

Genetic dissection of autoimmune type I diabetes in the BB rat

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Nature Genetics 2, 56–60 (1992)

Originally we reported that *Iddm1* is responsible for causing lymphopenia and is tightly linked to the *Npy* locus (0.7 cM from *Npy* in the direction of *Igk*) on rat chromosome 4, the exact position based on three recombinants out of 429 animals. Subsequently, we have determined that the three apparent recombinants were due to sample and phenotype errors: there were no actual recombinants between *Npy* and *Iddm1*. Studies of more than 500 additional animals have shown that *Iddm1* still maps very close to *Npy*, but towards the locus *D4Mit6* (see Fig. 1). These findings do not affect the conclusions of the paper, apart from repositioning *Iddm1* by approximately 1.5 cM.

Automated construction of genetic linkage maps using an expert system (MultiMap): a human genome linkage map

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Nature Genetics 6, 384–390 (1994)

Information about MultiMap can be obtained from the following revised e-mail addresses:
aravinda@chimera.gene.cwru.edu or
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erratum

A null mutation in the human *CNTF* gene is not causally related to neurological diseases

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Table 1 was inadvertently omitted from the final layout of this paper. Table 1 appears below.

Table 1 Distribution of *CNTF* genotypes in healthy subjects and patients with neurological diseases

Subjects	Number of subjects (%)		
	Genotype		
	N/N	N/M	M/M
Healthy volunteer	95 (62.9)	52 (34.4)	4 (2.6)
ALS	27 (57.4)	18 (38.3)	2 (4.3)
Alzheimer disease	17 (56.7)	13 (43.3)	0 (0)
Parkinson disease	30 (57.7)	21 (40.4)	1 (1.9)
Miscellaneous disease	73 (65.8)	36 (32.4)	2 (1.8)
Total	242 (61.9)	140 (35.8)	9 (2.3)



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