Reversible metamorphosis in a bacterium

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ABSTRACT

The cell wall is a shape-defining and protective structure that envelops virtually all bacteria. Wall-less variants, called L-forms, have been generated in laboratories for many decades under highly specialized conditions, invariably aimed at interrupting cell wall synthesis. As such, the relevance of these cells has remained obscure. Here we show that the filamentous actinomycete *Kitasatospora viridifaciens* has the natural ability to switch between a wall-less state and the canonical mycelial mode-of-growth. We show that this organism thrives in a cell wall-less form, and identify the polar growth determinant DivIVA as an essential regulator required for reversible metamorphosis. This is the first report of a reversible metamorphosis in a bacterium that includes wall-less cells as a natural stage in bacterial development.
INTRODUCTION

The cell wall, an essential component of virtually all bacteria, is a dynamic structure that largely determines bacterial cell shape\(^1\). It provides structural rigidity and protection to osmotic stresses and forms the barrier between the bacterium and its environment\(^2,3\). Given its protective role, the cell wall and its biosynthetic enzymes are targets of some of the best known antibiotics, including penicillin and vancomycin\(^4,5\). The synthesis of its major constituent, peptidoglycan (PG), involves the activity of large protein complexes that cooperatively build and incorporate new PG precursors into the growing glycan strands at the cell surface\(^3,6,7\). These strands are then cross-linked to form in essence a single, giant sacculus\(^8\). Major differences in growth and cell division are seen between the planktonic firmicutes that grow via lateral cell-wall growth, and the Actinobacteria, which include many multicellular genera that form complex mycelia and grow via apical extension\(^9,10\). A large number of genes required for PG synthesis and cell division are organized in the highly conserved dcw gene cluster (for division and cell wall synthesis), which contains the cell division scaffold FtsZ\(^11,12\). Some genes in this cluster have gained species-specific functions, or have been lost during evolution. A well-known example is DivIVA: in *Bacillus subtilis*, this protein is involved in septum-site localization by preventing accumulation of the cell division initiator protein FtsZ\(^13\). In contrast, in filamentous actinomycetes, DivIVA is a crucial component of the polarisome complex that drives polar growth\(^14,15\). As a consequence, *divIVA* is dispensable in *B. subtilis* but essential in actinomycetes\(^14,16\). Conversely, cell division is not essential for growth in streptomycetes, and uniquely, *ftsZ* and various other genes that encode members of the divisome can be deleted\(^17,18\). These examples imply a major evolutionary diversification that has separated Actinobacteria from other bacteria.
Although the cell wall is a vital structure for bacteria, several species can be manipulated to produce cells that propagate without their wall, which are known as L-forms\textsuperscript{19-22}. Typically, this was done in an artificial way, by growing strains on osmo-protective media in the presence of high levels of antibiotics targeting the enzymes required for cell-wall synthesis\textsuperscript{23-25}. More recently, L-form growth was also achieved genetically by turning off PG synthesis\textsuperscript{20,26}. These approaches often yield unstable L-form variants that quickly revert to the walled state when the inducing agents are omitted. By contrast, stable L-forms can propagate indefinitely without the cell wall and require two kinds of mutations\textsuperscript{27}. The first class of mutations leads to an increase in membrane synthesis, either directly by increasing fatty acid biosynthesis, or indirectly by reducing cell wall synthesis\textsuperscript{28}. The second class of mutations reduces oxidative damage caused by reactive oxygen species, which are detrimental to formation of L-forms\textsuperscript{29}. Notably, proliferation of L-forms is independent of the FtsZ-based division machinery\textsuperscript{20,30}. Instead, their proliferation can be explained solely by biophysical processes, in which an imbalance in the cell surface area to volume ratio leads to spontaneous blebbing and the generation of progeny cells\textsuperscript{28}. This purely biophysical mechanism of L-form proliferation is independent of the prokaryotic lineage. This observation has led to the hypothesis that early life forms propagated in a similar fashion well before the cell wall had evolved\textsuperscript{20,28,31}. Until now, no examples are known of bacteria that generate L-form cells in the absence of the commonly used inhibitors of cell-wall synthesis and that propagate without accumulating mutations.

Here we present evidence that filamentous actinomycetes have the natural ability to form such wall-less cells. These cells can propagate in the wall-less state, but can also revert to mycelial growth. Our work provides the first direct evidence that L-form cells are a natural stage in bacterial life.
Transitions between filamentous and L-form growth

One of the key requirements for growth of L-form cells is the presence of an osmotically balanced medium to prevent cell lysis. Unexpectedly, when the actinomycete *Kitasatospora viridifaciens* (previously called *Streptomyces viridifaciens*\(^{32}\)) is grown in liquid LPB (containing high levels of sucrose) in the absence of any cell wall synthesis inhibitor, spherical vesicles were visible at low frequency in the culture broth and in between the mycelial networks (Fig. 1A). Replacing sucrose by an equimolar amount of NaCl also leads to the formation of such vesicles, although they are less abundant and smaller than those formed in LPMA containing sucrose (Extended Data Fig. S1B). In the absence of either osmolyte, we could not detect any vesicles (Extended Data Fig. S1C). The spherical vesicles often contained smaller vesicles inside and are surrounded by a membrane. Importantly, staining with SYTO9 indicated that these vesicles contained DNA (Fig. 1B), while transmission EM indicated that they contained no cell wall (Fig. 1C, D), which make them very reminiscent of L-form-like cells. Comparable DNA-containing vesicles were also observed during growth of *Kitasatospora* strains MBT63 and MBT66\(^{32}\), as well as *Streptomyces venezuelae*. However, they were not detected in *Streptomyces coelicolor*, *Streptomyces griseus* or in *Streptomyces lividans* (Extended Data Fig. S2).

To determine the location where these L-form-like cells originate from in the context of the hyphae, we performed live imaging of growing colonies (Fig. 1E; Extended Data Video S1). Approximately 7 hr after the visible emergence of germ tubes, we detected a transient arrest in tip extension of the leading hypha (Fig. 1E, t=430 mins). By that time, L-form-like cells became visible, which were extruded from the hyphal tip (see arrow in Fig. 1E). After the extrusion process, the leading
hypha resumed elongation. Subapically, new branches became visible about 210 minutes after the first appearance of these cells (Extended Data Video S1, t= 640 min). Notably, such branches frequently also extruded L-form-like cells, similarly to the leading hypha (Extended Data Video S2). This showed that the L-form-like cells are produced at hyphal tips.

To study the viability of these L-form-like cells, we separated them from the mycelia by filtration, and used the filtrate to inoculate fresh LPB medium. Strikingly, these cells were able to proliferate in a manner reminiscent of the extrusion-resolution mechanism that was previously described for proliferation of B. subtilis L-forms (Fig. 1F; Extended Data Video S3). Apparently, these wall-less cells are natural L-forms that can proliferate without the need for any genetic mutations. To discriminate these natural L-forms from other stable and unstable L-forms generated by inducing agents, we hereinafter refer to them as N-forms (for natural L-forms).

