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## AP52471PU-N

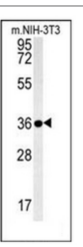
## IPU-N Polyclonal Antibody to LGALS9B (Center) - Aff - Purified

Quantity:	0.4 ml
Concentration:	lot specific
Uniprot ID:	<u>Q3B8N2</u>
NCBI:	<u>NP_001036150</u>
GenelD:	<u>284194</u>
Host / Isotype:	Rabbit / Ig
Immunogen:	KLH conjugated synthetic peptide between 167-196 amino acids from the Central region of Human LGALS9B
Format:	<b>State:</b> Liquid purified Ig fraction <b>Purification:</b> Protein A column, followed by peptide affinity purification <b>Buffer System:</b> PBS containing 0.09% (W/V) Sodium Azide as preservative
Applications:	ELISA: 1/1000. Western Blot: 1/100-1/500. Flow Cytometry: 1/10-1/50. Other applications not tested. Optimal dilutions are dependent on conditions and should be determined by the user.
Specificity:	This antibody recognizes Human and Mouse LGALS9B (Center).
Add. Information:	Molecular Weight: 39660 Da
Storage:	Store undiluted at 2-8°C for one month or (in aliquots) at -20°C for longer. Avoid repeated freezing and thawing. Shelf life: one year from despatch.
Pictures:	Western blot analysis of LGALS9B293Antibody (Center) CatNo AP52471PU-95N in 293 cell line lysates (35ug/lane).72This demonstrates the LGALS9B antibody55detected the LGALS9B protein (arrow).55
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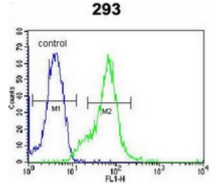
For research and in vitro use only. Not for diagnostic or therapeutic work. Material Safety Datasheets are available at www.acris-antibodies.com or on request.



Western blot analysis of LGALS9B Antibody (Center) Cat.-No AP52471PU-N in mouse NIH-3T3 cell line lysates (35ug/lane). This demonstrates the LGALS9B antibody detected the LGALS9B protein (arrow).



Flow cytometric analysis of 293 cells using LGALS9B Antibody (Center) Cat.-No AP52471PU-N (right histogram) compared to a negative control cell (left histogram). FITC-conjugated goat-antirabbit secondary antibodies were used for the analysis.



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