De novo biosynthesis of garbanzol and fustin in Streptomyces albus based on a potential flavanone 3-hydroxylase with 2-hydroxylase side activity

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Summary
Flavonoids are important plant secondary metabolites, which were shown to have antioxidant, anti-inflammatory or antiviral activities. Heterologous production of flavonoids in engineered microbial cell factories is an interesting alternative to their purification from plant material representing the natural source. The use of engineered bacteria allows to produce specific compounds, independent of soil, climatic or other plant-associated production parameters. The initial objective of this study was to achieve an engineered production of two interesting flavanones, garbanzol and fustin, using Streptomyces albus as the production host. Unexpectedly, the engineered strain produced several flavones and flavonols in the absence of the additional expression of a flavone synthase (FNS) or flavonol synthase (FLS) gene. It turned out that the heterologous flavanone 3-hydroxylase (F3H) has a 2-hydroxylase side activity, which explains the observed production of 7,4′-dihydroxyflavone, resokaempferol, kaempferol and apigenin, as well as the biosynthesis of the extremely rare 2-hydroxylated intermediates 2-hydroxyliquiritigenin, 2-hydroxyxarigenin and probably licodione. Other related metabolites, such as quercetin, dihydroquercetin and eriodictyol, have also been detected in culture extracts of this recombinant strain. Hence, the enzymatic versatility of S. albus can be conveniently exploited for the heterologous production of a large diversity of plant metabolites of the flavonoid family.

Introduction
Flavonoids comprise a large family of polyphenols, representing about 9,000 compounds ubiquitously distributed in plants (Kumar and Pandey, 2013; Li et al., 2014; Shah et al., 2019). They are one of the largest families of natural products and share a generic chemical structure consisting of 15 carbon atoms (C6-C3-C6). The core structure harbours two aromatic rings (rings A and B) connected by a heterocyclic tetrahydropyran (ring C) (Verma and Pratap, 2010; Ravishankar et al., 2013; Yao et al., 2014), and this molecule’s backbone can carry multiple substituents, such as hydroxyl or methyl groups, as well as sugars (Crozier et al., 2009). Flavonoids have received significant attention due to their medicinal value as nutraceuticals and cancer chemopreventive agents, acting against different types of chronic diseases related to the cardiovascular system (Sánchez et al., 2019) or cancer (Fernández et al., 2016; George et al., 2017; Redondo-Blanco et al., 2017). Recently, the flavonoid herbacetin was shown to inhibit 3C-like protease (3Cpro) of coronaviruses (CoVs) (Jo et al., 2020). Outside of the field of medicine, flavonoids find potential applications in food preservation (Gutiérrez-del-Río et al., 2018).

Similar to other flavonoids, fustin (sometimes called dihydrofisetin) and garbanzol are valuable nutraceuticals and have been found to possess antioxidant (Chen et al., 2017), anticancer (Park et al., 2004; Fotso et al., 2017),...
antibacterial (Jang et al., 2018), antiviral (Kang et al., 2012), anti-inflammatory (Li et al., 2018) and neuroprotective (Park et al., 2007) properties. Nevertheless, very little is known about the biological activities of licidione or 2-hydroxyliquiritigenin, and only two licidione derivatives (2′-methoxy-3′-prenyl-licidione and 2′-methoxy-3′,3′-diprenyl-licidione) have been shown to possess isoprenylcysteine carboxyl methyltransferase (Icmnt) inhibitory activity, proving their potential as anticancer agents (Buchanan et al., 2008). The compound 7,4′-dihydroxyflavone was demonstrated to be a potent aromatase inhibitor (Pelissero et al., 1996; Ta and Walle, 2007) and additionally inhibits expression of mucin 5AC (MUC5AC), a key mucin in obstructive lung disease (Liu et al., 2007) and additionally inhibits expression of mucin 5AC (Lee et al., 2010; Yu et al., 2016).

In plants, the initial steps of flavonoid biosynthesis are part of the phenylpropanoid pathway, in which L-phenylalanine is converted to p-coumaroyl-CoA in three enzymatic steps, which are catalysed by phenylalanine ammonia-lyase (PAL, EC 4.3.1.24), cinnamate 4-hydroxylase (C4H, EC 1.14.14.91) and p-coumaroyl CoA ligase (4CL, EC 6.2.1.12; Falcone Ferreyra et al., 2012). The activity of six enzymes (TAL, 4CL, CHS, CHR, CHI and F3H) is required to produce garbanzol. In this work, the engineered production of putative garbanzol has been previously achieved by combinatorial biosynthesis using *Escherichia coli* and *Saccharomyces cerevisiae* (Stahlhut et al., 2015; Rodriguez et al., 2017), but we report de novo biosynthesis of resokaempferol and garbanzol in *Streptomyces albus*. We also report de novo biosynthesis of fustin, 2-hydroxyliquiritigenin, licidione (tenta
give identification) and 7,4′-dihydroxyflavone in a microorganism. Additionally, we report for the very first time the biosynthesis of flavones and flavonols from flavanones and flavanones, respectively, through 2-hydroxy intermediates and without the need for expression of an exogenous gene coding for an FNS or FLS. This fact suggests a possible catalytic F2H side activity of F3H, followed by an endogenous *S. albus* dehydratase activity; however, further studies are needed to unequivocally corroborate the assumption in this bacterium.

