# **Mouse IL-6 ELISA-KIT**

Ref.: mIL-6-EIA-5-KIT

MabTag's ELISA for mouse Interleukin-6 (mIL-6) contains appropriate reagents sufficient for processing of 5 microplates (5 x 96 wells; 100  $\mu$ l/well)

For research only. Not for use in diagnostic or therapeutic procedures.

Specificity: mouse Interleukin-6 (mIL-6)

Typical standard curve range: 16 – 1000 pg/ml

**Detection limit**: 12 pg/ml

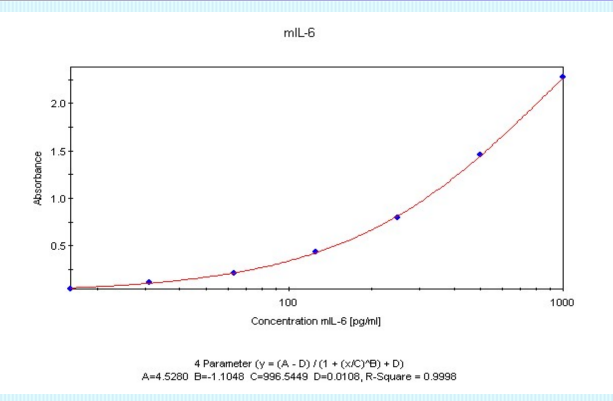
**Samples:** Culture supernatants, serum, plasma and other body fluids.

For serum and plasma a dilution of  $\geq$  1:10 is recommended.



Content	Working dilution	Storage
1 x vial 500 μl liquid anti-mIL-6 Capture-Antibody (red cap)	1:100	-20°C
1 x vial 500 μl liquid anti-mIL-6 Detection-Antibody (yellow cap)	1:100	-20°C
1 x vial 50 ng lyophilized rmIL-6 Standard (white cap)	customer specific	-20°C
1 x vial 50 μl Poly-HRP-Streptavidin (blue or green cap)	1:1000	-20°C
5 x 96well-Microplate	-	room temperature
1 x 60 ml Coating-Buffer	ready-to-use	2-8°C
3 x 100 ml Blocking-Buffer / Reagent-Diluent	ready-to-use	2-8°C
3 x 100 ml Wash-Buffer (10x concentrated)	100 ml must be filled up to 1L with aqua dest.	2-8°C
1 x 4 ml TMB-Solution A	10 ml TMB-B + 0.5 ml TMB-A	2-8°C
1 x 60 ml TMB-Solution B	TO IIII TIVIB-B + 0.5 MI TIVIB-A	protect from light
1 x 30 ml Stop-solution ACID → wear gloves & protective glasses	ready-to-use	2-8°C

## Typical standard curve



Step	Incubation	Procedure		
Coating Capture-antibody	≥ OVERNIGHT	Dilute capture-antibody 1:100 in COATING-BUFFER		
	at room	(100 μl capture-antibody in 10 ml COATING-BUFFER).		
	temperature	Subsequently transfer 100 µl of this working-solution to each well and incubate.		
Remove capture-antibody completely by inverting the microplate and blotting it <i>vigorously</i> against clean paper towels.				
Blocking	≥ 1 Hour	Add 200 of DEOCKING DUFFFD to cook well and involves		
	at room	Add 300 μl BLOCKING-BUFFER to each well and incubate.		
Davis DLOC	temperature	ala da circa de la comitación de la contrata de la		
Remove BLOCKING-BUFFER completely by inverting the microplate and blotting it <i>vigorously</i> against clean paper towels.				
Standard & Sample		Dilute standard & samples in REAGENT-DILUENT and transfer 100 μl in the respective		
	≥ 2 Hours at room temperature	wells in duplicates. Standard: Make serial dilutions in REAGENT-DILUENT and begin		
		with a high standard and end with blanks. The standard vial of this set contains <b>50 ng</b>		
		lyophilized standard. Reconstitute this in exactly 1 ml REAGENT-DILUENT (stock solution = 50 ng/ml) and choose a sufficient high standard working solution for your		
		assay (e.g. prepare a 1:20 dilution for a standard curve beginning with 2500 pg/ml).		
Wash Ex <b>vigoro</b>	refurmith MASHING BII			
vvasii ox <b>vigorot</b>	Wash 5x <i>vigorously</i> with WASHING-BUFFER and remove resting buffer completely by inverting the microplate and blotting it			
	≥ 2 Hours	vigorously against clean paper towels.  Dilute detection-antibody 1:100 in REAGENT-DILUENT		
Detection- antibody	≥ 2 ⊓ours at room	(100 μl detection-antibody in 10 ml REAGENT-DILUENT).		
	temperature	Subsequently transfer 100 µl of this working-solution to each well and incubate.		
Wash 5x <b>vigorously</b> with WASHING-BUFFER and remove resting buffer completely by inverting the microplate and blotting it				
Wash SA Vigorot	vigorously against clean paper towels.			
Poly-HRP- Streptavidin	20-30 Min	Dilute Poly-HRP-Streptavidin 1:1000 in REAGENT-DILUENT		
	at room	(10 μl in 10 ml REAGENT-DILUENT).		
	temperature	Subsequently transfer 100 μl of this working-solution to each well and incubate.		
Wash 5x <i>vigorously</i> with WASHING-BUFFER and remove resting buffer completely by inverting the microplate and blotting it				
vigorously against clean paper towels.				
Substrate solution	Up to 60 Min*	Optionally warm the solution to room temperature before use.		
	at room	Add 100 µl of the SUBSTRATE-SOLUTION to each well and incubate.		
	temperature	Control the development of the colour reaction continuously and stop at an		
	in the dark	appropriate time point.		
Stop solution		When the enzymatic colour reaction is sufficiently proceeded stop the reaction by		
	<u>_</u>	adding of 50 μl stop solution. Read the microplate at the substrate-depending		
		wavelength. (e.g. <b>450</b> nm for TMB substrate)		
		(if wavelength correction is available, set to 540 nm, 570 nm or 630 nm as reference)		

<sup>\*</sup>The speed of enzymatic colour development is influenced by many customer-specific factors. Therefore the incubation time is variable und specific for each test system.

#### Note:

All incubation steps except <u>TMB substrate</u> could be optionally carried out over-night. Do not use sodium azide-containing solutions, nor add sodium azide to the supplied reagents. Sodium azide inactivates the peroxidase.

#### Storage:

Specific storage conditions in the table above.

Reconstituted reagents should be stored at -20°C. Please prevent repeated freeze- thaw cycles. Stable for up to 6 months after opening when stored at -20° C. The performance of the unopened reagents is guaranteed until one year after point of delivery.

### **Precautions for use:**

!The stop solution is an <u>acid solution</u>. TMB-Solution A contain  $\underline{H_2O_2}$  and <u>tetramethylbenzidine</u> (TMB). All Buffers and liquid antibody solutions contain 0.045% (v/v) <u>Proclin®950</u> as preservative. All these compounds are harmful and cause respiratory, skin and eye irritation. Do not swallow any components of the set/kit (R22). Wear face, eye and hand clothing protection when using this material (S36). Keep out of reach of children (S2). Keep away from food, drink and animal feeding stuff (S13). !These reagents are offered for research purposes only! For *in vitro* use only. MabTag will not be held responsible for patent infringement or other violations that may occur with the use of our products.

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