

## Produktinformation



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Zellkultur & Verbrauchsmaterial



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## Lieferung & Zahlungsart

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# GSS (m): 293T Lysate: sc-126926



The Power to Question

#### **BACKGROUND**

GSS (glutathione synthetase) is a 474 amino acid protein encoded by the gene located at human chromosome 20q11.22. GSS consists of three loops projecting from an antiparallel  $\beta$ -sheet, a parallel  $\beta$ -sheet and a lid of antiparallel sheets, which provide access to the ATP-binding site. Although Southern blot and gene analysis suggest that GSS may be the only member of a unique family, the crystal structure indicates that GSS belongs to the ATP-GRASP superfamily. GSS is expressed in hemocytes and nucleated cells, including the brain. GSS occurs as a homodimer. There are two steps in the production of glutathione, begining with γ-GCS and ending with GSS. In an ATP-dependent reaction, GSS produces glutathione from γ-glutamylcysteine and glycine precursors. Partial hepatectomy, diethyl maleate, buthionine sulfoximine, tert-butylhaydroquinone and thioacetamide increase the expression of GSS, which causes an increase in glutathione levels. An inherited autosomal recessive disorder, 5-oxoprolinuria (pyroglutamic aciduria), is caused by GSS deficiencies, which leads to central nervous system damage, hemolytic anemia, metabolic acidosis and urinary excretion of 5-oxoproline. A missense mutation in the gene encoding GSS leads to a GSS deficiency restricted to erythrocytes, which causes only hemolytic anemia.

#### **REFERENCES**

- Webb, G.C., Vaska, V.L., Gali, R.R., Ford, J.H. and Board, P.G. 1995. The gene encoding human glutathione synthetase (GSS) maps to the long arm of chromosome 20 at band 11.2. Genomics 30: 617-619.
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- Shi, Z.Z., Habib, G.M., Rhead, W.J., Gahl, W.A., He, X., Sazer, S. and Lieberman, M.W. 1996. Mutations in the glutathione synthetase gene cause 5-oxoprolinuria. Nat. Genet. 14: 361-365.
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  Molecular basis of glutathione synthetase deficiency and a rare gene permutation event. EMBO J. 18: 3204-3213.
- 5. Huang, Z.A., Yang, H., Chen, C., Zeng, Z. and Lu, S.C. 2000. Inducers of  $\gamma$ -glutamylcysteine synthetase and their effects on glutathione synthetase expression. Biochim. Biophys. Acta 1493: 48-55.

#### CHROMOSOMAL LOCATION

Genetic locus: Gss (mouse) mapping to 2 H1.

#### **PRODUCT**

GSS (m): 293T Lysate represents a lysate of mouse GSS transfected 293T cells and is provided as 100  $\mu$ g protein in 200  $\mu$ l SDS-PAGE buffer.

#### **STORAGE**

Store at -20° C. Repeated freezing and thawing should be minimized. Sample vial should be boiled once prior to use. Non-hazardous. No MSDS required.

#### **PROTOCOLS**

See our web site at www.scbt.com for detailed protocols and support products.

#### **APPLICATIONS**

GSS (m): 293T Lysate is suitable as a Western Blotting positive control for mouse reactive GSS antibodies. Recommended use: 10-20 µl per lane.

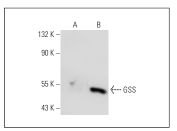
Control 293T Lysate: sc-117752 is available as a Western Blotting negative control lysate derived from non-transfected 293T cells.

GSS (H-7): sc-166882 is recommended as a positive control antibody for Western Blot analysis of enhanced mouse GSS expression in GSS transfected 293T cells (starting dilution 1:100, dilution range 1:100-1:1,000).

#### **RECOMMENDED SUPPORT REAGENTS**

To ensure optimal results, the following support reagents are recommended: 1) Western Blotting: use m-lgG $\kappa$  BP-HRP: sc-516102 or m-lgG $\kappa$  BP-HRP (Cruz Marker): sc-516102-CM (dilution range: 1:1000-1:10000), Cruz Marker<sup>TM</sup> Molecular Weight Standards: sc-2035, UltraCruz<sup>®</sup> Blocking Reagent: sc-516214 and Western Blotting Luminol Reagent: sc-2048.

#### **DATA**



GSS (H-7): sc-166882. Western blot analysis of GSS expression in non-transfected: sc-117752 (**A**) and mouse GSS transfected: sc-126926 (**B**) 293T whole cell lysates.

#### **RESEARCH USE**

For research use only, not for use in diagnostic procedures.

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