

MICROBIAL TRANSFORMATION OF 2'-PROPOXY ANALOGS OF (-)- AND (+)-
DEHYDROGRISOFULVIN AND (+)-2'-DEMETHOXYDEHYDROGRISOFULVIN BY
STREPTOMYCES CINEROCROCATUS

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Abstract----- By the fermentation of Streptomyces cinereocrocatus, 2'-propoxy analogs of (-)- and (+)-dehydrogrisofulvin were both converted into the corresponding analog of (+)-grisofulvin and the same treatment of (+)-2'-demethoxydehydrogrisofulvin afforded (+)-2'-demethoxygrisofulvin and (+)- and (-)-2'-demethoxy-2',3'-dihydrodehydrogrisofulvin, indicating that the (+)-substrates were isomerized into the corresponding (-)-enantiomers and subsequently transformed to the reduction products.

The microbial transformation of (-)-dehydrogrisofulvin to (+)-grisofulvin was initially investigated by Andres and his co-workers¹ using Streptomyces cinereocrocatus NRRL 3443. Since then, we have demonstrated that (-)- and (+)-dehydrogrisofulvin are both transformed mainly into (+)-grisofulvin by Streptomyces species including Streptomyces cinereocrocatus and the stereochemistry of the microbial reduction is successfully elucidated by ²H NMR spectroscopy.^{2,3}

We describe the following studies which clarify that the microbial transformations of 2'-propoxy analogs (1 and 2) of (-)- and (+)-dehydrogrisofulvin and (+)-2'-demethoxydehydrogrisofulvin (3) by Streptomyces cinereocrocatus take place directly or after isomerizations with hydrogenations depending on 2'-substituents of (-)- and (+)-dehydrogrisofulvin analogs. 2'-Propoxy analogs (1 and 2) were used as the substrates in place of 2'-ethoxy analogs, since in the microbial transformation the former will give more informations connected with the replacement of the 2'-methoxy group of (-)- and (+)-dehydrogrisofulvin than the latter.

Firstly, the substrates (1 and 2) were synthesized as follows. Reaction of 2'-propoxy analog (4)⁴ of (+)-grisofulvin in the presence of selenium dioxide in

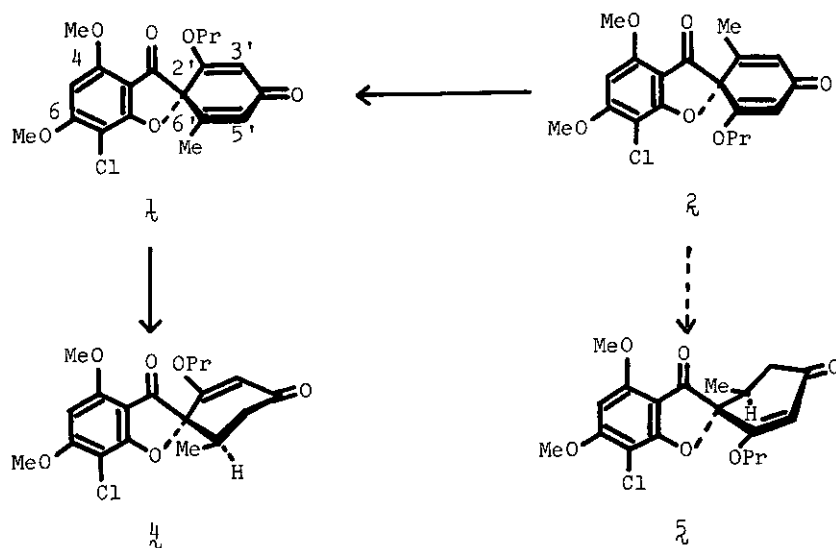
tert-butanol, followed by silica gel column chromatography afforded 2'-propoxy analog (λ) of (-)-dehydrogriseofulvin [PMR δ (CDCl₃) 0.79 (3H, t, J = 7 Hz, 2'-OCH₂CH₂CH₃), 1.60 (2H, sextet, J = 7 Hz, 2'-OCH₂CH₂CH₃), 1.79 (3H, bs, 6'-CH₃), 3.79 (2H, t, J = 7 Hz, 2'-OCH₂CH₂CH₃), 4.00 (3H, s, 4-OCH₃), 4.07 (3H, s, 6-OCH₃), 5.66 (1H, bs, 3'-H), 6.18 (1H, s, 5-H), 6.20 (1H, bs, 5'-H)].⁵ The same dehydrogenation reaction of 2'-propoxy analog of (+)-epigriseofulvin which was obtained by alkylation of (+)-epigriseofulvic acid with diazopropane afforded 2'-propoxy analog (ρ) of (+)-dehydrogriseofulvin, which showed the same PMR spectrum but exhibited opposite optical properties⁶ compared with those of λ . Catalytic hydrogenation of ρ yielded 2'-propoxy analog (ξ)⁷ of (-)-griseofulvin. In connection with the previous studies³, the microbial treatment of λ by *S. cinereocrocatus* under previously described conditions gave η as the reduction product and the recovered material, which were separated by silica gel column chromatography. On the other hand, the same microbial treatment of ρ was performed and its results were compared with those of the enantiomer (λ). The results indicate that reactions of λ proceed more rapidly than those of ρ by comparisons of the yields of the reduction product and the recovered material(s). Moreover, the comparisons of susceptibilities to microbial transformations between (-)- and (+)-dehydrogriseofulvin and their 2'-propoxy analogs indicate that the propoxy analogs are less

