

Phosphoserine Detection Kit

Order No.: 0701/PSER-KIT



www.nanotools.de

orders & support:

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02/080507

Background and Specificity

Phosphorylation and dephosphorylation of cellular proteins are central steps in transducing extracellular signals to the nucleus. Phosphorylated epitopes may serve as docking sites for the assembly of protein complexes or may alter the 3-dimensional protein structure thus modulating enzymatic activity or the ability to undergo protein-protein interactions.

Modification of proteins on tyrosine residues is mediated by protein tyrosin kinases. Tyrosine phosphorylation may alter the biological activity or mediate the assembly of protein complexes via the interaction of phosphotyrosine residues with SH2 or PTB domains.

Antibodies direct against phosphorylated epitopes recognize the phosphorylated amino acid in the context of the surrounding amino acid sequence. Recognition is therefore dependent on 2 criteria: 1) phosphorylation and 2) the surrounding amino acid motif. If one of the two criteria is not fulfilled, the antibody will not detect the phosphorylation site. Since the amino acid sequence varies between different phosphorylation sites, certain proteins - though phosphorylated - may not be detected by the antibody. Phosphorylation patterns in a given cell extract may differ when probed with different antibodies due to sequence specificity.

The Phosphoserine Detection Kit contains 6 different phosphotyrosine specific monoclonal antibodies.

Do not use Milk or Casein based blocking and incubation buffers.

clone	isotype	order number
1C8	IgM	0018-025
4A3	IgM	0019-025
4A9	IgM	0020-025
4H4	IgM	0021-025
7F12	IgG1	0022-025
16B4	IgM	0023-025

Postive control

This product contains the following positive control for immunoblot applications:

#0901-PSRECO phosphoproteins from rabbit muscle

Mouse Monoclonal Antibody to

Phosphoserine

clone 1C8

Order No.: 0018-025/PSER-1C8

Size (µg) 25

Lot No.: 0018S



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03/160307F

Isotype	Species Reactivity	Applications	Mol. Weight	Ref.Cell Line	Epitope	Immunogen
IgM	human, mouse, rat, dog	WB, ELISA, IP	pattern			phosphoserine conjugated to KLH

Background and Specificity:

Phosphorylation and dephosphorylation of cellular proteins are central steps in transducing extracellular signals to the cell nucleus. Phosphorylated epitopes may serve as docking sites for the assembly of protein complexes or may alter the 3-dimensional protein structure thus modulating enzymatic activity or the ability to undergo protein-protein-interactions. Modification of proteins on serine residues is mediated by serine/threonine kinases.

Please note that phosphoserine detection by monoclonal antibodies is always dependent on the surrounding amino acid sequence!

Mab PSER-1C8 recognizes a broad range of serine-phosphorylated proteins in crude cell extracts, preferring positively charged amino acids directly neighboured to phosphoserine.

Related Products

mab against Phosphoserine

#0019-100/pSer-4A3

#0020-100/pSer-4A9

#0021-100/pSer-4H4

#0022-100/pSer-7F12

#0023-100/pSer-16B4

mab against Phosphothreonine

#0024-100/pThr-1E11

#0025-100/pThr-4D11

#0026-100/pThr-14B3

Purification: The antibody was purified from serum-free cell culture supernatant by subsequent thiophilic adsorption and size exclusion chromatography.

Formulation: lyophilized from 1 ml 2 x PBS / 0.09 % Na-azide / PEG and Sucrose.

Reconstitution: Reconstitute with 1 ml H₂O (15 min, RT).

Stability: For long-term storage, freeze lyophilizate upon arrival (-20°C). Upon reconstitution, aliquote and freeze in liquid nitrogen; reconstituted antibody can be stored frozen at -80°C up to 1 year. Thaw aliquots at 37°C. Thawed aliquots may be stored at 4°C up to 1 week.

Avoid repeated freeze / thaw cycles.

