

Type XIII RTKs: Ephrin receptor family in GtoPdb v.2025.3

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Abstract

Ephrin receptors are a family of 15 RTKs - the largest family of RTKs - with two identified subfamilies (EphA and EphB), which have a role in the regulation of neuronal development, cell migration, patterning and angiogenesis. Their ligands are membrane-associated proteins, thought to be glycosylphosphatidylinositol-linked for EphA ([ephrin-A1](#) , [ephrin-A2](#), [ephrin-A3](#), [ephrin-A4](#) and [ephrin-A5](#)) and transmembrane proteins for Ephrin B ([ENSFM0025000002014: ephrin-B1](#), [ephrin-B2](#) and [ephrin-B3](#)). Ephrin-A3 and ephrin-B3 have also been shown to interact with heparan sulphate proteoglycans [22].

Contents

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Database links

Type XIII RTKs: Ephrin receptor family

<https://www.guidetopharmacology.org/GRAC/FamilyDisplayForward?familyId=327>

Introduction to Type XIII RTKs: Ephrin receptor family

<https://www.guidetopharmacology.org/GRAC/FamilyIntroductionForward?familyId=327>

Receptors

[EphA1\(EPH receptor A1\)](#)

<https://www.guidetopharmacology.org/GRAC/ObjectDisplayForward?objectId=1821>

[EphA2\(EPH receptor A2\)](#)

<https://www.guidetopharmacology.org/GRAC/ObjectDisplayForward?objectId=1822>

[EphA3\(EPH receptor A3\)](#)

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[EphA4\(EPH receptor A4\)](#)

<https://www.guidetopharmacology.org/GRAC/ObjectDisplayForward?objectId=1824>

[EphA5\(EPH receptor A5\)](#)

<https://www.guidetopharmacology.org/GRAC/ObjectDisplayForward?objectId=1825>

[EphA6\(EPH receptor A6\)](#)

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[EphA7\(EPH receptor A7\)](#)

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[EphA8\(EPH receptor A8\)](#)

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