When N-forms were plated on solid LPMA medium, we noticed that some of the cells generated colonies consisting of both mycelia and N-form cells, implying that these N-forms can revert to mycelial growth (Fig. 1G; Extended Data Video S4). To exclude that mycelial growth was caused by outgrowing spores present in the filtrate, we also analysed a non-sporulating ΔssgB mutant of K. viridifaciens (Extended Data Fig. S3A-C). Like the wild-type strain, the N-forms of the ΔssgB mutant propagated in the wall-less state in liquid, and reverted to mycelial growth on solid medium (Extended Data Fig. S3D, E; Extended Data Video S5).

Altogether, this work thus presents the first example of a bacterium with the natural ability to generate wall-less cells as a natural stage in bacterial life. These cells can propagate without their cell wall, but can also revert to mycelial growth (Fig. 1H).
Generation of the stable L-form lineage \textit{alpha}

Although N-forms could propagate in the wall-less state, their phenotype was reversible leading to mixtures of both hyphae and N-forms. To understand the mechanism of this reversible metamorphosis, we aimed to generate a strain that could reproducibly switch between an all-walled and completely wall-less state. To this end, we exposed the wild-type strain to penicillin and lysozyme following a weekly sub-culturing regime to obtain a derivative strain, which we designated \textit{alpha}. As intended, in sucrose-based media \textit{alpha} exclusively proliferated in the L-form-like state and showed no signs of reversion (Fig. 2A; Extended Data Video S6).

Characterization using transmission electron microscopy (TEM) confirmed that \textit{alpha} contained no PG-based cell wall (Fig. 2B). Excitingly, while \textit{alpha} cells preferred the L-form state, they still possessed the capacity of reversible metamorphosis. When \textit{alpha} was streaked onto plates lacking osmotic protectants, mycelial colonies were obtained (Fig. 2C). Conversely, when mycelial colonies were transferred to plates containing sucrose, the formation of L-form cells was readily induced, which, like in the parental wild-type strain, emerged from the hyphal tip (Extended Data Video S7).

However, unlike the N-forms generated by the wild-type strain, the L-forms formed by \textit{alpha} indefinitely propagated in the wall-less state and did not revert to mycelial growth on osmo-protective media.

To see if we could identify the mutations that predisposed proliferation in the L-form state, we performed SNP analysis by comparing the genome sequence of \textit{alpha} to that of the parent DMS40239\textsuperscript{33}. This revealed that \textit{alpha} had lost the 1.7MB linear plasmid KVP1 that is present in the wild-type strain. In addition, three SNPs were identified in the genome (Extended Data Table 1). One of the mutations mapped to a non-coding DNA region between two genes encoding a putative glutamine-
fructose-6-phosphate transaminase and an ABC transporter, respectively. A second
mutation was identified in one of the two \textit{lysX} genes (BOQ63_22710), and which
encodes a bifunctional phosphatidylglycerol lysyltransferase\textsuperscript{34,35}. This protein is
involved in lipid modification by transferring L-lysine from lysyl-tRNA to
phosphatidylglycerol. This mutation resulted in a T183I mutation in LysX. The third
SNP was in \textit{uppP} (BOQ63_21575), resulting in an L58M change in its gene product
undecaprenyl-diphosphate phosphatase. This phosphatase is involved in the recycling
pathway of the carrier lipid undecaprenyl phosphate, which transports glycan
biosynthetic intermediates for cell wall synthesis\textsuperscript{36}. While their function related to
cell-wall or membrane synthesis is suggestive at least, and they may therefore
facilitate proliferation in the L-form state, none of them blocked reversible
metamorphosis.

\textbf{DivIVA is essential for reversible metamorphosis}

DivIVA is required for polar growth in \textit{Streptomyces} and \textit{Corynebacteria}\textsuperscript{14,16}. We
therefore wondered if DivIVA has a role in the formation or proliferation of \textit{K. viridifaciens} L-forms, which lack any obvious polarity. To address this question, we
first created plasmid pKR3, which leads to the constitutive expression of a C-terminal
eGFP fusion to DivIVA, which was partially active (see below). Analysis of a
transformant constitutively expressing DivIVA-eGFP revealed that the chimeric
protein localized to hyphal tips (Fig. 2D). This apical localization pattern is consistent
with earlier findings in streptomycetes\textsuperscript{14}. Notably, some hyphae expressing the fusion
protein were wider and showed hyphal tip-splitting, which may be due to a polar
effect or deregulated expression of DivIVA. To localize DivIVA in the L-form state,
we made use of the fact that the generated strain reverts to the L-form mode-of-
growth on osmoprotective media. Despite the absence of polarity, L-forms containing pKR3 typically showed one or two DivIVA foci per cell, which invariably were localized to the membrane (Fig. 2D). In contrast, no foci were detected in L-form cells constitutively expressing eGFP (pGreen) or those containing the empty plasmid (pKR1).

We then constructed plasmids pKR4 to delete divIVA and pKR5 to delete a large part of the dcw gene cluster, which includes divIVA\textsuperscript{11} (Fig. 3A). Introduction of these plasmids into alpha by transformation and subsequent screening yielded the desired divIVA and dcw mutants. Western blot analysis using antibodies against DivIVA of Corynebacterium glutamicum confirmed the absence of DivIVA in both the divIVA and the dcw mutant (Fig. 3B). Analysis of growth in LPB medium or on solid LPMA plates indicated that the L-form cells proliferated normally in the absence of divIVA or part of the dcw gene cluster (Fig. 3C, D). However, when L-form cells were plated on MYM medium, only the alpha strain could grow due to its ability to form mycelial cells, while the divIVA and dcw mutants could not grow due to their failure to undergo reversible metamorphosis (Fig. 3D). Introduction of plasmid pKR6 containing sepF-sepG-divIVA under control of their natural promoters restored growth to the divIVA null mutant on MYM medium (Fig. 3D). To exclude that this was caused by the additional sepF and sepG genes, we also introduced plasmid pKR7, which expresses divIVA from the constitutive gap1 promoter. This plasmid also complemented growth of the divIVA mutant on MYM medium, thus establishing that divIVA alone was sufficient to complement the divIVA null mutant (Fig. 3D).

Importantly, the DivIVA-eGFP fusion protein also restored filamentous growth to the divIVA mutant, implying that the chimeric protein is largely functional \textit{in vivo}. However, some hyphae showed apical branching or tip-splitting indicative of polar
effects on growth and development (Extended Data Fig. S4). Altogether, these results identify DivIVA as a key protein that is essential for reversible metamorphosis.