**Results**

**Heterologous biosynthesis of garbanzol**

The activity of six enzymes (TAL, 4CL, CHS, CHR, CHI and F3H) is required to produce garbanzol. In this work, six synthetic genes coding for the enzymes required for garbanzol biosynthesis (with codon usage adapted to the translation characteristics of *Streptomyces*) were cloned into a replicative high-copy number *E. coli*–*Streptomyces* shuttle vector under the control of P_{ermC} (see Experimental Procedures section; Fig. 2A). The P_{ermC} promoter is one of the most widely used promoters for heterologous gene expression in streptomycetes (Lombó et al., 2006; Park et al., 2009; Wang et al., 2012; Takano et al., 2017; Liu et al., 2018), and its correct activity for
controlling flavonoid biosynthetic genes has been confirmed in previous studies of this research group (Marín et al., 2017, 2018; García-Gutiérrez et al., 2020). Genes were selected from different bacteria (Rhodobacter capsulatus and S. coelicolor) and plants (Glycine max, Petroselinum crispum and Arabidopsis thaliana) as they were already tested in the laboratory for these enzymatic steps (Marín et al., 2017, 2018; García-Gutiérrez et al., 2020). In bacteria, the use of TAL is preferred for heterologous biosynthesis purposes, as starting from L-tyrosine, the need for the C4H activity (a plant membrane-bound enzyme) does not longer exist. Because almost all the TALs show activity towards both phenylalanine and tyrosine, TAL enzyme from R. capsulatus was chosen based on its high affinity for tyrosine ($K_m = 160$), instead of phenylalanine ($K_m = 560$) (Xue et al., 2007). The following enzymes 4CL, CHS, CHR and CHI, common to both garbanzol and fustin biosynthesis, were chosen from S. coelicolor and Glycine max as they were already functioning properly in other experiments (Marín et al., 2017, 2018; García-Gutiérrez et al., 2020). The final plasmid pGR was used for the transformation of S. albus J1074, yielding S. albus pGR. The recombinant strain was then tested for garbanzol production.

To confirm the heterologous production of garbanzol, the constructed strain S. albus pGR was cultivated, and extracts of the culture were analysed by HPLC-HRESIMS. Simultaneously, a negative control strain carrying the empty vector pIAGO, S. albus pIAGO, was analysed under the same conditions and compared with S. albus pGR. The obtained base peak chromatograms (BPCs) were extracted for the mass peak $m/z$ 271.06118 [M-H]⁻ (calculated for C₁₅H₁₂O₅), with a mass error range of 0.005 mmu (millimass units). The obtained extracted ion chromatograms (EICs) revealed the presence of six peaks (retention times: 4.7, 5.0, 5.1, 5.3, 6.5 and 6.7 min) for the proposed $m/z$ that were all absent in the negative control S. albus pIAGO (Fig. 3A). The retention time and the observed ions for two (peak 2 and peak 6, Fig. 3B) of the five detected peaks were consistent with those observed in the garbanzol and

![Proposed biosynthetic pathway for de novo production of flavonoids and chalcone licidione from naringenin chalcone (A) and isoliquiritigenin (B). 4CL, p-coumarate CoA ligase; CHI, chalcone isomerase; CHR, chalcone reductase; CHS, chalcone synthase; F2H, flavanone 2-hydroxylase; F3'H, flavonoid 3'-hydroxylase; F3H, flavanone 3-hydroxylase; FNS, flavone synthase; TAL, tyrosine ammonia-lyase.](A)
peaks 1, 3, 4 and 5 (Fig. 3A) could not be assigned to any of the flavonoids taking part in the proposed biosynthetic pathway; hence, further studies were carried out to identify those unknown compounds, focusing on peak 1.

A time-course cultivation of the garbanzol-producing strain *S. albus* pGR and the negative control strain *S. albus* pIAGO was carried out. In this experiment, biomass concentrations and garbanzol titres were monitored over 168 h of incubation (Fig. 4), showing that the growth of the garbanzol-producing strain is similar to the control strain as no statistically significant differences were observed between both strain growth curve values, except for the biomass value at 72 h. Also, there are not statistically significant differences between the final dry cell weight at 168 h between both strains (Fig. 4). This

![Fig. 2](image)

Fig. 2. Designed gene clusters for heterologous expression of garbanzol (A) and fustin (B) in *S. albus*. Both DNA constructions were finally subcloned as a *Bgl*II-*Bam*HI fragment into the pIAGO plasmid. Blocked restriction enzymes sites (after DNA polymerase I Klenow fragment treatment during subcloning) are represented in brackets.

![Fig. 3](image)

Fig. 3. EICs at *m/z* 271.06118 ± 0.005 [M-H]⁻ of *S. albus* pGR and *S. albus* pDF crude extract (A) and commercial flavonoids standards of NRG (naringenin) and GRB (garbanzol) (B). Peak 1 corresponds to 2-hydroxyliquiritigenin (probably two isomers based on the stereochemistry of the 2-hydroxyl group), peak 2 corresponds to the two commercial garbanzol enantiomers, peaks 3 and 4 probably correspond to different tautomeric forms of licodione (tentative identification) and peak 6 corresponds to naringenin. GRB, garbanzol; NRG, naringenin.
rules out a possible toxic effect of some of the pathway intermediates or the end-products associated with the cloned biosynthetic pathway. Garbanzol titres increased during the exponential phase (until 72 h in R5A medium), and a maximum was detected at the beginning of the stationary phase (96 h). Then, the garbanzol titres started to decline in the days following incubation (Fig. 4). Interestingly, this drop from the maximum production is accompanied by an increase in the relative intensity of peak 1 (see Section Identification of liquiritigenin derivatives: 2-hydroxyliquiritigenin, licodione and 7,4'-dihydroxyflavone produced by the S. albus pGR strain), the maximum of which was reached at 144–168 h. This fact led to the belief that there might be a biosynthetic relationship between garbanzol and peak 1 (Fig. S5).

A comparison between the culture supernatant and cellular pellet was carried out to detect the main flavonoid compounds, both after solvent extraction (Fig. S4). Interestingly, the compound distribution was not the expected one as in the culture supernatant, mostly, p-coumaric acid, garbanzol and 2-hydroxyliquiritigenin were present. However, naringenin was mainly present in cellular pellet extracts. Isoliquiritigenin and liquiritigenin were similarly distributed between both organic extracts. This could imply the presence of specific membrane transporters in S. albus for some of these nutraceutical polyphenols.

Identification of liquiritigenin derivatives: 2-hydroxyliquiritigenin, licodione and 7,4'-dihydroxyflavone produced by the S. albus pGR strain

With the objective of deciphering the identity of the most abundant isomers of garbanzol under negative ionization conditions (peak 1), the engineered S. albus pGR strain was regrown (6 l laboratory scale) and submitted to HRESIMS-guided fractionation with different consecutive fractionation steps (SPE fractionation and semipreparative HPLC fractionation; see Section Experimental procedures). After purification, 0.8 mg of an almost pure compound was obtained, and the putative elucidation of the structure was performed on the basis of HPLC-HRESIMS/MS, LC-UV/vis and 1D-2D-NMR methods using the analysis of MS-, MS²-, UV/vis absorbance-, COSY- and HSQC-generated spectra. An authentic standard of (2R,3R)-garbanzol was also submitted to the same spectrometric and spectroscopic analyses, and all spectra were added to the pool of data available in our research group for commercially available flavonoids.

