Sexual Activation of Carotenogenesis in *Phycomyces blakesleeanus*

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Sexual activation increases the β-carotene content of the fungus *Phycomyces blakesleeanus*. The same effect is observed in single cultures exposed to natural and synthetic trisporates and in intersexual heterokaryons. Synthetic racemic (9Z)-methyl trisporate B, a trisporate precursor made by cultures of the (+) mating type, stimulates carotenogenesis only in (−) cultures. Synthetic racemic (9Z)-methyl trisporate C is less effective and the corresponding all-(E) isomers and other related compounds are inactive. Sexual stimulation of carotenogenesis is additive with the stimulations induced by light, retinol, dimethyl phthalate, and mutation of the gene *carS*.

INTRODUCTION

Several external stimuli accentuate the yellow colour of the fungus *Phycomyces blakesleeanus* by increasing its β-carotene content (review by Cerda-Olmedo & Torres-Martinez, 1979). The main active agents are light (Bergman et al., 1973; Jayaram et al., 1979; López-Díaz & Cerda-Olmedo, 1980), sexual interaction, and various chemicals, including β-ionone (Mackinney et al., 1952), retinol (Eslava et al., 1974), dimethyl phthalate and others (Cerda-Olmedo & Hüttermann, 1986).

There are several kinds of mutants with defective β-carotene synthesis. Some of them block specific reactions of the pathway: *carR* mutants are red and accumulate lycopene, *carB* mutants are white and accumulate phytoene. A high constitutive level of β-carotene is found in *carS* mutants. Most *carA* mutants are white and contain only little β-carotene, but become yellow on addition of retinol. Mutations having little or no effect on the basal β-carotene content but rendering the mycelia insensitive to the action of retinol and other chemicals occur at genes *carI* and *carA*. For references on these mutants see Table 1 and the review by Cerda-Olmedo (1985).

There are two mating types or sexes in *Phycomyces*, called (+) and (−) (Blakeslee, 1904). Hyphae of opposite sex growing near each other have a pronounced yellow colour (Blakeslee, 1904; Hocking, 1965). Trisporic acids, found in mated cultures of the Mucorales, including *Blakeslea trispora* (Caglioti et al., 1966; Sutter & Whitaker, 1981) and *P. blakesleeanus* (Sutter, 1977), stimulate carotene production and sexual morphogenesis when added to single cultures. Trisporic acids are made from β-carotene (Austin et al., 1969) via sex-specific precursors (Sutter et al., 1974). The methyl trisporates are produced by the (+) mycelia and converted to trisporic acids by the (−) mycelia (Sutter & Whitaker, 1981). Intersexual heterokaryons, whose mycelia mix nuclei of both sexes (Burgeff, 1914), form many bizarre morphological structures, called pseudophores, but few regular sporangiophores. Intersexual heterokaryons have more β-carotene than comparable homokaryons whose nuclei belong all to the same sex (Murillo et al., 1978).

We have investigated the genetic requirements for the sexual activation of carotenogenesis and the relations between sexual and other stimuli. Sexual stimulation was achieved by adding...
synthetic methyl trisporates or natural trisporic acids to single (−) cultures of various genotypes or by constructing intersexual heterokaryons.

METHODS

The strains of Phycomyces blukesleeanus Burgeff used in this work are listed in Table 1. The auxotrophic strains A87 and B36 were given to us by Professor A. P. Eslava, Universidad de Salamanca, Spain; they require L-lysine (200 mg l⁻¹) and nicotinic acid (1 mg l⁻¹), respectively. C strains came from the collection of the late Professor M. Delbrück at the California Institute of Technology; the mutation mad-107 affects the late reactions of phototropism and has presumably no effect on carotenogenesis. S strains originated in this laboratory. Blakelea\n
activation trispora NRRL 2895, mating type (+), and NRRL 2896, mating type (−), and the standard P. blukesleeanus wild-type NRRL 1555 came from the Agricultural Research Service, Department of Agriculutre, Peoria, Ill., USA. Heterokaryons are designated by placing an asterisk (*) between the symbols of the constituent homokaryons.