**Functional complementation of the *K. viridifaciens dcw* mutant by the *S. coelicolor dcw* gene cluster**

The unique capacity of *alpha* to switch back-and-forth between a walled, filamentous mode-of-growth and a wall-less L-form state provides a unique platform to apply an engineering approach to cell morphology design. As a first step towards that goal, we introduced the *dcw* gene cluster from *S. coelicolor* into the *K. viridifaciens dcw* mutant (Fig. 3D, E). Excitingly, introduction of this gene cluster restored filamentous growth and reversible metamorphosis (Fig. 3D, E), demonstrating that the *S. coelicolor dcw* cluster functionally replaces the *dcw* cluster of *K. viridifaciens*. This is an important observation as it also shows that the ability of *K. viridifaciens* to undergo reversible metamorphosis is not dictated by specific adaptation in this organism of genes within its *dcw* cluster. Staining of nascent PG synthesis by fluorescent vancomycin (Van-FL) revealed a ladder-like staining pattern in the complemented *dcw* mutant, which could not be discriminated from that observed for the *alpha* strain growing as a mycelium (Fig. 3E). These ladders represent sites where PG-based septa are formed in an FtsZ-dependent manner. Altogether, this demonstrates that the mycelium formed by this 'hybrid' bacterium is established by the activity of the macromolecular machinery of two different bacteria belonging to separate genera. This paves the way for extending this principle with *dcw* gene clusters of morphologically distinct bacteria, such as the unicellular actinobacteria *Corynebacterium glutamicum* and *Mycobacterium tuberculosis*. 
DISCUSSION

Filamentous actinomycetes have been studied for more than 50 years as a model for multicellular bacterial development. We here provide compelling evidence that the formation of wall-less cells is a natural, and previously unnoticed developmental stage in these organisms. These natural wall-less cells, which we have dubbed N-forms, are extruded from the hyphal tips, contain DNA and are viable as concluded from their ability to form colonies. This work thus presents the first example of a bacterium with the natural ability to generate wall-less cells and provides a major step in our ability to reveal the ecological relevance of these enigmatic wall-less cells. It will be of key interest to dissect how this unique morphogenetic switch is regulated and to disclose how widespread this phenomenon is.

The spontaneous release of N-forms provides an important step forward towards characterising where such cells are found in nature. Actinomycetes are abundantly present in soil and marine environments and are known to interact with many eukaryotic organisms, such as insects and plants. Interestingly, the phloem of plants is rich in sucrose and provides a supportive environment to sustain wall-less cells. Given their presence in soil environments, we envision that these actinomycetes may associate with plant roots in their filamentous state, followed by a switch to the wall-less state allowing them to quickly travel throughout the plant tissue or via the phloem. In this way, they may be able to traverse long distances to the sites where their activity is required, for instance in providing protection to the plant mediated by their large repertoire of secondary metabolites they produce.

However, for other species this transient mode-of-growth, during which the wall-less cells may escape from the plant immune system, could also help to cause damage to plants. In this context it is interesting to note that we have been able to induce L-form
growth in the potato pathogen *Streptomyces scabies*, which, as a non-sporulating
strain, is known to traverse long distances within the plant (R. Loria, personal
communication). More generally, we suppose that wall-less cells may be more
commonly present in nature, but have simply been overlooked due to their specific
requirements for growth.

N-forms are released from hyphal tips, which contain extended membrane
structures \(^3\). One of the key requirements for L-form proliferation is an upregulation
of membrane synthesis \(^2\). We think that the presence of these extended membranes at
the hyphal tip may facilitate the formation of N-forms in these actinomycetes. Their
release from the hyphal tip coincides with an arrest in tip extension, indicating that
PG synthesis at those sites is transiently blocked. Notably, sequencing of the genome
of the stable L-form lineage *alpha* identified two mutations that may exert their effect
on PG synthesis. One mutation was identified in the non-coding region of a gene
encoding a protein controlling the flux of glucose in the hexosamine pathway,
ultimately leading to synthesis of PG precursors. A second mutation was found in
*uppP*, which is involved in PG precursor recycling. While these mutations may
regulate cell-wall precursor accumulation, none of them was essential for mycelial
growth. This can be derived from the fact that both N-forms as well as the L-forms
formed by *alpha* may revert to mycelial growth. However, the fact that the mutations
do not prevent mycelial growth does not rule out that they may have been essential for
the formation of a previous lineage that finally evolved into *alpha*. The role of the
mutations and their contribution to the formation of N-forms and proliferation of
*alpha* is under current investigation.

Summarising, our work on reversible metamorphosis provides the first
evidence for the existence of natural wall-less cells, and thereby broadens the large
diversity in bacterial cell shapes\textsuperscript{1}. Finally, it will enable us to design ‘hybrid’ bacteria with dramatically changed cell morphologies by replacing some of the most fundamental shape-defining genes. Ultimately, this should allow us to convert a truly multicellular bacterium into a walled, unicellular derivative, which is not only exciting from an evolutionary point-of-view, but which may also prove valuable as an industrial workhorse for the biosynthesis of natural products. The functional replacement of the \textit{dcw} gene cluster between bacteria from distinct genera provides an important first step towards that goal.

\section*{METHODS}

\subsection*{Strains and media}

Bacterial strains used in this study are shown in Extended Data Table 2. To obtain sporulating cultures, \textit{Streptomyces} and \textit{Kitasatospora} species were grown at 30°C for 4 days on MYM medium, containing (w/v) 0.4% maltose, 0.4% yeast extract, 1% malt extract, 2% Iberian agar, 0.2 (v/v) R5 trace elements\textsuperscript{43}. To support growth of wall-less cells, strains were grown in liquid L-Phase Broth (LPB), containing (w/v) 0.15% yeast extract, 0.25% bacto-peztone, 0.15% oxoid malt extract, 0.5% glucose, 22% sucrose, 1.5% oxoid tryptone soya broth powder and 25 mM MgCl\textsubscript{2}. Alternatively, wall-less cells were propagated in liquid L-Phase Medium (LPM), containing (w/v) 0.5% glucose, 0.5% yeast extract, 0.5% peptone, 20% sucrose, 0.01% MgSO\textsubscript{4} \cdot 7H\textsubscript{2}O, and 25 mM MgCl\textsubscript{2}. Cultures were inoculated with $10^6$ spores ml\textsuperscript{-1} and grown in 250 ml flasks. Cultures were incubated at 30°C, while shaking at 100 rpm. For propagation of wall-less cells on solid medium, we used LPMA (i.e. LPM supplemented with (w/v) 0.75% Iberian agar and 5% (v/v) horse serum).
Generation of the stable *K. viridifaciens* L-form lineage alpha was performed in LPB, following the cultivation regime previously described\(^4\). 50 ml cultures were grown in 250 ml flasks in an orbital shaker at 100 rpm.