The MS spectrum for this new molecule showed a deprotonated ion at m/z 271.0613 [M-H]⁻, consistent with the molecular formula C₁₅H₁₂O₅ (calculated m/z 271.0613 [M-H]⁻). Examination of UV/vis and MS² data-sets led to similar absorbance peaks (Fig. 5) and identical fragmentation patterns compared with authentic commercial garbanzol (Fig. 6A), revealing the existence of a putative isomer of garbanzol. Although the chromatographic analysis showed a major peak with minor contaminants, the NMR analysis failed because no major signals arose and a vast mixture of minor compounds was detected. Interestingly, the HSQC spectrum did not show any oxygenated methine carbon. This fact not only excluded this new molecule to be a stereoisomer of garbanzol but also indicated the presence of a potential hydroxylation in the C2 position of the C ring, which would in fact correspond to 2-hydroxyliquiritigenin. However, stronger evidence about a positional isomer differing in the location of the hydroxyl group was needed to validate this hypothesis. Additionally, commercial (2S,3S)-garbanzol, which is a naturally occurring enantiomer of the (2R,3R)-garbanzol produced in the present study (Hashida et al., 2014), was purchased, but it showed an identical retention time as (2R,3R)-garbanzol (Fig. 3B). To gain an insight into this proposed C2 hydroxylation, the HPLC-HRESIMS/MS fragmentation profile of the new molecule, possibly 2-hydroxyliquiritigenin, was studied (Fig. 6B and C, see Appendix S1). The presence of a shoulder peak on the left side of peak 1 could point to the existence of two stereoisomers in relation to the position of the 2-hydroxy moiety at C2 of the heterocycle (Fig. 3A).

Fig. 4. Cultivation data (dry cell weight) of S. albus pGR (red) and negative control S. albus pAGO (green) grown on RSA medium sampled every 24 h over 168 h. Garbanzol titres in µg L⁻¹ have been overlayed (dashed line). The data are expressed as mean value ± standard error of mean (SEM). * indicate statistically significant differences between both strains (S. albus pGR and negative control S. albus pAGO) after two-way ANOVA.
To confirm the existence of an alternative biosynthetic pathway leading to licodione/7,4'-dihydroxyflavone, the BPC obtained from the crude organic extraction of \textit{S. albus} pGR was re-analysed in detail for the presence of ions corresponding to licodione (calculated \(m/z\) 271.06118 [M-H]−; \(\text{C}_{13}\text{H}_{12}\text{O}_{3}\)) and 7,4'-dihydroxyflavone (calculated \(m/z\) 253.05062 [M-H]−; \(\text{C}_{13}\text{H}_{13}\text{O}_{4}\)). The EIC for the mass peak 271.06118 [M-H]− showed the six peaks previously described (peaks 1–6) (Fig. 3A), with three of them representing known compounds: garbanzol (peak 2), 2-hydroxyliquiritigenin (peak 1, tentative identification) and naringenin (peak 6). The unidentified peaks Fig. 5. UV/vis spectra of several commercial flavonoid standards and purified putative 2-hydroxyliquiritigenin with \(m/z\) 271.0613 [M-H]− (peak 1 in Fig. 3A). 2OH-LIQ, 2-hydroxyliquiritigenin; BUT, butin; ILQ, isoliquiritigenin; LIQ, liquiritigenin; NRG, naringenin.

Fig. 6. HPLC-HRESIMS/MS analysis of authentic garbanzol and purified 2-hydroxyliquiritigenin. According to the nomenclature of Yang et al., each fragment is denoted by the combined use of \(^{i,i}\)A or \(^{i,i}\)B. A and B represent the flavonoid intact ring, and the superscript on the left indicates the broken bounds of the deprotonated molecule (Yang \textit{et al.}, 2012). A, Proposed retrocyclization cleavages of the C ring. B, MS\(^2\) fragmentation products for the parent ion \(m/z\) 271.0535 [M-H]−. C, MS\(^3\) fragmentation products for the in-source-formed species \(m/z\) 243.0591 [M-H]−. Product ions resulting from losses of H\(_2\)O, CO and CO\(_2\) have also been included.
3, 4 and 5 could belong to different tautomeric forms of liciodone (keto- and enol forms). When the \( m/z \) 253.05062 [M-H]⁻ ion was extracted, an unknown peak in the chromatogram at 5.5 min was observed (Fig. S2B). For both EICs, a mass error range of 0.005 mmu was selected, and they were not present in the negative control S. albus pGR (data not shown). The presence of both compounds could be confirmed using authentic liciodone (keto form) and 7,4′-dihydroxyflavone standards, but only the latter was commercially available and could be analysed under HPLC-HRESIMS conditions. The standard proved the presence of 7,4′-dihydroxyflavone (Fig. S2B) in the recombinant strain S. albus pGR constructed in this work. Thus, the existence of this molecule confirmed the presence of an unexpected additional pathway that generates 7,4′-dihydroxyflavone from liquiritigenin with 2-hydroxyl liquoritigenin as potential intermediate (Fig. 1B).

**Identification of the flavononol dihydrokaempferol and other flavone and flavonol derivatives: apigenin, resokaempferol and kaempferol produced by the S. albus pGR strain**

