

P2X receptors (version 2019.4) in the IUPHAR/BPS Guide to Pharmacology Database

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Abstract

P2X receptors (nomenclature as agreed by the [NC-IUPHAR Subcommittee on P2X Receptors \[46, 134\]](#)) have a trimeric topology [[118](#), [132](#), [177](#)] with two putative TM domains, gating primarily Na⁺, K⁺ and Ca²⁺, exceptionally Cl⁻. The Nomenclature Subcommittee has recommended that for P2X receptors, structural criteria should be the initial criteria for nomenclature where possible. X-ray crystallography indicates that functional P2X receptors are trimeric and three agonist molecules are required to bind to a single receptor in order to activate it [[132](#), [88](#), [96](#), [161](#)]. Native receptors may occur as either homotrimers (e.g. P2X1 in smooth muscle) or heterotrimers (e.g. P2X2:P2X3 in the nodose ganglion [[251](#)], P2X1:P2X5 in mouse cortical astrocytes [[146](#)], and P2X2:P2X5 in mouse dorsal root ganglion, spinal cord and mid pons [[50](#), [207](#)]). P2X2, P2X4 and P2X7 receptors have been shown to form functional homopolymers which, in turn, activate pores permeable to low molecular weight solutes [[229](#)]. The hemi-channel pannexin-1 has been implicated in the pore formation induced by P2X7 [[188](#)], but not P2X2 [[38](#)], receptor activation.

Contents

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