abcam

Product datasheet

Recombinant Human Beta Arrestin 2 protein ab132309

1 Image

Description		
Product name	Recombinant Human Beta Arres	in 2 protein
Expression system	Wheat germ	
Accession	<u>P32121</u>	
Protein length	Full length protein	
Animal free	No	
Nature	Recombinant	
Species	Human	
Sequence		 MGEKPGTRVFKKSSPNCKLTVYLGKRDFVDHLDKVDPV DGVVLVDPDYLK DRKVFVTLTCAFRYGREDLDVLGLSFRKDLFIATYQAFPP VPNPPRPPTR LQDRLLRKLGQHAHPFFFTIPQNLPCSVTLQPGPEDTGKA CGVDFEIRAF CAKSLEEKSHKRNSVRLVIRKVQFAPEKPGPQPSAETTR HFLMSDRSLHL EASLDKELYYHGEPLNVNVHVTNNSTKTVKKIKVSVRQYA DICLFSTAQY KCPVAQLEQDDQVSPSSTFCKVYTITPLLSDNREKRGLAL DGKLKHEDTN LASSTIVKEGANKEVLGILVSYRVKVKLVVSRGGDVSVEL PFVLMHPKPH DHIPLPRPQSAAPETDVPVDTNLIEFDTNYATDDDIVFEDF ARLRLKGMK DDDYDDQLC
Predicted molecular weight	73 kDa including tags	
Amino acids	1 to 409	

Specifications

Our Abpromise guarantee covers the use of ab132309 in the following tested applications.

The application notes include recommended starting dilutions; optimal dilutions/concentrations should be determined by the end user.

