Production of 11α-hydroxy steroids from sterols in a single fermentation step by *Mycobacterium smegmatis*

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Summary

11α-hydroxylated steroid synthons are one of the most important commercially pharmaceutical intermediates used for the production of contraceptive drugs and glucocorticoids. These compounds are currently produced by biotransformation using fungal strains in two sequential fermentation steps. In this work, we have developed by a rational design new recombinant bacteria able to produce 11α-hydroxylated synthons in a single fermentation step using cholesterol (CHO) or phytosterols (PHYTO) as feedstock. We have designed a synthetic operon expressing the 11α-hydroxylating enzymes from the fungus *Rhizopus oryzae* that was cloned into engineered mutant strains of *Mycobacterium smegmatis* that were previously created to produce 4-androstene-3,17-dione (AD), 1,4-androstadiene-3,17-dione (ADD) from sterols. The introduction of the fungal synthetic operon in these modified bacterial chassis has allowed producing for the first time 11α-OH-AD and 11α-OH-ADD with high yields directly from sterols in a single fermentation step. Remarkably, the enzymes of sterol catabolic pathway from *M. smegmatis* recognized the 11α-hydroxylated intermediates as alternative substrates and were able to efficiently funnel sterols to the desired hydroxylated end-products.

Introduction

Steroids represent one of the most widely used drugs for multiple clinical purposes (e.g. anti-inflammatory, immunosuppressive, anti-allergic, anti-cancer) (Fernández-Cabezón et al., 2018). The oxidation state of steroid rings and the presence of different attached functional groups determine the specific physiological function of each steroid (Lednicer, 2011), and therefore, structural modifications of steroids, such as hydroxylation, dehydrogenation or esterification, highly affect their biological activity (Donova and Egorova, 2012; Szaleniec et al., 2018). Among them, hydroxylation is one of the most important steroid modifications introduced by the pharmaceutical industry, since it introduces deep changes not only in their clinical activities, but also in their physico-chemical properties (e.g. solubility, adsorption).

Fungi have been reported to carry out hydroxylations at almost all stereo core centres of the steroid molecule (Kristan and Rižner, 2012), and they have been widely applied at industrial scale for the production of hydroxylated steroids with a broad range of biological activities (Donova, 2017). This oxyfunctionalization of steroids is mainly catalysed by cytochromes P450 (CYPs) acting as monooxygenases by inserting a single oxygen atom into a non-activated C–H bond of the substrate with a concomitant reduction of other oxygen atom to water (Bernhardt, 2006). Fungal hydroxylations are usually carried out by two-component enzyme systems consisting of a CYP monooxygenase and a NAD(P)H CYP-reductase (CPR) (Crešnar and Petrič, 2011; Kristan and Rižner, 2012).

In particular, 11α-hydroxylated steroid synthons are one of the most important commercially steroid intermediates used for the production of contraceptive drugs and glucocorticoids. Important fungal species for steroid 11α-hydroxylation reactions include *Rhizopus nigricans*, *Aspergillus oryzae*, *Aspergillus oryzae* (Wang et al., 2017; Kollerov et al., 2020). In this sense, the 11α-hydroxylation system of *R. oryzae* has been used for the production of hydroxyprogesterone (Fernandes et al., 2003; Petric et al., 2010). This fungal system consists in the CYP509C12, one of the 48 CYPs encoded in its genome, and its redox partner RoCPR1, a NAD(P)H-dependent CPR. By expressing CYP509C12...
in yeast, it has been demonstrated that this CYP hydroxylates predominantly at 11α and 6β positions of steroids (Petric et al., 2010), although 11α-hydroxylation of steroids by fungi is the synthetic procedure currently used by the steroid industry (Petric et al., 2010). The current method for obtaining 11α-OH-ADD at industrial level is a two-step one-pot bioconversion from phytosterols based on the specific biochemical activities of two microbial strains without separation and purification of the intermediate product (Dovbnya et al., 2017). However, the large number of CYPs contained in the fungal strains often generates during the in vivo hydroxylation a multiplicity of oxygenated steroid by-products, which not only are difficult to separate, but also contribute to reduce the bioconversion yields (Fernandes et al., 2003). Therefore, the design of alternative and more specific hydroxylating microbial cell factories created by recombinant DNA technologies has been considered an industrial challenge (Donova, 2017; Fernández-Cabezón et al., 2018).

