

ift88^{tz288}

Nature of the mutation

The *tz288* allele contains a single T-to-A point mutation that introduces a premature stop codon in exon 11 of the *ift88* transcription unit (Tsujikawa and Malicki, Neuron 42(5): 703-716, 2004).

Genotyping assay

Genotyping of the *tz288* allele is based on the RFLP assay (**R**estriction Fragment Length **P**olymorphism; Botstein *et al.*, Am. J. Hum. Genet. 32: 314-331, 1980). This method is used to detect a mutation that either creates or abolishes a site recognized by a specific restriction enzyme. In the RFLP assay, a sequence of interest is first PCR-amplified and then the PCR product is subjected to restriction enzyme digestion. The presence or absence of the mutation is determined by the resulting restriction pattern. The *tz288* mutation creates a site recognized by the Ddel restriction enzyme.

Primers:

OVL_01: 5' GCA AAT TAG TGC ATA TAA CGC CTC 3' OVL_02: 5' CCA ATG TTC TGC ATG ATC TTT ATC C 3'

PCR program (53_30_30):

- 1. 94°C for 3 min
- 2. 94°C for 30 sec
- 3. **53°**C for **30** sec
- 4. 72°C for **30** sec
- 5. Go to step 2 (above) for 39 cycles
- 6. 72°C for 5 min
- 7. 8.0°C hold
- 8. END

Product size: 401 bp

Digestion of the PCR product with the DdeI restriction enzyme:

Product type	Product digestion	DNA fragments after digestion (bp)
PCR product derived from the WT template	unaffected	401 bp
PCR product containing the mutation	cleaved	259 bp and 142 bp