

(For scientific research use only, not for clinical diagnosis	(For	· scientific re	esearch use	only, not	for clinical	l diagnosis
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Human Cell Globulin (CYGB)

Quantitative Detection Kit (ELISA)

Instructions for Use Specification:

96T/48T Catalog Number: SYP-H2415

Purpose: Used to detect human cell globulin in serum, plasma, cell culture supernatant and other samples

(CYGB) concentration.

Website: www.upingbio.com

Official hotline: 400-999-8863

Supervision phone number:

Page 1 of 20

Please read the instructions carefully before use. If you have any questions, please contact us via:

Official hotline: 400-999-8863

Technical phone number: 18358180525

Email: UpingBio@163.com

Company website: www.upingbio.com For specific shelf life, please refer to the outer packaging label of the kit. Please use the kit within the shelf life. When contacting us, please provide the product number and production date (see box label) so that we can serve you more efficiently.

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Supervision phone number:

Page 2 of 20

[Kit performance]

Physical properties: Each liquid component is clear and transparent, with no sediment or floc. Microplate aluminum foil bags should be vacuum packed without damage or leakage.

Calibration curve linearity: The correlation coefficient r value of the calibrator dose-response curve is greater than or equal to 0.9900. Precision: intra-batch variation coefficient CV% is less than 10%; inter-batch variation coefficient CV% is less than 15%. Sensitivity: The

Recovery rate: The recovery rate is between 85%-115%.

lowest detectable dose is less than 0.078 ng/ml.

Sensitivity: This kit recognizes natural human cell globulin (CYGB) and has no overlap with structural analogs.

Stability: Stored at 2°C-8°C, validity period is 6 months.

Detection range: 0.312 ng/ml - 10 ng/ml.

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Official hotline: 400-999-8863

Supervision phone number:

Page 3 of 20

Experimental principle

This kit uses a double-antibody sandwich enzyme-linked immunosorbent assay (ELISA). In the microwell enzyme plate pre-coated with anti-human cell globulin (CYGB) antibody (solid-phase antibody), add human cell globulin (CYGB) calibrator and sample to be tested, then add biotin-labeled antibody, and after warming After incubation and sufficient washing, HRP-coupled avidin is added. After incubation and sufficient washing, unbound components are removed, and a solid-phase antibody-antigen-biotin-labeled antibody-avidin is formed on the solid surface of the microplate. Sandwich complexes of enzymes. Add TMB chromogenic solution to produce a blue product. Under the action of the stop solution, it is finally converted into yellow. The absorbance (OD value) is measured on a microplate reader at a wavelength of 450nm. The absorbance (OD value) is related to the human cell globulin in the sample to be tested. (CYGB) concentration is positively related. By fitting the calibrator curve, the concentration of human cell globulin (CYGB) in the sample can be calculated.

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Official hotline: 400-999-8863

Supervision phone number:

Page 4 of 20

[Kit components and storage]

Comp	onents	quantity	Main
Calibrator	High Standard	2 vial	Calibrator freeze-
Calibration solution	Reconstitution	2 vial	PBS
Calibrators & Sample Diluents	Standard & Sample Diluent	25mL	PBSTN
coated microplate	Microelisa Stripplate	96T/48T	Pre-coated solid phase
biotin antibody	Bio-Antibody	10mL	biotin antibody
HRP labeled avidin	HRP- Conjugate	10mL	HRP labeled avidin
TMB chromogenic	TMB	10mL	TMB
stop solution	Stop Solution	6mL	acidic solution
20×concentrated	20X Wash Solution	25mL	0.05%Tween20
manual	manual	1 serving	
ziplock bag	ziplock bag	1	
Self-adhesive	Self-adhesive	4 pieces	

Note: 1. Before use, please check whether the label and quantity of the reagents in the kit are consistent with the table.

- 2. The test kit should be stored at 2-8°C. Expired test kits must not be used.
- 3. If the coated microplate is not used up in a single time, remember to seal it and store it at 2-8°C.
- 4. The reconstituted calibrator can only be used on the same day.

5. If the components of the kit need to be used again, please ensure that they have not been contaminated since the last use.

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Page 5 of 20

Prepare your own test equipment required for the

test (not provided, but can assist in purchasing) 1.

Standard specification microplate reader.

- 2. Automatic plate washing machine.
- 3. Oscillator.
- 4. A series of adjustable pipettes and tips. When testing a large number of samples at one time, it is best to use a multi-channel pipette.

