



Recombinant Human Liver X receptor β - LBD (Ligand Binding Domain)

Human recombinant protein expressed in *Nicotiana benthamiana*.

Cat. No. RF0019-1

RF0019-5

RF0019-10

RF0019-25

RF0019-100

Molecular formula:

C₁₄₂₂H₂₂₄₉N₄₀₉O₄₁₅S₈

Molecular weight:

31.9 kDa (201-461 amino acids)

p.I:

6,4

Extinction coefficient :

E_{0.1%} = 1,27 (A 280 nm)

Purity assay:

> 97% by SDS-PAGE gel

Endotoxin level:

< 0.04 EU/ μ g protein (LAL method)

Animal-free product*

Sequence:

HHHHHHSSGIEGRGRGLIKHMTPGGSEAGSQSGEGEGVQLTAAQELMIQQLVAAQL
QCNKRSFSDQPKVTPWPLGADPQSRDARQQRFAHFTELAIISVQEIVDFAKQVPGFLQ
LGREDQIALLKASTIEIMLLETARRYNHETECITFLKDFTYSKDDFHRAGLQVEFINPIF
EFSRAMRRLLGLDDAEYALLIAINIFSADRPNVQEPGRVEALQQPYVEALLSYTRIKRP
QDQLRFPRMLMKLVSLRTLSSVHSEQVFALRLQDKKLPPLLSEIWDVHE

Description:

Liver X Receptors (LXRs) are nuclear receptors that regulate the metabolism of cholesterol and bile acids. There are two subtypes of LXRs, LXR α and LXR β . The LXRs are ligand-dependent transcription factors that form permissive heterodimers with the retinoid X receptor (RXR). LXR- member of Nuclear Receptor Family is activated by certain oxysterol derivatives of cholesterol. They play an important role in cholesterol, lipid, and carbohydrate metabolism. LXR α is highly expressed in liver tissue. They respond to elevated cholesterol levels via transactivation of genes involved in sterol transport (ABCA1, ABCG1, ABCG5, and ABCG8), cholesterol efflux and high-density lipoprotein (HDL) metabolism, and sterol catabolism (CYP7A1). They also play a central role in regulating cellular lipid content through activation of SREBP-1c, which is the master regulator of de novo lipogenesis. LXRs were found to upregulate angiopoietin- like protein 3 (Angpf13), a member of the family of vascular endothelial growth factors that is also a key regulator of lipid metabolism. The liver's X receptors are critical for the control of lipid homeostasis. LXRs serve as cholesterol sensors that regulate the expression of multiple genes involved in the efflux, transport, and excretion of cholesterol. Synthetic LXR agonists inhibit the development of atherosclerosis in murine models. These observations identify the LXR pathway as a potential target for therapeutic intervention in human cardiovascular disease.

Formulation:

Lyophilized from a Tris HCl 20mM buffer at pH 8, 0.1% SDS.

Source:

Produced by transient expression of LXR in non-transgenic plants. Recombinant human LXR contains a 6-His-tag at the N-terminal end and is purified by sequential chromatography (FPLC). Contains no animal-derived components or impurities.

Reconstitution recommendation:

Lyophilized protein should be reconstituted in water to a concentration of 50 ng / μ l.

Storage and Stability:

This lyophilized preparation is stable at 2-8° C. For long storage should be kept at -20° C and it is recommended to add a carrier protein (0.1% HSA or BSA). Repeated freezing and thawing is not recommended.

Purity and Serological Identification:

The protein was resolved by SDS polyacrylamide gel electrophoresis and the gel was stained with Coomassie blue.

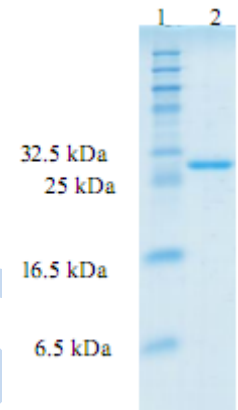
Figure 1. SDS-PAGE analysis of recombinant LXR. Samples were loaded in 15% SDS-polyacrylamide gel and stained with Coomassie blue. Lane 1: Molecular weight marker (kDa); lane 2: contains 0.3 ug of recombinant LXR.

Recombinant Human Liver X receptor β

Serological Identification:

The protein was electrophoresed under reducing condition on a 15% SDS-polyacrylamide gel, transferred by electroblotting to a NC membrane and visualized by immune-detection with specific LXR antibody.

Figure 2. Western Blot analysis of recombinant LXR. Lane 1: Molecular weight marker (kDa). Lane 2: 0.2 ug of LXR.



References

-Toresson G., Schuster G.U., Steffensen K.R., Bengtsson M., Ljunggren J., Dahlman-Wright K., Gustafsson J.A. 2004. Purification of functional full-length liver X receptor beta produced in Escherichia coli. *Protein Expr. Purif.* 35(2):190-8.

-Quinet E.M., Savio D.A., Halpern A.R., Chen L., Schuster G.U., Gustafsson J.A., Basso M.D., Nambi P. 2006. Liver X receptor (LXR)-beta regulation in LXRalpha-deficient mice: implications for therapeutic targeting *Mol. Pharmacol.* 70(4):1340-9.

-Patel M. B., Oza N. A., Anand I. S., Deshpande S. S., Patel C. N. 2008. Liver X Receptor: A Novel Therapeutic Target. *Indian J Pharm Sci.*; 70(2): 135–144.

For R+D purposes only. Purchaser must determine the suitability of the product(s) for their particular use.

***Agrenvec products are expressed in a plant system and intrinsically have extremely low endotoxin levels and are Animal-free.**

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Marienbongard 20
52062 Aachen Deutschland