CREB binding protein (catalytic domain) (human), (recombinant)

Although originally termed histone acetyltransferases (HATs) for their lysine acetylation activity on histone N-terminal tails, CBP and its paralogue, p300, have been shown to acetylate a variety of non-histone proteins including p53, DNA polymerase β and nuclear import factors. p300/CBP acetylations play regulatory roles in transcription, DNA repair and replication, the cell cycle, p53 turnover, and nuclear import. In addition to functions which overlap with those of p300, CBP has a number of unique functions, including a role in p27Kip1 induction.

Citations: 5

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Ordering Information

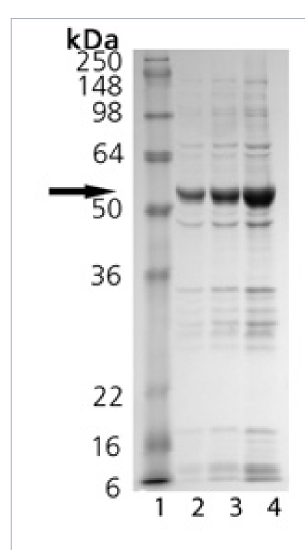
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BML-SE452-0100

100µg

Manuals, SDS & CofA

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SDS-PAGE Analysis: Lane 1: MW Marker, Lane 2: 1 μg ,

Lane 3: 2 µg, Lane 4: 5 µg CBP.

Handling & Storage

Handling Avoid freeze/thaw cycles. After opening, prepare aliquots and store at -80°C.

Long Term Storage -80°C

Shipping Dry Ice

Regulatory Status RUO - Research Use Only

Product Details

Alternative Name CBP

Application NotesCan be used to study CBP kinetics and regulation, for

screening for inhibitors or activators and for radiolabeling of proteins or peptides with, for example [³H]-Acetyl CoA.

Formulation Liquid. In 50mM TRIS-HCl, pH 8.0, containing 0.1mM

EDTA, 10% glycerol.

MW 45.4 kDa

Purity Detail Partially purified by single-step affinity chromatography

and gel filtration.

Source Produced in *E. coli*. Catalytic domain (aa 1319-1710) of

human CBP.

Specific Activity ≥250 pmol/min/µq assayed as production of CoA-SH from

AcCoA in the presence of a peptide comprising human p53 residues 368-386 (Prod. No. BML-P198). CoA-SH is determined colorimetrically by reaction with DTNB (5,5'-

Dithiobis(2-nitrobenzoic acid)).

UniProt ID Q92793

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ENZO LIFE SCIENCES, INC. Phone: 800.942.0430 infousa@enzolifesciences.com European Sales Office ENZO LIFE SCIENCES (ELS) AG Phone: +41 61 926 8989 infoeu@enzolifesciences.com Belgium, The Netherlands & Luxembourg Phone: +32 3 466 0420 infobe@enzolifesciences.com

France
Phone: +33 472 440 655
infofr@enzolifesciences.com

Germany Phone: +49 7621 5500 526 infode@enzolifesciences.com UK & Ireland
Phone (UK customers):
0845 601 1488
Phone: +44 1392 825900
infouk@enzolifesciences.com