[Kit limitations]

- 1. For scientific research use only and not for clinical diagnosis.
 - 2. Use within the validity period marked on the kit. Expired products must not be used.
 - 3. Do not mix with kits or components from other manufacturers.
 - 4. Use the sample diluent provided with the kit.
 - 5. If the sample value is higher than the highest calibrator concentration value,

please dilute the sample appropriately and then re-measure. 6. Human anti-mouse

and other heterophilic antibodies present in the sample to be tested will interfere with the test results. Please eliminate this factor before testing.

7. The test results obtained by other methods are not directly comparable to the test results of this kit.

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Page 6 of 20

[Precautions]

- 1. This kit is for in vitro research only and not for clinical diagnosis.
- 2. Please wear a lab coat and latex gloves for protection during the test.

 Especially when testing blood or other body fluid samples, please follow the national biological laboratory safety protection regulations.
- 3. Incubate strictly according to the specified time and temperature to ensure accurate results. All reagents must reach room temperature 20-25°C before use. Store reagents refrigerated immediately after use.
- 4. Incorrect plate washing may lead to inaccurate results. Make sure to absorb as much liquid as possible from the wells before adding substrate. Do not allow the microwells to dry out during incubation.
 - 5. Eliminate residual liquid and fingerprints on the bottom of the plate, otherwise it will affect the OD value.
 - 6. The substrate chromogenic solution should be colorless or very light in color.
 - 7. Avoid cross-contamination of reagents and specimens to avoid erroneous results.
 - 8. Avoid direct exposure to strong light during storage and incubation.

- 9. After balancing to room temperature, open the sealed bag to prevent water droplets from condensing on the cold slats.
- 10. Any reaction reagents must not come into contact with bleaching solvents or strong gases emitted by bleaching solvents. Any bleaching ingredients will destroy the biological activity of the reagents in the kit.
 - 11. The microplate reader used for detection needs to be equipped with a filter capable of detecting a wavelength of 450±10nm.

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Page 7 of 20

Optical density ranges from 0-3.5. It is recommended to preheat 15 minutes in advance before use.

- 12、请勿使用其他批号或其他来源的试剂混合或替代本试剂盒中的试剂。
- 13、试验中所用的 EP 管和吸头均为一次性使用,严禁混用。
- 14、请勿使用过期的试剂。

【样品的准备和保存】

以下只是列出样品采集和保存的一般指南。所有样本采集保存过程中, 不得使用叠氮钠做为防腐剂。样品如果不立即分析,应分装后冷冻保存, 且避免反复冻融。

细胞培养上清: 离心去除沉淀, 立即分析或分装后-20℃冷冻保存。

血清:用干净试管收集血液,室温凝固 30 分钟,离心 2000×g 20 分钟,收集血清。立即分析或分装后-20℃冷冻保存。

血浆:采用肝素、柠檬酸盐或 EDTA 抗凝,抽血后 30 分钟内在 2-8℃ 离心 2000×g 20 分钟。为消除血小板的影响,建议在 2-8℃进一步离心 10000×g 10 分钟。立即分析或分装后-20℃冷冻保存。

细胞裂解液:对于贴壁细胞,去除培养液,用PBS、生理盐水或无血

清培养液洗一遍。加入适量裂解液,用移液器吹打数下,使裂解液和细胞充分接触。通常 10 秒后,细胞就会被裂解。对于悬浮细胞,离心收集细胞,用 PBS、生理盐水或无血清培养液洗一遍。加入适量裂解液,用移液器吹打把细胞吹散,用手指轻弹以充分裂解细胞。充分裂解后,10000—14000×g 离心 3-5 分钟,取上清。立即分析或分装后-20℃冷冻保存。

尿液:用无菌管收集,离心 2000×g 20 分钟。仔细收集上清。如有 沉淀形成,应再次离心。

【试剂准备】

- 1、使用前,所有的组分都要至少复温 120min,确保充分复温到室温。
- 2、浓缩洗涤液: 从冰箱取出的浓缩洗涤液, 会有结晶产生, 这属于正常现象, 水浴加热使结晶完全溶解。浓缩洗涤液与蒸馏水, 按 1:20 稀释, 即 1 份的浓缩洗涤液, 添加 19 份的蒸馏水。

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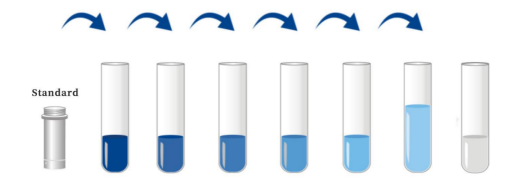
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第9页共20页

【校准品稀释方法】

Steps for redissolving the calibrator: Redissolve the calibrator with the calibrator reconstituted solution. Add all the liquid in one bottle of the calibrator reconstituted solution to the bottle of calibrator freeze-dried powder. Vortex gently to ensure thorough mixing., the concentration of the mother solution of the calibrator after redissolution is 20 ng/ml, mix thoroughly before dilution.