Positive Control: #0901: phosphoserine/phosphothreonine positive control

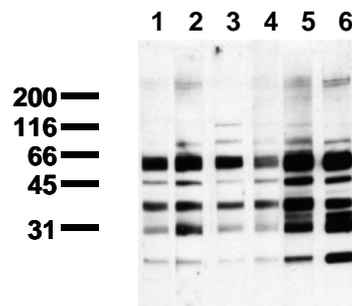
Immunoblotting: 1 µg/ml for HRPO/ECL detection
Recommended blocking buffer: BSA/Tween 20 based blocking buffer.
DO NOT USE MILK OR CASEIN FOR BLOCKING!

Immunoprecipitation: use at 1 - 10 µg per 10⁶ pervanadate-treated A431 cells

Immunocytochemistry: ND

ELISA: use at 0.05 µg/ml

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Phosphoserine Detection

Phosphoprotein Positive Control was probed with

lane 1: mab 1C8 (IgM), 1 µg/ml

lane 2: mab 4A3 (IgM), 1 µg/ml

lane 3: mab 4A9 (IgM), 1 µg/ml

lane 4: mab 4H4 (IgM), 1 µg/ml

lane 5: mab 7F12 (IgG), 1 µg/ml

lane 6: mab 16B4 (IgM), 1 µg/ml

Mouse Monoclonal Antibody to

Phosphoserine

clone 4A3

Order No.: 0019-025/PSER-4A3

Size (µg) 25

Lot No.: 0019S



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03/160307F

Isotype	Species Reactivity	Applications	Mol. Weight	Ref.Cell Line	Epitope	Immunogen
IgM	human, mouse, rat, dog	WB, ELISA, IP	pattern			phosphoserine conjugated to KLH

Background and Specificity:

Phosphorylation and dephosphorylation of cellular proteins are central steps in transducing extracellular signals to the cell nucleus. Phosphorylated epitopes may serve as docking sites for the assembly of protein complexes or may alter the 3-dimensional protein structure thus modulating enzymatic activity or the ability to undergo protein-protein-interactions. Modification of proteins on serine residues is mediated by serine/threonine kinases.

Please note that phosphoserine detection by monoclonal antibodies is always dependent on the surrounding amino acid sequence!

Mab PSER-4A3 recognizes a broad range of serine-phosphorylated proteins in crude cell extracts, preferring positively charged amino acids directly neighbored to phosphoserine.

Related Products

mab against Phosphoserine

#0018-100/pSer-1C8

#0020-100/pSer-4A9

#0021-100/pSer-4H4

#0022-100/pSer-7F12

#0023-100/pSer-16B4

mab against Phosphothreonine

#0024-100/pThr-1E11

#0025-100/pThr-4D11

#0026-100/pThr-14B3

Purification: The antibody was purified from serum-free cell culture supernatant by subsequent thiophilic adsorption and size exclusion chromatography.

Formulation: lyophilized from 1 ml 2 x PBS / 0.09 % Na-azide / PEG and Sucrose.

Reconstitution: Reconstitute with 1 ml H₂O (15 min, RT).

Stability: For long-term storage, freeze lyophilizate upon arrival (-20°C). Upon reconstitution, aliquote and freeze in liquid nitrogen; reconstituted antibody can be stored frozen at -80°C up to 1 year. Thaw aliquots at 37°C. Thawed aliquots may be stored at 4°C up to 1 week.

Avoid repeated freeze / thaw cycles.

Positive Control: #0901: phosphoserine/phosphothreonine positive control

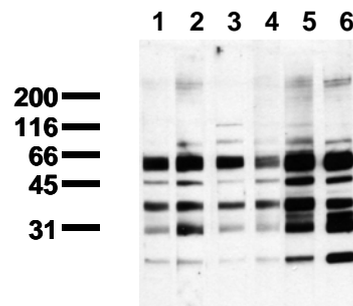
Immunoblotting: 1 µg/ml for HRPO/ECL detection
Recommended blocking buffer: BSA/Tween 20 based blocking buffer.
DO NOT USE MILK OR CASEIN FOR BLOCKING!

