

Quantitative proteome and phosphoproteome analyses of *Streptomyces coelicolor* reveal proteins and phosphoproteins modulating differentiation and secondary metabolism

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Running title: The *Streptomyces* proteome and phosphoproteome

Key words: *Streptomyces* / phosphoproteomics / differentiation / sporulation / TMT/LC-MS/MS

Abbreviations: LC/MS/MS, Liquid Chromatography-Mass Spectrometry-Mass Spectrometry; PCD, programmed cell death; CPP, calcium phosphate precipitation; TMT, tandem mass tag

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SUMMARY

Streptomyces are multicellular bacteria with complex developmental cycles. They are of biotechnological importance as they produce most bioactive compounds used in biomedicine, e.g. antibiotic, antitumoral and immunosuppressor compounds. *Streptomyces* genomes encode a large number of Ser/Thr/Tyr kinases, making this genus an outstanding model for the study of bacterial protein phosphorylation events. We used mass spectrometry based quantitative proteomics and phosphoproteomics to characterize bacterial differentiation and activation of secondary metabolism of *Streptomyces coelicolor*. We identified and quantified 3461 proteins corresponding to 44.3% of the *S. coelicolor* proteome across three developmental stages: vegetative hypha (MI); secondary metabolite producing hyphae (MII); and sporulating hyphae. A total of 1350 proteins exhibited more than 2-fold expression changes during the bacterial differentiation process. These proteins include 136 regulators (transcriptional regulators, transducers, Ser/Thr/Tyr kinases, signalling proteins), as well as 542 putative proteins with no clear homology to known proteins which are likely to play a role in differentiation and secondary metabolism. Phosphoproteomics revealed 85 unique protein phosphorylation sites, 58 of them differentially phosphorylated during differentiation. Computational analysis suggested that these regulated protein phosphorylation events are implicated in important cellular processes, including cell division, differentiation, regulation of secondary metabolism, transcription, protein synthesis, protein folding and stress responses. We discovered a novel regulated phosphorylation site in the key bacterial cell division protein FtsZ (pSer319) that modulates sporulation and regulates actinorhodin antibiotic production. We conclude that manipulation of distinct protein phosphorylation

events may improve secondary metabolite production in industrial streptomycetes, including the activation of cryptic pathways during the screening for new secondary metabolites from streptomycetes.

INTRODUCTION

Streptomyces is a genus of Gram-positive soil bacterium of great importance for biotechnology given their ability to produce a large array of bioactive compounds, including antibiotics, anticancer agents, immunosuppressants and industrial enzymes (1). *Streptomyces* has a complex morphogenesis which modulates secondary metabolism (2). After spore germination, a fully compartmentalized mycelium (MI) initiates development (3) until it undergoes an ordered programmed cell death (PCD) (4) and differentiates to a second multinucleated mycelium (substrate mycelium, early MII) which activates secondary metabolism (2) before it starts to express the chaplin and rodlin genes encoding the proteins constituting the hydrophobic coats necessary for growth into the air (aerial mycelium; late MII) (5). At the end of the cycle, there is hyphal septation and sporulation (6).

Reversible protein phosphorylation at serine, threonine and tyrosine is a well-known dynamic post-translational modification with stunning regulatory and signalling potential in eukaryotes (7). In contrast, the extent and biological function of Ser/Thr/Tyr protein phosphorylation in bacteria are poorly defined. Phosphorylation in bacteria is dramatically lower than in eukaryotes, making bacterial phosphoproteomics challenging. Several large-scale Ser/Thr/Tyr phosphoproteome studies in bacteria have been reported (8-24) (summarised in Table I), using, in most cases, large amounts of protein (milligrams) obtained during the vegetative growth to detect the relatively low number of phosphopeptides (Table I). This precludes application of isobaric mass

tagging quantitation strategies, since labelling such large amount of peptides is prohibitively expensive. This makes quantitative phosphoproteomics more difficult and to our knowledge, there are only six quantitative phosphoproteomic studies reported for bacteria. Some studies used stable isotope labelling by amino acids in cell culture (SILAC) (25-27), others used the scheduled reaction monitoring analyses (28, 29) and we performed a phosphopeptide profiling (label-free) quantitative proteomics study (16) (see Table I).

Streptomyces coelicolor, the model *Streptomyces* strain (30), has 47 predicted eukaryotic-like protein kinases, twice the number predicted from genomes of other well characterized bacteria, including *E. coli* and *B. subtilis*, but 10-times less than human cells (31). *S. coelicolor* constitutes an outstanding model for the study of bacterial Ser/Thr/Tyr phosphorylation (16). The *S. coelicolor* chromosome was predicted to encode for 30 secondary metabolite biosynthetic pathways, 14 of them not yet experimentally observed (cryptic pathways) (32). None of these secondary metabolites has clinical applications, but *S. coelicolor* still is one of the best models to search for conserved pleiotropic secondary metabolism regulators in streptomycetes.

Here we investigate dynamic regulation of the *S. coelicolor* proteome and phosphoproteome during differentiation in solid sporulating cultures at three developmental time points, 16-, 30- and 65-hours, corresponding to vegetative hyphae (MI), antibiotic producing hyphae (MII) and sporulating hyphae, respectively. This study reports the first isobaric-tag labelling quantitative phosphoproteomic study to identify Ser/Thr/Tyr phosphorylation implicated in secondary metabolism and differentiation in *Streptomyces*. Many of the proteins and phosphorylation sites differentially regulated during the vegetative (MI) and secondary metabolite producing mycelia (MII) are uncharacterised, but conserved in the *Streptomyces* genus. They are

potential modulators of secondary metabolism and differentiation. Genetic manipulation of the genes encoding these biomolecules might improve bioactive compound production in streptomycetes.

EXPERIMENTAL PROCEDURES

Bacterial Strains and Media – The *S. coelicolor* M145 strain was used in this study (30). Solid cultures were grown in Petri dishes (85 mm diameter) with 25 mL of solid GYM medium (glucose, yeast/malt extract) (33) that were covered with cellophane disks, inoculated with 100 μ L of a spore suspension (1×10^7 viable spores/mL) and incubated at 30 °C. *FtsZ* mutants were grown in solid SFM and MM (supplemented with mannitol) media (30). Liquid cultures were performed in sucrose-free R5A (34) in 100 ml flasks, containing 20 ml media, at 200 rpm and 30°C).

Experimental design and statistical rationale – Three key developmental stages were analysed: vegetative hypha (MI_{16h}), secondary metabolite producing hyphae (MII_{30h}) and sporulating hyphae (MII_{65h}) (Fig. 1). Quantitative proteomics and phosphoproteomics were performed by means of TMT-10-plex using 4:3:3 biological replicates from MI_{16h}, MII_{30h} and MII_{65h} respectively (Fig. 1). In the case of the *FtsZ* mutants, samples were collected from sucrose-free R5A liquid cultures at 93-hours, the developmental time-point preceding the maximum phenotypical differences (see results), and TMT-11-plex was used (4:3:4 biological replicates from Ser-Ser, Glu-Glu and Ala-Ala *FtsZ* alleles respectively). As detailed below, abundance changes were statistically significant if they passed the differential expression test between the different developmental stages (q-value less than 0.01).

Sampling of S. coelicolor cells throughout the differentiation cycle - The mycelial lawns of *S. coelicolor* M145 grown on cellophane disks were scraped off at different time points (16- 30- and 65-h; MI, MII and sporulating MII) (Fig. 1) using a plain spatula. In the case of the FtsZ mutants, samples were collected from sucrose-free R5A liquid cultures at 93-hours by centrifugation at 20,000 g for 5 min. Samples were resuspended in 2% SDS, 50 mM pH 7 Tris-HCl, 150 mM NaCl, 10 mM MgCl₂, 1 mM EDTA, 7 mM β-mercaptoethanol, EDTA-free Protease Inhibitor Cocktail Tablets from Roche and 1% phosphatase inhibitor mixtures 1 and 2 (Sigma) and lysed by boiling for 10 minutes; sample viscosity was reduced by sonication (MSE soniprep 150, in four cycles of 10 s, on ice); cell debris were discarded after centrifugation at 20,000 g for 10 min; the sample was cleaned by precipitation with acetone/ethanol (sample/EtOH/acetone 1:4:4 [v/v/v] overnight at -20°C; washed with EtOH/acetone/H₂O 2:2:1 [v/v/v]); resuspended in water; dialyzed against large volumes of water (1 h at 4°C with four water changes); quantified as previously described (35); lyophilized in aliquots of 100 µg; and stored at -80°C.

