

by widespread microglial activation across large areas of white matter, associated with reduced myelin staining and axonal injury (“dirty white matter”). Activated astrocytes induced by microglia may also contribute to tissue damage ([Chap. 425](#)). These observations imply that ongoing inflammation occurs behind an intact BBB in many patients with progressive MS, and this feature could explain the failure of immunotherapies not capable of crossing the BBB to benefit patients with progressive MS.

■ Physiology

Nerve conduction in myelinated axons occurs in a saltatory manner, with the nerve impulse jumping from one node of Ranvier to the next without depolarization of the axonal membrane underlying the myelin sheath between nodes ([Fig. 444-2](#)). This produces faster conduction velocities (~ 70 m/s) than the slow velocities (~ 1 m/s) produced by continuous propagation in unmyelinated nerves. Conduction block occurs when the nerve impulse is unable to traverse the demyelinated segment. This can happen when the resting axon membrane becomes hyperpolarized due to exposure of voltage-dependent potassium channels that are normally buried underneath the myelin sheath. A temporary conduction block often follows a demyelinating event before sodium channels (originally concentrated at the nodes) redistribute along the naked axon ([Fig. 444-2](#)). This redistribution ultimately allows continuous propagation of nerve action potentials through the demyelinated segment. Conduction block may be incomplete, affecting high- but not low-frequency volleys of impulses. Variable conduction block can occur with raised body temperature or metabolic alterations and may explain clinical fluctuations that vary from hour to hour or appear with fever or exercise. Conduction slowing occurs when the demyelinated segments of the axonal membrane are reorganized to support continuous (slow) nerve impulse propagation.