

RPC984Hu01 100µg Recombinant Poly ADP Ribose Polymerase 4 (PARP4) Organism Species: *Homo sapiens (Human) Instruction manual*

FOR RESEARCH USE ONLY

NOT FOR USE IN CLINICAL DIAGNOSTIC PROCEDURES

12th Edition (Revised in Aug, 2016)

Coud-Clone Corp.

[PROPERTIES]

Source: Prokaryotic expression Host: E.coli Residues: Val338~Thr609 Tags: N-terminal His Tag Subcellular Location: Nucleus, Cytoplasm **Purity:** > 95% Traits: Freeze-dried powder Buffer formulation: 20mM Tris, 150mM NaCl, pH8.0, containing 0.01% SKL, 5% Trehalose. Original Concentration: 120µg/mL Applications: Positive Control; Immunogen; SDS-PAGE; WB. (May be suitable for use in other assays to be determined by the end user.) Predicted isoelectric point: 5.9 Predicted Molecular Mass: 34.2kDa Accurate Molecular Mass: 40kDa as determined by SDS-PAGE reducing conditions. Phenomenon explanation: The possible reasons that the actual band size differs from the predicted are as follows: 1.Splice variants: Alternative splicing may create different sized proteins from the same gene. 2. Relative charge: The composition of amino acids may affects the charge of the protein. 3. Post-translational modification: Phosphorylation, glycosylation, methylation etc. 4. Post-translation cleavage: Many proteins are synthesized as pro-proteins, and then cleaved to give the active form. 5. Polymerization of the target protein: Dimerization, multimerization etc. [USAGE] Reconstitute in 20mM Tris, 150mM NaCI (pH8.0) to a concentration of 0.1-1.0 mg/mL. Do not

[STORAGE AND STABILITY]

vortex.



Storage: Avoid repeated freeze/thaw cycles.

Store at 2-8°C for one month.

Aliquot and store at -80°C for 12 months.

Stability Test: The thermal stability is described by the loss rate. The loss rate was determined

by accelerated thermal degradation test, that is, incubate the protein at 37°C for 48h, and no

obvious degradation and precipitation were observed. The loss rate is less than 5% within the

expiration date under appropriate storage condition.

[SEQUENCE]

VNL GLLAKKADLC QLIRDMVNVC ETNLSKPNPP SLAKYRALRC KIEHVEQNTE EFLRVRKEVL QNHHSKSPVD VLQIFRVGRV NETTEFLSKL GNVRPLLHGS PVQNIVGILC RGLLLPKVVE DRGVQRTDVG NLGSGIYFSD SLSTSIKYSH PGETDGTRLL LICDVALGKC MDLHEKDFSL TEAPPGYDSV HGVSQTASVT TDFEDDEFVV YKTNQVKMKY IIKFSMPGDQ IKDFHPSDHT ELEEYRPEFS NFSKVEDYQL PDAKTSSST

[IDENTIFICATION]

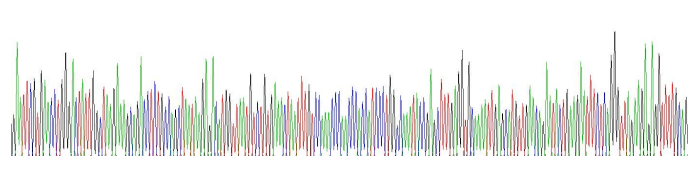
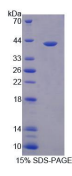


Figure . Gene Sequencing (extract)



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[<u>IMPORTANT NOTE</u>]

The kit is designed for research use only, we will not be responsible for any issue if the kit was used in clinical diagnostic or any other procedures.