McDougall, K. J. and V. W. Woodword. Suppression of pyr-3 mutants by arg-12 mutants. Until recently the arg-|2 locus, the locus which structures ornithine transcarbamylase (OTCase), was represented by a single mutant, $arg-|2^{5}$ (37301), $arg-|2^{5}$ possesses about 3%

of wild-type OTCore activity and is capable of suppressing the pyrimidine requirement of pyr-3 mutants characterized by in vitro aspartic transcarbamylase (ATCase) activity. (The pyr-3 mutants used here are denoted by the KS-prefix. KS16 and KS20 ore ATCase⁺; KS23 and KS43 are ATCase⁻. The arg-12 mutants ore designated as 6-1, 6-2, 6-3, 6-8 and 7.0.) The mechanism of suppression is thought to be due to metabolic cross-feeding of carbamyl phosphate (CAP), a substrate common to both pathways. Independent efforts by us and by Davis and Thwaites (1963 Genetics 48: 1551) resulted in the isolation of OTCoreless mutants phenotypically distinguishable from arg-12s by reduced growth rates and by 99% or more reduction of OTCore activity. There mutants were found to be located near arg-5 on linkage group II (Wood word and Schwarz 1964 Genetics 49: B45). It was not possible to demonstrate suppression of pyr-3 ATCase⁺ mutants by the new arg-12 mutants, since the required arginine supplement offsets suppression, possibly by repression or inhibition of carbomyl phosphokinase, shutting off the remaining source of CAP. The effect of exogenous arginine on pyr-3 ATCase⁺; arg-12 double mutants can be overcome, however, by adding lysine to the culture medium (Houldan and Mitchell 1947 Proc. Nat1, Acad. Sci. U. S. 33; 223), This procedure was employed to demonstrate that the new arg-12 isolates are capable of suppressing pyr-3 ATCase⁺ mutants.

	at g cond	centration of	0.3 mg/mi.						
Strain	Time (days)	Minimal	Aroinine	Uridine	Medium Arainine + Uridine		Lvsine + Aroinine		
K\$20	4	0	0	84.4	79.5	0	0	0	
	8	0	0	110.3	110.4	0	0	0	
b-9	4	0	85.4	0	86.7	97.7	90.1	97.2	
	8	0	75.9	0	78.9	99.	102.0	103.8	
KS43	4	0	0	90.]	87.3	0	0	0	
	8	0	0	119.4	112.1	0	0	0	
K\$43;6-	-94	0	0	0	91.3	0	0	0	
	8	0	0	0	79.9	0	0	0	
KS20;6-	-94	0	0	0	90.2	0	tr	4.1	
	8	0	4.9	0	77.6	12.4	24.5	30.0	

Table]. Dry weights, in mg, from 125 ml stationary flask cultures at 30°C containing 40 ml of medium; supplements were Used at g concentration of 0.3 mg/ml.

The data in Table 1 illustrate suppression of a <u>pyr-3</u> ATCase⁺ mutant (KS20) by an <u>arg-12</u> isolate (6-9). It is seen that the <u>pyr-3</u> ATCase⁻ strain (KS43) is not suppressed, which is in agreement with earlier findings (Davis and Woodward 1962 Genetics 47: 1075). Suppression is also observed 0n arginine medium providing the double mutant is cultured for a prolonged period, in much the same way as the RU-suppressors of pyr-3 mutants (McDougall and Woodward 1965 Genetics 50: 397). Although data are presented for only two double mutants, suppression was observed in the following combinations involving an ATCase⁺ <u>pyr-3</u> mutant and an <u>arg-12</u> mutant: KS16;7-0, KS20;6-2, KS20;6-8, KS20;7-0. In addition, all of there double mutants: KS23;6-1, KS23;6-3, KS43;6-1, KS43;6-3.

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