Protein digestion - Lyophilized proteins were dissolved in 8 M urea in 100 mM TEAB buffer. Protein concentration was measured using bicinchoninic acid assay and adjusted to 2 mg of protein per mL. Samples were reduced with 5 mM DTT at room temperature for 1 h; S-alkylated with 15 mM iodoacetamide (15-30 min at RT in darkness); the alkylation reaction was stopped by adding DTT up to 15 mM; 300 µg of protein were digested using a combined trypsin/LysC digestion protocol (36); samples were desalted using Oasis HLB reverse-phase columns (Waters).

TMT labelling - Samples were labelled using TMT-10-plex or TMT-11-plex reagent (ThermoFisher). The 30- and 65 h samples were analysed in biological triplicate and the 16 h samples were analysed in quadruplicate. The samples from the strains harbouring the FtsZ Ser-Ser (wild-type) and the Ala-Ala alleles were analysed in quadruplicate, while the sample from the strain harbouring the FtsZ Glu-Glu allele in triplicate. Peptide concentration was quantified by amino acid analysis (37). For each condition, 60 µg of peptides was labelled with 0.5 mg of TMT-10-plex reagent following the manufacturer's protocol. The samples were then combined (Fig. 1).

Phosphoenrichment - Phosphopeptides were pre-enriched using calcium phosphate precipitation (CPP) as previously described (16, 38). One-hundred micrograms of TMT labelled peptides were precipitated using CPP and desalted using reverse phase chromatography (POROS R3 resin), before TiO₂ enrichment (16) (Fig 1). The optimal proportion between TiO₂ beads and peptides was tested using 100 µg of TMT labelled peptides (before CPP enrichment) and different amounts of TiO₂ beads. The best phosphopeptide enrichment efficiency, producing an enriched sample containing $48 \pm 1.8\%$ phosphopeptide identifications, was obtained for 0.15 mg TiO₂ beads per 100 µg of TMT labelled peptide sample, vs. the $26 \pm 2.3\%$ and $20 \pm 2.6\%$ obtained with 0.6 and 1.2 mg TiO₂ beads of TMT labelled peptides.

LC-MS/MS analysis of the proteome and phosphoproteome temporal dynamics – Fifty micrograms of TMT-labelled peptides were fractionated by hydrophilic interaction liquid chromatography (HILIC) to generate twelve fractions. Each of the fractions was analysed by LC-MS/MS on EasyLC system (Thermo) coupled to Orbitrap-Fusion-LUMOS. The LC aqueous mobile phase contained 0.1% (v/v) formic acid in water and

the organic mobile phase contained 0.1% (v/v) formic acid in 95% (v/v) acetonitrile. Before injection the trap and analytical columns were pre-equilibrated with 15 and 3.5 μ l buffer A, respectively. The samples were injected on a custom 3-cm trap column (100- μ M internal diameter silica tubing packed with Reprosil 120 C18 5- μ M particles) and desalted with 18 μ l of buffer A. Separation was performed on a home-made 20-cm column (75- μ M internal diameter silica tubing packed with Reprosil 120 C18 3- μ M particles) with a pulled emitter at 250 nL min⁻¹. For total proteome analysis, separation was performed with a 1-8% gradient over 3 min, 8 to 28% over 80 min, 28 to 40% over 10 min and 40 to 100% over 5 min, the column was then kept at 100% buffer B for 8 min. For phosphopeptide analysis, peptides were separated with 1 to 34% buffer B over 60 min and 34 to 100% over 5 min and the column was then kept at 100% for 8 min.

The eluted peptides were analysed on an Orbitrap Fusion mass spectrometer in data-dependent mode. The MS1 spectrum was acquired on an Orbitrap mass analyser at the 400-1600 mass range 120,000 resolution with an AGC target of 5e5 (60 ms maximum injection time). For MS2 scans, peptides were isolated with a quadrupole using a 1.2-Da isolation window and fragmented at 40% normalised collision energy. AGC was set at 5e4 (maximum injection time 120 ms) and dynamic exclusion was 20 s.

Raw data are available via ProteomeXchange with identifier PXD005558.

LC-MS/MS of the S. coelicolor FtsZ mutants - Fifty μ g of TMT labelled peptides were fractionated by high pH LC into thirty fractions. Each of the fractions was analysed by LC-MS/MS using a Dionex Ultimate 3000 system (Thermo) interfaced to a Thermo Orbitrap Fusion Lumos electrospray ionization (ESI) tandem mass spectrometer. The LC mobile phases contained 0.1% (v/v) formic acid in water (A) and 0.1% (v/v) formic acid in 100% (v/v) acetonitrile (B). The samples were injected on a

commercial trap column (300 μ M \times 5mm C18 PepMap 100) and desalted with 18 μ l of solvent A. Separation was performed at 1.1 μ L/min on a home-made 50-cm column (150- μ M internal diameter silica tubing packed with Inertsil C18 1.9- μ M particles) at 35°C, equipped with 10 μ M ID emitter from PepSep (PSFSE10). The gradient was run from 2 to 10% B in 1 minute, from 10 to 12% in 2 minutes, from 12 to 28% in 44 minutes, from 28 to 35% in 10 minutes, from 35 to 47% in 3 minutes. The column was then washed with 95% B for 3 minutes.

The Lumos MS instrument was operated with the recommended settings for SPS analysis in data-dependent MS/MS mode. The MS1 spectrum was acquired using the Orbitrap mass analyzer at m/z 375-1500 at mass resolution 120.000, AGC target of 4e5 (50 ms maximum injection time). For MS2 scans, peptides were isolated with the quadrupole using a 0.7 Da isolation window, fragmented at 35% normalized collision energy. Centroid spectra were collected in the ion trap in turbo mode. For MS3 scans, 10 notches were selected in a 2 Da isolation window and fragmented at 65 NCE by HCD and reporter ions were detected at mass resolution 50.000 in the m/z 100-500 range.

Raw data are available via ProteomeXchange with identifier PXD005558.

Data analysis - Data were analysed in Proteome Discoverer 2.1 software. A database search was performed with Mascot 2.3.2 using *Streptomyces coelicolor* UniProt database (retrieved on 06.03.15, 8038 entries). Methionine oxidation was set as variable modification, while Cys carbamidomethylation, TMT6plex (K) and TMT6plex (N-term) were specified as fixed modifications. A maximum of two trypsin missed cleavages were permitted. The precursor and fragment mass tolerances were 10 ppm

and 0.02 Da respectively. Contaminants from maxQuant website (retrieved May 2015) were excluded from the final analysis. Peptides were validated by Mascot Percolator with a 0.01 posterior error probability (PEP) threshold. For phosphopeptide analysis, the phosphorylation position was validated using the ptmRS algorithm. Peptide scores, precursor charge, peptide m/z, ptmRS scores, annotated mass labelled spectra for all phosphopeptides and other details are available via ProteomeXchange with identifier PXD005558. Peptide–spectrum matches, peptides per protein and protein coverage are indicated in supplemental Table S1. For phosphopeptides, we report the charge, m/z and mascot score for the highest confident PSM (that showing the highest Mascot score in our dataset) in supplemental Table S2. Peptide spectrum matches with total summed reporter intensities of less than $4e5$ were not considered for quantitation due to the high level of noise in the quantitation data. The quantification results of peptide spectrum matches were converted to peptide-level quantitation, which in turn was converted into protein quantitation using an R script (39). Only proteins with two or more quantified peptides were considered. Proteins with similar temporal abundance profiles were clustered using a fuzzy c-means algorithm with a Euclidean distance matrix (40).