Applications

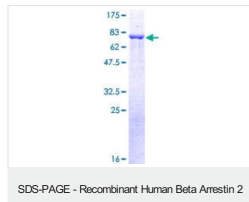
ELISA

SDS-PAGE

	Western blot
Form	Liquid
Preparation and Storage	
Stability and Storage	Shipped on dry ice. Upon delivery aliquot and store at -80°C. Avoid freeze / thaw cycles.
	pH: 8.00
	Constituents: 0.31% Glutathione, 0.79% Tris HCI
General Info	
Function	Eurotions in regulating agapted mediated Constain equilled recenter (CRCR) signaling by
Function	Functions in regulating agonist-mediated G-protein coupled receptor (GPCR) signaling by mediating both receptor desensitization and resensitization processes. During homologous
	desensitization, beta-arrestins bind to the GPRK-phosphorylated receptor and sterically preclude
	its coupling to the cognate G-protein; the binding appears to require additional receptor
	determinants exposed only in the active receptor conformation. The beta-arrestins target many
	receptors for internalization by acting as endocytic adapters (CLASPs, clathrin-associated sorting
	proteins) and recruiting the GPRCs to the adapter protein 2 complex 2 (AP-2) in clathrin-coated
	pits (CCPs). However, the extent of beta-arrestin involvement appears to vary significantly
	depending on the receptor, agonist and cell type. Internalized arrestin-receptor complexes traffic
	to intracellular endosomes, where they remain uncoupled from G-proteins. Two different modes of
	arrestin-mediated internalization occur. Class A receptors, like ADRB2, OPRM1, ENDRA, D1AR
	and ADRA1B dissociate from beta-arrestin at or near the plasma membrane and undergo rapid
	recycling. Class B receptors, like AVPR2, AGTR1, NTSR1, TRHR and TACR1 internalize as a
	complex with arrestin and traffic with it to endosomal vesicles, presumably as desensitized
	receptors, for extended periods of time. Receptor resensitization then requires that receptor-
	bound arrestin is removed so that the receptor can be dephosphorylated and returned to the plasma membrane. Mediates endocytosis of CCR7 following ligation of CCL19 but not CCL21.
	Involved in internalization of P2RY1, P2RY4, P2RY6 and P2RY11 and ATP-stimulated
	internalization of P2RY2. Involved in phopshorylation-dependent internalization of OPRD1 and
	subsequent recycling or degradation. Involved in ubiquitination of IGF1R. Beta-arrestins function
	as multivalent adapter proteins that can switch the GPCR from a G-protein signaling mode that
	transmits short-lived signals from the plasma membrane via small molecule second messengers
	and ion channels to a beta-arrestin signaling mode that transmits a distinct set of signals that are
	initiated as the receptor internalizes and transits the intracellular compartment. Acts as signaling
	scaffold for MAPK pathways such as MAPK1/3 (ERK1/2) and MAPK10 (JNK3). ERK1/2 and
	JNK3 activated by the beta-arrestin scaffold are largely excluded from the nucleus and confined to
	cytoplasmic locations such as endocytic vesicles, also called beta-arrestin signalosomes. Acts as
	signaling scaffold for the AKT1 pathway. GPCRs for which the beta-arrestin-mediated signaling
	relies on both ARRB1 and ARRB2 (codependent regulation) include ADRB2, F2RL1 and PTH1R.
	For some GPCRs the beta-arrestin-mediated signaling relies on either ARRB1 or ARRB2 and is
	inhibited by the other respective beta-arrestin form (reciprocal regulation). Increases ERK1/2
	signaling in AGTR1- and AVPR2-mediated activation (reciprocal regulation). Involved in CCR7-
	mediated ERK1/2 signaling involving ligand CCL19. Is involved in type-1A angiotensin II
	receptor/AGTR1-mediated ERK activity. Is involved in type-1A angiotensin II receptor/AGTR1- mediated MAPK10 activity. Is involved in dopamine-stimulated AKT1 activity in the striatum by
	disrupting the association of AKT1 with its negative regulator PP2A. Involved in AGTR1-mediated
	chemotaxis. Appears to function as signaling scaffold involved in regulation of MIP-1-beta-
	stimulated CCR5-dependent chemotaxis. Involved in attenuation of NF-kappa-B-dependent
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	transcription in response to GPCR or cytokine stimulation by interacting with and stabilizing CHUK. Suppresses UV-induced NF-kappa-B-dependent activation by interacting with CHUK. The function is promoted by stimulation of ADRB2 and dephosphorylation of ARRB2. Involved in p53/TP53-mediated apoptosis by regulating MDM2 and reducing the MDM2-mediated degradation of p53/TP53. May serve as nuclear messenger for GPCRs. Upon stimulation of OR1D2, may be involved in regulation of gene expression during the early processes of fertilization. Also involved in regulation of receptors others than GPCRs. Involved in endocytosis of TGFBR2 and TGFBR3 and down-regulates TGF-beta signaling such as NF-kappa-B activation. Involved in endocytosis of low-density lipoprotein receptor/LDLR. Involved in endocytosis of smoothened homolog/Smo, which also requires ADRBK1. Involved in endocytosis of SLC9A5. Involved in endocytosis of ENG and subsequent TGF-beta-mediated ERK activation and migration of epithelial cells. Involved in Toll-like receptor and IL-1 receptor signaling through the interaction with TRAF6 which prevents TRAF6 autoubiquitination and oligomerization required for activation of NF-kappa-B and JUN. Involved in insulin resistence by acting as insulin-induced signaling scaffold for SRC, AKT1 and INSR. Involved in regulation of inhibitory signaling of natural killer cells by recruiting PTPN6 and PTPN11 to KIR2DL1.
Sequence similarities	Belongs to the arrestin family.
Domain	The [DE]-X(1,2)-F-X-X-[FL]-X-X-R motif mediates interaction the AP-2 complex subunit AP2B1.
Post-translational modifications	Phosphorylated at Thr-382 in the cytoplasm; probably dephosphorylated at the plasma membrane. The phosphorylation does not regulate internalization and recycling of ADRB2, interaction with clathrin or AP2B1. The ubiquitination status appears to regulate the formation and trafficking of beta-arrestin-GPCR complexes and signaling. Ubiquitination appears to occurr GPCR-specifc. Ubiquitinated by MDM2; the ubiquitination is required for rapid internalization of ADRB2. Deubiquitinated by USP33; the deubiquitination leads to a dissociation of the beta-arrestin-GPCR complex. Stimulation of a class A GPCR, such as ADRB2, induces transient ubiquitination and subsequently promotes association with USP33. Stimulation of a class B GPCR promotes a sustained ubiquitination.
Cellular localization	Cytoplasm. Nucleus. Cell membrane. Membrane > clathrin-coated pit. Cytoplasmic vesicle. Translocates to the plasma membrane and colocalizes with antagonist-stimulated GPCRs.

Images



protein (ab132309)

12.5% SDS-PAGE stained with Coomassie Blue showing ab132309.

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