Table I. Yields of Reduction Products and Relative Ratios of (+)- and (-)-Enantiomers of Recovered Substrates

Substrates	Reduction products:	Recovered substrates:		
	Yields (%)	Yields (%)	Relative ratios (%) of	
			(+)-	(-)-
(-)-Dehydrogriseofulvin*	88	0	—	—
2'-Propoxy analog of (-)-dehydrogriseofulvin	17	24	0	100
(+)-Dehydrogriseofulvin*	31	15	100	0
2'-Propoxy analog of (+)-dehydrogriseofulvin	3	57	96	4
2'-Propoxy analog of (λ)-dehydrogriseofulvin	20	25	72	28

* The experiments using these substrates were performed as controls for the corresponding (-)- and (+)-2'-propoxy analogs. And their reduction products were the same optically pure (+)-griseofulvin.³

transformed by *S. cinereocrocatus* (see Table I).⁸ Furthermore, the relative ratios of these compounds clearly demonstrate that both substrates were transformed into the optically pure 2'-propoxy analog of (+)-griseofulvin in spite of a fact that recovered dehydrogriseofulvin analogs were a mixture of (+)- and (-)-enantiomers in the microbial treatment of 2'-propoxy analog ($\hat{2}$) of (+)-dehydrogriseofulvin. Further, it is of importance to notice that in the microbial treatment by *S. cinereocrocatus* 2'-propoxy analog ($\hat{2}$) of (+)-dehydrogriseofulvin was not transformed into the corresponding hydrogenated product ($\hat{5}$). These results are summarized in Scheme 1.