The crude organic extract of S. albus pGR and the different SPE (solid-phase extraction) fractions (I to VI) obtained during the purification process of 2-hydroxyl liquoritigenin were screened using HPLC-HRESIMS for the presence of flavonoids, for which MS data was available. Interestingly, clear signals of the deprotonated molecule mass peaks at \( m/z \) 269.0467/269.0457 [M-H]⁻ (two signals with different retention times), \( m/z \) 287.0563 [M-H]⁻ and \( m/z \) 285.0402 [M-H]⁻ were detected which eluted at the same retention time and were consistent with apigenin (calculated \( m/z \) 269.04553 [M-H]⁻; \( C_{15}H_{10}O_{5} \)); resokaempferol (calculated \( m/z \) 269.04553 [M-H]⁻; \( C_{15}H_{10}O_{5} \)); 2-hydroxykaempferol (calculated \( m/z \) 287.05610 [M-H]⁻; \( C_{15}H_{12}O_{6} \)) and kaempferol (calculated \( m/z \) 285.04045 [M-H]⁻; \( C_{15}H_{10}O_{5} \)) respectively (Fig. S2B). Besides, the EIC at \( m/z \) 287.05610 [M-H]⁻ showed one additional distinct peak (\( m/z \) 287.0569 [M-H]⁻) at a different retention time, which belongs to a molecule with the proposed molecular formula \( C_{15}H_{12}O_{6} \). In addition to dihydrokaempferol, the identified peak corresponds to 2-hydroxykaempferol, a coeluent of the commercial 2-hydroxykaempferol standard (Fig. S2B). In relation to this 2-hydroxy-derivative, a residual signal at \( m/z \) 303.0512 [M-H]⁻ consistent with the molecular formula \( C_{15}H_{12}O_{7} \) (calculated \( m/z \) 303.05101 [M-H]⁻) was detected, indicating the presence of 2-hydroxydihydrokaempferol, which could explain the unexpected detection of kaempferol. However, the assignment could not be confirmed because no authentic 2-hydroxydihydrokaempferol standard was commercially available. In addition, very low signals possibly belonging to 2-hydroxygarbanzol (no commercial standard available) were detected. None of these signals were present in the negative control S. albus pAGO.

Quantification of the intermediates and final products obtained with S. albus pGR led to the identification of p-coumaric acid (694.65 ± 9.52 µg l⁻¹), as well as lower concentrations of dihydrokaempferol (53.95 ± 0.84 µg l⁻¹), liquiritigenin (40.48 ± 1.81 µg l⁻¹), naringenin (38.52 ± 0.62 µg l⁻¹), isoliquiritigenin (19.081 ± 0.17 µg l⁻¹), garbanzol (3.74 ± 0.55 µg l⁻¹), kaempferol (1.74 ± 0.08 µg l⁻¹), apigenin (0.36 ± 0.01 µg l⁻¹), 2-hydroxyliquiritigenin (0.32 ± 0.05 µg l⁻¹), resokaempferol (0.14 ± 0.016 µg l⁻¹) and 7,4′-dihydroxyflavone (0.002 ± 0.0003 µg l⁻¹) (Fig. S3).

When the S. albus pGR strain was supplemented with 0.1 mM of naringenin, the garbanzol yield was drastically reduced (0.41 ± 0.04 µg l⁻¹). The same was true for liquiritigenin (12.16 ± 1.12 µg l⁻¹), 7,4′-dihydroxyflavone (LOQ, limit of quantification) and resokaempferol (LOQ), whereas titres of apigenin (2.82 ± 0.09 µg l⁻¹) and kaempferol (2.45 ± 0.41 µg l⁻¹) increased; titres of p-coumaric acid (753.40 ± 10.23 µg l⁻¹), isoliquiritigenin (14.35 ± 0.29 µg l⁻¹) and dihydrokaempferol (46.91 ± 2.26 µg l⁻¹) remained unaltered (Fig. S3).

**Heterologous biosynthesis of fustin**

Fustin is a hydroxylated derivative of garbanzol, and its biosynthesis requires the activity of one additional enzyme (F3′H) able to hydroxylate garbanzol at C3′. After having checked that the S. albus host was able to produce garbanzol, the gene coding for the F3′H enzyme was cloned into pGR (Fig. 2B). The new plasmid pDF was used for the transformation of S. albus, and the crude extract obtained after cultivation of the recombinant strain was subjected to HPLC-HRESIMS analysis.

As expected, all final products and the intermediates of the pathway that have been previously described for S. albus pGR were also detected in S. albus harbouroing pDF, including the 2-hydroxy intermediates 2-hydroxyliquiritigenin, 2-hydroxyliquiritigenin (Fig. S2C) and putative 2-hydroxydihydrokaempferol. Regarding the differential peaks from S. albus pGR, clear signals arose at \( m/z \) 287.0565/287.0558 [M-H]⁻ (two signals with different retention times), \( m/z \) 285.0402 [M-H]⁻, \( m/z \) 271.0615 [M-H]⁻, \( m/z \) 303.0505 [M-H]⁻ and \( m/z \) 301.0350 [M-H]⁻, which had the same retention time as authentic fustin (calculated \( m/z \) 287.05610 [M-H]⁻; \( C_{15}H_{12}O_{6} \)); eriodictyol (calculated \( m/z \) 287.05610 [M-H]⁻; \( C_{15}H_{12}O_{6} \)); fisetin (calculated \( m/z \) 285.04045 [M-H]⁻; \( C_{15}H_{12}O_{6} \)); pachypodiumin (calculated \( m/z \) 271.06118 [M-H]⁻; \( C_{15}H_{12}O_{6} \)).

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diydroquercetin (calculated m/z 303.05101 [M-H]–
C15H12O7) and quercetin (calculated m/z 301.03536 [M–
H]–; C15H10O7) respectively (Fig. S2C). All these differ-
tential peaks result from the activity of F3'H, which can use naringenin, dihydrokaempferol, kaempferol, liquiditi-
genin and garbanzol as substrates, and its activity was described in a previous study by our group (Marin et al., 2018).

For S. albus pDF, product titres were similar to S. albus pGR: β-coumaric acid (672.80 ± 19.34 μg l–1), dihydrokaempferol (47.32 ± 4.73 μg l–1), liquiritigenin (35.12 ± 0.007 μg l–1), naringenin (34.41 ± 2.17 μg l–1), isoliquiritigenin (17.22 ± 0.31 μg l–1), garbanzol (2.69 ± 0.19 μg l–1), kaempferol (1.30 ± 0.23 μg l–1), apigenin (0.25 ± 0.02 μg l–1), 2-hydroxynaringenin (0.19 ± 0.09 μg l–1), resokaempferol (0.12 ± 0.02 μg l–1) and 7,4'-dihydroxyflavone (0.002 ± 0.0003 μg l–1). Also, fustin (0.45 ± 0.10 μg l–1), dihydroquercetin (3.57 ± 1.27 μg l–1) and quercetin (0.07 ± 0.04 μg l–1) were detected due to F3'H activity (Fig. S3).

