

Fig. 1. Curves showing average weekly gains of rats fed on opaque-2 maize and Indiana hybrid 453.

Fortification Mixture; diet D, the same as diet C except containing 20 percent soybean meal (8), no casein, and 65 percent corn starch.

The results of feeding the rats on

Table 2. Gains in weight, protein consumption, and protein efficiency ratios (28 days).

Animal No.	Gain (g)	Protein eaten (g)	PER*	
	Diet conta	ining casein		
C-1	70	20.3	3.4	
C-2	54	17.2	3.1	
C-3	72	31.6†	2.3	
C-4	73	21.0	3.5	
C-5	43	1 7.4 ‡	2.5:	
C-6	85	26.7	3.2	
Average	66		3.1	
Die	t containing	opaque-2 ma	ize	
A-7	86	30.0	2.9	
A-8	102	38.0†	2.7	
A-9	134	41.6	3.2	
A-10	81	30.7	2.6	
A-11	96	37.0	2.6	
A-12	85	38.2†	2.2	
Average	97		2.8	
Diet	containing	Indiana hybrid	453	
B-13	32	38.6†	0.8	
B-14	33	19.8	1.7	
B-15	23	21.5†	1.1	
B-16	23	14.3	1.6	
B-17	23	19.7†	1.2	
B-18	30	19.5	1.5	
Average	27		1.6	
D	iet containii	ng soybean med	d i a	
D-19	67	35.6†	1.9	
D-20	95	34.0	2.8	
D-21	95	36.0	2.6	
D-22	101	37.2‡	2.7	
D-23	78	28.7	2.7	
D-24	105	37.3	2.8	
Average	90		2.7	

* Protein efficiency ratio (grams gained divided by grams protein eaten). The percentage of pro-tein (N \times 6.25) in the diets was as follows: diet C, 8.1; diets A and B, 9.5; and diet D, 10.0. \dagger Value not included in average because of food scattering. \ddagger Rat scattered the food slightly, but value was included in average.

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these diets are summarized in Table 2. In spite of special feed cups, scattering was a problem in all groups, with greatest incidence in the group fed on hybrid maize (diet B). Protein efficiency ratios (PER) were calculated for all animals, but the average PER values include only animals which did not scatter food, or scattered only slightly. Howe et al. (9) surveyed the PER values of nine unsupplemented, ground, whole cereal grains fed to rats on diets containing 7.8 to 10 percent protein, and found oats to have the highest value (1.8) when compared with casein (2.5). Barley, rice, maize, bulgar, rye, wheat, sorghum, and millet were below oats in PER. Our data indicate that in young rats opaque-2 maize proteins have a food value equal to that of heat-treated soybean meal, and superior to any cultivated cereal grain.

Diets A and B were labeled with 1 percent metallic oxide, and the digestibility of the proteins was measured as described by Mertz et al. (10). White rats weighing approximately 100 g were used, four animals being fed on each diet. Protein digestibility values of 80, 83, 86, and 87 percent, respectively, were obtained with the rats on diet A (opaque-2) and values of 84, 84, 87, and 87 percent, respectively, were obtained with the rats on diet B (Indiana hybrid 453). Better digestibility of proteins therefore does not appear to be a factor contributing to the growth-promoting properties of opaque-2 maize.

In a previous feeding test with the same number of rats (3), the average gain in weight for 28 days was 86 g for rats fed a different strain of opaque-2 maize and 23 g for rats fed Indiana hybrid 257, a 3.7-fold difference. In the present test, a 3.6-fold difference was observed. Figure 1 shows average weekly gains of the animals on diets A and B. Similar curves were obtained in the previous feeding test.

The greater efficiency of opaque-2 proteins in rats provides a basis for assuming that opaque-2 proteins would also be superior to ordinary maize proteins in the diet of man and domestic animals. Loci homologous to opaque-2 probably exist in other cereals, and methods for detection and use of mutants should be developed.

Note added in proof: Opaque kernels in high protein background gave endosperms containing 19.2 percent protein, of which 2.7 percent was lysine. Nonopaque kernels on the same ear yielded endosperms with 19.6 percent protein, of which 1.3 percent was lysine. Thus, the opaque gene exerts its effect in a high protein background, and selection is feasible for lines that are high in both lysine and protein. Such lines would provide more lysine per gram of corn than diet A.

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Isoantigens of

Gamma Globulin in Pigs

Abstract. Two gamma globulin isoantigens in pigs have been identified by hemagglutination-inhibition tests. Two codominant autosomal alleles, Gla and determine three phenotypes, Gl^b. Gl(a+b-), Gl(a+b+), and Gl(a-b+).

Genetic differences in types of γ -globulins have been described for a number of mammalian species (1-3). The hemagglutination-inhibition system used to identify isoantigens of γ -globulins in pigs is modeled after that used for typing Gm and Inv, the hereditary antigenic determinants of γ -globulin in man (2).

Antibodies against the serum factor designated Gl_a were first produced by isoimmunization of pigs according to a procedure similar to that described for rabbits (3), but repeated immunizations were required to produce antibodies, and inhibition of agglutination was not easily interpreted. Wilson and Steinberg

(4) found a high frequency of antibodies to Gm and Inv in children aged 6 months to 5 years, and serums from young pigs may similarly have a high frequency of isoantibodies to y-globulins without deliberate immunization. There were antibodies to Gl_a in 2 out of 20 serum samples from pigs which were approximately 4 months old, and in another there were antibodies to a complementary serum factor, designated $Gl_{\rm b}$ (5). One of the serums with antibodies to Gl_a and the single serum with antibodies to Gl_h were used as agglutinators in all of the tests reported here.

