



Rat APOA1 Elisa Kit

- 货号: LV20031
- 种 属: 大鼠
- 规 格: 48T/96T
- 有效期: 6 个月
- 批 号: 详见试剂盒外包装
- 检测范围: 0.156 – 10ng/ml
- 灵 敏 度: 0.031ng/ml
- 保存温度: 2-8℃

- 标准曲线浓度为: 10.0、5.0、2.5、1.25、0.62、0.31、0.156、0 ng/ml
- 特异性: 可检测样本中大鼠的 APOA1, 且与其



一、检测原理:

上海爱萌优宁生物生产的 ELISA 试剂盒采用“夹心法”:将捕获抗体包被于酶标板上, 捕获样品及标准品中的靶蛋白, 生物素化的检测抗体与靶蛋白结合, SABC 复合物与生物素化检测抗体结合, 形成免疫复合物, 加入 TMB 显色液后, 若反应孔中有靶蛋白则显蓝色, 加入终止液变黄色, 检测过程中未结合的成分均被洗去, 用酶标仪在 **450 nm** 处测 OD 值, 靶蛋白浓度与 OD 值之间呈正比, 从而对检测样品进行定性或相对定量分析。

二、试剂盒组分: (保存温度 4℃)



2. 进口品牌可调节移液器及吸头, 0.5-10 μ L, 2-20 μ L, 20-200 μ L, 200-1000 μ L。一次检测样品较多时, 最好用多通道移液器。
3. 干净的试管和 Ep 管。
4. 双蒸水或去离子水。

三、样本收集方法:

- 血清:** 室温血液自然凝固 10-20 分钟后, 离心 20 分钟左右 (2000-3000 转/分), 收集上清-20℃或者-80℃ 冷冻保存。
- 血浆**



3. 血清标本采集时，应注意避免溶血，红细胞溶解时会释放出具有过氧化物酶活性的物质，以 HRP 为标记的 ELISA 测定中，溶血标本可能会增加非特异性显色。
4. 为了保证尿液检测结果的准确性，必须正确收集尿液标本和保存，盛尿容器要清洁干燥。最好使用一次性的容器（如塑料尿杯），避免因用药并清洗不干净而造成的污染，影响检测结果，尿液标本必须新鲜，留取后，应及时检测或保存，以免细菌繁殖，因在室温（尤其是夏季）中久置后尿中的磷酸盐等可析出结晶而干扰检测。
5. 冻结标本融解后，蛋白质局部浓缩，分布不均，应充分轻缓混匀，避免气泡，可上下颠倒混和，不要在混匀器上强烈振荡。
6. 混浊或有沉淀的标本应先离心或过滤，澄清后再检测。
7. 反复冻融会使蛋白效价降低，所以待测标本如需保存作多次检测，宜少量分装冰



七、检测前试剂准备:

1. 试剂与样品预处理

实验前 30 min, 将全部试剂、样品置于室温平衡, 严禁加热促融; 已取出的试剂禁止倒回原瓶, 防止整瓶污染; 试剂配制、样品稀释后必须充分混匀。

2. 标准曲线与样品稀释

每次实验均需绘制标准曲线; 根据预估待测蛋白浓度, 对样品进行梯度稀释, 保证目的蛋白浓度



八、洗板方法:

● 手工洗板方法:

吸去（不可触及板壁）或甩掉酶标板内的液体；在实验台上铺垫几层吸水纸，酶标板朝下用力拍几次；将 $1 \times$ 洗涤缓冲液至少 $300 \mu\text{l}$ 注入孔内，浸泡 1-2 分钟。根据需要，重复此过程数次。

● 自动洗板:

全自动洗板机的使用应注意以下几点:



十一、结果判断与计算:

1. 所有 OD 值建议减除空白孔值后再进行计算, 如空白孔 OD 低于 0.1, 也可以直接计算。
2. 以标准品浓度作横坐标, OD 值作纵坐标, 手工绘制或用软件绘制标准曲线, 根据样品 OD 值计算出相应含量, 再乘以稀释倍数即可。

十二、声明:

1、本公司只对试剂盒本身负责, 不对因使用该试剂盒所造成的样本消耗负责, 请使用者使用前充分考虑到样本的可能使用量, 预留充足的样本。



问题分析:

若实验效果不好, 请及时对显色结果拍照, 保存实验数据, 保留所用板条及未使用试剂, 然后 联系我公司技术支持为您解决问题。同时您也可以参考以下资料:

标准曲线较差

原因	解决方案
标准品溶液配置有误	确认是否进行正确稀释。
标准品复溶不当	开盖前进行离心; 检查复溶后是否存在不溶物。
标准品已	

试剂盒没有充分平衡	试剂室温平衡至少 20 分钟，确保所有试剂已平衡至室温。
-----------	------------------------------

变异系数 (CV) 较大

原因	解决方案
孔中有气泡	读板前，确保不存在气泡。
孔洗涤不均/未充分洗涤	检查洗板机的所有管口是否畅通。使用推荐方法进行洗涤。
试剂混匀不充分	确保所有试剂充分混匀。
移液量不一致	正确使用经过校准的移液器
边缘效应	确保孔板和所有试剂处于室温。
样品制备或保存条件不一致	确保样品制备保持一致