Discussion

Previous studies of our research group have demonstrated de novo production of several flavonoids derived from naringenin chalcone in S. albus (Fig. 1A; Marin et al., 2017, 2018; García-Gutiérrez et al., 2020). However, the present work describes de novo biosynthesis of flavonoids from a 6’-deoxychalcone produced by the activity of CHR. To our knowledge, flavonoid biosyn-
sis in the genus Streptomyces has focussed on naringenin or pinocembrin chalcone derivatives (Park et al., 2009, 2010, 2011), and heterologous production of isoliquiritigenin derivatives such as garbanzol, fustin or resokaempferol has not yet been described in these acti-
nomycetes (Rozmer and Perjási, 2016; Fig. 1B). To achieve an engineered production of garbanzol and fustin without supplementation of intermediates, the different genes that configure the biosynthetic pathway of these flavanones were cloned and expressed in the recombinant strains S. albus pGR and S. albus pDF, but different pathway intermediates and shunt products were hitherto detected (see Appendix S1; Fig. S3).

Six peaks consistent with a molecular formula of C15H12O5 were identified (Fig. 3A), with peaks 2 and 6 assigned as garbanzol and naringenin respectively. Hence, we hypothesized that peaks 1, 3, 4 and 5 could be isomers related to the biosynthetic pathway of gar-
banzol, and peak 1 (Fig. 3A) was putatively identified as 2-hydroxyliquiritigenin, an extremely rare natural product suggested to be an intermediate in the conversion of fla-
vanones to flavones (Jiang et al., 2019; Yonekura-
Sakakibara et al., 2019; Fig. 1B) The close relationship between garbanzol and putative 2-hydroxyliquiritigenin was evidenced when S. albus pGR strain was monitored over time because maximum intensities of both peaks switched over time (72 h and 168 h respectively; Fig. S5).

It was not possible to perform the structural elucidation of putative 2-hydroxyliquiritigenin by NMR due to problems related to its instability under the tested conditions. In fact, it is said that 2-hydroxyflavanones exist together with dibenzoylmethanes as tautomeric mixtures (keto-
and enol forms) in specific solvents such as DMSO and acetone, generating overlapping signals that cannot be assigned to one or the other tautomer with certainty (Stevens et al., 1999; Ti et al., 2011). Although NMR analysis generated a broad spectrum with more signals than would be expected for an individual molecule structure, peak 1 (in Fig. 3A) could be traced back based on the absence of signals from oxygenated methines in the mixture, as well as on its UV/vis and fragmentation spec-
trum (see Appendix S1). Unfortunately, 2-
hydroxyliquiritigenin was not commercially available, and its fragmentation profile was published in metabolomic databases such as FlavonoidSearch (Akimoto et al., 2017).

The existence of 2-hydroxyliquiritigenin in the crude extracts of S. albus pGR and S. albus pDF was completely unexpected because the introduced biosynthetic pathway lacks the enzymes needed to introduce a hydroxyl group in a position other than C3. In order to understand these surprising results, it is necessary to introduce the biosynthetic pathway of 2-
hydroxyliquiritigenin, which is produced from liquiritigenin by the activity of the cytochrome P450-dependent enzyme flavanone 2-hydroxylase (F2H, EC 1.14.14.162), which is unstable and generates licodione by spontaneous hemiacetal opening (Aoki et al., 2000). Alterna-
tively, 2-hydroxyliquiritigenin gives rise to 7,4'-
dihydroxyflavone, either by acid treatment or by the action of a dehydratase (Fig. 1B; Akashi et al., 1998; Du et al., 2010). The hypothesis on the presence of this alternative route branching off from licorinitigenin was ver-
ified because 7,4'-dihydroxyflavone and licodione (ten-
tative identification, no commercial standard available) were detected in the crude extracts of S. albus pGR and S. albus pDF (Fig. S2B and C). Additionally, new flavo-
nols (resokaempferol and kaempferol) were identified as well as 2-hydroxy intermediates other than 2-
hydroxyliquiritigenin (2-hydroxy naringenin and putative 2-
hydroxydihydrokaempferol; Fig. S2A). The synthesis of flavones or 2-hydroxyflavanones was surprising because no F2H-, FNS- or flavonol synthase (FLS, EC 1.14.20.6)-encoding genes were expressed in either of these strains.

In order to discard the potential presence of a gene coding for a native F2H enzyme in the chromosome of
S. albus wild-type strain, a feeding experiment with 0.1 mM naringenin was carried out, but no 2-hydroxynaringenin was observed, ruling out a possible endogenous F2H activity in this bacterium. Moreover, when S. albus pGR was supplemented with naringenin, a 10-fold decrease of final garbanzol titre was observed and the 2-hydroxyliquiritigenin signal almost disappeared, whereas the amount of dihydrokaempferol and its derivatives increased. As it will be further discussed, we think that F3H has a F2H side activity because F3H is acting on naringenin substrate under feeding conditions, rather than on liquiritigenin to generate 2-hydroxyliquiritigenin (Figs 1 and S3).

Interestingly, F2H activity can also be obtained by other enzymes such as flavone syntheses (FNSs): FNSI (soluble Fe³⁺/2-oxoglutarate-dependent dioxygenase, EC 1.14.20.5) and FNSII (NADPH-dependent cytochrome P450 membrane-bound monoxygenase, EC 1.14.19.76; Martens and Mithöfer, 2005). They are responsible for catalysing the conversion of flavanones to flavones by introducing a double bond between C2 and C3 in the heterocycle (Britsch, 1990; Zhang et al., 2007; Jiang et al., 2019), but some FNSII can also exhibit F2H activity, being required the activity of an additional unknown dehydratase to generate the flavone in planta (Akashi et al., 1999; Du et al., 2010; Wu et al., 2016). This said, the attention then focused on F3H as it is a 2-oxoglutarate-dependent oxygenase closely related to FNSI, being the only enzyme in the cloned pathway being able to catalyse the hydroxylation of the heterocycle. We believe that there is a relationship between F3H and FNSI, which could explain an additional unknown dehydratase to generate the flavone by introducing a double bond between C2 and C3.

