

Catcheside, D. E. A. A new carotenoid

mutant of *Neurospora*.

A mutant, MN58p, found amongst the survivors of UV-irradiated conidia, is pigmented somewhat like ad-3 strains grown on limiting adenine. The 'purple' pigmentation of MN58p is due to a lock of neurosporaxanthine and at least two yellow carotenoids which are, as yet, unidentified. The new mutant is epistatic to y1o-1 (Y30539y).

Crosses of MN58p to markers on linkage groups II (pe), III (try-1), IV (lys-5, try-2) and VII (nit-3), failed to detect linkage (Table 1). However, MN58p proved to be closely linked to nic-1, aur, al-2 and alY on linkage group IR. Failure to detect a single recombinant amongst 5,742 progeny of a cross of al-2 (15300) suggests that MN58p may be on allele of al-2.

A forced heterocaryon (A, MN58p; me-5 (86304)) + (A, al-2 (15300); cot-1 (C102); am (47305). hist-1 (K83)) was found to be phenotypically MN58p, although the purple pigmentation is less intense than in the MN58p parent, and not wild type. A heterocaryon (A, MN58p; lys-5 (DS6-85) y1o-1 (Y30539y); nit-3 (Y31881)) + (fr (B110), A, our (34508)), phenotypically fr⁺ and able to grow on minimal medium, has a normal distribution of pigment, unlike the our parent, which has pigmented conidia but locks colored polyenes in the mycelium, and is wild type in color. MN58p complements our (34508) but not al-2 (15300).

The ability of MN58p to complement various mutants deficient in carotenoids has also been investigated by examining the color phenotype of the escape growth of the initially slow-growing progeny, containing a duplication, of crosses between normal sequence and a translocation T(IR:VL) AR190 (Table 2). Wild type pigmentation was observed amongst the progeny of crosses of MN58p with alY (ALS4) and our (34508), but not with al (al^s), al (K96) or al-2 (15300).

As MN58p complements and recombines with our (34508) and alY (ALY), it is probable that MN58p is in a gene separate from these two mutations. However, as no complementation is observed with al-2 (15300), al (al^s) and al (K96), it would seem that MN58p is allelic to these mutations, although it has a phenotype distinct from each of these albino mutants, which fail to produce colored polyenes. This suggests that the al-2 gene product is concerned in two steps in carotenoid biosynthesis; the conversion of colorless polyenes and a later step concerned in the conversion of colored polyenes. The MN58p mutation would affect only the latter function, whilst al-2 (15300) affects both. However, this interpretation should be considered in conjunction with Huang's findings (1964 *Genetics* 49:453) that complementation may not be observed between albino mutants which, on the basis of recombination frequency, could be considered to be in different cistrons. Although some cases of complementation failure can be considered as due to unfavorable nuclear ratios in forced heterocaryons, the phenomenon is not inconsistent with the hypothesis that all of the mutations affecting polyene biosynthesis and located in this region of linkage group IR result in an alteration in the structure of one functionally integrated unit possessing heterocatalytic sites, analogous to the products of the grom gene cluster (Giles, Case, Partridge and Ahmed 1967 *Proc. Natl. Acad. Sci. U. S.* 58: 1453) and the hist-3 region of *Neurospora* (Minson and Creaser 1969 *Biochem. J.* 114:49; Catcheside, D. G. 1965 *Biochem. Biophys. Res. Commun.* 18:648; Ahmed, Core and Giles 1964 *Brookhaven Symp. Biol.* 17:55). The properties of the new mutant, MN58p, are consistent with this and similar suggestions which have been made by Huang (op. cit.) and by Subden and Threlkeld (1968 *Con. J. Genet. Cytol.* 10:351).

Table 1. Two-point linkage data for MN58p.

Cross No.	Locus	Allele	Linkage Group	Progeny parental	Progeny recombinant	Map Distance upper limit ^g
898	pe	L	II R	45	17 ^a	-
1046	lys-5	DS6-85	VI L	134	124	-
1046	try-2	75001	VI R	133	125	-
1110	try-1	A10	III R	65	60	-
1110	cot-1	C102	IV R	66	59	-
1116	nic-3	Y31881	V L	59	67	-
1106	nic-1	3416	I R	119	6	9.2
1105	al-2	15300	I R	5742 ^b	0 ^a	0.1
2242	aur	34508	I R	208	1 ^{a,c}	4.5
2348	aur	34508	I R	35	1 ^{a,d}	e
3771 f	alY	ALS4	I R	49	1 ^{a,d}	e

a - only one recombinant class distinguishable from parental classes; b - 2430 albino, 2812 purple. Ascospores were germinated and grown 3 days at 25°C in the dark on sorbose:glucose:fructose (1: 0.025 : 0.025 %) medium and then transferred to 4°C under fluorescent lighting for up to 14 days (Huang op. cit.) Progeny of all other crosses were scored, following total isolation, in 4" by 1/2" tubes incubated at 25°C for 3 days in the dark and subsequently for 3-7 days at ca. 23°C in diffused daylight; c = not pseudowild; d = not tested for pseudowild; e = contains translocation T(IR:VL) AR190. Scores of non-duplicated progeny only; f - D.D. Perkins data and cross number; g - 95% confidence level.

Table 2. Complementation between carotenoid mutants in the escape growth of duplications from crosses containing T(IR:VL) AR190.

Allele crossed to MN58p	Presumptive duplications examined*	Number showing wild type color in escape growth
al _y (ALS4)	26	19**
al (al ^s)	17	0
al (K96)	20	0
al-2 (15300)	28	0
aur (34508)	40	22

* The escape growth may have either type of parental pigmentation and is frequently sectored with both. Conidial re-isolates normally include both parental phenotypes, but the nuclear ratio appears to be unstable and may favor either parental type.

• * Data of D. D. Perkins.

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