灵敏度偏低

原因	解决方案
ELISA 试剂盒保存不当	按推荐方式保存所有试剂。请注意，各试剂的保存条件可能有所不同。
靶标不足	浓缩样品或降低样品稀释度。
检测试剂失活	确保报告酶/荧光素具有预期的活性。
酶标仪设置不正确	在检测中，确保酶标仪设置为正确的吸收波长或激发/发射波长。
测定方法不够灵敏	更换更灵敏的检测系统（例如从比色检测转变为化学发光/荧光检测）。更换更灵敏的测定方法（例如从直接 ELISA 方法转变为夹心 ELISA 方法）。延长孵育时间或升高温度。
微量滴定板吸附靶标的	



Rat APOA1 Elisa Kit

- **Cat NO.:** LV20031
- **Reactivity:** Rat
- **Size:** 48T/96T
- **Validity:** 6 months
- **Lot NO.:** Please refer to the exterior package
- **Range:** 0.156–10ng/ml
- **Sensitivity:** 0.031ng/ml
- **Storage:** 2-8℃
- **Standard curve concentration:** 10.0、5.0、2.5、1.25、0.62、0.31、0.156、0 ng/ml
-



Test principle

The ELISA kit produced by Shanghai Aimin Youning Biotechnology adopts the "sandwich method": The capture antibody is coated on the microplate. The target protein in the sample and standard solution binds to the capture antibody. The biotinylated detection antibody combines with the target protein, and the SABC complex combines with the biotinylated detection antibody to form an immune complex. After adding the TMB color development solution, if there is the target protein in the reaction well, it will turn blue. After adding the stop solution, it will turn yellow. During the detection process, all unbound components are washed away. The OD value is measured at 450 nm using an enzyme detector. The concentration of the target protein is proportional to the OD value, thereby



5. Clean tubes and Eppendorf tubes
6. Deionized or distilled water

Sample Collection and Storage (universal)

- **Serum:** after 10-20 minutes of natural coagulation of blood at room temperature, centrifugation for about 20 minutes (2000-3000 rpm/min). Collect and clear- 20 ° C or -80 ° C cryopreservation.
- **Plasma:** Collect plasma using (EDTA-Na₂ or heparin as an anticoagulant. Centrifuge samples for 15 minutes at 1000×g at 2

retention, should be timely detection or preservation, in order to avoid bacterial reproduction. It interferes with the detection because of the precipitation of phosphate in urine after long-term storage at room temperature (especially in summer).

6. After thawing the frozen sample, the protein is locally concentrated and unevenly distributed. It should be fully mixed gently to avoid bubbles. It can be mixed upside down, and do not vibrate strongly on the mixer. Repeated freezing and thawing of samples should be avoided.
7. Turbid or precipitated samples should be centrifuged or filtered before detection.
8. Repeated freezing and thawing will reduce the potency of the protein, so if the samples to be tested need to be preserved for multiple tests, they should be stored in a small amount of ice. Appropriate preservatives can also be added.
9. Aprotinin should be added to hormone samples.

Sample preservation:



Reagent Preparation and Storage

1. Reagent and Sample Pretreatment

All reagents and samples were equilibrated to room temperature for **30** minutes before the experiment. **Heating thawing is prohibited**. The removed reagents shall not be poured back into the original bottle to avoid contamination. All reagents and diluted samples must be fully mixed during preparation.

2. Standard Curve and Sample Dilution



Sample Dilution

The user should estimate the concentration of target protein in the test sample, and select a proper dilution factor to make the diluted target protein concentration fall in the optimal detection range of the kit. Dilute the sample with the provided dilution buffer, and several trials may be necessary. The test sample must be well mixed with the dilution buffer. And also standard curves and sample should be making in pre-experiment. If samples with very high concentrations, dilute samples with PBS first and then dilute the samples with Sample Dilution.

The matrix components in the sample will affect the test results, which it need to be diluted at least 1/2 with Sample Dil



4. **Wash:** The washing steps are the same as above.
5. **Avidin-Biotin-Peroxidase Complex (SABC) :** Add 100 μ l of SABC Working Solution into each well, cover the plate and incubate at 37°C for 30 minutes.
6. **Wash:** The washing steps are the same as above.
7. **TMB Solution:** Add 100 μ l TMB Solution into each well, cover the plate and incubate at 37°C in dark within 10-20 minutes. (**Note:** The reaction time can be shortened or extended according to the actual color change, but not more than 30 minutes. You can terminate the reaction when apparent gradient appeared in standard wells.)
8. **Stop:** Add 50 μ l stop solution into each well, mix well, and measure the absorbance at 450nm within 30 minutes.<



Problem analysis:

If the experimental effect is not good, please take photos of the color results in time, save the experimental data, keep the used strips and unused reagents, and then contact our technical support to solve the problem for you. You can also refer to the following information:

Poor standard curve



The coefficient of variation (CV) is large

reason	Solution
Before reading the board	make sure that there is no bubble in the hole.
Uneven / insufficient pore washing	Check whether all pipe orifices of washing machine are unblocked. Wash with recommended method.
Insufficient mixing of reagents	Ensure that all reagents are fully mixed.
Inconsistent fluid transfer	Correct use of calibrated pipettes
edge effect	Make sure that the pore plate and all reagents are at room temperature.



Low sensitivity

reason	Solution
Improper storage of Elisa Kit	Save all reagents as recommended. Please note that the storage conditions of each reagent may be different.
Insufficient targets	Concentrate the sample or reduce the dilution of the sample.
Deactivation of detection reagent	Ensure that the reporter enzyme / fluorescein has the desired activity.
Incorrect setting of microplate reader	In the detection, make sure that the microplate