

bs-0924R**[Primary Antibody]****Bioss**
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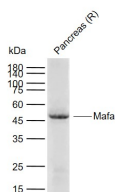
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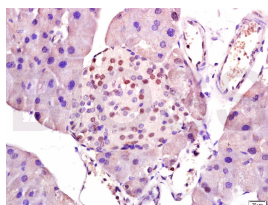
400-901-9800

Mafa Rabbit pAb**— DATASHEET —**

Host: Rabbit	Isotype: IgG	Applications: WB (1:500-2000)
Clonality: Polyclonal		IHC-P (1:100-500)
GeneID: 378435	SWISS: Q8CF90	IHC-F (1:100-500)
Target: Mafa		IF (1:100-500)
Immunogen: KLH conjugated synthetic peptide derived from mouse Mafa: 265-359/359.		Reactivity: Rat (predicted: Human, Mouse)
Purification: affinity purified by Protein A		
Concentration: 1mg/ml		Predicted MW.: 37 kDa
Storage: 0.01M TBS (pH7.4) with 1% BSA, 0.02% Proclin300 and 50% Glycerol. Shipped at 4°C. Store at -20°C for one year. Avoid repeated freeze/thaw cycles.		Subcellular Location: Nucleus
Background: Insulin gene expression is regulated by several islet-enriched transcription factors. However, MAFA is the only beta cell-specific activator. MAFA selectively induces endogenous insulin transcription in non-beta cells. MAFA was also first detected in the insulin-producing cells formed during the second and predominant phase of beta cell differentiation, and absent in the few insulin-positive cells found in Nkx6.1(-/-) pancreata, which lack the majority of second-phase beta cells. These results demonstrate that MAFA is a potent insulin activator that is likely to function downstream of Nkx6.1 during islet insulin-producing cell development.		

— VALIDATION IMAGES —

Sample: Lane 1: Rat Pancreas tissue lysates
Primary: Anti- Mafa (bs-0924R) at 1/1000 dilution
Secondary: IRDye800CW Goat Anti-Rabbit IgG at 1/20000 dilution Predicted band size: 37 kDa
Observed band size: 47 kDa



Tissue/cell: rat pancreas tissue; 4% Paraformaldehyde-fixed and paraffin-embedded; Antigen retrieval: citrate buffer (0.01M, pH 6.0), Boiling bathing for 15min; Block endogenous peroxidase by 3% Hydrogen peroxide for 30min; Blocking buffer (normal goat serum, C-0005) at 37°C for 20 min; Incubation: Anti-Mafa Polyclonal Antibody, Unconjugated(bs-0924R) 1:400, overnight at 4°C, followed by conjugation to the secondary antibody(SP-0023) and DAB(C-0010) staining

— SELECTED CITATIONS —

- **[IF=7.5]** Xie Tianqin. et al. Dysregulated lncRNAs regulate human umbilical cord mesenchymal stem cell differentiation into insulin-producing cells by forming a regulatory network with mRNAs. STEM CELL RES THER. 2024 Dec;15(1):1-17 IF ;Human. 38273351