

**bs-8513R****[ Primary Antibody ]****BioSS**  
ANTIBODIES

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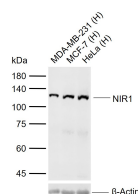
sales@bioss.com.cn

techsupport@bioss.com.cn

400-901-9800

**NIR1 Rabbit pAb****— DATASHEET —**

<b>Host:</b> Rabbit	<b>Isotype:</b> IgG	<b>Applications:</b> WB (1:500-2000)
<b>Clonality:</b> Polyclonal		<b>Reactivity:</b> Human (predicted: Mouse, Rat, Pig, Cow, Horse)
<b>GeneID:</b> 83394	<b>SWISS:</b> Q9BZ71	
<b>Target:</b> NIR1		<b>Predicted MW.:</b> 106 kDa
<b>Immunogen:</b> KLH conjugated synthetic peptide derived from human NIR1/RDGBA3: 131-250/974.		<b>Subcellular Location:</b> Cell membrane
<b>Purification:</b> affinity purified by Protein A		
<b>Concentration:</b> 1mg/ml		
<b>Storage:</b> 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.		
<b>Background:</b> Catalyzes the transfer of phosphatidylinositol and phosphatidylcholine between membranes (in vitro) (By similarity). Binds calcium ions. Involvement in disease: Defects in PITPNM3 are the cause of cone-rod dystrophy type 5 (CORD5) . CORDs are inherited retinal dystrophies belonging to the group of pigmentary retinopathies. CORDs are characterized by retinal pigment deposits visible on fundus examination, predominantly in the macular region, and initial loss of cone photoreceptors followed by rod degeneration.		

**— VALIDATION IMAGES —**

Sample: Lane 1: Human MDA-MB-231 cell lysates  
Lane 2: Human MCF-7 cell lysates Lane 3: Human HeLa cell lysates  
Primary: Anti-NIR1 (bs-8513R) at 1/1000 dilution  
Secondary: IRDye800CW Goat Anti-Rabbit IgG at 1/20000 dilution  
Predicted band size: 106 kDa  
Observed band size: 120 kDa