Therefore, the close evolutionary relationship between F3H and FNSI, along with the F2H activity described for an FNSI, leads us to hypothesize that our F3H (encoded in pGR and pDF recombinant vectors) can have an F2H side activity allowing the formation of 2-hydroxyliquiritigenin and 2-hydroxynaringenin (spectrum signals coming from putative 2-hydroxydihydrokaempferol were also detected, but no commercial standard was available). Finally, an endogenous S. albus dehydratase would be responsible for generating the corresponding flavones (apigenin and 7,4′-dihydroxyflavone) or flavonols (kaempferol or resokaempferol) from its 2-hydroxy precursors because FNSI (as well as FNSII) does not accept 2-hydroxy derivatives as substrates in vitro (Zhang et al., 2007; Han et al., 2014; Fig. 1). Our hypothesis involving a native S. albus dehydratase is similar to what has been observed in cultures of Saccharomyces sp. supplemented with naringenin because the heterologous expression of a gene encoding the Medicago truncatula FNSII led to flavone production in vivo, but it only yielded 2-hydroxynaringenin in vitro (Akashi et al., 1999; Zhang et al., 2007). The side activity of some flavonoid enzymes is not entirely new; in fact, it has been described that some FLSs present F3H activity, despite the low homology that exists between both enzymes (Prescott et al., 2002; Duan et al., 2017; Park et al., 2019; Sun et al., 2019).

The strain S. albus J1074 has been widely used as a workhorse for heterologous production of a wide range of natural products over the last two decades (Lombó et al., 2006; Myronovskyi et al., 2016; Liu et al., 2018; Fazal et al., 2020) and could be regarded as a promising platform for heterologous flavonoid expression, but multiple genome improvements are needed. Firstly, improvements focused on increasing the intracellular availability of the main bottleneck in flavonoid biosynthesis (malonyl-CoA) would be necessary (Takamura and Nomura, 1988; Kurth et al., 2009; Santos et al., 2011), as well as limiting its use by other native biosynthetic clusters (Myronovskyi et al., 2018). On the other hand, the fact that a negative control strain supplemented with naringenin generated apigenin and eriodictyol (see Appendix S1) makes necessary to identify and to silence the promiscuous activities (Verdel-Aranda et al., 2015; Esnault et al., 2017). Deletion of endogenous catabolic pathways responsible for the degradation of L-Tyr would also increase the cytoplasmic pool of building blocks for flavonoid biosynthesis in this strain (Jiang et al., 2005; Kim et al., 2018). All these efforts will consolidate the use of S. albus as a highly attractive host for production of flavonoids.
In conclusion, our experiments demonstrated the functional introduction of 4 different biosynthetic pathways branching off from naringenin or isoliquiritigenin into the actinomycete S. albus. When using naringenin as a substrate, apigenin (2-hydroxynaringenin as an intermediate), kaempferol (dihydrokaempferol and 2-hydroxydihydrokaempferol as intermediates) and quercetin (kaempferol as an intermediate) were obtained. In a related approach, resokaempferol (garbanzol as an intermediate) were biosynthesized from liquiritigenin (Fig. 1). The functional introduction of a pathway leading to putative licodione, fustin (garbanzol as an intermediate) were biosynthesized from liquiritigenin (Fig. 1). The functional introduction of a pathway leading to putative licodione, 2-hydroxyliquiritigenin or 7,4′-dihydroxyflavone has never been reported in a microbial host before. The competitive effects between the four branches of this complex biosynthetic pathway, in which naringenin and liquiritigenin both serve as substrates of F3H, can explain the high structural diversity of flavonoids achieved in this bacterial factory as well as the low titres of the expected end-products.

Experimental procedures

Bacterial strains, plasmids and culture conditions

All strains and plasmids used in this study are listed in Table 1. E. coli TOP10 (Invitrogen), pSL1180 (Brosius, 1989) and pUC57 (Fermentas) were used for routine subcloning. The strain Streptomyces albus J1074, a mutant of the S. albus G strain that lacks an active SalI restriction modification system, was used for the production of flavonoids (Chater and Wilde, 1980). The high copy number E. coli–Streptomyces shuttle vector pIAGO, which harbours the strong constitutive promoter of ermE* (PermE*), was chosen as the expression plasmid (Aguirrezabalaga et al., 2000) as this assures high expression levels in the host. This vector harbours the plJ101 origin of replication from S. lividans ISP5434, which assures around 300 copies per chromosome, or 30% of the total strain DNA (Kieser et al., 2000). The promoter PermE is derived from the promoter region of the erythromycin resistance gene (ermE) of the erythromycin producer Streptomyces erythraeus (Bibb et al., 1985), which actually contains two different promoters, PermE1 and PermE2, and the PermE* promoter is a stronger variant containing the PermE2 and a TGG deletion in the -35 region of the PermE1 part.

E. coli strains were grown in tryptic soy broth (TSB, VWR) or on TSB agar plates, supplemented with the corresponding antibiotic (ampicillin 100 μg ml⁻¹, Sigma Aldrich, Madrid, Spain) for maintenance of the plasmid. S. albus J1074 was grown at 30°C in yeast extract–malt extract (YEME) 17% (w/v) sucrose for the preparation of protoplasts. It was sporulated in Bennett medium (Kieser et al., 2000) and supplemented with the corresponding antibiotics, when necessary (thiostrepton 50 μg ml⁻¹, Cayman Chemical, Ann Arbor, MI, USA).

For flavonoid production, S. albus spores were quantified, and a pre-inoculum of 10⁷ spores ml⁻¹ was transferred into Bennett medium which was first incubated for 72 h until sporulation and subsequently transferred to 30 ml of liquid RSA medium (Fernández et al., 1998). The cultures were incubated for 84 h at 30°C and 250 revolutions per minute (rpm). In the feeding experiments, 0.1 mM of naringenin was added after 24 h of incubation. When higher culture volumes for 2-hydroxyliquiritigenin purification were needed, the volume was scaled up to 6 l. In these cases, a pre-inoculum was made in liquid RSA medium with 10⁷ spores ml⁻¹, incubated for 45 h at 30°C and 250 revolutions per minute.
250 r.p.m., and a 10% (v/v) inoculum size was finally used for 6 l final culture volume and incubated for 112 h at 30°C and 250 r.p.m.

For *S. albus* pGR growth and production curves, triplicate cultures of *S. albus* pGR were grown in shake flasks with 30 ml of RSA over a duration of 168 h. Samples were taken every 24 h per duplicate to determine biomass as dry cell weight as well as production titres of garbanzol, liquiritigenin and isoliquiritigenin.

