



UK Office

Everest Biotech Ltd

Cherwell Innovation Centre
77 Heyford Park
Upper Heyford
Oxfordshire
OX25 5HD
UK

Enquiries:

info@everestbiotech.com

Sales:

sales@everestbiotech.com

Tech support:

support@everestbiotech.com

Tel: +44 (0)1869 238326

www.everestbiotech.com

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diagnostic or therapeutic use.**

EB11539 - Goat Anti-Hsp31p (yeast, aa50-64) Antibody

Size: 100µg specific antibody in 200µl



Target Protein

Principal Names: Hsp31p, YDR533C

Official Symbol: HSP31

Accession Number(s): NP_010822.1

Immunogen

Peptide with sequence C-ETGKFGWDEHSLAKD, from the internal region of the protein sequence according to NP_010822.1.

Please note the [peptide](#) is available for sale.

Purification and Storage

Purified from goat serum by ammonium sulphate precipitation followed by antigen affinity chromatography using the immunizing peptide.

Supplied at 0.5 mg/ml in Tris saline, 0.02% sodium azide, pH7.3 with 0.5% bovine serum albumin.

Aliquot and store at -20°C. Minimize freezing and thawing.

Applications Tested

Peptide ELISA: antibody detection limit dilution 1:128000.

Western blot: Approx 24kDa band observed in lysates of postdiauxic (overnight grown yeast cells, and no signal in KO grown under identical conditions. Cells grown in logarithmic phase show diminished signals as expected. (calculated MW of 25.7kDa according to NP_010822.1). Recommended concentration: 0.02-0.06µg/ml. This antibody has been successfully used in the following paper: Natka█ska et al (2018) PMID: 29264711.

Species Reactivity

Tested: *S. cerevisiae*

Expected from sequence similarity: *Saccharomyces cerevisiae* S288c

Specific Reference

This antibody has been successfully used in Redox WB on Yeast:

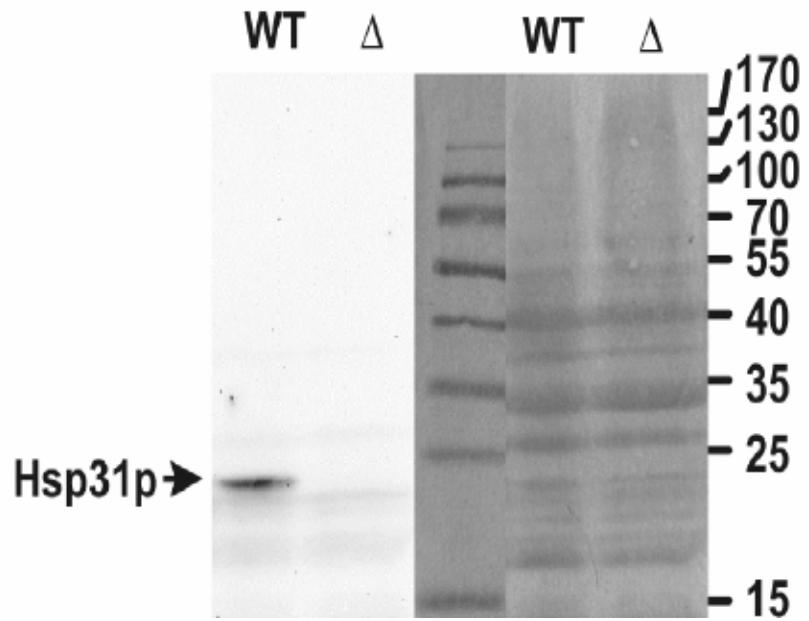
Urszula Natka█ska, Adrianna Skoneczna and Marek Skoneczny

Oxidative stress triggers aggregation of GFP-tagged Hsp31p, the budding yeast environmental stress response chaperone, and glyoxalase III

Cell Stress and Chaperones (2018) 23:595–607

PMID: 29264711

Postdiauxic (overnight grown) cells



EB11539 (0.05 μ g/ml) staining of yeast lysate (wt) and the KO (delta) in the left panel and the Ponceau stain in the right panel (35 μ g protein in RIPA buffer). Primary incubation was 1 hour. Detected by chemiluminescence. Data obtained by Dr. M. Skoneczny, IBB, Warsaw, Poland.