In this sense, the cloning and expression of hydroxylating CYPs and their CPRs partners in heterologous hosts to be used as alternative fungal bio-catalysts has been explored in yeasts and bacteria (Szczepański et al., 2003; Schmitz et al., 2014; Schiffer et al., 2015; Ichinose et al., 2015; Hull et al., 2017; Uno et al., 2017; Wang et al., 2017; Lu et al., 2018; Felpeto-Santero et al., 2019). However, all these bio-catalysers described so far are only able to transform pre-synthesized steroid synthons and, therefore, the complete synthesis requires two fermentation steps, one to produce the first synthons and other to hydroxylate them. Up to now, none of the developed hydroxylating recombinant bio-catalysts provide the possibility of producing hydroxylated steroids in a single fermentation step starting from natural sterols.

Some actinobacteria (e.g. Mycobacterium, Mycolicibacterium, Gordonia, Rhodococcus) that are able to metabolize cholesterol and other natural sterols are currently used by the industry to obtain steroid synthons (e.g. 4-androstene-3,17-dione (AD), 1,4-androstanediene-3,17-dione (ADD), 9-hydroxy-4-androstene-3,17-dione (9OH-AD), testosterone (TS) and C22 steroids (e.g. 22-hydroxy-23,24-bisnorchole-4-en-3-one (4-HBC), 3-oxo-23,24-bisnorchole-1,4-dien-22-oic acid (1,4-HBC), 9,22-dihydroxy-23,24-bisnorchole-4-en-3-one (9OH-4-HBC) from sterols (phytosterols (PHYTO) or cholesterol (CHO)) (Fig. 1) (Donova and Egorova, 2012; García et al., 2012; Galán et al., 2017). Although most of the strains used at industrial scale are natural isolated or random mutagenized strains, in the last few years a number of new steroid-producing strains have been rationally designed by metabolic engineering approaches (Wei et al., 2010, 2014; Xiong et al., 2017; He et al., 2018; Liu et al., 2018; Zhang et al., 2018). In this way, we have developed several mutant strains of Mycolicibacterium smegmatis (formerly Mycobacterium smegmatis) able to produce à la carte a large number steroid synthons (Galán et al., 2017; García et al., 2017).

In this work, we have tested the possibility of rationally design new recombinant bacteria able to produce 11α-hydroxylated steroids in a single fermentation step using sterols as feedstock. To this aim, we have designed a synthetic operon containing the 11α-hydroxylating enzymes from R. oryzae that was cloned and expressed into two previously engineered mutant strains of M. smegmatis created to produce AD or ADD from sterols. The expression of the synthetic operon in these modified bacterial chassis has allowed us to produce for the first time 11α-hydroxylated compounds directly from sterols in a single fermentation step.

Results

Production of 11αOH–ADD in M. smegmatis

To test the possibility of producing 11α-hydroxylated steroids from natural sterols (CHO and PHYTO) in a single step in M. smegmatis, we designed the synthetic bacterial operon FUN (RoCPR59830-CYP509C12), harboring the CYP 11α-hydroxylase from R. oryzae and its redox partner CRP (Fig. S1). The synthetic operon was cloned into the mycobacterial replicative plasmid pMV261, creating the pMVFUN plasmid, that was transformed into the M. smegmatis MS6039 mutant strain. This mutant strain has been previously engineered and accumulates ADD from CHO or PHYTO (Table 1) (García et al., 2012; García et al., 2017). The expression of the enzymes RoCPR59830 and CYP509C12 in the MS6039 (pMVFUN) recombinant strain was determined by SDS-PAGE (Fig. S2A) and the ability to transform sterols into 11αOH-ADD was analysed by HPLC-MS along the growth curve using steroids (CHO or PHYTO) as feedstock.