For general cloning purposes, *E. coli* strains DH5α and JM109 were used, while *E. coli* ET12567 and SCS110 were used to obtain unmethylated DNA (Extended Data Table 2). *E. coli* strains were grown at 37 °C in LB medium, supplemented with chloramphenicol (25 \(\mu\)g ml\(^{-1}\)), ampicillin (100 \(\mu\)g ml\(^{-1}\)), apramycin (50 \(\mu\)g ml\(^{-1}\)), kanamycin (50 \(\mu\)g ml\(^{-1}\)), or viomycin (30 \(\mu\)g ml\(^{-1}\)), where necessary.

### Construction of plasmids

All plasmids and primers used in this work are shown in Extended Data Table 3 and Extended Data Table 4, respectively.

#### Construction of the DivIVA localization construct pKR3

To localize DivIVA, we first created plasmid pKR2 containing a viomycin resistance cassette cloned into the unique NheI site of pIJ8630\(^45\). To this end, the viomycin resistance cassette was amplified from pIJ780\(^46\) with the primers *vph*-FW-NheI and *vph*-RV-NheI. Next, we amplified the strong constitutive gap1 promoter as a 450 bp fragment from the genome of *S. coelicolor* with the primers P\(_{Gap1}\)-FW-BglII and P\(_{Gap1}\)-RV-XbaI. We also amplified the *divIVA* coding sequence (the +1 to +1335 region relative to the start codon of *divIVA* (BOQ63_31065) from the chromosome of *K. viridifaciens* using primers *divIVA*-FW-XbaI and *divIVA*-no stop-RV-NdeI.

Finally, the promoter and *divIVA* coding sequence were cloned into pKR2 as a BglII/XbaI and XbaI/NdeI fragment respectively, yielding plasmid pKR3.
Construction of the deletion constructs pKR1, pKR4 and pKR5

The \textit{ssgB} mutant was created in \textit{K. viridifaciens} using pKR1, which is a derivative of the unstable plasmid pWHM\textsuperscript{47}. In the \textit{ssgB} mutant, nucleotides +20 to +261 relative to the start codon of \textit{ssgB} were replaced with the \textit{loxP-apra} resistance cassette as described\textsuperscript{48}. A similar strategy was used for the deletion of the \textit{divIVA} gene or part of the \textit{dcw} cluster, using plasmids pKR4 and pKR5, respectively. In the \textit{divIVA} deletion mutant, nucleotides +205 to +349 relative to the start codon of \textit{divIVA} (BOQ63_31065) were replaced with the apramycin resistance cassette, whereas in the \textit{dcw} mutant the chromosomal region from +487 bp relative to the start of the \textit{ftsW} gene (BOQ63_31025) until +349 relative to the start of the \textit{divIVA} gene was replaced with the apramycin resistance marker.

Construction of the complementation constructs pKR6 and pKR7

The constructs pKR6 and pKR7 were created to complement the \textit{divIVA} mutant. For complementation of \textit{divIVA} under control of its native promoter, the genomic sequence from -1565 to +1357 relative to the translational start site of \textit{divIVA} was amplified from the chromosomal DNA of \textit{K. viridifaciens} using the primers \textit{sepF-sepG-divIVA-FW-BglII} and \textit{sepF-sepG-divIVA-RV-XbaI}. This product contained the coding sequences of \textit{sepF}, \textit{sepG} and \textit{divIVA} including the native \textit{divIVA} promoter, and was cloned into the integrative vector pIJ8600\textsuperscript{45}.

The pKR7 plasmid was constructed for the expression of \textit{divIVA} from the constitutive \textit{gap1} promoter. To this end, the promoter region of \textit{gap1} was amplified with the primers P\textsubscript{gap1}-FW-BglIII and P\textsubscript{gap1}-RV-XbaI using \textit{S. coelicolor} genomic DNA as the template. The \textit{divIVA} coding sequence was amplified from the genome of \textit{K. viridifaciens} with the primers \textit{divIVA-XbaI-FW} and \textit{divIVA-NdeI-RV}. Finally, both
the gapI promoter and divIVA coding sequence were cloned as BglII/XbaI and XbaI/NdeI fragments into the integrative vector pIJ8600\textsuperscript{45}. All constructs were sequenced for verification.

For the orthologous complementation of the \textit{K. viridifaciens} dcw mutant, the \textit{S. coelicolor} cluster from cosmid ST4A10\textsuperscript{49} was used. The ST4A10 cosmid was cut with BglII and Scal, followed by gel extraction of a 13,268 bp BglII fragment, encompassing the partial \textit{S. coelicolor} dcw cluster. This fragment was subsequently ligated into BglII-digested pIJ8600, yielding pKR8.

**Filtration of N-forms**

An individual colony of \textit{K. viridifaciens} DSM40239 that had been grown on MYM medium for 4 days was used to inoculate 50 ml LPB medium. The culture was grown for 7 days at 30°C in an orbital shaker at 100 rpm. To separate the mycelium from the N-forms, the culture was passed through a sterile filter made from an EcoCloth\textsuperscript{TM} wiper. A subsequent filtration step was done by passing the N-forms through a 5 µm Isopore\textsuperscript{TM} membrane filter. The filtered cells were centrifuged at 1,000 rpm for 40 min, after which the supernatant was carefully removed by decantation to avoid disturbance of the cells.

**Transformation of L-forms**

Transformation of \textit{alpha} essentially followed the protocol for the rapid small-scale transformation of \textit{Streptomyces} protoplasts\textsuperscript{50}, with the difference that 50 µl cells from a mid-exponential growing L-form culture were used instead of protoplasts.

Typically, 1 µg DNA was used for each transformation. Transformants were selected by applying an overlay containing the required antibiotics in P-buffer after 20 hours.
Further selection of transformants was done on LPMA medium containing the appropriate antibiotics.

**Microscopy**

Strains grown in LPB or LPMA were imaged using a Zeiss Axio Lab A1 upright microscope equipped with an AxioCam Mrc. A thin layer of LPMA (without horse serum) was applied to the glass slides to immobilize the cells prior to the microscopic analysis.

**Fluorescence microscopy**

Fluorescence microscopy pictures were obtained with a Zeiss Axioskope A1 upright fluorescence microscope with an AxioCam Mrc5 camera. Live cells were stained with SYTO 9 (0.05 mM) and BODIPY vancomycin (1 µg ml\(^{-1}\)) for 10 min to detect DNA and nascent PG, respectively. Stains were obtained from Molecular Probes\textsuperscript{TM}. The fluorescent images were obtained using a 470/40 nm band pass excitation and a 505/560 band pass detection, using an 100x N.A. 1.3 objective. Stained live cells where immobilized on microscope slides using a thin layer of LPMA (without horse serum). To obtain a sufficiently dark background, the background of the images was set to black. These corrections were made using Adobe Photoshop CS5.