Reagents and biochemicals

All solvents used for solid-phase extraction, HPLC analysis/purification and MS-based experiments were LC-MS grade from either Sigma-Aldrich or VWR Chemicals. Authentic standards for LC-HRESIMS quantification and molecule identification through LC-HRESIMS and LC-UV/vis were purchased from different suppliers: *p*-coumaric acid (Sigma Aldrich, USA), naringenin (Sigma Aldrich, USA), dihydrokaempferol (Sigma Aldrich, USA), kaempferol (Cayman Chemical, USA), apigenin ( Extrasynthese, Genay, France), 2-hydroxynaringenin (Ambinter, Orléans, France), isoliquiritigenin (Sigma Aldrich, USA), liquiritigenin (Tocris Bioscience, Bristol, UK), (2R,3R)-garbanzol (BioBioPha, Kunming, Yunnan, China), (2S,3S)-garbanzol (AnalytiCon Discovery, Potsdam, Germany), resokaempferol ( Extrasynthese), fustin (Synthems, France), isoliquiritigenin (Sigma Aldrich, USA), liquiritigenin (Tocris Bioscience, Bristol, UK), (2R,3R)-garbanzol (BioBioPha, Kunming, Yunnan, China), (2S,3S)-garbanzol (AnalytiCon Discovery, Potsdam, Germany), resokaempferol (Extrasynthese), fustin (Synthems, France), quercetin (Cayman Chemical) and dihydroquercetin (Sigma Aldrich).

Genes and enzymes

Recombinant DNA techniques were performed following standard protocols (Sambrook and Russell, 2001). Restriction enzymes were purchased from Takara Biochemicals, T4 DNA ligase from Thermo Scientific and DreamTaq DNA Polymerase from Thermo Scientific. Synthetic genes for the following ORFs were synthesized by GenScript after codon optimization: TAL from *Rhodobacter capsulatus* (Genbank accession no. WP_013066811), 4CL from *S. coelicolor* (Genbank accession no. NP_628552), CHS from *Glycine max* (Genbank accession no. L07647.1), CHR from *G. max* (Genbank accession no. X55730.1), CHI from *G. max* (Genbank accession no. AY959413.1), F3H from *Petroselinum crispum* (Genbank accession no. YR23248) and F3H from *A. thaliana* (Genbank accession no. Q9SD85). Genbank accession numbers LT629805.1, LT629806.1, LT629807.1, MW404307, LT629808.1, LT629809.1 and MG748610 belong to the synthetic genes encoding TAL, 4CL, CHS, CHR, CHI, F3H and F3H respectively. Compatible restriction sites were added at each the gene cassette end and ribosome-binding sites at the 5'-ends. The RBS sequence (in bold) for all genes was 5’-TCCCGTGAGGAGCAGC-3’.

All constructed plasmids described later were verified by restriction enzyme digestions and also by sequencing of the cloned regions. Recombinant *Streptomyces* clones were confirmed by PCR. Primers used to amplify the first two common genes included 5’-GTATCGAGCTGGA CATGAA-3’ as the forward primer and 5’-GGCGTCC ACGAGGTGC-3’ as the reverse primer.

Construction of pGR

The plasmid pGR contains the *ermE* promoter (PermE*) and the six genes encoding enzymes responsible for garbanzol biosynthesis. All synthetic gene cassettes were independently cloned in pUC57, and plasmids were named pLMF1 (pUC57 containing TAL gene), pLMF2 (4CL), pLMF3 (CHS), pLMF4 (CHR), pLMF5 (CHI) and pLMF-F3H (F3H) (Table 1). Additionally, the TAL gene was subcloned into vector pSL1180 using HindIII-BamHI (pLMF7) in the first step of the cloning strategy. The 4CL gene (from pLMF2) was cloned into pLMF7 as *Pst*I-BamHI gene cassette, generating pLMF8. The next step involved subcloning of the CHR gene cassette from pLMF4 into pLMF3 using EcoRI, yielding pLMF11. The correct orientation of each DNA fragment was confirmed after each step by restriction enzyme digestions and sequencing. The two gene cassettes from pLMF11 (CHS and CHR) were subcloned together into pLMF8 using *SacI*-BamHI in order to assemble the first four genes in a plasmid (pLMF14). Finally, the F3H gene was subcloned into pLMF5 (cut with EcoRV-BamHI) as an EcoRI (blunt-ended)-BamHI gene cassette, and the ligation product (harbouring the genes CHI and F3H) was subcloned into pLMF14 using XbaI-BamHI, resulting in pLMF18, which contains the six genes required for garbanzol biosynthesis. As the expression host was *Streptomyces*, a further subcloning step was required, in which the *BglII*-BamHI DNA fragment carrying the six genes was finally subcloned into the pIAGO plasmid (Fig. 2A), a derivative of the bifunctional replicative vector pWHM3, which contains the *ermE* promoter, giving rise to the final plasmid pGR.

Construction of pDF

The plasmid pDF contains the *ermE* promoter (PermE*) and the seven genes required for the biosynthesis of fustin. To obtain this plasmid, it was required to add one more gene to the previous plasmid pLMF18 constructed to produce garbanzol. This last gene, F3H, was cloned into pLMF18 as Dral-BamHI gene cassette, giving rise to pLMF19. The DNA fragment containing seven genes...
was subcloned into the vector plAGO to be further expressed in Streptomyces (Fig. 2B). The gene cassette was cloned as a BgIII-BamHI DNA fragment to obtain plasmid pDF (Table 1).

**Flavonoid extraction and LC-HRESIMS analysis**

*S. albus* J1074 clones harbouring pGR, pDF or plAGO (negative control) were incubated (three replicates for each strain) as described earlier. The recovery of flavonoids from the recombinant strains developed in this study was achieved by an organic extraction with acetone (cellular pellet) and ethyl acetate (culture supernatant). Briefly, 30 ml of culture medium was centrifuged at 10 000 r.p.m. for 5 min, and the biomass and the supernatant were extracted separately. Firstly, an equal volume of acetone was added to the pellet to facilitate the breaking of the mycelium and then a second extraction was performed with an equal volume of ethyl acetate. For the supernatant, two extractions were performed with the same volume of ethyl acetate. Finally, the organic phases were dried sequentially under vacuum in a rotavapor (RV 10 Digital, IKA) equipped with a vertical condenser maintained at −10°C (RC-10 Digital Chiller, VWR) and brought together in a single 10 ml glass vial.