Protein and phosphopeptide abundances of the reproductive stages (MII) were normalised against the vegetative stage (MI). The fold change of the MII stages (30- and 65- h) compared to the MI was estimated using the average TMT abundances from three or four biological replicates. Abundances were analysed for differential expression using the limma package in R (41). P- values for differential expression were adjusted for multiple comparisons using the R stats package. Phosphopeptide abundances were further normalized against the abundance of the non-phosphorylated proteins. Three thousand and eighty-three proteins exhibited significant expression changes (q-value less than 0.01) at least in one of the two MII stages compared to MI (supplemental

Table S1); seventy-eight phosphopeptides passed the abundance expression test (q-value < 0.01) at least in one of the MII stages analysed compared to MI (supplemental Table S2).

Analysis of protein functions - Proteins were classified into functional categories according to their annotated functions in the Gene Bank database, publications and by homology/functions according to the Gene Ontology (42), the Conserved Domain (<https://www.ncbi.nlm.nih.gov/Structure/cdd/cdd.shtml>) and the KEGG Pathway (http://www.genome.jp/kegg-bin/show_organism?org=sco) databases. When a protein was involved in the synthesis of a specific secondary metabolite, it was classified in the secondary metabolism group, even if it matched additional categories. When a protein was involved in cell division regulation, it was included in cell division, instead of the category of regulatory proteins.

FtsZ site-directed mutagenesis – Ser 319 (supplemental Table S2) and Ser 387 (16) were replaced by Glu (to mimic phosphate group) and by Ala (to mimic nonphosphorylation). The synthesis of *FtsZ* mutated alleles was ordered to GeneCust. They included the *FtsZ* ORF and an upstream region (380 nucleotides) large enough to contain the three promoter regions described by Flårdh et al. (43). Genes were cloned in the integrative plasmid pNG3 (44) and introduced into *S. coelicolor* by conjugation (30). The native *FtsZ* ORF was mutated using the CRISPR-Cas9 system developed by Tong et al. (45). The 20-nt target sequence was selected inside the *FtsZ* gene and amplified using primers sgRNA-F (CATGCCATGGCGATGACTTTGATGACTGCGGTTTTAGAGCTAGAAATAGC) and sgRNA-R (ACGCCTACGTAAAAAAGCACCGACTCGGTGCC). The product

of 110 bps was digested with *NcoI/SnaBI* and cloned in *NcoI/SnaBI* digested pCRISPR-Cas9. The *FtsZ* surrounding regions were amplified by PCR with 2082LeftF (AGGCCTA GACCGACCACCGCCGAG) / 2082LeftR (CCTATCACTTCAGGAAGTCCGTGATGACTGCGAGGTAGTTCTG) and 2082RightF (CAGAACTACCTCGCAGTCATCACGGACTTCCTGAAGTGATAGG) / 2082RightR (AGGCCTAGTAACCGACCACGGAACGCA) primers. The two DNA fragments were combined by overlap extension PCR (46) using primers 2082LeftR and 2082RightF. The PCR product was cloned and sequenced in pCR™-Blunt II-TOPO®. The insert was released with *EcoRI*, filled with the Klenow enzyme and cloned into pCRISPR-Cas9 (harbouring the 20-nt target sequence) digested with *StuI*. Mutations were confirmed by PCR using primers 2082LeftF and 2082RightR after plasmid clearance (45).

Antibiotic quantification - Undecylprodigiosin and actinorhodin were quantified according to the spectrophotometric assays described by Tsao et al. (47) and Bystrykh et al. (48). Cells were ruptured in the culture medium by adding 0.1 N KOH. After vortexing and centrifugation, actinorhodin was quantified in the supernatant (ϵ_{640} 25,320). Undecylprodigiosin was measured after vacuum drying of the mycelium, followed by extraction with methanol and acidification with HCl (to 0.5 M) (ϵ_{530} 100,500). Analyses were performed in three biological replicas.

Hypha and cell wall staining –Hypha and cell wall were stained using the LIVE/DEAD BacLight Bacterial Viability Kit (Invitrogen, L-13152) or Texas Red WGA (Invitrogen W21405) respectively as previously reported (49). Samples were observed under a Leica TCS SP8 laser scanning microscope as described before (3).

RESULTS

1. MI and MII show distinctive proteomes and phosphoproteomes.

A total of 3461 proteins (44.3% of *S. coelicolor* proteome) were identified and quantified (supplemental Table S1). Protein abundances were highly consistent across the biological replicates analysed (average correlation coefficient of 0.95; median coefficient of variation of 6.5%) (Fig. 2 and supplemental Table S1). Five clusters of proteins with similar temporal abundance profiles were identified (Fig. 3, supplemental Table S1): cluster 1 includes 345 proteins up-regulated at the MI stage; cluster 2 includes 241 proteins up-regulated at the MI and MII_{30h} stages; clusters 3-5 include 974 proteins up-regulated at the MII stages. Most proteins demonstrating significant changes (q-value less than 0.01) in the MII_{30h} and MII_{65h} stages (compared to MI) showed similar trends i.e., they were up- or down-regulated in both (see heat maps in Figs. 4A and 5A) (supplemental Table S1). In order to focus on the most outstanding changes, we focused on the function of the 1350 proteins with significant expression changes and at least 2-fold change (\log_2 ratio MII/MI lower than -1 or higher than 1). The abundances of key proteins are presented as histograms (Figs 4B and 5B), detailed below and outlined in Figure 6.

Eighty-five unique phosphorylation sites (48% Ser, 52% Thr) from 92 phosphopeptides and 48 phosphoproteins were identified (supplemental Table S2). All the phosphoproteins detected, were also identified and quantified in the proteomics set, which allowed us to normalise the change in phosphopeptide abundance against the change in protein, avoiding fluctuations due to variations in protein amount. To prevent artificial changes due to ratio suppression (underestimation of the relative change due to co-fragmentation of co-selected ions with similar m/z) (50) we focused on

phosphopeptides that passed the abundance expression test (q -value < 0.01) and showed at least two-fold increase or decrease in abundance (MII vs. MI) (58 phosphopeptides) (Figs. 7 and 8 and Table III). There is a clear and striking up-regulation of phosphorylation levels at MII, especially at 65 h (red bars in Fig. 7).

2. Proteins responsible for biosynthesis of secondary metabolites are upregulated at MII stages.

Proteins involved in secondary metabolite, such as actinorhodin, CPK, deoxysugar and coelichelin biosynthetic proteins, were up-regulated at both MII stages (clusters 3-5 in Fig. 3) (Figs. 4B and 6, Table II).

We did not detect any Ser/Thr/Tyr phosphorylation in proteins involved in secondary metabolite biosynthesis.

3. Proteins controlling aerial mycelium formation and sporulation showed different abundances at the MI and MII stages.

Proteins involved in the regulation of the aerial mycelium formation and sporulation (Bld, Whi, Wbl, Sap proteins) showed different abundances between MI and MII (Figs. 4B and 6, Table II). Most of them were up-regulated at the MII stages as SapA (51), WhiE ORFs VI and VIII (52), BldB (53), WhiH (51), SCO4301/SCO6476/SCO6638 (potential BldA targets; they harbour Leu encoded by TTA) (54), BldN (55) and WblA (56). Others were down-regulated at the MII stages as BldD/BldC/BldKA (57), WblE (58), SCO3897/SCO6717 potential BldA targets and SCO3424, an uncharacterized BldB homologue.

We did not detect any Ser/Thr/Tyr phosphorylation in proteins controlling aerial mycelium formation and sporulation.

4. *Transcriptional regulators, transducers, Ser/Thr/Tyr kinases, and signalling proteins showed different abundances and phosphorylation at the MI and MII stages.*

One-hundred-thirty-six regulatory proteins (transcriptional regulators, transducers, Ser/Thr/Tyr kinases and signalling proteins) had different abundances in at least one of the MII stages compared to MI (supplemental Table S1). Well-characterised regulatory proteins were included here (Figs. 4B and 6, Table II). However, the regulatory role of most of these proteins has been inferred from amino acid sequence similarity to previously characterised regulators.