Activated (48 °C, 15 min) P. blukesleeanus spores were inoculated onto plates containing 25 ml glucose minimal medium (Sutter, 1975) with L-asparagine. H₂O (2 g l⁻¹) and agar (15 g l⁻¹) and incubated at 22 °C for 4 d in the dark (if not otherwise stated). Experiments with heterokaryons, and their constitutive homokaryons as controls, were started with mycelial pieces, about 2 mm in width, taken from 2-d-old colonies. Heterokaryons were made by the microsurgical technique of Ootaki (1973).

Methyl trisporates and their analogues (Compounds I–VIII, Fig. 4) were dissolved in ethanol and spread over agar plates preincubated and preincubated for 24 h. Each place received 0.1 ml ethanol. Dimethyl phthalate (dimethyl ester of 1,2-benzenedicarboxylic acid) was dissolved in ethanol and added to the molten medium before the plates were poured; the final concentrations were 2 mmol dimethyl phthalate l⁻¹ and 1% (v/v) ethanol. Retinol acetate (Sigma) was dissolved in a mixture of absolute ethanol and polyoxyethylene sorbitan mono-oleate (Tween 80, Sigma), then added to 1 l agar medium before the plates were poured; the final concentrations were: retinol acetate, 0.75 mmol l⁻¹ (250 µg ml⁻¹); ethanol, 0.1% (v/v); and Tween 80, 0.4% (v/v).

The methods for the production, purification and assay of trisporates from mated B. trispora cultures were described by Govind & Modi (1981) and Sutter (1970), respectively. Most of the trisporic acids used in these experiments came from cultures started with fragments of mycelia of the two sexes and grown for 6 d in liquid malt medium; the yield was about 15.7 mg trisporic acids l⁻¹. In minimal medium the yield was considerably smaller.

For the determination of carotenoids, 4-d-old mycelia were peeled from the plates, freed from any visible bits of agar, and lyophilized. A weighed sample was extracted with petroleum ether (b.p. 40–60 °C), vacuum-dried, and resuspended in n-hexane. β-Carotene, lycopene and phytoene were quantified from their absorption coefficients (Davies, 1965). Phytoene was previously purified chromatographically on aluminum oxide (Davies, 1965). The results shown are means of three independent experiments, unless otherwise stated.

RESULTS

Activation of the wild-type by trisporates and related compounds

The trisporate preparation from B. trispora activated carotenogenesis (Fig. 1). The maximal effect was observed with 10 µg trisporates ml⁻¹; higher amounts were less effective. The dry weight of 4-d-old mycelia – 317 ± 23 mg (SD) per Petri plate - was not affected by the addition of the preparation.

Synthetic (9Z)-methyl trisporates stimulated carotene accumulation in P. blukesleeanus cultures of the (−) mating type (Figs 2 and 3; structural formulae and names in Fig. 4). The B form was clearly more effective than the C form. Little or no effect was shown by the corresponding all-(E) isomers and other synthetic compounds (V–VIII) structurally related to the trisporic acids. These eight compounds were not toxic to P. blukesleeanus under the conditions of the experiments: the mycelial dry weight was not affected by concentrations of up to 40 µg ml⁻¹ of any of these compounds.

While the natural trisporates from B. trispora stimulated carotenogenesis in both (+) and (−) wild-types, compound I did not appear to be effective on the (+) strain B36 (Table 1).

Activation of car mutants

Representative strains carrying various car mutations were exposed to natural trisporates from B. trispora and to racemic (9Z)-methyl trisporate B (Table 1). Carotene production by the wild-type was activated by both agents, as expected from results given above. The same was found with the carS mutant. The chemoinensitive mutants S119 and S144 responded to neither
Activation of carotenogenesis in Phycomyces

Fig. 1. Effect of trisporates from mated B. trispora cultures on the accumulation of β-carotene in the standard P. blakesleeanus wild-type. The bars indicate standard errors.

Fig. 2. Effect of methyl trisporates (compounds I–IV, see Fig. 4) on the accumulation of β-carotene in the standard P. blakesleeanus wild-type. ●, Compound I; ○, compound II; △, compound III; Δ, compound IV.

Fig. 3. Effect of compounds V–VIII (see Fig. 4) on the accumulation of β-carotene in the standard P. blakesleeanus wild-type. ●, Compound V; ○, compound VI; △, compound VII; Δ, compound VIII.