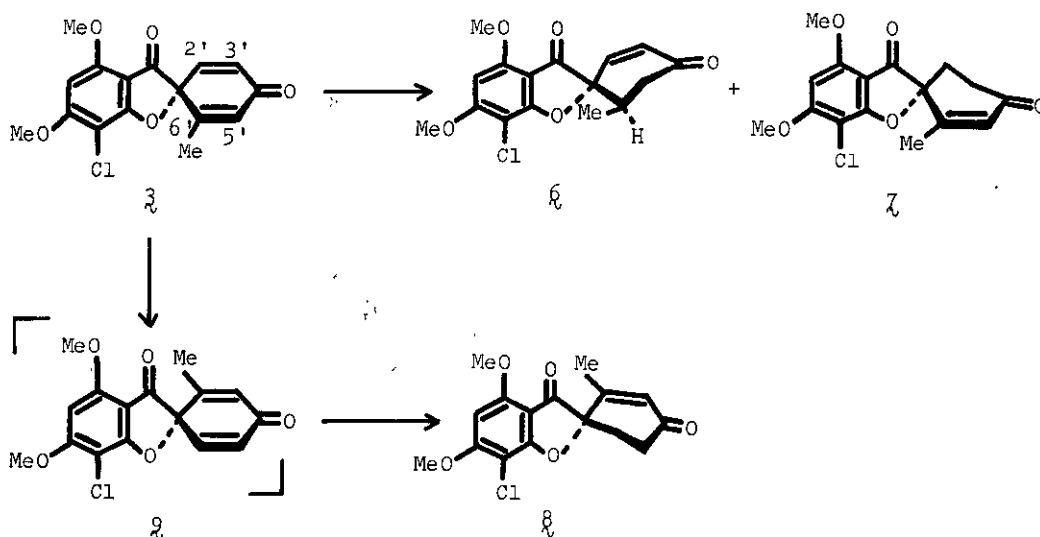


Scheme 1

In order to extensively elucidate the microbial transformation, (+)-2'-demethoxydehydrogriseofulvin was synthesized as follows. A solution of (+)-2'-demethoxygriseofulvin ($\hat{6}$)⁹ and pyridinium hydrobromide perbromide¹⁰ in chloroform was reacted under reflux for 2 hr to give a 40:60 mixture (g.l.c.) of 3'-bromo-2'-demethoxygriseofulvin¹¹ and 5'-bromo-2'-demethoxygriseofulvin¹², which was separated by repeated recrystallization from methanol and silica gel column chromatography. Subsequent dehydrobromination¹³ of the 5'-bromo derivative with LiCl and Li₂CO₃ in DMF containing pyridine at 100°C for 24 hr yielded (+)-2'-demethoxydehydrogriseofulvin ($\hat{3}$)¹⁴ [PMR δ (CDCl₃) 1.82 (3H, bs, 6'-CH₃), 3.98 (3H, s, 4-OCH₃), 4.08 (3H, s, 6-OCH₃), 6.18 (1H, s, 5-H), 6.29 (1H, m, 3'-H), 6.42 (1H, d, J = 2 Hz, 5'-H), 6.53 (1H, d, J = 10 Hz, 2'-H)].

The microbial treatment of (+)-2'-demethoxydehydrogriseofulvin ($\hat{3}$) by *S. cinereo-*

crocatus for 12 hr under the same conditions described above afforded (+)-2'-demethoxygriseofulvin (**6**) (12%) and a mixture of (-)- and (+)-2'-demethoxy-2',3'-dihydrodehydrogriseofulvin (**7** and **8**)¹⁵ (8%), whose relative ratio was calculated as 19:81 from the value¹⁶ of its circular dichroism. When the incubation period was shortened by 3 hr, **6** and a mixture¹⁷ of **7** and **8** were obtained in 3 and 8% yields, respectively, with 52% yield of the recovered **3**. These results are summarized in Scheme 2, which indicates that the microbial reductions of **3** proceed more preferentially in 5',6'- than 2',3'-double bond. Furthermore, the formation of (+)-2'-demethoxy-2',3'-dihydrogriseofulvin (**8**) suggests that the microorganism has the abilities of the isomerization of the substrate (**3**) into the enantiomer (**9**) and of the subsequent reduction of the latter.



Scheme 2

Hence, we conclude that in the treatment with S. cinereocrocatus, the analogs of (-)- and (+)-dehydrogriseofulvin which have or have not substituents at 2'-position are reduced directly or after isomerization into the corresponding enantiomers, yielding (+)- and/or (-)-dihydro compound(s) as the transformation products.