Immunoprecipitation: use at 1 - 10 µg per 10⁶ pervanadate-treated A431 cells

Immunocytochemistry: ND

ELISA: use at 0.05 µg/ml

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Phosphoserine Detection

Phosphoprotein Positive Control was probed with

lane 1: mab 1C8 (IgM), 1 µg/ml

lane 2: mab 4A3 (IgM), 1 µg/ml

lane 3: mab 4A9 (IgM), 1 µg/ml

lane 4: mab 4H4 (IgM), 1 µg/ml

lane 5: mab 7F12 (IgG), 1 µg/ml

lane 6: mab 16B4 (IgM), 1 µg/ml

Mouse Monoclonal Antibody to

Phosphoserine

clone 4A9

Order No.: 0020-025/PSER-4A9

Size (µg) 25

Lot No.: 0020S



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03/160307F

Isotype	Species Reactivity	Applications	Mol. Weight	Ref.Cell Line	Epitope	Immunogen
IgM	human, mouse, rat, dog	WB, ELISA, IP	pattern			phosphoserine conjugated to KLH

Background and Specificity:

Phosphorylation and dephosphorylation of cellular proteins are central steps in transducing extracellular signals to the cell nucleus. Phosphorylated epitopes may serve as docking sites for the assembly of protein complexes or may alter the 3-dimensional protein structure thus modulating enzymatic activity or the ability to undergo protein-protein-interactions. Modification of proteins on serine residues is mediated by serine/threonine kinases.

Please note that phosphoserine detection by monoclonal antibodies is always dependent on the surrounding amino acid sequence!

Mab PSER-4A9 recognizes a broad range of serine-phosphorylated proteins in crude cell extracts, preferring positively charged amino acids directly neighboured to phosphoserine.

Related Products

mab against Phosphoserine

#0018-100/pSer-1C8

#0019-100/pSer-4A3

#0021-100/pSer-4H4

#0022-100/pSer-7F12

#0023-100/pSer-16B4

mab against Phosphothreonine

#0024-100/pThr-1E11

#0025-100/pThr-4D11

#0026-100/pThr-14B3

Purification: The antibody was purified from serum-free cell culture supernatant by subsequent thiophilic adsorption and size exclusion chromatography.

Formulation: lyophilized from 1 ml 2 x PBS / 0.09 % Na-azide / PEG and Sucrose.

Reconstitution: Reconstitute with 1 ml H₂O (15 min, RT).

Stability: For long-term storage, freeze lyophilizate upon arrival (-20°C). Upon reconstitution, aliquote and freeze in liquid nitrogen; reconstituted antibody can be stored frozen at -80°C up to 1 year. Thaw aliquots at 37°C. Thawed aliquots may be stored at 4°C up to 1 week.

Avoid repeated freeze / thaw cycles.

Positive Control: #0901: phosphoserine/phosphothreonine positive control

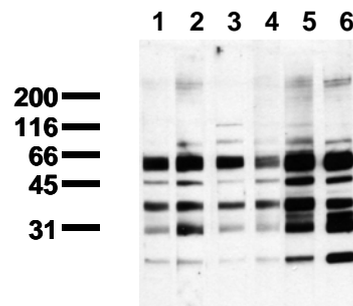
Immunoblotting: 1 µg/ml for HRPO/ECL detection
Recommended blocking buffer: BSA/Tween 20 based blocking buffer.
DO NOT USE MILK OR CASEIN FOR BLOCKING!