Table 1. Carotene content of various P. blakesleeanus strains grown for 4 d in the presence of synthetic racemic (9Z)-methyl trisporate B (compound I, 40 μg ml⁻¹) or natural trisporates from B. trispora (10 μg ml⁻¹)

The table gives the concentration (μg per g dry weight) of the major carotene in each strain, which was β-carotene except in strains C9 (lycopene) and C173 (phytoene) (results marked †).

<table>
<thead>
<tr>
<th>Strain</th>
<th>Genotype and references</th>
<th>Control</th>
<th>With natural trisporates</th>
<th>With compound I</th>
</tr>
</thead>
<tbody>
<tr>
<td>NRRL 1555</td>
<td>Standard wild-type (―)</td>
<td>54</td>
<td>296</td>
<td>300</td>
</tr>
<tr>
<td>C2</td>
<td>carA5 (―) (Murillo et al., 1981)</td>
<td>2</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>C9</td>
<td>carR21 (―) (de la Guardia et al., 1971)</td>
<td>1723†</td>
<td>2748†</td>
<td>2006†</td>
</tr>
<tr>
<td>C173</td>
<td>carB32 carR21 (―) (Presti et al., 1977)</td>
<td>1503†</td>
<td>1535†</td>
<td>1447†</td>
</tr>
<tr>
<td>C115</td>
<td>carS42 mad-107 (―) (Murillo &amp; Cerdá-Olmedo, 1976)</td>
<td>2609</td>
<td>4277</td>
<td>4417</td>
</tr>
<tr>
<td>S119</td>
<td>carA113 (―) (Roncero &amp; Cerdá-Olmedo, 1982)</td>
<td>28</td>
<td>30</td>
<td>31</td>
</tr>
<tr>
<td>S114</td>
<td>car1131 (―) (Roncero &amp; Cerdá-Olmedo, 1982)</td>
<td>29</td>
<td>27</td>
<td>38</td>
</tr>
<tr>
<td>S218 • S219</td>
<td>See Murillo et al. (1978)</td>
<td>14062</td>
<td>14137</td>
<td>15317</td>
</tr>
<tr>
<td>B36</td>
<td>nicA101 (―) (Alvarez &amp; Eslava, 1983)</td>
<td>52</td>
<td>105</td>
<td>38</td>
</tr>
<tr>
<td>A87</td>
<td>lysA401 (―)</td>
<td>64</td>
<td>201</td>
<td>223</td>
</tr>
<tr>
<td>B36 • A87</td>
<td>nicA101 (―) • lysA401 (―)</td>
<td>269</td>
<td>252</td>
<td>249</td>
</tr>
</tbody>
</table>
Fig. 4. Structural formulae of compounds used in this work: racemic (9Z)-methyl trisporate B (I), racemic (9Z)-methyl trisporate C (II), racemic all-(E)-methyl trisporate B (III), racemic all-(E)-methyl trisporate C (IV), 4-oxo-β-ionone (V), racemic 3-acetoxy-4-oxo-β-ionone (VI), 4-hydroxy-4-[(3Z)-5-hydroxy-3-methyl-1,3-pentadienyl]-3,5,5-trimethyl-2-cyclohexen-1-one (VII), and 8-[2,6,6-trimethylcyclohexenyl]-2,6-dimethyl-octatrien(2,4,6)-al(1) (VIII). Mated B. trispora cultures contain mainly trisporates B (IX) and C (X) (Sutter et al., 1974).

Table 2. Synergism between the effects of intersexual heterokaryosis and those of light, retinol, and dimethyl phthalate on carotenogenesis

The table gives the concentration of β-carotene (μg per g dry weight) in 4-d-old cultures. The tests of the chemicals and their controls were incubated in the dark.