First, MS6039 (pMVFUN) and MS6039 (pMV261) strains were grown in the biotransformation medium in the presence of CHO. During the exponential phase, the recombinant strain MS6039 (pMVFUN) showed a slight delay in growth compared to the control strain MS6039 (pMV261), but both cultures reached similar biomass at the stationary phase (Fig. 2A). After 24 h, 11αOH-ADD was detected in the culture supernatant of MS6039 (pMVFUN) and the maximum bioconversion of 99.6 ± 0.3 % was obtained after 60 h of growth with a production yield of 65.8 ± 3.9 % for 9φ11αOH-ADD/CHO (Fig. 2B). Interestingly, the HPLC-UV/DAD-MS monitoring allowed us to identify and quantify small amounts of some by-products in the culture medium, such as ADD, 11αOH-AD and trace amounts of 1,4-HBC (Fig. 2B). Production yields for the main by-products were 25.4 ± 4.3 % for ADD/CHO and 8.6 ± 0.5 % for 11αOH-
AD/CHO (Fig. 2B, Fig. S3A and B). We have detected one additional compound eluting at 4.92 min (Fig. S3A and B). Its m/z of 345 coincides with the molecular mass of 1,4-HBC increased by 16, suggesting that it could correspond to 11α-OH-1,4-HBC.

As expected, MS6039 (pMV261) control strain did not produce 11α-OH-ADD from CHO (Fig. S3A) and ADD was detected as the main biotransformation product with a conversion rate of 96.2 ± 5.9% and a production yield of 99.3 ± 0.3% for ADD/CHO.

Taking into account that PHYTO are used in the steroid industry as the preferred low-cost raw material to produce steroid synthons, we tested it as feedstock to produce 11α-OH-ADD in the MS6039 (pMVFUN) recombinant strain. To this aim, MS6039 (pMVFUN) and MS6039 (pMV261) strains were grown in the biotransformation medium in the presence of PHYTO and monitored by HPLC DAD-MS as performed for CHO cultures (Fig. 2C). The MS6039 (pMVFUN) strain successfully achieved the transformation of PHYTO into 11α-OH-ADD. The conversion rate was 67.5 ± 0.3%, and the 11α-OH-ADD production yield 11α-OH-ADD/PHYTO was 33.3 ± 0.2% (Fig. 2D and Fig. S4). Some by-products as ADD, 11α-OH-ADD, 1,4-HBC and 11α-OH-1,4-HBC were detected when PHYTO was used as feedstock (Fig. S4).

The yields for these by-products were 59.5 ± 0.4% for ADD/PHYTO and 3.4 ± 0.1% for 11α-OH-ADD/PHYTO. Derived compounds 11α-OH-1,4-HBC and 1,4-HBC could not be quantified because they were present at very low concentrations.

As expected, MS6039 (pMV261) control strain only produced ADD from PHYTO with a conversion rate of 67.5 ± 0.3% and a transformation yield of 95.7 ± 0.9% for ADD/PHYTO (Fig. S4).

Production of 11α-OH-AD in M. smegmatis

To achieve the conversion of sterols into 11α-OH-AD in a single fermentation step, we used as chassis the M. smegmatis MS6039-5941 mutant strain that was engineered to produce AD from CHO or PHYTO (Table 1) (García et al., 2012; García et al., 2017). The MS6039-5941 strain was transformed with the pMVFUN plasmid as described above. The production of CYP509C12 and RoCPR59830 proteins in the recombinant MS6039-5941 (pMVFUN) strain was confirmed by SDS-PAGE (Fig. S2B), and the ability to transform sterols (CHO and PHYTO) into 11α-OH-AD was analysed by HPLC-MS.

