

Phosphodiesterases, 3',5'-cyclic nucleotide (PDEs) in GtoPdb v.2025.1

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Abstract

3',5'-Cyclic nucleotide phosphodiesterases (PDEs, 3',5'-cyclic-nucleotide 5'-nucleotidohydrolase), [E.C. 3.1.4.17](#), catalyse the hydrolysis of a 3',5'-cyclic nucleotide (usually [cyclic AMP](#) or [cyclic GMP](#)). [isobutylmethylxanthine](#) is a nonselective inhibitor with an IC₅₀ value in the millimolar range for all isoforms except PDE 8A, 8B and 9A. A 2',3'-cyclic nucleotide 3'-phosphodiesterase ([E.C. 3.1.4.37](#) CNPase) activity is associated with myelin formation in the development of the CNS.

Contents

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Database links

Phosphodiesterases, 3',5'-cyclic nucleotide (PDEs)

<https://www.guidetopharmacology.org/GRAC/FamilyDisplayForward?familyId=260>

Introduction to Phosphodiesterases, 3',5'-cyclic nucleotide (PDEs)

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Enzymes

PDE1A(phosphodiesterase 1A)

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PDE1B(phosphodiesterase 1B)

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PDE1C(phosphodiesterase 1C)

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PDE2A(phosphodiesterase 2A)

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PDE3A(phosphodiesterase 3A)

<https://www.guidetopharmacology.org/GRAC/ObjectDisplayForward?objectId=1298>

PDE3B(phosphodiesterase 3B)

<https://www.guidetopharmacology.org/GRAC/ObjectDisplayForward?objectId=1299>

PDE4A(phosphodiesterase 4A)

<https://www.guidetopharmacology.org/GRAC/ObjectDisplayForward?objectId=1300>

PDE4B(phosphodiesterase 4B)

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[PDE4C\(phosphodiesterase 4C\)](https://www.guidetopharmacology.org/GRAC/ObjectDisplayForward?objectId=1302)
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[PDE4D\(phosphodiesterase 4D\)](https://www.guidetopharmacology.org/GRAC/ObjectDisplayForward?objectId=1303)
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[PDE10A\(phosphodiesterase 10A\)](https://www.guidetopharmacology.org/GRAC/ObjectDisplayForward?objectId=1310)
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[PDE11A\(phosphodiesterase 11A\)](https://www.guidetopharmacology.org/GRAC/ObjectDisplayForward?objectId=1311)
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