**Time-lapse microscopy**

For visualization of N-form formation, spores of *K. viridifaciens* were pre-germinated in TSBS (i.e. TSB containing 10% sucrose) for 5 hours. Aliquots of 10 µl of the recovered germlings were placed in an ibiTreat 35 mm low imaging dish (ibidi\textsuperscript{®}), after which an LPMA patch was placed on top of the germlings. For visualization of
N-form growth, 50 µl aliquots of filtered N-forms were used. All samples were imaged for ~15 hours using an inverted Zeiss Axio Observer Z1 microscope equipped with a Temp Module S (PECON) stage-top set to 30°C. Z-stacks with a 1 µm spacing were taken every six minutes using a 40x water immersion objective. Average intensity projections of the in-focus frames were used to compile the final movies. Light intensity over time was equalised using the correct bleach plugin of ImageJ (version 1.51f).

To visualize the proliferation of alpha or filtered N-forms, cells were collected and resuspended in 300 µl LPB (containing 4-22% sucrose) and placed in the wells of a chambered 8-well µ-slide (ibidi®). Cells were imaged on a Nikon Eclipse Ti-E inverted microscope equipped with a confocal spinning disk unit (CSU-X1) operated at 10,000 rpm (Yokogawa), using a 40x Plan Fluor Lens (Nikon) and illuminated in bright-field. Images were captured every 2 minutes for 10-15 hours by an Andor iXon Ultra 897 High Speed EM-CCD camera (Andor Technology). Z-stacks were acquired at 0.2-0.5 µm intervals using a NI-DAQ controlled Piezo element. During imaging wall-less cells were kept at 30 °C using INUG2E-TIZ stage top incubator (Tokai Hit).

Electron microscopy

For transmission electron microscopy, L-forms obtained from a 7-day-old liquid culture of the K. viridifaciens alpha strain were trapped in agarose blocks prior to fixation with 1.5% glutaraldehyde. In contrast, cultures of the wild-type strain forming N-forms were immediately fixed for an hour with 1.5% glutaraldehyde before being filtered (see above). Filtered N-forms were then washed twice with 1X PBS prior to embedding in 2% low melting agarose. A post-fixation step with 1% OsO₄ was performed on both L- and N-forms. Samples were embedded in Epon and sectioned.
into 70 nm slices. Samples were stained using uranyl-acetate (2%) and lead-citrate (0.4%) if necessary. Samples were imaged using a Jeol 1010 or a Fei 12 BioTwin transmission electron microscope.

**Genome sequencing and SNP analysis**

For genomic DNA isolation, the stable L-form cell line *alpha* was grown in LPB medium for 3 days. After harvesting of the cells by centrifugation, genomic DNA was isolated as described\(^5\). Whole-genome sequencing and SNP analyses were performed by BaseClear (Leiden, The Netherlands), using the sequence of the parental strain as a reference\(^3\).

**DivIVA detection using Western analysis**

To detect DivIVA using Western analysis, biomass of L-form strains was harvested after 7 days of growth in LPB medium (100 rpm), while biomass of mycelial strains was obtained from TSBS medium (200 rpm) after 17 hours. Cell pellets were washed twice with 10% PBS, after which they were resuspended in 50mM HEPES pH 7.4, 50 mM NaCl, 0.5% Triton X-100, 1 mM PFMS and P8465 protease inhibitor cocktail (Sigma). The cells and mycelia were disrupted with a Bioruptor Plus Sonication Device (Diagenode). Complete lysis was checked by microscope, after which the soluble cell lysate was separated by centrifugation at 13,000 rpm for 10 min at 4°C. The total protein concentration in the cell lysates was quantified by BCA assay (Sigma-Aldrich). Equal amounts of total proteins were separated with SDS-PAGE using 12.5% gels. Proteins were transferred to polyvinylidene difluoride (PVDF) membranes (GE Healthcare) with the Mini Trans-Blot® Cell (Bio-Rad Laboratories) according to the manufacturer’s instructions. DivIVA was detected using a 1:5,000
dilution of polyclonal antibodies raised against *Corynebacterium glutamicum* DivIVA, and kindly provided by Professor Bramkamp (Ludwig-Maximilian-Universität München). The secondary antibody, anti-rabbit IgG conjugated to alkaline phosphatase (Sigma), was visualized with the BCIP/NBT Color Development Substrate (Promega).

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Author Contributions

K.R., J.W., E.U., A.J.W., A.M., A.B., D.H., G.P.W. and D.C. collected the data and aided in data analysis. D.C. and G.P.W. designed the experiments and supervised the research. K.R., G.P.W. and D.C. wrote the manuscript with input from all co-authors.

Competing Financial Interests

The authors declare there are no competing financial interests

LEGENDS

Figure 1. Wall-less cells are a natural stage in bacterial development.

(A) In media containing high levels of sucrose, the filamentous bacterium *K. viridifaciens* forms spherical vesicles (arrowheads), which stain with the dye SYTO9. (B). Transmission electron micrographs indicate that these vesicles are cells that lack the cell wall (C, D). The arrowheads indicate the cell membrane. (E) These wall-less cells (arrows) are extruded from the hyphal tip. Arrowheads indicate new branches, while “S” designates the spore. Images were taken from Extended Data Video S1. (F) Phase-contrast images from Extended Data Video S3 showing proliferation of wall-less cells in LPB medium. The times are indicated in HH:MM. The cell indicated with
an asterisk had entered the field after 38 mins. (G) Reversion of wall-less cells to hyphal growth yields colonies consisting of hyphae and (newly-formed) wall-less cells. Images were taken from Extended Data Video S4. (H) Model indicating that wall-less cells (or N-forms) are a natural stage in bacterial development. On osmo-protective media, vegetative hyphae generate N-form cells that either proliferate transiently in the wall-less state, or immediately revert to mycelial growth (right). Reversion to mycelial growth allows the strain to complete its canonical life cycle (left). Scale bars represent 1 µm (D), 5 µm (C, F), 10 µm (A, B, E) or 20 µm (G).