First, MS6039-5941 (pMVFUN) and control MS6039-5941 (pMV261) strains were cultured in the
biotransformation medium in the presence of CHO. As observed with MS6039 (pMVFUN), during the exponential phase, the recombinant strain MS6039-5941 (pMVFUN) showed a delay in growth compared to the control strain MS6039-5941 (pMV261), but this effect was partially restored at stationary phase (Fig. 2E). CHO started to be consumed after 48 h of growth and completely depleted at 96 h. The product yield reached was 74.0 ± 0.2% 11α-OH-AD (CHO) (Fig. 2F). Some by-products as AD, 4-HBC, 11α-OH-ADD, ADD, and 1,4-HBC were produced during the biotransformation (Fig. 2F, Fig. S5A and B). The more relevant detected by-products were AD (AD/CHO = 9.0 ± 10%), 4-HBC (4-HBC/CHO = 8.3 ± 0.4%) and 11α-OH-ADD (11α-OH-ADD/CHO = 7.9 ± 0.4%). We have detected two additional compounds eluting at 2.2 and 5.7 min (Fig. S5A and B). Based on their m/z, they could be assigned to the double hydroxylated compound 11α, 6β-diOH-4-HBC (m/z of 319) and 11α-OH-4-HBC (m/z of 319) (Fig. S5A and B). It is worth to mention that the control strain MS6039-5941 (pMV261) did not produce 11α-OH-AD (Fig. S5) and, as expected, AD is the main product, having a yield of 76.7 ± 3.7% with a conversion of 99.1 ± 0.2%. Curiously, a significant amount of 4-HBC was detected in the control strain (4-HBC/CHO = 18.3 ± 2.9%) when compared to the strain carrying the FUN operon. A small amount of ADD was also detected in the control strain (ADD/CHO = 4.9 ± 0.9%). On the other hand, the MS6039-5941 (pMVFUN) strain was grown in the presence of PHYTO to check its ability to convert this raw material into 11α-OH-AD. During the biotransformation, the recombinant strain MS6039-5941 (pMVFUN) showed also a slight decrease in growth compared to the control strain MS6039-5941 (pMV261) (Fig. 2G). The HPLC monitoring revealed that this strain transformed PHYTO into 11α-OH-AD (Fig. 2H). As it is shown, phytosterols reached a conversion of 73.0 ± 1.7% at 96 h. At this time, the product yield was...
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57.3 ± 5.3% for 11α-OH-AD/PHYTO. In addition, other by-products, such as AD (AD/PHYTO = 17.9 ± 1.3%), 4-HBC (4-HBC/PHYTO = 12.1 ± 2.3%), 11α-OH-ADD (11α-OH-ADD/PHYTO = 2.9 ± 0.2%) and ADD (ADD/PHYTO = 1.6 ± 0.5%) were detected in the culture supernatant (Fig. 2H and Fig. S5). As expected, MS6039-5941 (pMV261) control strain did not render any 11α-OH-AD (Fig. S6) and only produced AD achieving a conversion of 52.7 ± 12.9 % yielding 77.6 ± 2.0 % AD/PHYTO. A significant amount of 4-HBC (17.0 ± 1.2% 4-HBC/PHYTO) and a small amount of ADD (ADD/PHYTO = 3.3 ± 0.4%) were produced as by-products (Fig. S6).

Discussion

In the pharmaceutical sector, steroid hydroxylation plays an important role to produce new functionalized steroids, because it usually introduces deep changes in their physicochemical and pharmaceutical properties. In particular, the 11α- or 11β-hydroxylation of steroids are essential functionalization steps to develop commercially important intermediates to synthesize glucocorticoids and contraceptive drugs. Current methods of production of hydroxylated steroids mainly rely on biotransformations using wild-type fungal whole cells that harbour these enzymatic activities. The production of the hydroxylated steroids is carried out in at least two fermentation steps exhibiting in most cases some drawbacks such as low selectivity and reduced conversion yield. Therefore, the design of alternative fermentation processes by using recombinant DNA technologies has been proposed in recent years. In this sense, several fungal hydroxylases have been successfully expressed in yeasts (Petric et al., 2010; Hull et al., 2017; Lu et al., 2018) demonstrating their potential biotechnological applications. However, to the best of our knowledge these recombinant yeasts have not been implemented at industrial scale yet.

In this work, we have advanced one-step forward in the direction of creating alternative processes to produce the 11-hydroxylated steroids directly from sterols in a single fermentation step. To fulfil this aim, we have used a rationally designed synthetic operon (named FUN operon) carrying the 11α-hydroxylating system from R. oryzae and cloned them in M. smegmatis generating new recombinant bacterial strains able to produce 11-hydroxylated derivatives in a single fermentation directly from natural sterols such as CHO and PHYTO.