Most of these regulatory proteins were up-regulated at the MII stages (most of them grouped into clusters 4 and 5) (Fig. 3): OhkA, a regulator of secondary metabolism and morphological differentiation (59); RarA-C “restoration of aerial mycelium formation” proteins (60); SarA, a protein involved in the regulation of sporulation and secondary metabolism (61); AtrA, an activator of the actinorhodin biosynthetic gene expressions (62); DevA, a GntR-Like transcriptional regulator controlling aerial mycelium development (63); NsdA, a BldD target regulating morphological differentiation and secondary metabolism (64); and ScbR, a γ -butyrolactone binding protein regulating morphological differentiation and secondary metabolism (65). By contrast, some regulatory proteins were down-regulated at the MII stages: CspB, a repressor of secondary metabolism (66) and NusG, a transcription antiterminator repressing undecylprodigiosin production (67).

As introduced above, the *Streptomyces coelicolor* genome encodes 47 Ser/Thr/Tyr kinases. In this work we identified 40 of these kinases. Most of them were up-regulated at the MII_{65h} stage (supplemental Table S2). The variation of eight Ser/Thr/Tyr kinases was over the 2-fold threshold (Fig. 4B): PkaE, an inhibitor of

actinorhodin production (68), PkaI, one of the proteins involved in spore wall synthesis (69) and six uncharacterised putative kinases (SCO1468, SCO2666, SCO3360, SCO3621, SCO3821, SCO4776).

Seven regulatory proteins were differentially phosphorylated during development, especially at the MII_{65h} stage (Table III) (Figs. 7A and 8): SCO0204, an orphan response regulator whose expression is affected by the sporulation-specific cell division activator SsgA (70); DevA, a GntR-like transcriptional regulator required for aerial mycelium development (63); DasR, a pleiotropic regulator of differentiation (71); SCO5357, rho transcription terminator involved in RNA degradation (KEGG pathway sco03018); SCO5544, part of the *cvn1* conservon, a set of genes involved in regulating antibiotic production and aerial mycelium formation (72); SCO5704, a putative NusA transcription elongation factor; and SCO7463, a putative histidine kinase.

5. Cell division proteins are differentially expressed and phosphorylated in the compartmentalised MI and the multinucleated MII hyphae.

Forty-seven cell division proteins showed significant differences in their abundances between the MI and MII (supplemental table S1). Cell division proteins do not present large differences in abundance during *Streptomyces* differentiation (3). However, they are crucial in differentiation, controlling the compartmentalisation of the MI hyphae (3) and sporulation (73). Hence and only for cell division proteins, we discuss here all the well-characterised proteins showing significant variation, including those that did not pass the 2-fold threshold defined above (Figs. 4 and 6, Table II). Some cell division proteins that were reported to be involved in sporulation were up-regulated at the MII stages (clusters 4 and 5 in Fig. 3): SffA (74), ParJ (75), CrgA (76), FemX (77), SCO4114 (78) and SmC (SCO5577) (79). Other well characterised cell

division proteins were down-regulated at the MII stages: DivIVA (80), FtsZ (81), MreC (69), ParA, ParB (82), the SCO4439 DD-CPase (83) and the nucleoid-associated HupS protein (84).

Four proteins involved in cell division and cell wall synthesis were differentially phosphorylated during development (Table III) (Figs. 7A and 8). Three cell division proteins were phosphorylated at the MII_{65h} stage: DivIVA, one of the proteins regulating apical growth in *Streptomyces*, which was demonstrated to be regulated by phosphorylation (80); SepF one of the proteins included in the *Streptomyces* division and cell wall (dcw) cluster (85) and FtsZ, the key tubulin-like bacterial division protein (81). SCO4439, a D-alanyl-D-alanine carboxypeptidase involved in sporulation (83), was phosphorylated at the MII_{30h} and MII_{65h} stages.

6. Proteins involved in primary metabolism are differentially expressed and phosphorylated during development.

Most proteins involved in protein synthesis (translation/protein folding) were up-regulated at the MI (cluster 1 in Fig. 3) (Figs. 5 and 6, Table II). Proteins involved in energy production (krebs cycle, oxidative phosphorylation), lipid metabolism and catabolism were mostly up-regulated at the MII stages (Figs. 5 and 6, Table II). Other proteins involved in primary metabolic pathways were not clearly overrepresented in the MI or MII developmental stages (supplemental Table S1).

Nine proteins involved in primary metabolism (translation, glycolytic enzymes, catabolism, nucleotide metabolism, folate biosynthesis) were differentially phosphorylated during development. Most of them were phosphorylated at the MII_{65h} stage (Table III) (Figs. 7 and 8).

7. Stress proteins and chaperones are regulated by phosphorylation.

Stress proteins and chaperones were not clearly overrepresented in the MI or MII developmental stages (supplemental Table S1). However, some of them were differentially phosphorylated during development, mostly at the MII_{65h} (Figs. 7B and 8, Table III): PspA, a phage shock protein (86); chaperonin GroES; SCO5419, a putative thioredoxin; and SCO1836, a putative stress-like protein.

8. Proteins and phosphoproteins of unknown function showed different abundances at the MI and MII stages.

Five-hundred-forty-two uncharacterized proteins without any clear homology showed different abundances in at least one of the MII stages analysed compared to MI. As discussed below, these proteins constitute an outstanding database of potential regulators and effectors of differentiation (supplemental Table S1). For example, SCO5191 was up-regulated 11- and 8-fold at the MI_{16h} stage compared to the MII_{30h} and MII_{65h} respectively, whereas SCO6650 was up-regulated 28-fold at the MII_{65h} (Table II).

Eight proteins with unknown functions were differentially phosphorylated during development. Interestingly, SCO2668 and SCO3859 were multiphosphorylated at eight and nine positions respectively (Fig. 7C) (supplemental Table S2) (Fig. 8) (Table III). SCO2668 has a “von willebrand factor type A” domain (vWA, conserved domain database accession cd00198) which is involved in a wide variety of important cellular functions (basal membrane formation, cell migration, cell differentiation, adhesion, homeostasis); SCO3859 has a “helix-turn-helix” domain (HTH, conserved domain database accession cl22854) which is present in several transcriptional regulators and other DNA binding proteins. Interestingly, phosphorylation in both

proteins do not look to be random, as it is localised into small regions (58 and 13 amino acids respectively), outside the conserved domains (Fig. 7C).

9. Phosphorylation of FtsZ affects sporulation and secondary metabolism.

As mentioned above, we identified a novel FtsZ phosphorylation at Ser 319 (supplemental Table S2), which is significantly up-regulated at the MII_{30h} and MII_{65h}, 1.25- and 4.1-fold respectively. In our previous label-free study, we identified another FtsZ phosphorylation at Ser 387 (16). In order to study the effect of these phosphorylations, we replaced the native *FtsZ* gene with two mutated alleles, one mimicking phosphorylation (replacing phosphoserines by Glu) and another one mimicking non-phosphorylation (replacing phosphoserines by Ala), a well-established methodology to analyse the effect of Ser phosphorylation (see some examples in (87-89)).

FtsZ is crucial for cell division and sporulation (90) and sporulation is affected in the FtsZ mutants. It is highly reduced in the strain harbouring the “Glu-Glu” allele and delayed in the strain harbouring the “Ala-Ala” allele (Fig. 9A). However, we did not observe differences in hypha septation between the FtsZ mutants and the wild-type strain during the vegetative stage (Fig. 9B). The Ala-Ala mutant shows a delayed growth at the early development compared to the Glu-Glu mutant and the wild-type strains (Fig. 9C). The most surprising phenotype, was the effect in secondary metabolism. Actinorhodin, purple pigmented polyketide antibiotic, was overproduced (2.9-fold) in the Ala-Ala mutant and repressed (0.53-fold the wild-type value) in the “Glu-Glu” mutant (Fig. 9C), in all the culture media (SFM, MM, GYM and R5A) and conditions (solid and liquid cultures) tested (Fig. 9C,D). The production of the tripyrrole antibiotic undecylprodigiosin, was slightly diminished (0.77-fold the wild-type value) in

the Ala-Ala mutant and highly repressed (0.24-fold the wild-type value) in the Glu-Glu mutant (Fig. 9C).

