea, I. M. Focus on membrane differentiation and membrane domains in the prokaryotic cell. *J. Mol. Microbiol. Biotechnol.* **23**, 345–356 (2013).

34. Schlimpert, S. et al. General protein diffusion barriers create compartments within bacterial cells. *Cell* **151**, 1270–1282 (2012).

35. Celler, K., Koning, R. I., Koster, A. J. & van Wezel, G. P. Multidimensional view of the bacterial cytoskeleton. *J. Bacteriol.* **195**, 1627–1636 (2013).

36. Wolanski, M. et al. Replisome trafficking in growing vegetative hyphae of *Streptomyces coelicolor* A3(2). *J. Bacteriol.* **193**, 1273–1275 (2011).

37. Verkade, P. Moving EM: the Rapid Transfer System as a new tool for correlative light and electron microscopy and high throughput for high-pressure freezing. *J. Microsc.* **230**, 317–328 (2008).

38. Koning, R. I. et al. MAVIS: an integrated system for live microscopy and vitrification. *Ultramicroscopy* **143**, 67–76 (2014).

39. Kisser, T., Bibb, M. J., Buttner, M. J., Chater, K. F. & Hopwood, D. A Practical Guide to *Streptomyces* Genetics (John Innes Foundation, 2000).

40. Kremer, J. R., Mastronarde, D. N. & McIntosh, J. R. Computer visualization of three-dimensional image data using IMOD. *J. Struct. Biol.* **116**, 71–76 (1996).

41. Frangakis, A. S. & Hegerl, R. Noise reduction in electron tomographic reconstructions using nonlinear anisotropic diffusion. *J. Struct. Biol.* **135**, 239–250 (2001).

**Acknowledgements**

We are grateful to Dennis Claessen for the gift of pGreen and for discussions. *S. coelicolor* M145 was obtained from the John Innes Centre strain collection, and *S. albus* G. ATCC 25426 was obtained from the American Type Culture Collection strain collection. This work was supported in part by Cytrron II (Life Sciences Health framework FES0908); the Netherlands Centre for Electron Nanoscopy, Leiden; the Netherlands Organization for Scientific Research (NWO VICI grant 10379; grant 175.010.2009.001) and the European Regional Development Fund of the European Commission (21Z,014). We would like to acknowledge support of R.K. from the European Union through the Horizon 2020 Programme grant iNEXT (grant agreement No. 653706).

**Author contributions**

K.C. and R.I.K. performed the cryo-correlative light and electron microscopy. K.C. and J.W. performed the (live) light imaging. G.P.v.W. and A.J.K. supervised the work. K.C., R.I.K. and G.P.v.W. wrote the manuscript. All authors discussed the results and commented on the manuscript.

**Additional information**

Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications

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

Reprints and permission information is available online at http://npg.nature.com/reprintsandpermissions/

How to cite this article: Celler, K. et al. Cross-membranes orchestrate compartmentalization and morphogenesis in *Streptomyces*. *Nat. Commun.* **7**:11836 doi: 10.1038/ncomms11836 (2016).

This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/