

## **Product Datasheet**

HT-1080 Cell Nuclear Extract (orb348707)



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**Description**nts. HT-1080 Cell Nuclear Extract

Conjugation Unconjugated

Tested
Applications

ChIP, IP, WB

**Preservatives** Preservative: None. Stabilizer: None. 1X

RIPA Buffer with HALT Protease and

**Phosphatase Inhibitors** 

Form/Appearance Liquid (sterile filtered)

**Concentration** 4.0 mg/ml

**Storage** Store vial at -70° C or COLDER. For

extended storage, aliquot Nuclear Extract

to minimize freeze/thaw cycles.

**Note** For research use only

**Application notes** Multi-purpose HT-1080 nuclear extracts are

especially prepared as positive control for multiple assays including western blot, immunoprecipitation (IP), capture ELISA or other assays requiring native protein sample. For separation by SDS-PAGE and subsequent western blot analysis, lysates should be diluted by user to desired concentration in SDS-PAGE buffer with 2-mercaptoethanol or dithiothreitol as the reducing agent and heated to 95° C for 5 minutes. Sample is ready for use in immunoprecipitation and ELISA experiments, conditions should be optimized by the user. Rockland recommends its TrueBlot IP reagents for

immunoprecipitation experiments.

**Purity** HT-1080 cells were grown in Eagle's

Minimum Essential Medium supplemented with 10% fetal bovine serum. Cells were washed with PBS and then incubated on ice in modified RIPA buffer, containing 150 mM sodium chloride, 50 mM Tris HCl, pH 7.4, 1 mM EDTA, 1.0% NP-40, 0.5% sodium deoxycholic acid , 0.1% SDS and 0.01% (w/v) sodium azide to lyse the cells. Protein integrity was 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  $\mu$ M Aprotinin, 5  $\mu$ M Bestatin, 1.5  $\mu$ M E-64, 2

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Biorbyt Ltd.

7 Signet Court, Swann's Road, Cambridge, CB5 8LA, United Kingdom
Email: info@biorbyt.com | Phone: +44 (0) 1223 859-353 | Fax: +44 (0)1223 280

Biorbyt LLC.

68 TW Alexander Drive<br>Research Triangle Park<br>Durham, North Carolina<br/>br>27709. United States

 ${\sf Email: info@biorbyt.com \mid Phone: +1 (415) \ 906-5211 \mid Fax: +1 \ (415) \ 651-8558}$