M. smegmatis is a model microorganism that has been used for the study of steroid catabolism due to some strengths: (i) it is a fast-growing bacterium capable to grow in natural sterols (CHO and PHYTO) as a sole carbon and energy source; (ii) its genome sequence has been accessible since 2006; (iii) it can be manipulated genetically with many genetic tools; (iv) it is a robust chassis resistant to stressing industrial production conditions; and (v) it has a very effective transport system for sterols. Hence, M. smegmatis has been proposed and used before as an efficient platform to produce steroidal drugs (Fernández-Cabezón et al., 2017; Galán et al., 2017).

One of the most important achievements of this work is the demonstration that the synthetic fungal operon was fully functional in M. smegmatis. Only few fungal CYPs have been successfully produced in its active form in bacteria so far (Barnes et al., 1991; González and Korzekwa, 1995; Hannemann et al., 2006; Felpeto-Santero et al., 2019). In our case, we achieved significant levels of CYP and CRP expression as determined by SDS-PAGE (Fig. S1). Apparently, depending on the substrate and the mutant strain used, the production of the hydroxylating system generates some cellular stress in the host as deduced by comparing the corresponding growth curves (Fig. 2).

The detection of the 11α-hydroxylated steroids in the culture medium of the recombinant bacteria at high yields demonstrated that the fungal 11α-hydroxylating enzymes have been integrated in the sterol metabolism in M. smegmatis strains creating a new expanded pathway. Such an efficient integration was in fact a surprising result, because one likely outcome could be the hydroxylation of different metabolic intermediates that might block some of the multiple reactions required to accumulate the desired compound due to the specificity of the sterol catabolic enzymes. However, MS6039 (pMVFUN) and MS6039-5941 (pMVFUN) recombinant strains successfully accumulated 11α-OH-AD and 11α-OH-ADD, respectively, using CHO and PHYTO as precursors. In this sense, the detection of 11α-OH derivatives of AD, ADD, 4-HBC and 1,4-HBC suggests, on the one hand that the enzymes of R. oryzae are not highly specific for a particular steroid. However, on the other hand, this also means that although several hydroxylated intermediates can be produced, none of them blocks the pathway, and fortunately, can be finally funnelled to the final products 11α-OH-AD or 11α-OH-ADD, depending on the mutant host used. Therefore, we can conclude that the enzymes of the M. smegmatis sterol degradation pathway can also recognize as substrates the 11α-OH derivatives of their own natural substrates.

It is worth to mention that when the 11α-hydroxylating system of R. oryzae was expressed in yeast, it was able to hydroxylate steroids, both at 11α and 6β positions, with good yields (Petric et al., 2010). However, in our case the main products obtained were hydroxylated at 11α position. We have detected trace amounts of a compound of m/z 319 compatible with the dihydroxylated...
derivative of 4-HBC in the biotransformation of CHO by the MS6039-5941 (pMVFUN) strain. Although more experiments are required to confirm the presence of small amounts of 6β-OH derivatives in M. smegmatis, this finding suggests that 6β-hydroxylation is not relevant in our recombinant strain. This interesting result suggests that, most probably, the specificity of CYP enzymes depends on their metabolic environment (location, pH, substrates, other reductases, etc.), and the hydroxyl group destination (1α or 6β position) can be favoured in one selective direction. This characteristic has been also previously observed in the case of heterologous expression of the same R. oryzae hydroxylating system in yeast where the distribution between 11α- and 6β-hydroxylated derivatives, and the appearance of non-identified monohydroxysteroids depended on the substrate that was used (Petric et al., 2010).

Although the whole process has to be further optimized at industrial scale to improve the yield of the 11α-hydroxylated compounds and to reduce the amount and number of by-products, our results open a new avenue for searching effective biocatalysts for producing hydroxylated steroids from sterols in a single fermentation step. In addition, they reinforce the assumption that engineered M. smegmatis strains represent a new generation of biocatalysts with a great potential to be applied for industrial processes. Based on the increased knowledge on the steroid metabolism in M. smegmatis, we uphold this bacterium as an exceptional bacterial chassis to implement à la carte metabolic engineering strategies based on synthetic biology for the industrial production of other valuable pharmaceutical steroids directly from sterols.

