

Bile acid receptor in GtoPdb v.2025.3

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Abstract

The bile acid receptor (GPBA) responds to bile acids produced during the liver metabolism of [cholesterol](#). Selective agonists are promising drugs for the treatment of metabolic disorders, such as type II diabetes, obesity and atherosclerosis.

Contents

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Database links

Bile acid receptor

<https://www.guidetopharmacology.org/GRAC/FamilyDisplayForward?familyId=8>

Introduction to Bile acid receptor

<https://www.guidetopharmacology.org/GRAC/FamilyIntroductionForward?familyId=8>

Receptors

GPBA receptor

<https://www.guidetopharmacology.org/GRAC/ObjectDisplayForward?objectId=37>

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