**Figure 2. Characterization of the stable L-form cell line alpha.** (A) Phase-contrast images from Extended Data Video S6, which demonstrate the proliferation of alpha in the L-form state. The arrowhead points to the dividing mother cell. Please note that after the generation of new progeny the mother cell lysis (indicated with an asterisk). (B) Transmission electron micrograph showing an alpha cell lacking the cell wall. (C) On plates containing sucrose and MgCl₂, alpha forms colonies exclusively containing L-forms, while it forms mycelial colonies on plates without these supplements. (D) L-forms of alpha (alpha⁵) transformed with pGreen, leading to the constitutive expression of eGFP, show a diffuse cytosolic GFP signal. By contrast, L-form cells containing pKR3, thereby expressing eGFP fused to DivIVA, show distinct foci localized to the membrane. When the strains are grown as a mycelium (alpha⁴), DivIVA-eGFP localized to the hyphal tips (arrowheads). No fluorescence is observed in cells or mycelium containing the control plasmid pKR1. Scale bars represent 500 nm (B), 5 µm (A), 10 µm (D) or 20 µm (C).
Figure 3. Reversible metamorphosis depends on DivIVA. (A) Illustration of the dcw clusters in alpha and the derivative strains lacking divIVA or part of the dcw cluster. (B) Western analysis confirming the absence of DivIVA in the ΔdivIVA and Δdcw mutants. Introduction of divIVA under control of its own (pKR6) or the gap1 (pKR7) promoter restores the presence of DivIVA. Likewise, introduction of the S. coelicolor dcw gene cluster (on pKR8) in the ΔdivIVA or Δdcw mutant strains restores the presence of DivIVA. MW is the molecular weight in kDa. (C)

Comparable growth between alpha (solid black line) and the ΔdivIVA (dashed line) and the Δdcw mutants (dotted line) in liquid LPB medium. (D) The mutant strains lacking divIVA can no longer grow on MYM plates lacking sucrose and MgCl2. (D) Reintroduction of the S. coelicolor dcw gene cluster (on pKR8) in the Δdcw mutant restores mycelial growth, including septal PG synthesis, as revealed by staining with fluorescent vancomycin (vanFL). Scale bar represents 5 µm.

Extended Data Figure S1. Osmolytes are required for the formation of wall-less cells. The addition of 0.6M sucrose (A) or NaCl (B) to LPB medium is required for the formation of wall-less cells. No wall-less cells are detected without the addition of osmolytes (C). Scale bars represent 20 µm.

Extended Data Figure S2. Formation of wall-less cells in Streptomyces and Kitasatospora species. Unlike Streptomyces coelicolor (A), Streptomyces lividans (B) and Streptomyces griseus (C), wall-less (arrowheads) are evident in Streptomyces venezuelae (D), and Kitasatospora strains MBT63 (E) and MBT66 (F) grown in LPB medium. The inlays show magnified versions of wall-less cells in these species. (G-I).
Visualization of DNA in wall-less cells using SYTO9. Scale bars represent 5 µm (inlays), 10 µm (G-I) or 20 µm (A-F).

**Extended Data Figure S3. Reversible metamorphosis in the K. viridifaciens ΔssgB mutant.** (A) Deletion of ssgB in K. viridifaciens blocks the formation of grey spores. (B, C) Transmission electron micrographs of N-forms of the ΔssgB mutant strain indicates that they lack the cell wall. The arrowhead indicates the cell membrane (C). (D) Phase-contrast images from Extended Data Video S5 showing N-form proliferation in liquid medium. The times are indicated in HH:MM. (E) Plating of N-forms of the ssgB mutant on LPMA yields colonies consisting of mycelia and N-forms. Scale bars represent 200 nm (C), 2 µm (B), 10 µm (D) or 20 µm (E).

**Extended Data Figure S4. Functional complementation of the ΔdivIVA mutant by DivIVA-eGFP** The DivIVA-eGFP fusion expressed from pKR3 restores filamentous growth of the divIVA mutant, and localizes to the hyphal tips (arrowheads). Scale bars represent 10 µm.

**Extended Data Video S1. Apical extrusion of N-forms in K. viridifaciens.** N-forms are extruded from the hyphal tip after 430 min, coinciding with a transient arrest in tip growth. After extrusion of the N-forms, the hypha resumes elongation after 540 min, while subapically new branches become visible after 620 min. The times are indicated in min.

**Extended Data Video S2. Extrusion of N-forms from branches in K. viridifaciens.** N-forms are extruded from the tips of branches that are formed subapically.
Extended Data Video S3. Proliferation of N-forms. N-forms from K. viridifaciens proliferate in the wall-less state in liquid LPB medium. The arrowhead indicates the cells that are shown in Fig. 1F. The times are indicated in HH:MM. The scale bar represents 10 µm.

Extended Data Video S4. Proliferation and reversion of wild-type N-forms. Growth and reversion of N-forms on solid LPMA medium yields colonies consisting of both hyphae and N-forms. The times are indicated in HH:MM. The scale bar indicates 20 µm.

Extended Data Video S5. Proliferation of N-forms of the ΔssgB mutant strain. N-forms of the ΔssgB mutant strain proliferate in the wall-less state in liquid LPB medium. The arrowhead indicates the cell that is shown in Extended Data Fig. S3D. The times are indicated in HH:MM. The scale bar represents 10 µm.

Extended Data Video S6. Proliferation of alpha in the L-form state. The stable cell line alpha exclusively proliferates in the L-form state in medium containing high levels of sucrose. The arrowhead indicates the proliferating cell that is used to make Fig. 2A. The times are indicated in HH:MM. The scale bar represents 10 µm.

Extended Data Video S7. L-forms are released from hyphal tips in alpha. Transfer of mycelial colonies of alpha on plates containing high levels of sucrose leads to the extrusion of L-forms from the hyphal tips. The times are indicated in min.
Extended Data Table 1. Detected single nucleotide polymorphisms (SNPs) in the chromosome of alpha.

| SNP | Position in genome | Locus | Variant in wild-type | Variant in alpha | Effect on protein | Protein function |
|-----|--------------------|-------|----------------------|-----------------|------------------|------------------|
| 1   | 546 832            | NCR   | C                    | A               | None             | Unknown          |
| 2   | 3 549 271          | BOQ63_22710 | G          | A               | LysX^{T183I}     | Biosynthesis of lysyl-phosphatidyl glycerol |
| 3   | 3 297 354          | BOQ63_21575 | C                    | A               | UppP^{L55M}     | Dephosphorylation of undecaprenyl pyrophosphate |

NCR: non-coding region. LysX: phosphatidylglycerol lysyltransferase, UppP: undecaprenyl diphosphatase.
Extended Data Table 2. Strains used in this study.