<table>
<thead>
<tr>
<th>Strain</th>
<th>White light (W m⁻²)</th>
<th>Retinol acetate (mM)</th>
<th>Dimethyl phthalate (mM)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0</td>
<td>0.43</td>
<td>0</td>
</tr>
<tr>
<td>B36</td>
<td>66</td>
<td>321</td>
<td>62</td>
</tr>
<tr>
<td>A87</td>
<td>66</td>
<td>306</td>
<td>62</td>
</tr>
<tr>
<td>B36 * A87, mycelium 1</td>
<td>118</td>
<td>594</td>
<td>102</td>
</tr>
<tr>
<td>B36 * A87, mycelium 2</td>
<td>181</td>
<td>695</td>
<td>144</td>
</tr>
<tr>
<td>B36 * A87, mycelium 3</td>
<td>289</td>
<td>776</td>
<td>271</td>
</tr>
<tr>
<td>B36 * A87, mycelium 4</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
the natural nor the synthetic compounds. The stimulation of the carotene pathway by trisporic acids seemed to require the presence of β-carotene in the receiving cells: the amount of major carotene (phytoene) of strain C173, which is totally devoid of β-carotene, was not affected by trisporic acids, but the major carotenoids of strains C2 (β-carotene) and C9 (lycopene), which both produce some β-carotene, were appreciably increased.

**Regulation of carotenogenesis in intersexual heterokaryons**

Intersexual heterokaryons contained up to five times more β-carotene than either of the constituent homokaryons (Tables 1 and 2). The different carotene contents of different heterokaryotic mycelia depend presumably on their nuclear proportions. The intersexual heterokaryons S218 • S219 and B36 • A87 were not further stimulated by either trisporates or methyl trisporates (Table 1). Intersexual heterokaryosis thus appears to provide a saturating sexual stimulation. Cultures grown in the light or in the presence of retinol or dimethyl phthalate contained more β-carotene than those grown in the dark, irrespective of sex and intersexual heterokaryosis (Table 2). The emulsifiers needed for the application of the chemicals, present in the corresponding controls, had no effect by themselves.

**DISCUSSION**

Trisporates are relatively ineffective activators of carotene biosynthesis, if we compare inputs and outputs. Thus, to increase the carotene content of a Petri-dish culture from 17 to 90 μg one has to add 1 mg of the most active synthetic compound, (9Z)-methyl trisporate B (compound 1). The trisporate preparations from *B. trispora* produce a similar stimulation. The carotenogenic activity of the sex pheromones under physiological conditions does not appear to be very high either: mated cultures and intersexual heterokaryons (Murillo & Cerda-Olmedo, 1976; our results) contain about the same carotene concentrations as those obtained with exogenous trisporates.

The chemoinsensitive mutants (strains S119 and S144) are insensitive to stimulation by either synthetic methyl trisporates or natural trisporates (Table 1) and by retinol and dimethyl phthalate (unpublished). This establishes a common step in the regulation of carotenogenesis by all of these chemicals.

The sexual stimulation of carotenogenesis is additive with those induced by retinol, dimethyl phthalate, light, and mutation at gene *carS*. This synergism of *carS* mutations and sexual interaction had been observed by Murillo & Cerda-Olmedo (1976) and Murillo *et al.* (1978). The additivity indicates that at least the rate-limiting steps in the stimulation of carotenogenesis by sexual interaction are different from those of the other mechanisms of stimulation.

Sexual stimulation must thus have some common steps with the stimulation of carotenogenesis by retinol and dimethyl phthalate, but be separate from them at least at the respective limiting steps.

Particularly intriguing is the fact that the stimulatory action of trisporates apparently required β-carotene, as if the carotene itself or a product of its metabolism were involved in the chain of events leading to increased carotenogenesis. It may be that the trisporates by themselves have no effect on carotenogenesis, but only derepress the formation of trisporates from β-carotene (Bu'lock, 1976): an intermediate compound in the pathway from β-carotene to trisporate would then be the true activator of carotenogenesis.

The lack of activity of methyl trisporates on the (+) strain suggests that these compounds are only effective when converted to trisporates. Methyl trisporates are believed to be made only by the (+) strains and converted to trisporates by the (−) strains (Sutter & Whitaker, 1981).

The inactivity of compounds V–VIII (Table 1) is somewhat surprising, in view of their resemblance to the trisporic acids and to other aromatic compounds which have turned out to be potent activators of carotenogenesis. This is only a particular case of a more general problem: the relationships between chemical structure and activation of carotenogenesis are still unclear (Cerdá-Olmedo & Hüttermann, 1986).
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REFERENCES


