

# MAA Lysine Rabbit Polyclonal Antibody

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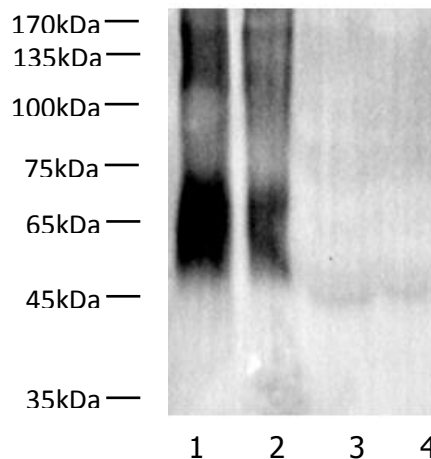
Cat#: 3090

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Only For Research. Not For Diagnosis.

<b>Attribute:</b>	Rabbit Polyclonal Antibody
<b>Isotype:</b>	IgG
<b>Purity:</b>	Antigen Affinity Purification
<b>Application:</b>	WB, IP, ELISA
<b>Reactivity:</b>	MAA adducts
<b>Immunogen:</b>	MAA protein
<b>Buffer:</b>	PBS with 0.1% sodium azide and 50% glycerol, pH 7.2
<b>Storage:</b>	Store at -20°C. Do not aliquot
<b>Recommended Dilution:</b>	WB: 1:1000-2000 IP: 1:5000-10000



1. MAA-HSA (10ug/lane)
2. MAA- HSA 5ug/lane)
3. HSA (10uIglane)
4. HSA (5ug/lane)

Western blot of anti-MAA Lysine Rabbit Polyclonal Antibody at dilution of 1:1000

## Background:

Malondialdehyde (MDA) and acetaldehyde (AA) react synergistically to form hybrid malondialdehyde–acetaldehyde (MAA) adducts . MAA adducts are unique among aldehyde–protein adducts in their stability, potent immunogenicity, ability to alter protein regulatory elements (i.e.,ε-amino lysine residue), and dose-dependent direct cellular toxicity . These properties of MAA adducts represent one pathogenic mechanism by which classical risk factors for cardiovascular disease(i.e., hypertension, hyperlipidemia, diabetes, and tobacco use) may result in inflammation and lead to specific clinical entities such as atherosclerosis.

We focus on precise protein quantification