Next we compared the proteomes of the FtsZ mutants and the wild-type strain (supplemental Table S3). Most of the proteins involved in secondary metabolism that showed significant variations, were up-regulated in the Ala-Ala mutant and down-regulated in the Glu-Glu mutant (Fig. 10A). When we focus on the proteins that passed the 2-fold threshold (Fig. 10B), we observe that all the actinorhodin biosynthetic proteins are up-regulated in the Ala-Ala mutant and not in the Glu-Glu mutant, while the undecylprodigiosin biosynthetic proteins are down-regulated in both mutants compared to the wild-type strain (Ser-Ser allele). These results corroborate that the actinorhodin overproduction observed in the Ala-Ala mutant and the undecylprodigiosin repression detected in the Ala-Ala and the Glu-Glu mutants (Fig. 9C), are regulated at the protein level.

DISCUSSION

Here we quantified the variation of 3461 proteins during the MI and MII stages in *Streptomyces coelicolor*. This corresponds to 44.3% of *S. coelicolor* theoretical proteome and 45% of the transcriptome identified by us in a previous work at the same developmental stages (91). Our results exceed any previous quantitative studies of the *S. coelicolor* proteome (92-96), quintuplicating the number of proteins quantified in our previous MI and MII quantitative proteomic study (16). We also quantified the variation of 92 phosphopeptides from 48 phosphoproteins during development, using the proteome abundances to normalise the phosphoproteome abundances and reporting the first quantitative dataset of Ser/Thr/Tyr phosphorylation variations during *Streptomyces* differentiation. We used MS2 TMT to profile proteome and phosphoproteome temporal

changes, which is a method known to suffer from ratio suppression (97). However, we and others have studied the issue of ratio compression in TMT based quantitative proteomics and have concluded that the effect on the overall outcome of proteomics studies can be neglected, unless high precision is needed (98-100). In order to reduce the interference of ratio suppression, we focused on the most reliable quantitative data, i.e. proteins and phosphopeptides that changed in abundance by at least two-fold. Despite this very stringent criterion, we found that 1350 proteins (supplemental Table S1) and 58 phosphopeptides (supplemental Table S2) were changing, indicating significant reorganisation of the *Streptomyces coelicolor* proteome and phosphoproteome throughout its life cycle. As described above, most of the 40 Ser/Thr/Tyr kinases identified, as well as Ser/Thr/Tyr phosphorylations were overrepresented at the sporulating stage (MII_{65h}), indicating a role of Ser/Thr/Tyr phosphorylation in the regulation of differentiation and sporulation.

The clearest physiological difference between the MI and MII is the activation of secondary metabolism at the MII (91, 95) and most proteins involved in secondary metabolism were highly up-regulated at the MII stages (CPK, ACT, deoxysugar synthases, coelichelin). By contrast, a majority of proteins involved in primary metabolism, particularly those involved in protein synthesis, translation and protein folding were highly up-regulated in the MI stage, which is not strange, considering that the MI corresponds to the exponential vegetative growth stage, while growth is arrested at the MII stage (101). Other primary metabolism proteins involved in energy production (krebs cycle, oxidative phosphorylation), lipid metabolism, amino acid metabolism and pentose phosphate pathway, were mostly up-regulated in MII stages, perhaps contributing the energy and precursors necessary for secondary metabolism. Secondary metabolism is associated to MII (4), but the expression of different

secondary metabolites need specific activators (elicitors) or repressors and not all secondary metabolites are produced simultaneously in the MII (5, 102). For instance, AtrA (SCO4118), an activator of actinorhodin biosynthetic genes (62) is slightly up-regulated at the MII_{65h} (2.1-fold), activating actinorhodin production, while PkaE, a repressor of actinorhodin production was also up-regulated at the MII_{65h} (2.8-fold), indicating that actinorhodin production is blocked in sporulating hyphae as reported before (91, 95). SCO6286, a CPK repressor (103), was up-regulated at the MII_{65h}, blocking CPK production during the aerial mycelium stage. SCO3225 and SCO3226, a two-component system repressing CDA biosynthesis showed similar abundances at the MI and MII stages (supplemental Table S1), repressing CDA production under the culture conditions used in this work.

The clearest morphological difference between MII_{30h} (substrate mycelium) and MII_{65h} (aerial sporulating mycelium), is the presence of hydrophobic coats in the aerial sporulating hyphae and proteins involved in the regulation of hydrophobic cover formation and sporulation (Bld, Whi, SapA, RarA-C) were up-regulated in the MII_{65h}.

None of the enzymes synthesising secondary metabolites, or the proteins forming the hydrophobic coats were detected as phosphorylated, indicating that modulation of these processes by Ser/Thr/Tyr phosphorylation is not at the final effectors. By contrast, some proteins and enzymes involved in central metabolic pathways (translation, glycolytic enzymes, catabolism, nucleotide metabolism, folate biosynthesis) were differentially phosphorylated in the MI and MII, suggesting that their activities are modulated by phosphorylation.

There are two developmental stages in which *Streptomyces* hyphae are compartmentalised, the MI and the sporulating MII, while the early stages of MII (substrate and aerial hyphae) consist in multinucleated hyphae with sporadic septa.

Streptomyces cell division studies focused on sporulation, while MI cell division remains poorly known. Recently, our group, was able to identify the existence of a new kind of cell division based on cross-membranes without detectable peptidoglycan in the MI hyphae (3, 104), whose regulation remains basically uncharacterised. Here, we detected several cell division proteins more abundant at the MI stage than at the sporulating MII stage. This result suggest a putative role of these proteins in the MI cross-membrane based cell division, as it happens with FtsZ, the key effector of bacterial cell division (3). These proteins constitute a good starting point to study the mechanisms controlling cross-membrane cell division in *Streptomyces*.

Cell division proteins such as DivIVA and FtsZ were more phosphorylated during the MII_{65h} stage, suggesting a role of phosphorylation in the regulation of cell-division accompanying sporulation. FtsZ phosphorylation has a direct effect in sporulation, that is highly reduced in the mutant mimicking double phosphorylation (“Glu-Glu” allele). This might indicate that phosphorylation affects FtsZ polymerization, reducing sporulation, as it happens in *Mycobacterium tuberculosis* (105). Interestingly, we detected a dramatic effect of FtsZ phosphorylation in actinorhodin production. *FtsZ* is highly conserved in the *Streptomyces* genus and the “Ala-Ala” allele, might be used in industrial streptomycetes to enhance the production of other antibiotics. Further work will be necessary to fully understand the biological role of FtsZ phosphorylation and its potential industrial application.

One-hundred-thirty-six regulatory proteins (transcriptional regulators, transducers, Ser/Thr/Tyr kinases and signalling proteins), most of them uncharacterised putative regulators identified by amino acid sequence homology, and 542 putative proteins (without any clear homology) showed differential abundances during MI and MII. Key regulatory proteins were differentially phosphorylated during *S. coelicolor*

development, such as DasR or DevA, two pleiotropic regulators controlling secondary metabolism and aerial mycelium formation. These results suggest a putative role of phosphorylation in the regulation of secondary metabolism, aerial mycelium formation and sporulation to be studied and characterised in the future. Eight proteins without known function were also differentially phosphorylated during development. For instance, SCO2668 is highly multiphosphorylated at the sporulating mycelium (MII_{65h}), and harbours a pleiotropic eukaryotic vWA domain which is involved in a wide variety of important cellular functions in eukaryotic cells (basal membrane formation, cell migration, cell differentiation, adhesion, homeostasis). This suggests a complex regulatory role of SCO2668 in sporulation, resembling eukaryotic regulatory pathways. *S. coelicolor* is the best characterised streptomycete, despite that it does not produce commercial secondary metabolites. However, many of the phosphorylated proteins, as well as regulatory proteins and proteins with unknown function differentially expressed during the vegetative (MI) and secondary metabolite producing mycelia (MII), are conserved in the *Streptomyces* genus. These proteins are potential modulators of secondary metabolism and differentiation to be studied and characterised in the future, as it happens with FtsZ. This knowledge can contribute to the optimisation of secondary metabolite production in industrial streptomycetes, including the activation of cryptic secondary metabolite pathways (pathways that are not activated in laboratory cultures) during the screening for new secondary metabolites from streptomycetes (106).

