

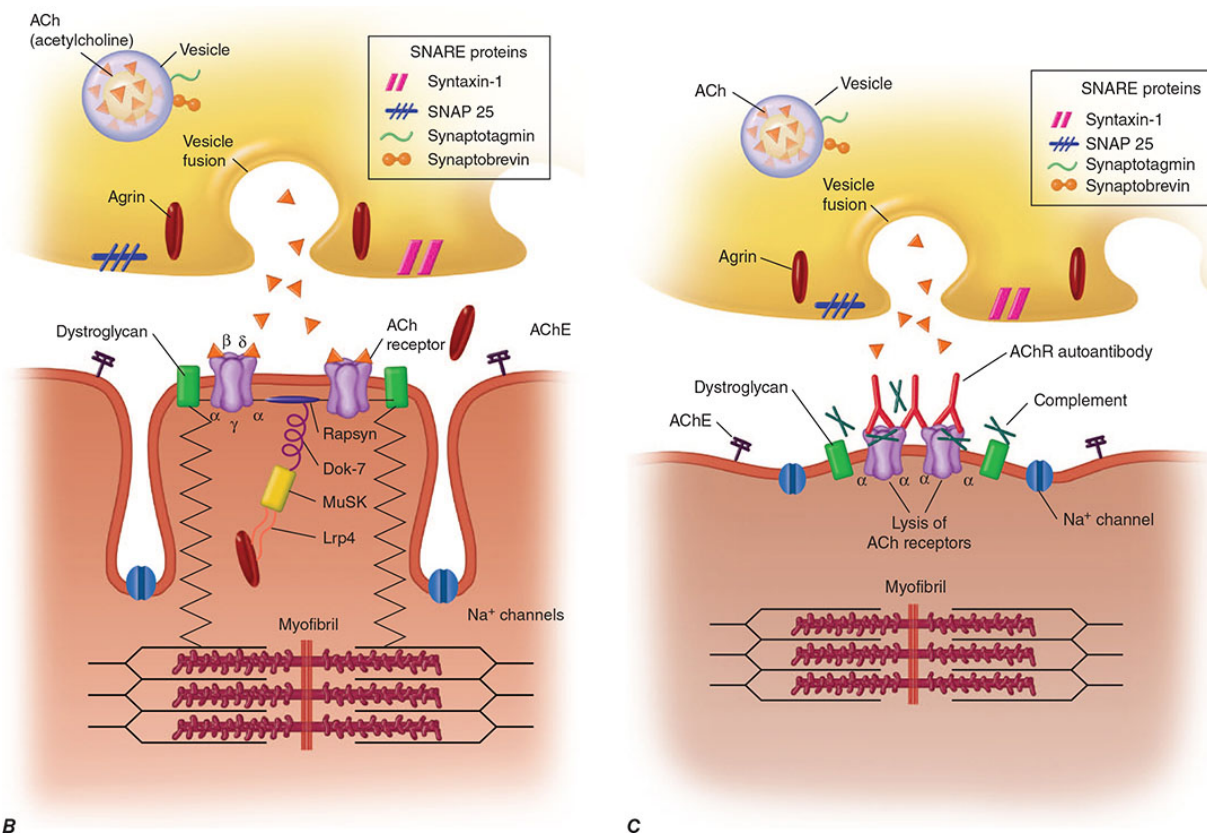


FIGURE 448-1 Illustrations of (A) a normal presynaptic neuromuscular junction, (B) a normal postsynaptic terminal, and (C) a myasthenic neuromuscular junction. AChE, acetylcholinesterase. See text for description of normal neuromuscular transmission. The myasthenia gravis (MG) junction demonstrates a reduced number of acetylcholine receptors (AChRs); flattened, simplified postsynaptic folds; and a widened synaptic space. See [Video 448-1](#) also. (From AA Amato, J Russell: *Neuromuscular Disorders*, 2nd ed. New York, McGraw-Hill, 2016, Figures 25-3 [p 588], 25-4 [p 589], and 25-5 [p 590]; with permission.)

In MG, the fundamental defect is a decrease in the number of available AChRs at the postsynaptic muscle membrane. In addition, the postsynaptic folds are flattened, or “simplified.” These changes result in decreased efficiency of neuromuscular transmission. Therefore, although ACh is released normally, it produces small end-plate potentials that may fail to trigger muscle action potentials. Failure of transmission results in weakness of muscle contraction.

The amount of ACh released per impulse normally declines on repeated activity (termed *presynaptic rundown*). In the myasthenic patient, the decreased efficiency of neuromuscular transmission