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4. L. A. Duncanson, J. F. Grove, and P. W. Jeffe, J. Chem. Soc., 1958, 2929.
5. The molecular ellipticity $[\theta]$ (c 1.0 mg/ml, CHCl_3): $[\theta]_{365}$ -560, $[\theta]_{335}$ -7000, $[\theta]_{328}$ -5600, $[\theta]_{298}$ -28350, $[\theta]_{290}$ 0, $[\theta]_{280}$ +19600, $[\theta]_{260}$ 0, $[\theta]_{257}$ -2100, $[\theta]_{255}$ 0, $[\theta]_{245}$ +18200, $[\theta]_{241}$ 0, $[\theta]_{235}$ -40600; $[\alpha]_D^{21}$ -57.5° (c 0.56, acetone).
6. The molecular ellipticity $[\theta]$ (c 1.0 mg/ml, CHCl_3): $[\theta]_{365}$ +490, $[\theta]_{335}$ +7000, $[\theta]_{328}$ +5600, $[\theta]_{298}$ +26600, $[\theta]_{290}$ 0, $[\theta]_{280}$ -19600, $[\theta]_{260}$ 0, $[\theta]_{257}$ +1750, $[\theta]_{255}$ 0, $[\theta]_{245}$ -17500, $[\theta]_{241}$ 0, $[\theta]_{235}$ +39900; $[\alpha]_D^{21}$ +56.0° (c 0.51, acetone).
7. PMR and mass spectra were identical with those of λ . However, the CD and optical rotation showed the opposite values.
8. The microbial transformations of (-)- and (+)-dehydrogriseofulvin and its 2'-propoxy analogs were performed in the incubation periods of 5 hr, one day, 3 days, and 5 days. Table I shows, however, the results of 3-day's incubations.
9. T. P. C. Mulholland, J. Chem. Soc., 1952, 3994.
10. C. Djerassi and C. R. Scholz, J. Am. Chem. Soc., 1948, 70, 417.
11. PMR (CDCl_3) δ 0.92 (3H, d, J = 6 Hz, 6'- CH_3), 2.5-3.3 (3H, m, 5' α , 5' β and 6' α -H), 3.97 (3H, s, 4- OCH_3), 4.02 (3H, s, 6- OCH_3), 6.13 (1H, s, 5-H), 7.01 (1H, s, 2'-H).
12. PMR (CDCl_3) δ 1.13 (3H, d, J = 6 Hz, 6'- CH_3), 3.03 (1H, d.d, J = 6 and 13 Hz, 6' α -H), 3.97 (3H, s, 4- OCH_3), 4.03 (3H, s, 6- OCH_3), 5.27 (1H, d, J = 13 Hz, 5' β -H), 6.16 (1H, s, 5-H), 6.29 (1H, d, J = 10 Hz, 3'-H), 6.63 (1H, d, J = 10 Hz, 2'-H).
13. R. P. Holysz, J. Am. Chem. Soc., 1953, 75, 4432.
14. The molecular ellipticity $[\theta]$ (c 1.0 mg/ml, CHCl_3): $[\theta]_{370}$ -480, $[\theta]_{343}$ -3360, $[\theta]_{333}$ -800, $[\theta]_{330}$ -900, $[\theta]_{327}$ 0, $[\theta]_{300}$ +9600, $[\theta]_{284}$ 0, $[\theta]_{270}$ -5440, $[\theta]_{263}$ 0, $[\theta]_{255}$ +6720, $[\theta]_{248}$ +4800, $[\theta]_{238}$ +43520.
15. Authentic sample of λ was synthesized by dehydrogenation of (-)-2'-demethoxy-dihydrogriseofulvin(cf. A. W. Dawkins and T. P. C. Mulholland, J. Chem. Soc., 1959, 1826) with selenium dioxide in tert-butanol. Compound λ : PMR (CDCl_3) δ 1.80 (3H, bs, 6'- CH_3), 2.3-2.8 (4H, m, 2'- and 3'-H), 4.03 (3H, s, 4- OCH_3),

4.07 (3H, s, 6-OCH₃), 6.14 (1H, bs, 5'-H), 6.20 (1H, s, 5-H); The molecular ellipticity [θ] (c 1.0 mg/ml, CHCl₃): [θ]₃₇₀ -190, [θ]₃₃₆ -21670, [θ]₃₃₃ -21410, [θ]₃₂₂ -31560, [θ]₃₁₁ -26890, [θ]₂₉₁ 0, [θ]₂₆₄ +17710, [θ]₂₄₂ 0, [θ]₂₃₄ -128800. The comparison of CD data of related compounds suggests that the conformation of λ is as shown in Scheme 2.

16. The molecular ellipticity [θ] (c 1.0 mg/ml, CHCl₃): [θ]₃₇₀ 0, [θ]₃₃₆ +12940, [θ]₃₃₃ +12690, [θ]₃₂₂ +18890, [θ]₃₁₁ +16100, [θ]₂₉₁ 0, [θ]₂₆₄ -700, [θ]₂₄₂ 0, [θ]₂₃₄ +40870.
17. The relative ratio of λ and δ was 28:72 and the recovered λ was optically pure on the basis of their CD data.

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