For the test for Gl_a, one volume of a 2 percent suspension of red cells (washed seven times) from a K_h-positive (6) pig of group O (7) was mixed with two volumes of an antiserum (diluted 1:16) which contained incomplete antibodies to K_b. Cells from a group-O pig were used since naturally occurring antibodies to pig O are rare (7) and would not be likely to interfere with the tests for Gl factors. The mixture was incubated for 21/2 hours at room temperature (24°C) to coat the red cells with antibody. The coated cells were then washed four times, and saline (0.91 percent NaCl) was added to make a 1-percent suspension. In each of two tubes one drop (1/30 ml) of the suspended red cells was added to a mixture of equal parts of antiserum to Gl_a (diluted 1:16) obtained from a pig, aged 4 months, and the diluted serum to be tested (1:4 and 1:8, respectively). The agglutinator and test serum were previously incubated at room temperature for 2 hours. The mixture of test serum, agglutinator, and coated cells was incubated at room temperature for 5 minutes, centrifuged gently (270g) for 1 minute, and examined macroscopically for agglutination. After an additional 2 hours incubation the tubes were again examined for agglutination, before and after centrifuging. The Gl(a+) test serums strongly inhibited agglutination of the coated cells by antibody to Gl_a; Gl(a-) serums inhibited weakly or not at all. For the test for Gl_b, the serum (diluted 1:8) obtained from a pig aged $4\frac{1}{2}$ months was used as the agglutinator, and incomplete antibodies to L_e were used for coating the L_e-positive (6) red cells of the same group-O pig used as a source of cells in the test for Gl_a. Controls consisted of: known Gl(a+b-) and Gl(a-b+) serums tested with each agglutinator and appropriate coated cells; diluted test serum, saline, and coated cells; saline, agglutinator, 25 JUNE 1965

Table	1.	Inheritance	of	Gl	types	in	pigs

Mating types	Mat- ings (No.)	Offspring			
		a+b-	a+b+	a-b+	
$\overline{a+b- \times}$					
a+b+	6	20	22	0	
$a+b- \times$					
a-b+	2	0	14	0	
$a+b+ \times$	-	-	06	0	
a+b+	3	1	26	8	
a+b+x a-b+	4	0	14	20	
$a-b+ \times a-b+$	4	0	0	30	

and coated cells; and saline plus coated cells. The red cells coated with incomplete antibodies to K_b or L_e can be agglutinated by the addition of rabbit antiserum to pig globulin as well as by the appropriate isoantibodies.

Porcine γ -globulins (Cohn Fraction II) and albumin (Cohn Fraction V) (8) were checked for freedom from other serum proteins by immunoelectrophoresis, and saline solutions of each fraction (20 mg/ml) were used in inhibition tests. There was no inhibition of agglutination by albumin, whereas the γ -globulin (diluted to 20 mg/ml) inhibited antibody to $\operatorname{Gl}_{\operatorname{a}}$ up to a further dilution of 1:8, and inhibited antibody to Gl_b up to 1:4096. These data suggest that most of the pigs from which the pool for preparation of the y-globulin was obtained were Gl(a-b+).

The distribution of Gl_a and Gl_b in serums from adult males and females (chosen to avoid including samples from pairs of full sibs) was: Duroc pigs, 2 Gl(a+b-), 10 Gl(a+b+), 2 Gl(a-b+); Yorkshire pigs, 1 Gl(a+b-), 3 Gl(a+b+), 12 Gl(a-b+).

No serums have been found to be Gl(a-b-), and it appears likely that allelic genes are responsible for factors Gl_a and Gl_b . The globulin types of 161 offspring, tested when 3 months old or older, from 21 matings are given in Table 1. These data are in accord with the hypothesis that Gl^a and Gl^b are codominant alleles, so that genotypes Gla Gla, Gla Glb, and Glb Glb determine phenotypes Gl(a+b-), Gl(a+b+), and Gl(a-b+). The existence of additional alleles, including $Gl^$ and Gl^{ab} , is not excluded by these limited data from two breeds of pigs.

Specific Gl types are not associated with specific haptoglobin, transferrin, prealbumin or amylase (9) types, nor are they associated with specific redcell antigens in the A-O, B, C, E, F, G, H, I, J, K, or L (6, 7, 10) systems. Results of segregation in offspring from matings of animals of known Gl and red-cell types have excluded sex linkage and close linkage between the Gl locus and the A, B, C, E, F, G, H, J, K, and L loci controlling red-cell antigens.

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Degeneration of the Eyes of **Tyrosine-Deficient Chick Embryos**

Abstract. Subjecting 4-day-old chick embryos to a yolk-sac perfusion medium lacking tyrosine resulted in arrest of retinal pigmentation and in degeneration of the neural retina. Phenylaldnine was ineffective in replacing tyrosine. Possibly retinal tyrosinase played a part in initiating the degenerative changes.

The first successful study of amino acid deficiencies in chick embryos was reported by Klein, et al. (1), who applied the explantation techniques of Spratt (2) and of Hayashi and Herrmann (3) in studies with defined