Acknowledgements - We thank the European Research Council (ERC Starting Grant; Strp-differentiation 280304) and the Spanish “Ministerio de Economía y Competitividad” (MINECO; BIO2015-65709-R) for financial support. The mass

spectrometry and proteomics infrastructure used in this study was provided by VILLUM Center for Bioanalytical Sciences at University of Southern Denmark supported by a generous grant from the VILLUM Foundation (O.N.J). P.S. was supported by a postdoctoral fellowship from the Lundbeck Foundation, Denmark. Nathaly Gonzalez-Quñonez was funded by a Severo Ochoa fellowship (FICYT, Consejería de Educación y Ciencia, Spain). Thanks to Beatriz Gutierrez Magan (Universidad de Oviedo, Dpto. Biología Funcional, Área de Microbiología) for her laboratory assistance.

Data availability - Raw data are available via ProteomeXchange with identifier PXD005558.

TABLES

Table I: Comparison of the *S. coelicolor* phosphoproteome with other published prokaryotic and human phosphoproteomes. Quantitative phosphoproteomic studies are indicated by an asterisk. N.r. Not reported.

	Bacterium	Protein (mg)¹	Phosphoproteins	Phosphopeptides	Phosphorylation sites	Reference
Gram +	<i>S. coelicolor</i>	0.1	48	92	85	This work
	<i>S. coelicolor</i>	0.3	127	260	289	(16)*
	<i>S. coelicolor</i>	50	40	44	46	(18)
	<i>S. erythraea</i>	10	88	109	n.r.	(28)*
	<i>B. subtilis</i>	10	78	103	78	(15)
	<i>B. subtilis</i>	12	139	177	144	(31)*
	<i>C. acetobutylicum</i>	2	61	82	107	(9)
	<i>L. lactis</i>	20	63	102	79	(22)
	<i>M. tuberculosis</i>	n.r.	301	380	500	(19)
	<i>S. pneumoniae</i>	1	84	102	163	(23)
	<i>L. monocytogenes</i>	10	112	155	143	(17)
	<i>L. monocytogenes</i>	8	191	256	242	(27)*
	<i>S. aureus</i>	50	108	n.r.	76	(10)
Gram -	<i>E. coli</i>	20	79	105	81	(14)
	<i>K. pneumoniae</i>	30	81	117	93	(13)
	<i>P. aeruginosa</i>	1.2	39	57	61	(20)
	<i>P. putida</i>	1.2	59	56	55	(20)
	<i>H. pylori</i>	n.r.	67	82	126	(11)
	<i>R. palustris (Ch)</i>	2	54	100	63	(12)
	<i>R. palustris (Ph)</i>	2	42	74	59	(12)
	<i>T. thermophilus</i>	100	48	52	46	(24)
	<i>E. coli</i>	9.8	133	n.r.	108	(25)*
	<i>E. coli</i>	10	n.r.	34	n.r.	(29)*
	<i>A. baumannii</i> Abh12O-A2	9	70	n.r.	80	(21)
	<i>A. baumannii</i> ATCC 17879	9	41	n.r.	48	(21)
Archaea	<i>H. salinarum</i>	20	26	42	31	(8)
Eukarya	<i>H. sapiens</i>	6	7832	>50000	38229	(108)

¹ The total amount of protein used for all the MS/MS or 2D gel experiments is indicated.

Table II: Averaged TMT protein abundances (from at least three biological replicates) of the substrate mycelium (MII_{30h}) and sporulating aerial mycelium (MII_{65h}) compared to the MI stage (16 h). The MII/MI ratio is shown in logarithm (log 2) and linear forms. Cl, clusters of proteins with similar abundance profiles. N.S. Not significant (q-value higher than 0.01). (-) The significant value that did not pass the 2-fold threshold. The proteins discussed in the text are indicated. Only changes of more than 2-fold up or down-regulated are shown, except for cell division proteins.

Category	SCO n°	Cl	Function	Log 2 (MII _{30h} /MI)	Log 2 (MII _{65h} /MI)	Fold-change (MII _{30h} /MI)	Fold-change (MII _{65h} /MI)
Secondary metabolism	SCO6286	4	CPK repressor	1	1.8	1.9	3.5
	SCO6276	3	CPK biosynthesis	2	n.s.	4	n.s.
	SCO6277	3		1.9	-	3.8	-
	SCO6278	3		1.4	n.s.	2.7	n.s.
	SCO6279	n.s.		2.6	1.6	6	3
	SCO6282	3		2.7	-	6.5	-
	SCO6283	3		2.6	-	5,9	-
	SCO0489			Coelichelin biosynthesis	1	-1.5	2
	SCO0492	3	1.1		n.s.	2.1	n.s.
	SCO0498	3	1.6		-	3	-
	SCO0499	3	1.2		n.s.	2.3	n.s.
	SCO5072	5	ACT biosynthesis	n.s.	3.6	n.s.	12.1
	SCO5073	5		n.s.	2.7	n.s.	6.5
	SCO5074	5		n.s.	4.6	n.s.	24.2
	SCO5075	5		n.s.	2.2	n.s.	4.6
	SCO5077	n.s.		-	2.1	-	4.3
	SCO5078	5		n.s.	3	n.s.	8
	SCO5079	5		-	3.4	-	10.5
	SCO5080	5		n.s.	3	n.s.	8
	SCO5083	5		n.s.	1.9	n.s.	3.7
	SCO5084	5		n.s.	3	n.s.	8
	SCO5086	n.s.		n.s.	1.6	n.s.	3
	SCO5087	5		n.s.	1.9	n.s.	3.7
	SCO5088	5		n.s.	1.6	n.s.	3
	SCO5089	n.s.		-	2.1	-	4.3
	SCO5090	5	n.s.	2.4	n.s.	5.3	
	SCO5091	4	-	1.6	-	3	
	SCO0381	5	Deoxysugar synthases	n.s.	2.5	n.s.	5.8
	SCO0382	n.s.		-	2.5	-	5.8
	SCO0383	5		-	2.7	-	6.3

	SCO0384	5		n.s.	2.4	n.s.	5.4
	SCO0385	5		n.s.	2.1	n.s.	4.3
	SCO0386	5		-	2.6	-	5.9
	SCO0387	n.s.		-	2.3	-	5
	SCO0388	5		-	2.8	-	7
	SCO0389	5		-	2.8	-	7
	SCO0391	5		n.s.	2.1	n.s.	4.2
	SCO0392	4		-	1.9	-	3.7
	SCO0393	n.s.		1	3.1	2	8.2
	SCO0394	5		-	3.3	-	9.6
	SCO0395	n.s.		1	3.1	2	8.7
	SCO0396	5		-	3.3	-	9.9
	SCO0398	5		-	2.7	-	6.4
	SCO0399	4		1.2	2.8	2.3	7
	SCO0400	n.s.		-	2.6	-	6.7
	SCO0401	n.s.		-	2	-	4
	SCO0409	5	SapA	-	1.7	-	3.2
	SCO1489	n.s.	BldD	-	-1.5	-	0.3
	SCO3323	n.s.	BldN	1.5	1.9	2.8	3.7
	SCO3424	1	BldB homologue	-1.8	-1.8	0.3	0.3
	SCO3579	n.s.	WblA	1.1	1.6	2.1	3
	SCO4091	n.s.	BldC	-	-1.8	-	0.3
	SCO5112	n.s.	BldKA	-1.1	n.s.	0.5	n.s.
	SCO5240	n.s.	WblE	-	-2.6	-	0.2
	SCO5315	5	WhiE ORFVI	n.s.	1.6	n.s.	3
	SCO5321	5	WhiE ORFVIII	n.s.	1.8	n.s.	3.5
	SCO5723	n.s.	BldB	-	1.5	-	2.8
	SCO5819	n.s.	WhiH	-	1.9	-	3.7
	SCO3897	1	Possible targets for bldA regulation	-1.4	-2.7	0.4	0.2
	SCO4301	4		-	1.5	-	2.8
	SCO6476	n.s.		-	1.4	-	2.6
	SCO6638	n.s.		1	1.3	2	2.5
	SCO6717	n.s.		-	-1.4	-	0.4
	SCO1416	5	SffA	n.s.	0.8	n.s.	1.7
	SCO1662	n.s.	ParJ	0.33	0.55	1.3	1.5
	SCO2077	n.s.	DivIVA	n.s.	-0.9	n.s.	0.5
	SCO2082	2	FtsZ	-0.2	-0.8	0.9	0.57
	SCO2610	2	MreC	n.s.	-0.7	n.s.	0.6
	SCO3854	5	CrgA	n.s.	1.4	n.s.	2.6
	SCO3886	n.s.	ParA	-0.3	-0.6	0.8	0.7
	SCO3887	2	ParB	n.s.	-0.3	n.s.	0.8
	SCO3904	4	FemX	n.s.	0.57	n.s.	1.48
	SCO4114	n.s.	Sporulation associated protein	0.4	0.2	1.3	1.1
	SCO4439	n.s.	DD-CPase	-0.4	-0.7	0.8	0.6
	SCO5556	1	HupS	-2	-1.7	0.25	0.3
	SCO5577	n.s.	SmC	n.s.	0.6	n.s.	1.5
Regulatory	SCO1468	5	Putative	n.s.	1.9	n.s.	3.8