For the identification of flavonoids using HPLC-HRESIMS, the dry extract obtained was reconstituted in 200 µl DMSO/MeOH 1:1 (v/v), and the samples were processed sequentially through 0.8 and 0.2 µm filters (Acrodisc®, Pall). Separation was performed in a UPLC system (Dionex Ultimate 3000, Thermoscientific, Madrid, Spain) equipped with an analytical RP-18 HPLC column (50 × 2.1 mm, Zorbax® Eclipse Plus, 1.8 µm, Agilent Technologies, Madrid, Spain) heated to 30°C, and a combination of distilled water (mobile phase A) and MeCN (mobile phase B), both acidified with 0.1% (v/v) of formic acid, was used. The analytes were eluted at a flow rate of 0.25 ml min⁻¹ in a 10–100% (v/v) gradient of MeCN under the following conditions: 0–1 min (10% B), 1–4 min (10–35% B), 4–5 min (35% B), 5–8 min (35–100% B), 8–10 min (100% B), 1011 min (100–10% B) and 11–15 min (10% B). The column effluent was directed to electrospray ionization mass spectrometry analysis (HPLC-ESI-MS) using an ESI-UHR-Qq-TOF Impact II spectrometer (Bruker España SA, Madrid, Spain) which acquired data in the negative ion mode, with a m/z range from 40 to 2000 Da. Data were analysed using Compass DataAnalysis 4.3 (Bruker). The obtained BPCs were extracted for the deprotonated ions of a set of flavonoids with a mass error range of 0.005 mmu (milli mass units), and the obtained EICs (extracted ion chromatograms) were compared with authentic commercial standards. When needed, flavonoids were quantified by comparing the peak area with that of a known amount of an authentic compound through a calibration curve. The production titres are given in µg/L, and the mean value was calculated from three biological replicates.

**Purification and elucidation of 2-hydroxyliquiritigenin**

In order to obtain sufficient amounts of 2-hydroxyliquiritigenin for structural elucidation, the production volume of *S. albus* pGR was scaled up, and the culture was subsequently extracted. The extraction method was followed as described earlier, with the only difference that the supernatant was extracted four times and a 2 l separating funnel was used to separate the organic from the aqueous phase. Finally, 4.03 g of dry residue were obtained, which was then subjected to HRESIMS-guided fractionation: a first phase of pre-fractionation by solid-phase extraction (SPE) was needed, followed by two HPLC-UV/vis purification rounds under semi-preparative conditions (see Appendix S1).

**Statistical methods**

A two-way ANOVA was used to compare cultivation data of *S. albus* pGR and negative control *S. albus* plAGO. Considering that each row represents a different incubation time, matched values were stacked into a sub-column, and a Sidák correction was used, which has a higher statistical power than Bonferroni correction. The alpha threshold and confidence level selected for data was of 0.05 (95% confidence level). The graphic representation of all the cultivation data was carried out using GraphPad Prism software (version 7, GraphPad Software, San Diego, CA, USA).

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Conflict of interest

The authors declare no conflict of interest.

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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.** MS spectra (first column), MS² spectra for the parent ion m/z 271.0535 [M-H]⁻ (second column) and MS² spectra for the in-source-formed species m/z 243.0591 [M-H]⁻ (third column). A-C, MS and MS² spectra of purified 2-hydroxyliquiritigenin. In B, daughter ions m/z' 253.0461 and m/z' 209.0554 were also detected but are not visible due to the scale; D-F, MS and MS² spectra of authentic garbanzol; and G-H, MS and MS² spectra of authentic butin.

**Fig. S2.** Overlaid EICs for several flavonoids, p-coumaric acid and chalcone liquiritigenin of S. albus pGR (B) and S. albus pDF crude extracts (C) compared with commercial standards (A). X-axis has been zoomed at 3.5–8 min. Y-axis has been modified to clearly show all peaks; hence, peak intensities cannot be compared. pCOU, p-coumaric acid; DHK, dihydrokaempferol; 7,4'-DHF, 7,4'-dihydroxyflavone; LIQ, liquiritigenin; ILQ, isoliquiritigenin; RSK, resokaempferol; 2OH-NAR, 2-hydroxynaringenin; API, apigenin; NRG, naringenin; KAE, kaempferol; ERI, eriodictyol; 2OH-LIQ, 2-hydroxyliquiritigenin (no commercial standard available); FUS, fustin; QRC, quercetin; DHQ, dihydroquercetin; GRB, garbanzol.

**Fig. S3.** Production titres in µg l⁻¹ of several flavonoids, chalcone isoliquiritigenin and p-coumaric acid in recombinant S. albus pGR and S. albus pDF strains as well as in S. albus plAGO-negative control strain. *: 7,4'-DHF was detected in S. albus pGR, S. albus pDF and S. albus pGR + 0.1 mM NRG but in LOQ (limit of quantification), as well as 2OH-NRG and RSK in S. albus pGR + 0.1 mM NRG. The mean value of compounds was calculated from three biological replicates. pCOU, p-coumaric acid; DHK, dihydrokaempferol; 7,4'-DHF, 7,4'-dihydroxyflavone; LIQ, liquiritigenin; ILQ, isoliquiritigenin; RSK, resokaempferol; 2OH-NAR, 2-hydroxynaringenin; API, apigenin; NRG, naringenin; KAE, kaempferol; FUS, fustin; QRC, quercetin; DHQ, dihydroquercetin; GRB, garbanzol.

**Fig. S4.** Comparison of main flavonoid intermediates as seen in the extracted supernatant (A) and cellular pellet (B) in S. albus pGR (overlaid EICs from HPLC-MS data). Same extracts concentrations were injected in the HPLC-MS equipment.

**Fig. S5.** EICs at m/z 271.06118 ± 0.005 [M-H]⁻ of S. albus pGR crude extract sampled every 24 h over 168 h. The scale of the Y-axis is fixed to the highest intensity value at 144 h.

**Appendix S1.** Supplementary material.