**Experimental procedures**

**Chemicals**

Commercial phytosterols from pine (provided by Gadea Biopharma, Spain) containing a mixture of different sterols (w/w percentage): β-sitosterol (83.61 %), stigmasterol (8.79 %) and campesterol (7.59 %), 22-hydroxy-23,24-bisnorchol-1,4-dien-22-oic acid (1,4-HBC kindly given by Gadea Pharma S.L., 4-androstene-3,17-ols (w/w percentage):

Commercial phytosterols from pine (provided by Gadea Genomics Stock Center, FGSC, University of Missouri; Petric et al., 2010) in bacteria, we designed the synthetic operon FUN that encodes the cytochrome CYP509C12 (EIE80372) with 11α-hydroxylase activity and its natural redox partner RoCPR1 cytochrome reductase (EIE89541). The codon usage was manually optimized for M. smegmatis, keeping a nucleotide identity percentage of approximately 83% for both genes. A consensus Shine–Dalgarno sequence of 6 bp (AAAGGGAG) was added upstream the respective start codons as well as some restriction sites to facilitate cloning (Fig. S1). Alanine was also included as the second amino acid of the resulting proteins to increase protein translation (Bivona et al., 2010). The operon engineered was chemically synthesized by ATG:biosynthetics GmbH and initially cloned into the pGH vector yielding plasmid pGH-FUN that was used to transform E. coli DH10B cells and check the sequence. Plasmid pGH-FUN was digested with BamHI-EcoRI to release the fragment containing the FUN operon that was further cloned under the control of the P_{rps} constitutive promoter, into the shuttle E. coli/Mycobacterium vector pMV261 (Stover et al., 1991) yielding pMVFUN. Plasmids pMV261 (empty vector, control plasmid) and pMVFUN were individually transformed into the M. smegmatis mutants by electroporation.

**Design and construction of the bacterial synthetic FUN (CYP509C12-RoCRP1) operon**

To achieve the heterologous production of the 11α-hydroxylase activity from R. oryzae RA 99-880 (Fungal Genetics Stock Center, FGSC, University of Missouri; Petric et al., 2010) in bacteria, we designed the synthetic operon FUN that encodes the cytochrome CYP509C12 (EIE80372) with 11α-hydroxylase activity and its natural redox partner RoCPR1 cytochrome reductase (EIE89541). The codon usage was manually optimized for M. smegmatis, keeping a nucleotide identity percentage of approximately 83% for both genes. A consensus Shine–Dalgarno sequence of 6 bp (AAAGGGAG) was added upstream the respective start codons as well as some restriction sites to facilitate cloning (Fig. S1). Alanine was also included as the second amino acid of the resulting proteins to increase protein translation (Bivona et al., 2010). The operon engineered was chemically synthesized by ATG:biosynthetics GmbH and initially cloned into the pGH vector yielding plasmid pGH-FUN that was used to transform E. coli DH10B cells and check the sequence. Plasmid pGH-FUN was digested with BamHI-EcoRI to release the fragment containing the FUN operon that was further cloned under the control of the P_{rps} constitutive promoter, into the shuttle E. coli/Mycobacterium vector pMV261 (Stover et al., 1991) yielding pMVFUN. Plasmids pMV261 (empty vector, control plasmid) and pMVFUN were individually transformed into the M. smegmatis mutants by electroporation.

**Analysis of the FUN operon expression by SDS-PAGE analysis**

The recombinant strains MS6039 and MS6039-5941 carrying plasmid pMVFUN (Table 1) were grown in bio-transformation media and conditions during 24 h. Cells
were harvested by centrifugation (15 min, 5000 × g, 4 °C) thawed on ice, washed twice with NaCl 0.9 % (w/v) and resuspended in 0.5 ml of Tris-HCl 50 mM pH 7.5. Cells were disrupted by sonication using a Branson sonicator 150 (6-8 pulses of 1 min at 90% power, with 30 s of cooling on ice between each). Cell debris was removed by centrifugation at 14,000 × g for 15 min at 4 °C. Soluble and insoluble fractions were analysed by SDS-PAGE to check FUN operon expression.

