

Tachykinin receptors in GtoPdb v.2023.1

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Abstract

Tachykinin receptors (**provisional nomenclature as recommended by NC-IUPHAR [91]**) are activated by the endogenous peptides [substance P](#) (SP), [neurokinin A](#) (NKA; previously known as substance K, neurokinin α , neuromedin L), [neurokinin B](#) (NKB; previously known as neurokinin β , neuromedin K), [neuropeptide K](#) and [neuropeptide \$\gamma\$](#) (N-terminally extended forms of neurokinin A). The neurokinins (A and B) are mammalian members of the tachykinin family, which includes peptides of mammalian and nonmammalian origin containing the consensus sequence: Phe-x-Gly-Leu-Met. Marked species differences in *in vitro* pharmacology exist for all three receptors, in the context of nonpeptide ligands. Antagonists such as [aprepitant](#) and [fosaprepitant](#) were approved by FDA and EMA, in combination with other antiemetic agents, for the prevention of nausea and vomiting associated with emetogenic cancer chemotherapy.

Contents

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NK₃ receptor

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