













## Axon degeneration in MS Direct or bystander immune-mediated attack...TNFa or Fas ligand Energy deficiency: inherent mitochondrial defects or due to damage by inflammatory mileu (e.g free radicals) Glutamate excitotoxicity Antibodies to neurofascin186, a component of the node/paranode (Mathy et al, 2007) Na<sup>+</sup> and Ca2<sup>+</sup> overloading in electrically active/ energy depleted axons (Trapp and Stys, 2009) Detected indirectly in patients by MRS for N-acetyl-aspartate (NAA). Reduction in NAA level correlates with increasing disability.



































































## Is multiple sclerosis a neurodegenerative disease ?

- Cortical pathology has a major impact on clinical progression in MS
- Presence of meningeal B-cell follicles leads to more extensive pathology and loss of neurons
- Cytotoxicity mediated by factors released by B-cells and/or CD8 Tcells and/or microglia
- How early during the disease course does this start?
- Different pathogenetic mechanisms may be involved in WM & GM pathology, suggesting novel treatment options

Multiple sclerosis is an inflammatory neurodegenerative disease

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![](_page_22_Figure_1.jpeg)

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## **Myelin repair in MS**

- remyelination leads to restoration of neurological function and may protect axons against damage
- The adult mammalian brain and spinal cord has an enormous capacity for repairing myelin damage

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