Bioinformatic tools

DNA and protein sequences from *R. oryzae* genome were obtained from Broad Institute Server (http://www.broad.mit.edu/annotation/genome/rhizopusoryzae/MultiHome.html) and National Center for Biotechnology Information (NCBI) Database (http://www.ncbi.nlm.nih.gov/). Nucleotide and amino acid sequences were compared to National Center for Biotechnology Information (NCBI) database using BLAST algorithms (http://www.ncbi.nlm.nih.gov/BLAST) to align and compared to local database. Local-BLAST (BioEdit) and HMMER algorithm at European Bioinformatics Institute (EBI) (http://www.ebi.ac.uk/Tools/hmmer/) were used. Multiple alignments of protein were carried out with the MUSCLE server programme at the European Bioinformatics Institute (EBI) (http://www.ebi.ac.uk/Tools/msa/muscle/).

Steroid biotransformation assay

Sterol biotransformation assays were carried out using *M. smegmatis* MS6039 (pMVFUN) and MS6039-5941 (pMVFUN) growing cells in shaken 100 ml flasks containing 20 ml of biotransformation medium having the following composition: 7H9 Broth supplemented with 18 mM glycerol as starter, 1 mM cholesterol or 1 mM phytosterols (previously dissolved in 3.6 % (v/v) Tyloxapol, 0.5 mM l-aminolevulinic acid (ALA) and kanamycin (20 µg ml⁻¹). First, a well-grown (48h) pre-culture of the MUSCLE server programme at the European Bioinformatics Institute (EBI) (http://www.ebi.ac.uk/Tools/msa/muscle/).

Steroids extraction

Aliquots of 10 µl of 5 mM TES in 10 % (v/v) tyloxapol were added to each 0.2 ml sample taken from the biotransformation experiments prior to extraction with chloroform, as an internal standard (ISTD). The samples were extracted using two volumes of chloroform. The aqueous fraction was discarded, and the chloroform fraction was dried at 60 °C using a Thermoblock and then dissolved in 0.5 ml of acetonitrile. Each sample was subjected to chromatographic analysis by HPLC-UV/DAD-MS (25 µl).

HPLC-UV/DAD-MS analysis

Experiments were carried out using a DAD detector and a LXiQ Ion Trap Mass Spectrometer, equipped with an atmospheric pressure chemical ionization source, electrospray ionization source and interfaced to a Surveyor Plus LC system (all from Thermo Electron, San Jose, CA, USA). Data were acquired with a Surveyor Autosampler and MS Pump and analysed with the Xcalibur Software (from Thermo Fisher Scientific, San Jose, CA, USA). High-purity nitrogen was used as nebulizer, sheath and auxiliary gas. MS analysis was performed both in full scan and in selected ion monitoring (SIM) mode by scanning all the daughter ions of the products in positive ionization mode. The quantification was performed from parent mass of compounds, and the specificity was obtained by following the specific fragmentations of all compounds.

The experiments were carried out with the following interface parameters: Ionization source APCI, volume to inject 25 µl, capillary temperature 275°C, vaporizing temperature 425°C, capillary voltage 39 V, corona discharge 6 kV, source power 6 µA and dissociation by collision energy 15 eV. Use high pure nitrogen as an auxiliary gas and sprayer. Chromatographic separation was performed on a *Tracer Excel* 120 ODSB C18 (4.6 mm x 150 mm, particle size 5 µm) (Teknokroma). The chromatography was performed using water containing 0.1 % (v/v) of formic acid, acetonitrile containing 0.1 % (v/v) of formic acid and isopropanol containing 0.1 % (v/v) of formic acid as mobile phases A, B and C, respectively (flow 1 ml min⁻¹). To monitor cholesterol (CHO) biotransformation, the examined compounds were as follows: TES (289 m/z), AD (287 m/z), ADD (285 m/z), 11αOH-ADD (301 m/z), 4-HBC (331 m/z) and 1,4-HBC (329 m/z). The HPLC gradient was as follows:
Quantification was performed by an ISTD (Internal Standard) method. The quantification of the compounds was calculated by the reaction yield and peak area regarding to IS. To monitor the biotransformation of PHYTO, that is a mixture of sitosterol (STIG), stigmasterol (STI), and campesterol (CAMP), the examined compounds were as follows: TES (289 m/z) used as ISTD; CAMP (383 m/z), STI (395 m/z), and STIG (397 m/z). The HPLC gradient used was as follows:

| Time (min) | % A | % B | % C |
|-----------|-----|-----|-----|
| 0         | 50  | 50  | 0   |
| 5         | 50  | 50  | 0   |
| 15        | 20  | 71  | 9   |
| 20        | 4   | 87  | 9   |
| 40        | 0   | 85  | 15  |
| 41        | 0   | 85  | 15  |
| 42        | 50  | 50  | 0   |
| 52        | 50  | 50  | 0   |

The concentration of sterols in 1 mM PHYTO is as follows: 0.84 mM STI, 0.08 mM STIG, and 0.08 mM CAMP. Therefore, the quantity of PHYTO represents the sum of the quantities of STI, STIG, and CAMP. The quantification of the compounds was calculated by the reaction yield and peak area regarding to IS.

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Conflict of interest
None declared.

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Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Fig. S1. Schematic representation of the genes contained in FUN operon (RoCPRI-CYP509C12). The sequences of the intergenic regions R1-R3 are indicated in the table. The sequence of restriction sites is underlined and RBS sequences are indicated in bold. The main restriction sites in the synthetic operon are also indicated.

Fig. S2. SDS-PAGE analysis of the 11α hydroxylating enzymes from R. oryzae in the M. smegmatis recombinant strains. (A) Lanes 1 and 6, molecular mass markers; lane 2, soluble (SN) control extract from MS6039 (pMV261); lane 3, soluble extract from MS6039 (pMFUN); lane 4, insoluble fraction from MS6039 (pMV261) and lane 5, insoluble fraction from MS6039 (pMV261). (B) Lanes 1 and 6, molecular mass markers; lane 2, soluble (SN) control extract from MS6039 (pMV261) and lane 5, insoluble fraction from MS6039 (pMV261) and lane 5, insoluble fraction from MS6039 (pMV261) and lane 5, insoluble fraction from MS6039 (pMFUN). The bands of CYP and CPR proteins are indicated by asterisks and arrows.

Fig. S3. CHO biotransformation by MS6039 (pMFUN) and MS6039 (pMV261) (Control) strains. A) HPLC-DAD chromatogram (50-600 nm) (green) and full scan mass spectra (m/z 150-400) at 0 h and 96 h of growth are shown. B) Magnification of the first 14 min of the HPLC-DAD chromatogram (50-600 nm) green, full scan mass spectra (m/z
150-400), mass spectra of m/z 301 and m/z 303 corresponding to 11αOH-ADD and 11αOH-AD are shown.

**Fig. S4.** PHYTO biotransformation by MS6039 (pMVFUN) (FUN) strain. HPLC-DAD chromatogram (50-600 nm) (green), full scan mass spectra (m/z 150-400), mass spectra of SITO (m/z 387), CAMP (m/z 383), STIG (m/z 386), 11αOH-ADD (m/z 301) and ADD (m/z 286) at 96 h are shown.

**Fig. S5.** CHO biotransformation by MS6039-5941 (pMVFUN) (FUN) and MS6039-5941 (pMV261) (Control) strains. A) HPLC-DAD chromatogram (50-600 nm) (green) and full scan mass spectra (m/z 150-400) at 0 h and 96 h of growth are shown. B) Magnification of the first 20 min of the HPLC-DAD chromatogram (50-600 nm) green, full scan mass spectra (m/z 150-400), mass spectra of m/z 303 corresponding to 11αOH-AD are shown.

**Fig. S6.** PHYTO biotransformation by MS6039-5941 (pMVFUN) (FUN) strain. HPLC-DAD chromatogram (50-600 nm) (green), full scan mass spectra (m/z 150-400), mass spectra of residual SITO (m/z 387), CAMP (m/z 383), STIG (m/z 386) and 11αOH-AD (m/z 303) at 96 h of growth are shown.