| Strains                        | Genotype                                                                 | Reference   |
|-------------------------------|-------------------------------------------------------------------------|-------------|
| **Escherichia coli strains**  |                                                                         |             |
| DH5α                          | F’ Φ80’lacZΔM15 Δ(lacZYA-argF)U169 recA1 endA1 hsdR17(K-, mK+) phoA      | 1           |
|                               | supE44 thi-1 gyrA96 relA1 λ  recA1 endA1 gyrA96 thi hsdR17 supE44 relA1 λ Δ(lac-proAB) [F’ traD36 proAB lacPZΔM15] | 2           |
| JM109                         |                                                                         |             |
| ET12567                       | F’ dam-13::Tn9 (ChlR) dcm-6 hsdM hsdR recF143 zji- 20I::Tn10 galK2 galT22 ara14 lacY1 xyl-5 leuB6 thi-1 tonA31 rpsL136 hisG4 tsx78 mil-1 glnV44 rpsL (StrR) thr leu endA thi-1 lacY galK galT ara tonA tsx dam dcm supE44 Δ(lac-proAB) [F’ traD36 proAB lacPZΔM15] | 3           |
| SCS110                        |                                                                         |             |
| **Streptomyces/Kitasatospora strains** |                                                                        |             |
| Streptomyces coelicolor A3(2) M145 | Wild-type                                                 | Lab collection |
| Streptomyces lividans 1326    | Wild-type                                                 | Lab collection |
| Streptomyces griseus          | Wild-type                                                 | Lab collection |
| Streptomyces venezuelae DIVERSA | Wild-type                                                 | Lab collection |
| Kitasatospora viridifaciens DSM40239 | Wild-type                                                 |            |
| Kitasatospora sp. MBT63       | Wild-type                                                 | 3           |
| Kitasatospora sp. MBT66       |Wild-type                                                 | 3           |
| K. viridifaciens ΔssgB        | K. viridifaciens DSM40239 derivative in which the ssgB gene has been replaced by an apramycin resistance marker | This work |
| **K. viridifaciens L-form strains** |                                                        |             |
| alpha                         | Stable K. viridifaciens L-form cell line                        | This work |
| ΔdivIVA                       | alpha derivative in which the divIVA gene has been replaced by an apramycin resistance marker | This work |
| Δdcw                          | alpha derivative in which part of the dcw gene cluster has been replaced by an apramycin resistance marker. The deleted genes include fisW, murG, fisQ, fisZ, ylmD, ylmE, sepG, sepF, and divIVA. | This work |

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**Extended Data Table 3. Plasmids used in this study.**

| Plasmid   | Description and relevant features                                                                 | Reference |
|-----------|---------------------------------------------------------------------------------------------------|-----------|
| pWHM3     | Unstable, multi-copy and self-replicating *Streptomyces* vector. Contains thiostrepton and ampicillin resistance cassette. | 1         |
| pIJ780    | Plasmid containing a viomycin (vph) resistance cassette.                                           | 2         |
| pIJ8600   | *E. coli–Streptomyces* shuttle vector containing the φC31 attP-int region for genomic integration. Confers resistance to apramycin and thiostrepton. | 3         |
| pIJ8630   | *E. coli–Streptomyces* shuttle vector containing the φC31 attP-int region for genomic integration. Confers resistance to apramycin. | 3         |
| pGreen    | pIJ8630 containing the eGFP gene under control of the constitutive gap1 promoter of *S. coelicolor.* | 4         |
| ST4A10    | Supercosmid fragment containing SCO2068-SCO2105 (43,147bp). Confers resistance to ampicillin and kanamycin. | 5         |
| pKR1      | pWHM3 containing the flanking regions of the *K. viridifaciens* sskB gene interspersed by the apra-loxP cassette. | This study|
| pKR2      | pIJ8630 derivative containing the viomycin resistance cassette from pIJ780 cloned into the unique NheI site. | This work |
| pKR3      | pKR2 derivative containing a C-terminal eGFP fusion to divIVA of *K. viridifaciens* under control of the gap1 promoter of *S. coelicolor.* | This work |
| pKR4      | pWHM3 containing the flanking regions of the *K. viridifaciens* divIVA gene interspersed by the apra-loxP cassette. | This work |
| pKR5      | pWHM3 derivative containing the flanking regions around the *K. viridifaciens* dcw gene cluster interspersed by the apra-loxP cassette. | This work |
| pKR6      | pIJ8600 containing divIVA of *K. viridifaciens* under control of its native promoter (which includes the coding sequences of sepF and sepG). | This work |
| pKR7      | pIJ8600 containing the divIVA gene of *K. viridifaciens* under control of the gap1 promoter of *S. coelicolor.* | This work |
| pKR8      | pIJ8600 containing a 13,268 bp BglII fragment encompassing the murD, ftsW, murG, ftsQ, ftsZ, ylmD, ylmE, sepF, sepG and divIVA genes of the *S. coelicolor* dcw gene cluster. | This work |

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### Extended Data Table 4. Primers used in this study.
Restriction sites are underlined.

| Primer           | Sequence (5’ – 3’) |
|------------------|--------------------|
| P1-ssgB-FW      | GACGAATTCAAGGGTCAGAAGAACGGGTATC |
| P2-ssgB-RV      | GAGTTATCCACACTCTAGAGCTACGGCTGGCTGGTTCATAAGC |
| P3-ssgB-FW      | GAGTTATACTGAGCTAGACTGACTGCTCTTGGAAGAGGAA |
| P4-ssgB-RV      | GACAGCTTTCACCTGAGCCTGGTGTTCAT |
| Delcheck-ssgB-FW| GGCCTGGTACTCCGTGATGATTC |
| Delcheck-ssgB-RV| AGCTTGCGAAGATGTTG |
| vph-FW-Nhel     | GACGCTAGGCGTCAGACCCGTGTTGATACCAAGG |
| vph-RV-Nhel     | GACGCTAGCAATGCTGAGCGAGGCCATCTCTAC |
| PGap-FW-BglIII  | GATTACAGATCCCGAGTTGCTGAGACC |
| PGap-RV-Xbal    | GATGACTTACACGAGTCCTCTCTTGTTGAC |
| divIVA-FW-Xbal  | GTCAAGCTTCTGAGATGCCACCCCGAGGAGGA |
| divIVA-Nostop-RV-Ndel  | GACCATATGGTTGCTCCGGCTCTCGTCAATTAGG |
| P1-divIVA-FW    | GACGACGAAATTCTGATGACCCGTGCTCCACTTG |
| P2-divIVA-RV    | GACGACTTACACGAGTCTAGCCTGAGCCCTGTTC |
| P1-dcw-FW       | GACGATTTCTCCGAGCTACGTACATC |
| P2-dcw-RV       | GACTCTAGAAGCAGAACGACCTGAGCTT |
| P3-dcw-FW       | GACTCTAGAAGCAGACGAGTAGGGAACCCAG |
| P4-dcw-RV       | GAATAGCTTCCGGCTACAACCTAGTTC |
| Delcheck-divIVA-FW | TGACCCGCCCAGGAGACTTTAC |
| Delcheck-divIVA-RV | GGCAGCCCCTCCAAACAC |
| Delcheck-dcw-FW | CCAGAACTGCGTTGATTTCG |
| Delcheck-dcw-RV | GTCTCCAGTACGACTCTAG |
| fisZ-int-FW     | ATGTGGCCGGTGAACCTACCC |
| fisZ-int-Rv     | GACCGCACCCGAAATGATG |
| fisQ-int-FW     | CTTGGGTGTGTCTTCTT |
| fisQ-int-Rv     | CTTGGGTGTGTCTTCTT |
| sepF-sepG-divIVA-FW-BglIII | GACGACAGATCTTCTGATGACCCGTGCTCCACTTG |
| sepF-sepG-divIVA-RV-Xbal | GACGACTTACAAACAGCCGCTACAACCTCAGTTGTC |
| divIVA-Xbal-FW  | GATCAGCTTACATGTGACCCGCTACAACCTCAGTTGTC |
| divIVA-Ndel-RV  | GATCGAATTCTATGCCCGCTACAACCTCAGTTGTC |
| divIVA seq1-FW  | AGCAGCAGATGGGGCAACACAG |
| divIVA seq2-FW  | CGCGCTTGAAGCTGCTACCTG |
| divIVA seq-RV   | ACCTCGTCTCTGCTCATAGC |
| SCO2079_F-520   | TACGGCGCTGTCGAAAGGACGGCC |
| SCO2079_R+1162  | CTCATCGAGGAAGCATCGAGCCTC |
| divIVA_Sco-FW   | AAGCTACCCGCTACCCAG |
| divIVA_Sco-RV   | AGATACGGGCTGCTGATGATTC |
Figure 1. Wall-less cells are a natural stage in bacterial development.

