

# **Product datasheet**

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# ARG80175 Human IgG4 ELISA Kit

Package: 96 wells Store at: 4°C

## Component

Cat. No.	Component Name	Package	Temp
ARG80175-001	Antibody-coated microplate	8 X 12 strips	4°C. Unused strips should be sealed tightly in the air-tight pouch.
ARG80175-002	Standard (Lyophilized)	3 X 0.7 μg/vial	4°C
ARG80175-003	Standard diluent buffer	40 ml	4°C
ARG80175-004	Antibody conjugate concentrate	400 μΙ	4°C
ARG80175-005	Antibody diluent buffer	16 ml	4°C
ARG80175-008	20X Wash buffer	50 ml	4°C
ARG80175-009	TMB substrate	12ml	4°C (Protect from light)
ARG80175-010	STOP solution	12ml	4°C
ARG80175-011	Plate sealer	4 strips	Room temperature

### Summary

Product Description	ARG80175 Human IgG4 ELISA Kit is an Enzyme Immunoassay kit for the quantification of Human IgG4 in Serum, Plasma, Cell culture supernatants.			
Tested Reactivity	Hu			
Tested Application ELISA				
Considerate.				

Specificity No significant cross-reactivity or interference with Human IgG1, IgG2, IgG3

Target Name IgG4
Conjugation HRP

Conjugation Note Substrate: TMB and read at 450 nm

Sensitivity 8 ng/ml

Sample Type Serum, Plasma, Cell culture supernatants

Standard Range 15.6 - 1000 ng/ml

Sample Volume  $100 \mu l$ 

Precision CV: less than10%

Alternate Names Immunoglobulin G4

## **Application Instructions**

Assay Time

4 hours

#### **Properties**

Form

96 well

Storage instruction

Store the kit at 2-8°C. Keep microplate wells sealed in a dry bag with desiccants. Do not expose test reagents to heat, sun or strong light during storage and usage. Please refer to the product user manual

for detail temperatures of the components.

Note

For laboratory research only, not for drug, diagnostic or other use.

#### Bioinformation

Background

IgG4 is present in very low levels in children younger than 10 years of age, so IgG4 deficiencies are not usually diagnosed before age 10. IgG4 may be undetectable in the serum of many "normal" adult individuals, and therefore low IgG4 alone is insufficient evidence of an antibody deficiency disorder requiring Ig replacement. IgG4—related systemic disease (IgG4-RSD), also known as hyper-IgG4 disease and IgG4-related disease is a disease in which inflammatory cells cause fibrosis, the deposition of connective tissue, in one or more organs. The disease is so named because the antibody subtype IgG4 can be detected on tissue samples and often at elevated levels in the bloodstream. The association with IgG4 is a relatively recent finding, and the condition has been described under numerous other names in the past. Diseases such asautoimmune pancreatitis, retroperitoneal fibrosis, mediastinal fibrosis, Riedel's thyroiditis, Mikulicz's syndrome, Kuttner's tumor and inflammatory pseudotumor are now regarded as forms of IgG4-RSD.

Highlight

Related products:

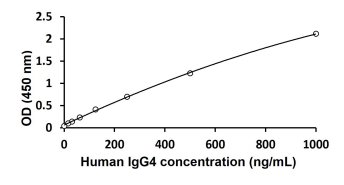
IgG4 ELISA Kits;

New ELISA data calculation tool: Simplify the ELISA analysis by GainData

Research Area

Immune System kit

#### **Images**



#### ARG80175 Human IgG4 ELISA Kit standard curve image

ARG80175 Human IgG4 ELISA Kit results of a typical standard run with optical density reading at 450 nm.