			Serine/Threonine protein kinase				
	SCO1596	n.s.	OhkA	-	1.5	-	2.8
	SCO1628	4	RarC	1.6	3.1	3	8.6
	SCO1629	n.s.	RarB	-	2.6	-	6.1
	SCO1630	5	RarA	-	3	-	8
	SCO2666	n.s.	Putative Serine/Threonine protein kinase	n.s.	1.3	n.s.	2.5
	SCO3102	5	PkaE	n.s.	1.5	n.s.	2.8
	SCO3360	n.s.	Putative Serine/Threonine protein kinase	-1.1	n.s.	0.4	n.s.
	SCO3621	5	Putative Serine/Threonine protein kinase	n.s.	1.1	n.s.	2.1
	SCO3821	n.s.	Putative Serine/Threonine protein kinase	-1.2	-0.9	0.4	0.5
	SCO4776	n.s.	Putative Serine/Threonine protein kinase	0.4	1.5	1.3	2.9
	SCO4778	n.s.	PkaI	0.9	1.5	1.9	2.8
	SCO4069	n.s.	SarA	-	1.5	-	2.8
	SCO4118	4	AtrA	-	1.1	-	2.1
	SCO4190	5	DevA	n.s.	1.3	n.s.	2.5
	SCO4325	n.s.	CspB	-	-2.2	-	0.2
	SCO4647	n.s.	NusG	-	-1.9	-	0.3
	SCO5582	4	NsdA	-	1.1	-	2.1
	SCO6265	n.s.	ScbR	1.6	1.5	3	2.8
Unknown	SCO5191	1	Hypothetical protein	-3.4	-3	0.09	0.12
	SCO6650	5	Hypothetical protein	n.s.	4.8	n.s.	27.8
Translation/ protein folding	SCO1491	n.s.	Elongation factor	-	-2.3	-	0.2
	SCO1804	5	tRNA ribosyltransferase -isomerase	n.s.	2.9	n.s.	7.4
	SCO2620	2	Trigger factor chaperone	-	-1.8	-	0.8
	SCO4648	1	Ribosomal protein	-1.3	-2.2	0.4	0.2
	SCO4725	2	Translation initiation	-	-1.9	-	0.3
	SCO4734	1	Ribosomal protein	-1.2	-2.2	0.4	0.2
Energy production	SCO0922	3	Succinate deshydrogenase	2.1	n.s.	4.2	n.s.
	SCO0923	3	Succinate	2.5	n.s.	5.5	n.s.

			deshydrogenase				
	SCO4563	5	NADH dehydrogenase	n.s.	2.4	n.s.	5.1
	SCO4564	5	NADH dehydrogenase	n.s.	3.2	n.s.	9.4
	SCO4566	5	NADH dehydrogenase	n.s.	2.6	n.s.	6.1
	SCO5107	5	Succinate deshydrogenase	n.s.	2.6	n.s.	6.2
Catabolism	SCO1640	4	Proteasome protein	1.2	2.8	2.6	7
	SCO1643	4	Proteasome protein	-	1.9	-	3.7
	SCO1646	n.s.	Proteasome protein	-	2.2	-	4.7
	SCO5443	5	Alpha-amylase	n.s.	2.2	n.s.	4.7
	SCO6411	5	Hydrolase	-	2	-	4
	SCO6414	4	Hydrolase	1.2	2.3	2.3	4.9

¹ For cell division proteins, variations inside the ± 1 interval showing significant variations (q-value <0.01), are shown.

Table III: Averaged TMT phosphopeptide abundances (from at least three biological replicates) of the substrate mycelium (MII_{30h}) and sporulating aerial mycelium (MII_{65h}) compared to the MI stage (16 h), of the phosphopeptides and phosphoproteins discussed in the manuscript. The MII/MI ratio is shown in the logarithm (log 2) and linear forms. N.S. Not significant (q-value higher than 0.01). (-) significant abundance values that did not pass the 2-fold threshold. Phosphorylation sites are highlighted in bold.

Category	PROTEIN	Phosphopeptide	Log 2 (MII _{30h} /MI)	Log 2 (MII _{65h} /MI)	Fold-change (MII _{30h} /MI)	Fold-change (MII _{65h} /MI)
Regulatory	SCO0204	AH p SEQGDNTGSPVR	n.s.	3.1	n.s.	8.5
	DevA (SCO4190)	ALQEDGLLTNV p SK	n.s.	2.1	n.s.	4.2
	DasR (SCO5231)	STDVSSAENEGG A pTVR	n.s.	2.1	n.s.	4.3
	SCO5357	VNASAEQAAPADD A pSER	n.s.	1.6	n.s.	3
	SCO5544	GHDEPD p SSRTDR	n.s.	3.2	n.s.	9.2
	SCO5544	GHDEPD S pSRTDRTPR	n.s.	1.9	n.s.	3.7
	SCO5544	GHDEPD p SpSRTDRTPR	n.s.	3.6	n.s.	12
	SCO5704	IDIRPDTEQPSDASPEQ S pSGGRGE	n.s.	4.8	n.s.	27.8
SCO7463	TSDTPGASSEGHEV p S	n.s.	2.8	n.s.	6.8	
Cell division	DivIVA (SCO2077)	QLETQADD p SLAPPR	n.s.	1.6	n.s.	3.1
	SepF (SCO2079)	IAEGGFFN Q pS	n.s.	1.1	n.s.	2.1
	SCO2082 (FtsZ)	VTVIAAGFDGGQ P pSK	n.s.	2	n.s.	4
	SCO4439	DGD ADGYDGGPPVD Q pTAVFK	1.3	1.9	2.4	3.7
Chaperones, thioredoxin and stress	SCO1836 ¹	SGQDTIET p SG p TAK	n.s.	-1	n.s.	0.49
	PspA (SCO2168 ¹)	QAIEGGQGQGE A p S pS S QSQPQDTPR	n.s.	1.72	n.s.	3.3
	GroES (SCO4761)	IVVQPLDAEQ p TTASGLVIPDTAK	-	2.2	-	4.5
	GroES (SCO4761)	IVVQPLDAEQ T A p SGLVIPDTAK	n.s.	2.5	n.s.	5.5
	GroES (SCO4761)	IVVQPLDAEQ T pTASGLVIPDTAK	n.s.	2.6	n.s.	5.9
	SCO5419	NVLAEEP G N p TEAK	n.s.	1.6	n.s.	3.1
Protein synthesis (translation)	RspA (SCO1998)	AAAEGVDTAGAAPAASGGGGGGSY SEGGDN p SGALASDEALALR	n.s.	2.1	n.s.	4.3
	RspA (SCO1998)	AAAEGVD p TAGAAPAASGGGGGGSY SSEGGDNSGALASDEALALR	n.s.	2.4	n.s.	5.4

