# abcam

## Product datasheet

## Human Apolipoprotein AI ELISA Kit (APOA1) ab108803

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Overview

Product name Human Apolipoprotein AI ELISA Kit (APOA1)

**Detection method**Colorimetric

Precision Intra-assay

Sample	n	Mean	SD	CV%	
Overall				4.6%	

Inter-assay

Sample	n	Mean	SD	CV%	
Overall				10%	

Sample type Saliva, Milk, Urine, Serum, Plasma, Cerebral Spinal Fluid

Assay type Sandwich (quantitative)

**Sensitivity** 0.95 ng/ml

**Range** 12.5 ng/ml - 100 ng/ml

Recovery 98 %
Assay time 4h 00m

**Assay duration** Multiple steps standard assay

Species reactivity Reacts with: Human

Product overview Abcam's Apolipoprotein AI (APOA1) Human in vitro ELISA (Enzyme-Linked Immunosorbent

Assay) kit is designed for the quantitative measurement of apolipoprotein AI concentrations in

plasma, serum, urine, saliva, milk, CSF and cell culture samples.

An Apolipoprotein AI specific antibody has been precoated onto 96-well plates and blocked. Standards or test samples are added to the wells and subsequently an Apolipoprotein AI specific biotinylated detection antibody is added and then followed by washing with wash buffer. Streptavidin-Peroxidase Complex is added and unbound conjugates are washed away with wash buffer. TMB is then used to visualize Streptavidin-Peroxidase enzymatic reaction. TMB is catalyzed by Streptavidin-Peroxidase to produce a blue color product that changes into yellow after adding acidic stop solution. The density of yellow coloration is directly proportional to the amount of Apolipoprotein AI captured in plate.

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Get results in 90 minutes with Human Apolipoprotein Al ELISA Kit ( $\underline{ab189576}$ ) from our SimpleStep ELISA $^{\circledR}$  range.

The entire kit may be stored at -20°C for long term storage before reconstitution - Avoid repeated freeze-thaw cycles.

Platform

Microplate

#### **Properties**

## Storage instructions

Store at -20°C. Please refer to protocols.

Components	1 x 96 tests
100X Streptavidin-Peroxidase Conjugate	1 x 80µl
10X Diluent N Concentrate	1 x 30ml
20X Wash Buffer Concentrate	2 x 30ml
50X Biotinylated Human Apolipoprotein Al Antibody	1 x 120µl
Apolipoprotein Al Microplate (12 x 8 well strips)	1 unit
Chromogen Substrate	1 x 7ml
Human Apolipoprotein Al Standard	1 vial
Sealing Tapes	3 units
Stop Solution	1 x 11ml

#### **Function**

Participates in the reverse transport of cholesterol from tissues to the liver for excretion by promoting cholesterol efflux from tissues and by acting as a cofactor for the lecithin cholesterol acyltransferase (LCAT). As part of the SPAP complex, activates spermatozoa motility.

#### Tissue specificity

Major protein of plasma HDL, also found in chylomicrons. Synthesized in the liver and small intestine.

#### Involvement in disease

Defects in APOA1 are a cause of high density lipoprotein deficiency type 2 (HDLD2) [MIM:604091]; also known as familial hypoalphalipoproteinemia (FHA). Inheritance is autosomal dominant.

Defects in APOA1 are a cause of the low HDL levels observed in high density lipoprotein deficiency type 1 (HDLD1) [MIM:205400]; also known as analphalipoproteinemia or Tangier disease (TGD). HDLD1 is a recessive disorder characterized by the absence of plasma HDL, accumulation of cholesteryl esters, premature coronary artery disease, hepatosplenomegaly, recurrent peripheral neuropathy and progressive muscle wasting and weakness. In HDLD1 patients, ApoA-I fails to associate with HDL probably because of the faulty conversion of pro-ApoA-I molecules into mature chains, either due to a defect in the converting enzyme activity or a specific structural defect in Tangier ApoA-I.

Defects in APOA1 are the cause of amyloid polyneuropathy-nephropathy lowa type (AMYLIOWA) [MIM:107680]; also known as amyloidosis van Allen type or familial amyloid polyneuropathy type

III. AMYLIOWA is a hereditary generalized amyloidosis due to deposition of amyloid mainly constituted by apolipoprotein A1. The clinical picture is dominated by neuropathy in the early stages of the disease and nephropathy late in the course. Death is due in most cases to renal amyloidosis. Severe peptic ulcer disease can occurr in some and hearing loss is frequent. Cataracts is present in several, but vitreous opacities are not observed.

Defects in APOA1 are a cause of amyloidosis type 8 (AMYL8) [MIM:105200]; also known as systemic non-neuropathic amyloidosis or Ostertag-type amyloidosis. AMYL8 is a hereditary generalized amyloidosis due to deposition of apolipoprotein A1, fibrinogen and lysozyme amyloids. Viscera are particularly affected. There is no involvement of the nervous system. Clinical features include renal amyloidosis resulting in nephrotic syndrome, arterial hypertension, hepatosplenomegaly, cholestasis, petechial skin rash.

Sequence similarities

Belongs to the apolipoprotein A1/A4/E family.

Post-translational modifications

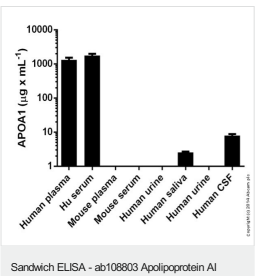
Palmitoylated.

Phosphorylation sites are present in the extracelllular medium.

Cellular localization

Secreted.

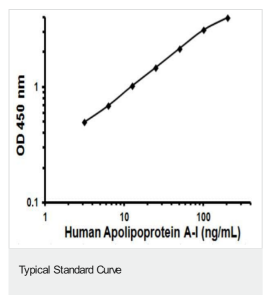
## **Images**



APOA1 measured in various samples showing quantity (microgram) per mL of tested sample.

(APOA1) Human ELISA Kit





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