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## IN BRIEF

#### **MULTIPLE SCLEROSIS**

# Risk of comorbid inflammatory diseases in MS might not be genetically determined

Patients with multiple sclerosis (MS) often present with other immune-mediated disorders, such as ulcerative colitis and type 1 diabetes. Researchers in Sweden analysed data from a large register of patients with MS and their parents to determine whether the risk of such comorbidity has a genetic component. They found that, in contrast to index cases, parents of patients with MS did not show an increased likelihood of having an immune-mediated disorder compared with control parents, arguing against a genetic influence.

**Original article** Roshanisefat, H. *et al.* Shared genetic factors may not explain the raised risk of comorbid inflammatory diseases in multiple sclerosis. *Mult. Scler.* doi:10.1177/1352458512438240

#### **EPILEPSY**

#### Immunotherapy could benefit treatment-resistant epilepsy

Accumulating evidence has suggested an autoimmune component to drug-resistant epilepsy. Now, immunotherapy in 27 patients with uncontrolled seizures with suspected autoimmune aetiology has shown striking results. At a median follow-up of 10 months, 22 patients (81%) exhibited substantial improvements in seizure status, and 18 patients (67%) were seizure-free. These results suggest that the treatment of autoimmunity could help to address the large unmet medical need of drug resistance in epilepsy.

**Original article** Quek, A. M. et al. Autoimmune epilepsy: clinical characteristics and response to immunotherapy. Arch. Neurol. doi:10.1001/archneurol.2011.2985

### **MOVEMENT DISORDERS**

# Interferon- $\gamma$ shows promise in a mouse model of Friedreich ataxia

Friedreich ataxia is a hereditary disease caused by mutation of the frataxin (FXN) gene and resultant low levels of the FXN protein. In a mouse model of the disease, Tomassini et al. found that subcutaneous injections of interferon- $\gamma$  for 14 weeks restored FXN levels in dorsal root ganglion neurons, which are particularly vulnerable in this disease, and improved motor coordination. The findings open therapeutic avenues for Friedreich ataxia, a disease for which no specific treatments are currently available.

 $\begin{tabular}{ll} \textbf{Original article} Tomassini, B.\ et\ al.\ Interferon\ gamma\ upregulates\ frataxin\ and\ corrects\ the\ functional\ deficits\ in\ a\ Friedreich\ ataxia\ model.\ \textit{Hum. Mol. Genet.}\ doi:10.1093/hmg/dds110\end{tabular}$ 

#### **ALZHEIMER DISEASE**

### Targeting apolipoprotein E improves AD phenotype in mice

Apolipoprotein E (APOE) is indirectly involved in the proteolytic degradation of amyloid- $\beta$ , and allelic variation in the APOE gene is the leading genetic risk factor for Alzheimer disease (AD). As APOE expression is transcriptionally regulated by retinoid X receptors (RXRs), Cramer et al. investigated the oral RXR agonist bexarotene in AD mice. The resultant increase in APOE levels was accompanied by increased clearance of amyloid- $\beta$ , rapid reduction in plaque volume and improvements in functional deficits associated with AD. Bexarotene is already approved as an anticancer drug.

**Original article** Cramer, P. E. *et al.* ApoE-directed therapeutics rapidly clear  $\beta$ -amyloid and reverse deficits in AD mouse models. *Science* **335**, 1503–1506 (2012)