Immunoprecipitation: use at 1 - 10 µg per 10⁶ pervanadate-treated A431 cells

Immunocytochemistry: ND

ELISA: use at 0.05 µg/ml

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Phosphoserine Detection

Phosphoprotein Positive Control was probed with

lane 1: mab 1C8 (IgM), 1 µg/ml

lane 2: mab 4A3 (IgM), 1 µg/ml

lane 3: mab 4A9 (IgM), 1 µg/ml

lane 4: mab 4H4 (IgM), 1 µg/ml

lane 5: mab 7F12 (IgG), 1 µg/ml

lane 6: mab 16B4 (IgM), 1 µg/ml

Mouse Monoclonal Antibody to

Phosphoserine

clone 4H4

Order No.: 0021-025/PSER-4H4

Size (µg) 25

Lot No.: 0021S



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Isotype	Species Reactivity	Applications	Mol. Weight	Ref.Cell Line	Epitope	Immunogen
IgM	human, mouse, rat, dog	WB, ELISA, IP	pattern			phosphoserine conjugated to KLH

Background and Specificity:

Phosphorylation and dephosphorylation of cellular proteins are central steps in transducing extracellular signals to the cell nucleus. Phosphorylated epitopes may serve as docking sites for the assembly of protein complexes or may alter the 3-dimensional protein structure thus modulating enzymatic activity or the ability to undergo protein-protein-interactions. Modification of proteins on serine residues is mediated by serine/threonine kinases.

Please note that phosphoserine detection by monoclonal antibodies is always dependent on the surrounding amino acid sequence!

Mab PSER-4H4 recognizes a broad range of serine-phosphorylated proteins in crude cell extracts, preferring positively charged amino acids directly neighboured to phosphoserine.

Related Products

mab against Phosphoserine

#0018-100/pSer-1C8

#0019-100/pSer-4A3

#0020-100/pSer-4A9

#0022-100/pSer-7F12

#0023-100/pSer-16B4

mab against Phosphothreonine

#0024-100/pThr-1E11

#0025-100/pThr-4D11

#0026-100/pThr-14B3

Purification: The antibody was purified from serum-free cell culture supernatant by subsequent thiophilic adsorption and size exclusion chromatography.

Formulation: lyophilized from 1 ml 2 x PBS / 0.09 % Na-azide / PEG and Sucrose.

Reconstitution: Reconstitute with 1 ml H₂O (15 min, RT).

Stability: For long-term storage, freeze lyophilizate upon arrival (-20°C). Upon reconstitution, aliquote and freeze in liquid nitrogen; reconstituted antibody can be stored frozen at -80°C up to 1 year. Thaw aliquots at 37°C. Thawed aliquots may be stored at 4°C up to 1 week.

Avoid repeated freeze / thaw cycles.

Positive Control: #0901: phosphoserine/phosphothreonine positive control

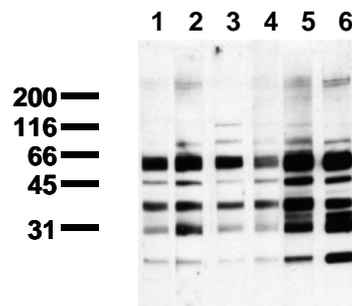
Immunoblotting: 1 µg/ml for HRPO/ECL detection
Recommended blocking buffer: BSA/Tween 20 based blocking buffer.
DO NOT USE MILK OR CASEIN FOR BLOCKING!

Immunoprecipitation: use at 1 - 10 µg per 10⁶ pervanadate-treated A431 cells

Immunocytochemistry: ND

ELISA: use at 0.05 µg/ml

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Phosphoserine Detection

Phosphoprotein Positive Control was probed with

lane 1: mab 1C8 (IgM), 1 µg/ml

lane 2: mab 4A3 (IgM), 1 µg/ml

lane 3: mab 4A9 (IgM), 1 µg/ml

lane 4: mab 4H4 (IgM), 1 µg/ml

lane 5: mab 7F12 (IgG), 1 µg/ml

lane 6: mab 16B4 (IgM), 1 µg/ml

Mouse Monoclonal Antibody to

Phosphoserine

clone 7F12

Order No.: 0022-025/PSER-7F12

Size (µg) 25

Lot No.: 0022S

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fax: +49-7641-455 671



03/160307F

Isotype	Species Reactivity	Applications	Mol. Weight	Ref.Cell Line	Epitope	Immunogen
IgG1	human, mouse, rat, dog	WB, ELISA, IP	pattern			phosphoserine conjugated to KLH