	SCO4703	AVD p TEGSEA	n.s.	3.3	n.s.	9.6
	SCO4711	SE p SNVTEETK	n.s.	3.3	n.s.	10.2
Catabolism	SCO1646 ¹	ApTRpSp TEEVVEEQAQDAQASEDLK	n.s.	1.1	n.s.	2.2
	SCO1646	p STEEVVEEQAQDAQASEDLK	n.s.	1.2	n.s.	2.3
	SCO1646	ApTR STEEVVEEQAQDAQASEDLK	n.s.	1.6	n.s.	3
	SCO1646 ¹	ApTRp STEEVVEEQAQDAQASEDLK	n.s.	3.1	n.s.	8.5
	SCO5864	KTDDDVDS p SLEELK	n.s.	3.3	n.s.	9.9
	SCO5864	TDDDV p SDSLEELK	n.s.	3	n.s.	8
	SCO5864	KTDDDV p SDSLEELK	n.s.	4.4	n.s.	21.1
Nucleotide metabolism	PurE (SCO3059)	EFQQDLNDQ Ap TEK	n.s.	3.2	n.s.	9
	SCO3542	ELPQIDPDQAPP p SR	n.s.	2	n.s.	4
	SCO3542	ELPQIDPDQAPP p SRR	n.s.	2.2	n.s.	4.5
Folate biosynthesis	SCO3401	SAPFAQGPSDPTVQPVPA S VIEQVDA AD Tp TL S NP K	n.s.	-1.5	n.s.	0.3
Glycolysis	SCO7443	TSGLADGVVV Tp SHNPPADGGFK	n.s.	1.4	n.s.	2.7
Unknown	SCO2668	AGDIDG Ap TAK	n.s.	1.1	n.s.	2.1
	SCO2668	VEEAD E MTLE TRpSp TK p TVR	n.s.	1.9	n.s.	3.8
	SCO2668	AVQLAGVSGNAD p TAK	1.62	1.3	3.1	2.5
	SCO3859	SQGGSVLSN T Tp T T Tp Sp SSGAP T V K	n.s.	3	n.s.	8.2
	SCO3859	SQGGSVLSN T T T T Tp T T Sp SSGAP T V K	n.s.	3.4	n.s.	10.4
	SCO3859	SQGGSVLSN T T T T Tp T T Sp SSGAP T V K	n.s.	3.3	n.s.	9.6
	SCO3859	SQGGSVLSN T T T T T Tp T Sp SSGAP T V K	n.s.	3.4	n.s.	10.4
	SCO3859	SQGGSVLSN p T T T T T Sp SSGAP T V K	n.s.	3.2	n.s.	9.3
	SCO3859	SQGG p SVLSN T T T T T Sp SSGAP T V K	n.s.	3.5	n.s.	11.2
	SCO3859	SQGGSVLSN T T T T T Tp Sp SSGAP T V K	n.s.	3.9	n.s.	14.8
	SCO3859	SQGGSVLSN T T T T T Tp T Sp SSGAP T V K	n.s.	5.3	n.s.	40.2
	SCO3859	SQGGSVLSN Tp T T T T Sp SSGAP T V K	n.s.	5.9	n.s.	61.1
	SCO3859	SQGGSVLSN T T T T T Tp T Sp SSGAP T V K	n.s.	3.4	n.s.	10.9
	SCO3859	SQGGSVLSN T T T Tp T T Sp SSGAP T V K	n.s.	3	n.s.	8
	SCO3859	SQGGSVLSN p Tp T T T Sp SSGAP T V K	n.s.	4.1	n.s.	17

¹ Only one phosphorylation site, at one of the positions marked, exists. See supplemental Table S2.

FIGURES

FIG. 1. **Sample preparation.** *S. coelicolor* hyphae stained with SYTO 9 and propidium iodide observed under the confocal microscope at the compartmentalised MI (16 h), multinucleated substrate mycelium (MII_{30h}) and sporulating aerial mycelium (MII_{65h}) stages are shown (upper panels). TMT labelling and the phosphopeptide enrichment workflows are outlined.

FIG. 2. **TMT abundance values distribution and correlation between biological replicates.** The distribution of the TMT abundances of two biological replicates of each developmental stage analysed (MI_{16h}, MII_{30h} and MII_{65h}) (panels in the diagonal), the correlation and coefficients of regressions of TMT abundances between biological replicates of the same developmental stage (marked in black) and the correlation and coefficients of regressions of TMT abundances between different developmental stages (marked in red), are shown. Notice the good correlation between replicates of the same developmental stage and the poor correlation between replicates of different developmental stages.

FIG. 3. **Clusters of the proteins with similar temporal abundance profiles.** Cluster 1, proteins up-regulated at the MI_{16h} stage; Cluster 2, proteins up-regulated at MI_{16h} and MII_{30h}; Clusters 3-5, proteins up-regulated at the MII stages.

FIG. 4. **Abundances (logarithm of TMT MII/MI) of proteins related with secondary metabolism, aerial mycelium formation, sporulation, cell division and regulation of cellular processes (transcriptional regulators, transducers Ser/Thr/Tyr kinases and signalling proteins).** Proteins involved in the synthesis of

specific secondary metabolites, were classified in the secondary metabolism group, even if they fitted additional categories; proteins involved in cell division regulation, were included in cell division (see Methods). *A*, Heat maps outlining the TMT abundances of proteins with significant variations in both MII stages with respect to MI. Asterisks indicate the few proteins showing opposite differences in both MII (i.e. up-regulated at MII_{30h} and down-regulated at MII_{65h} or vice versa). *B*, TMT abundances of the key proteins described in the text. Abundances have significant differences (q-value < 0.01) and passed the 2-fold threshold in at least one of the MII stages, except for cell division proteins, in which proteins below the 2-fold threshold, but showing significant differences, are also shown. The SCO numbers are indicated. Dashed lines indicate the 2-fold threshold. “K” indicates Ser/Thr/Tyr kinases.

FIG. 5. Abundances (logarithm of TMT MII/MI) of proteins related with central (primary) metabolism (aerobic energy production, catabolism, translation/protein folding). *A*, heat maps. *B*, TMT abundances of the key proteins described in the text. Statistics and labelling as in Fig. 4.

FIG. 6. Integrated *S. coelicolor* proteome variations during development (MI_{16h}, MII_{30h} and MII_{65h}). The key proteins discussed in the text are shown at the developmental stage in which they display significant up-regulation (q-value < 0.01).

FIG. 7. *S. coelicolor* phosphoproteome. *A-B*, Thirty-five peptides differentially phosphorylated during development (q-value < 0.01) that passed the 2-fold threshold at least in one of the MII stages grouped into functional categories. The SCO numbers are indicated. Dashed lines indicate the 2-fold threshold. The 23 peptides differentially

phosphorylated, belonging to proteins with unknown function, are not shown. *C*, Outline indicating the phosphorylation sites and conserved protein domains of two multiphosphorylated proteins with unknown function.

FIG. 8. *S. coelicolor* integrated phosphoproteome variations during development (MI_{16h}, MII_{30h} and MII_{65h}). The key proteins discussed in the text are shown at the developmental stage in which they display significant phosphorylation site up-regulation (q-value < 0.01).

FIG. 9. Phenotypes of the *S. coelicolor* harbouring the *FtsZ* wild-type gene (Ser-Ser), the *FtsZ* mutant mimicking double phosphorylation (“Glu-Glu”) and the *FtsZ* mutant mimicking no phosphorylation (“Ala-Ala”). *A*, macroscopic view of sporulated cultures 96-hours in SFM medium); the grey colour corresponds to spores. *B*, peptidoglycan staining (WGA staining) of vegetative hyphae (15-hours in GYM cultures). Arrows indicate septa. *C*, sucrose-free R5A liquid cultures; upper panel, growth curve (protein/ml); middle panel actinorhodin production; lower panel undecylprodigiosin production. *D*, macroscopic view of solid cultures; SFM and GYM at 72 h, MM at 96 h. Purple colour actinorhodin, red colour undecylprodigiosin (not visible at the time points shown).

FIG. 10. Abundances of proteins implicated in secondary metabolism in the *FtsZ* Glu-Glu and Ala-Ala mutants compared to the wild-type strain (Ser-Ser). *A*, heat maps. *B*, TMT abundances of the key proteins described in the text. Statistics and labelling as in Fig. 4.

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