(A) In media containing high levels of sucrose, the filamentous bacterium *K. viridifaciens* forms spherical vesicles (arrowheads), which stain with the dye SYTO9 (B). Transmission electron micrographs indicate that these vesicles are cells that lack the cell wall (C, D). The arrowheads indicate the cell membrane. (E) These wall-less cells (arrows) are extruded from the hyphal tip. Arrowheads indicate new branches, while “S” designates the spore. Images were taken from Extended Data Video S1. (F) Phase-contrast images from Extended Data Video S3 showing proliferation of wall-less cells in LPB medium. The times are indicated in HH:MM. The cell indicated with an asterisk had entered the field after 38 mins. (G) Reversion of wall-less cells to hyphal growth yields colonies consisting of hyphae and (newly-formed) wall-less cells. Images were taken from Extended Data Video S4. (H) Model indicating that wall-less cells (or N-forms) are a natural stage in bacterial development. On osmo-protective media, vegetative hyphae generate N-form cells that either proliferate transiently in the wall-less state, or immediately revert to mycelial growth (right). Reversion to mycelial growth allows the strain to complete its canonical life cycle (left). Scale bars represent 1 µm (D), 5 µm (C, F), 10 µm (A, B, E) or 20 µm (G).
Figure 2. Characterization of the stable L-form cell line alpha. (A) Phase-contrast images from Extended Data Video S6, which demonstrate the proliferation of alpha in the L-form state. The arrowhead points to the dividing mother cell. Please note that after the generation of new progeny the mother cell lysis (indicated with an asterisk). (B) Transmission electron micrograph showing an alpha cell lacking the cell wall. (C) On plates containing sucrose and MgCl₂, alpha forms colonies exclusively containing L-forms, while it forms mycelial colonies on plates without these supplements. (D) L-forms of alpha (alpha⁻) transformed with pGreen, leading to the constitutive expression of eGFP, show a diffuse cytosolic GFP signal. By contrast, L-form cells containing pKR3, thereby expressing eGFP fused to DivIVA, show distinct foci localized to the membrane. When the strains are grown as a mycelium (alpha⁺⁺), DivIVA-eGFP localized to the hyphal tips (arrowheads). No fluorescence is observed in cells or mycelium containing the control plasmid pKR1. Scale bars represent 500 nm (B), 5 µm (A), 10 µm (D) or 20 µm (C).
Figure 3. Reversible metamorphosis depends on DivIVA. (A) Illustration of the dcw clusters in alpha and the derivative strains lacking divIVA or part of the dcw cluster. (B) Western analysis confirming the absence of DivIVA in the ΔdivIVA and Δdcw mutants. Introduction of divIVA under control of its own (pKR6) or the gap1 (pKR7) promoter restores the presence of DivIVA. Likewise, introduction of the S. coelicolor dcw gene cluster (on pKR8) in the ΔdivIVA or Δdcw mutant strains restores the presence of DivIVA. MW is the molecular weight in kDa. (C) Comparable growth between alpha (solid black line) and the ΔdivIVA (dashed line) and the Δdcw mutants (dotted line) in liquid LPB medium. (D) The mutant strains lacking divIVA can no longer grow on MYM plates lacking sucrose and MgCl2. Reintroduction of the S. coelicolor dcw gene cluster (on pKR8) in the Δdcw mutant restores mycelial growth, including septal PG synthesis, as revealed by staining with fluorescent vancomycin (vanFL). Scale bar represents 5 μm.
Extended Data Figure S1. Osmolytes are required for the formation of wall-less cells. The addition of 0.6M sucrose (A) or NaCl (B) to LPB medium is required for the formation of wall-less cells. No wall-less cells are detected without the addition of osmolytes (C). Scale bars represent 20 µm.
**Extended Data Figure S2.** Formation of wall-less cells in *Streptomyces* and *Kitasatospora* species. Unlike *Streptomyces coelicolor* (A), *Streptomyces lividans* (B) and *Streptomyces griseus* (C), wall-less (arrowheads) are evident in *Streptomyces venezuelae* (D), and *Kitasatospora* strains MBT63 (E) and MBT66 (F) grown in LPB medium. The inlays show magnified versions of wall-less cells in these species (G-I). Visualization of DNA in wall-less cells using SYTO9. Scale bars represent 5 μm (inlays), 10 μm (G-I) or 20 μm (A-F).
Extended Data Figure S3. Reversible metamorphosis in the *K. viridifaciens* ΔssaB mutant.
(A) Deletion of *ssaB* in *K. viridifaciens* blocks the formation of grey spores. (B, C) Transmission electron micrographs of N-forms of the ΔssaB mutant strain indicates that they lack the cell wall. The arrowhead indicates the cell membrane (C). (D) Phase-contrast images from Extended Data Video S5 showing N-form proliferation in liquid medium. The times are indicated in HH:MM. (E) Plating of N-forms of the *ssaB* mutant on LPMA yields colonies consisting of mycelia and N-forms. Scale bars represent 200 nm (C), 2 µm (B), 10 µm (D) or 20 µm (E).
Extended Data Figure S4. Functional complementation of the ΔdivIVA mutant by DivIVA-eGFP. The DivIVA-eGFP fusion expressed from pKR3 restores filamentous growth of the divIVA mutant, and localizes to the hyphal tips (arrowheads). Scale bars represent 10 µm.