Background and Specificity:

Phosphorylation and dephosphorylation of cellular proteins are central steps in transducing extracellular signals to the cell nucleus. Phosphorylated epitopes may serve as docking sites for the assembly of protein complexes or may alter the 3-dimensional protein structure thus modulating enzymatic activity or the ability to undergo protein-protein-interactions. Modification of proteins on serine residues is mediated by serine/threonine kinases.

Please note that phosphoserine detection by monoclonal antibodies is always dependent on the surrounding amino acid sequence!

Mab PSER-7F12 recognizes a broad range of serine-phosphorylated proteins in crude cell extracts, preferring positively charged amino acids directly neighboured to phosphoserine.

Related Products

mab against Phosphoserine

#0018-100/pSer-1C8

#0019-100/pSer-4A3

#0020-100/pSer-4A9

#0021-100/pSer-4H4

#0023-100/pSer-16B4

mab against Phosphothreonine

#0024-100/pThr-1E11

#0025-100/pThr-4D11

#0026-100/pThr-14B3

Purification: The antibody was purified from serum-free cell culture supernatant by subsequent thiophilic adsorption and size exclusion chromatography.

Formulation: lyophilized from 1 ml 2 x PBS / 0.09 % Na-azide / PEG and Sucrose.

Reconstitution: Reconstitute with 1 ml H₂O (15 min, RT).

Stability: For long-term storage, freeze lyophilizate upon arrival (-20°C). Upon reconstitution, aliquote and freeze in liquid nitrogen; reconstituted antibody can be stored frozen at -80°C up to 1 year. Thaw aliquots at 37°C. Thawed aliquots may be stored at 4°C up to 1 week.

Avoid repeated freeze / thaw cycles.v

Positive Control: #0901: phosphoserine/phosphothreonine positive control

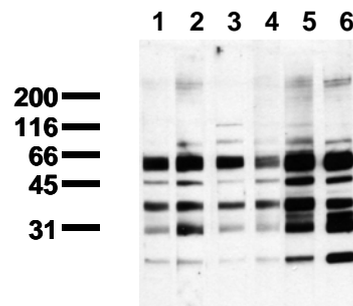
Immunoblotting: 1 µg/ml for HRPO/ECL detection
Recommended blocking buffer: BSA/Tween 20 based blocking buffer.
DO NOT USE MILK OR CASEIN FOR BLOCKING!

Immunoprecipitation: use at 1 - 10 µg per 10⁶ pervanadate-treated A431 cells

Immunocytochemistry: ND

ELISA: use at 0.05 µg/ml

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Phosphoserine Detection

Phosphoprotein Positive Control was probed with

lane 1: mab 1C8 (IgM), 1 µg/ml

lane 2: mab 4A3 (IgM), 1 µg/ml

lane 3: mab 4A9 (IgM), 1 µg/ml

lane 4: mab 4H4 (IgM), 1 µg/ml

lane 5: mab 7F12 (IgG), 1 µg/ml

lane 6: mab 16B4 (IgM), 1 µg/ml

Mouse Monoclonal Antibody to

Phosphoserine

clone 16B4

Order No.: 0023-025/PSER-16B4

Size (µg) 25

Lot No.: 0023S



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Isotype	Species Reactivity	Applications	Mol. Weight	Ref.Cell Line	Epitope	Immunogen
IgM	human, mouse, rat, dog	WB, ELISA, IP	pattern		...pSer - Pro...;...pSer - Lys	phosphopeptide conjugated to KLH

Background and Specificity:

Phosphorylation and dephosphorylation of cellular proteins are central steps in transducing extracellular signals to the cell nucleus. Phosphorylated epitopes may serve as docking sites for the assembly of protein complexes or may alter the 3-dimensional protein structure thus modulating enzymatic activity or the ability to undergo protein-protein-interactions. Modification of proteins on serine residues is mediated by serine/threonine kinases.

