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Datasheet for W09-001-A60

A431 Cell Nuclear Extract

Overview

| Description: | A431 Cell Nuclear Extract - W09-001-A60 |
|--------------|---|
| Item No.: | W09-001-A60 |
| Size: | 200 μg |
| Origin: | Human |

Product Details

| Product Details | |
|--------------------|--|
| Background: | Ready-to-use nuclear extracts produced by Rockland Immunochemicals are derived from cell lines or tissues using highly refined extraction protocols to ensure exceptionally high quality, protein integrity and lot-to-lot reproducibility. All extracts are tested by SDS-PAGE using 4-20% gradient gels and immunoblot analysis using antibodies to key cell signaling components to confirm the presence of both high molecular weight and low molecular weight proteins. |
| Synonyms: | A431 Cell Nuclear Extract, A431 Nuclear Lysate, A431 Lysate Nuclear Extract |
| Species of Origin: | Human |
| | |

Target Details

Purity/Specificity:

The cells were grown in DMEM supplemented with 10% FBS (Fetal Bovine Serum). The lysate was prepared by first washing the cells in PBS. Washed cells were then incubated on ice in lysis buffer containing 10 mM HEPES, 60 mM KCl, 1.0 mM EDTA, 0.075% (v/v) NP40 and 1.0 mM DTT, pH 7.6. Protein integrity is ensured using a cocktail of protease inhibitors with broad specificity for the inhibition of aspartic, cysteine, and serine proteases as well as aminopeptidases (0.1 mM AEBSF HCl, 0.08 μ M Aprotinin, 5 μ M Bestatin, 1.5 μ M E-64, 2 μ M Leupeptin Hemisulfate and 1 μ M Pepstatin A). Nuclei were then collected and washed in lysis buffer minus detergent. Nuclei were lysed by vortexing in extraction buffer containing 20 mM Tris-Cl, 1.5 mM MgCl2, 0.42 M NaCl, 0.2 mM EDTA, and 25% (v/v) glycerol, pH 8.0, supplemented with protease inhibitors (see above). The lysate was clarified by centrifugation. Protein concentration was determined by Lowry assay using a commercially available kit. The protein concentration was adjusted to 2.0 mg/ml and then an equal volume of 2X SDS-PAGE sample buffer was added.

Application Details

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| Application Note: | Ready-to-use nuclear extracts are especially prepared as positive controls for separation by SDS-PAGE and subsequent western blot analysis. Nuclear extracts are supplied in denaturing buffer without dissociating agents. Heat nuclear extract to 95° C for 5 minutes and rapidly cool. If dissociating conditions are desired add reducing agent prior to heating. The recommended loading volume per lane is 10-30 🛽 depending on the size format of your gel. |
|-------------------|--|
| Assay Dilutions: | All assays should be optimized by the user. Recommended dilutions (if any) may be listed below. |
| WB: | User Optimized |

Cell Line Data

| Cell Line: | Human A431 (epidermoid carcinoma) |
|-----------------------|-----------------------------------|
| Lysate Fractionation: | Nuclear Extract |
| Lysate Stimulation: | Not Stimulated |
| Culture Type: | Tissue Culture |
| Induction: | None (Control) |

Formulation

| Physical State: | Liquid (sterile filtered) |
|-----------------|--|
| Concentration: | 1.0 mg/mL by modified Lowry assay |
| Buffer: | 1X SDS-PAGE Sample Buffer (62.5 mM Tris HCl, 2% SDS, 10% Glycerol and 0.005% bromophenol blue, pH 6.8) |
| Preservative: | None |

Shipping & Handling

| Shipping Condition: | Dry Ice |
|----------------------------|--|
| Storage Condition: | Store vial at -70 $^{\circ}$ C or COLDER. For extended storage, aliquot contents to minimize freeze/thaw cycles. |
| Expiration: | Expiration date is three (3) months from date of receipt. |

Disclaimer

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