Please note that phosphoserine detection by monoclonal antibodies is always dependent on the surrounding amino acid sequence!

Mab PSER-16B4 recognizes a broad range of serine-phosphorylated proteins in crude cell extracts, preferring positively charged amino acids directly neighboured to phosphoserine.

Related Products

mab against Phosphoserine

#0018-100/pSer-1C8

#0019-100/pSer-4A3

#0020-100/pSer-4A9

#0021-100/pSer-4H4

#0022-100/pSer-7F12

mab against Phosphothreonine

#0024-100/pThr-1E11

#0025-100/pThr-4D11

#0026-100/pThr-14B3

Purification:	The antibody was purified from serum-free cell culture supernatant by subsequent thiophilic adsorption and size exclusion chromatography.
Formulation:	lyophilized from 1 ml 2 x PBS / 0.09 % Na-azide / PEG and Sucrose.
Reconstitution:	Reconstitute with 1 ml H ₂ O (15 min, RT).
Stability:	For long-term storage, freeze lyophilizate upon arrival (-20°C). Upon reconstitution, aliquote and freeze in liquid nitrogen; reconstituted antibody can be stored frozen at -80°C up to 1 year. Thaw aliquots at 37°C. Thawed aliquots may be stored at 4°C up to 1 week.

Avoid repeated freeze / thaw cycles.

Positive Control: #0901: phosphoserine/phosphothreonine positive control

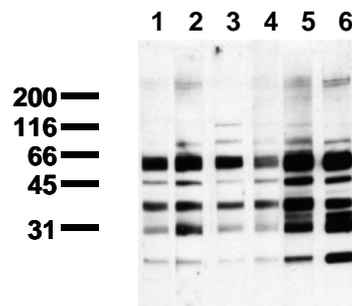
Immunoblotting: 1 µg/ml for HRPO/ECL detection
Recommended blocking buffer: BSA/Tween 20 based blocking buffer.
DO NOT USE MILK OR CASEIN FOR BLOCKING!

Immunoprecipitation: use at 1 - 10 µg per 10⁶ pervanadate-treated A431 cells

Immunocytochemistry: ND

ELISA: use at 0.05 µg/ml

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Phosphoserine Detection

Phosphoprotein Positive Control was probed with

lane 1: mab 1C8 (IgM), 1 µg/ml

lane 2: mab 4A3 (IgM), 1 µg/ml

lane 3: mab 4A9 (IgM), 1 µg/ml

lane 4: mab 4H4 (IgM), 1 µg/ml

lane 5: mab 7F12 (IgG), 1 µg/ml

lane 6: mab 16B4 (IgM), 1 µg/ml

pSer / pThr Molecular Weight Marker

Order No.: 0901/PSERCO
Lot: 0901
Size 20 Blots



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orders & support:

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03/100407F

Formulation

The pSer/pThr molecular weight marker contains rabbit muscle phosphoproteins isolated by Fe³⁺/IDA - affinity chromatography. Proteins are lyophilized from PBS/NaF/PEG/Sucrose/ Bromophenolblue and Na - azide. After reconstitution the solution contains 0.09% Na-azide.

Stability

Reconstitute by addition of 200 µl H₂O. After complete solubilization add 200 µl 2x SDS-PAGE sample buffer, mix and incubate at 90°C for 5 min.

Aliquote and store frozen. Avoid repeated freeze/thaw cycles.

Application

The pSer/pThr molecular weight marker is recommended for immunoblot applications. Use 20µl molecular weight marker per lane.

Note: Use BSA based blot incubation buffers. Milk, Casein and Blotto might interfere with antibody - antigen interaction.