Calibrator mother solution dilution steps: Let the calibrator working solution stand for 1-2 minutes before dilution, and dilute the calibrator mother solution with the calibrator & sample universal diluent. Double dilution method: Take 7 EP tubes, and Add 500 μ L calibrator & sample diluent, pipet 500 μ L from the 20 ng/ml calibrator mother solution into the first EP tube, mix well to prepare a 10 ng/ml calibrator working solution, and follow this step in sequence. Pipette and mix. As shown on the following page.



20ng/ml

Recommended dilution concentration of calibrator: It is recommended to prepare the following concentrations: 10, 5, 2.5, 1.25, 0.625,

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Page 10 of 20

0.312, 0 ng/ml, and used as the calibrator concentration value of the fitted standard curve.

Tip: Add the calibrator & sample universal diluent directly to the last tube as the 0 value. There is no need to draw liquid from the penultimate tube. The diluted calibrator working solution needs to be prepared and used immediately.

Coperating Procedure

Recommended sample dilution scheme: It is recommended that teachers conduct preliminary experiments to explore the optimal dilution ratio of samples before conducting formal experiments.

All reagents and components should be returned to room temperature first.

It is recommended to perform duplicate wells for calibrators, quality control materials and samples.

- 1. Prepare the working solution of various components of the kit according to the method described in the previous instructions.
 - 2. Take out the required slats from the aluminum foil bag, seal the remaining slats in a ziplock bag and return it to the refrigerator.

- 3. Set up the calibrator hole, sample diluent hole, blank hole and sample hole. Add 50 μL of calibrator of different concentrations to each of the calibrator holes. Add 50 μL of sample diluent to the sample diluent hole. Do not add any to the blank hole. Add the test well to the sample hole. Sample 50μL. Except for the blank wells, add 100μL of biotin antibody to each well, cover the reaction plate with a sealing film, and incubate in a 37°C water bath or incubator in the dark for 60 minutes.
 - 4. Uncover the sealing film, discard the liquid, pat dry on absorbent paper, fill each well with washing solution, and let it sit.

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Page 11 of 20

1 minute, shake off the washing liquid, pat dry on absorbent paper, repeat this 5 times. If using an automatic plate washer, please wash the plate according to the operating procedures of the plate washer. Adding a soaking program for 30 seconds can improve the detection accuracy. After washing the plate and before adding substrate, pat the reaction plate dry on clean, lint-free paper.

- 5. Except for the blank wells, add 100uL of HRP-labeled avidin to each well, cover the reaction plate with a sealing film, and incubate in a 37°C water bath or incubator in the dark for 20 minutes.
 - 6. Repeat step 4.
- 7. Add $100\mu L$ of TMB chromogenic solution to all wells. Cover the reaction plate with sealing film and incubate in a $37^{\circ}C$ water bath or incubator in the dark for 15 minutes.
- 8. Add 50 μ L of stop solution to all wells, and read the absorbance (OD value) of each well on a 450 nm wavelength microplate reader.

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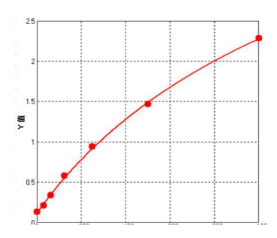
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Supervision phone number:

Page 12 of 20

Result calculation

1. Use the concentration of the calibrator as the abscissa and the corresponding absorbance (OD value) as the ordinate. Use computer software and four-parameter Logistic curve fitting (4-pl) to create a standard curve equation. Through the absorbance (OD value) of the sample value), use the equation to calculate the concentration value of the sample. [Calculation using ELISA Calc software] 2. If the sample is diluted, the concentration value measured by the above method must be multiplied by the dilution factor to obtain the final concentration of the sample.



(Schematic diagram, for reference only)

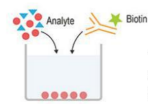
Website: www.upingbio.com

Official hotline: 400-999-8863

Supervision phone number:

Page 13 of 20

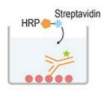
[Operation Summary]



1、反应板孔中加入50uL校准品工作液或样本后,立即每孔加入100uL生物素化抗体工作液,37℃孵育60分钟。



2、弃掉板内液体,洗板5次。



3、每孔加入100uL HRP酶结合物工作液 37℃孵育20分钟,弃掉板内液体,洗板 5次。



4、每孔加入100uL TMB显色液, 37℃孵育 15分钟。



5、每孔加入 50uL 终止液。



6、立即在 450nm波长下读数,处理数据。

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[problem analysis]

Problem Description	Possible Causes	Corresponding	
	Incorrect liquid	Check pipettes and tips	
Negative and positive	Equilibration time is too	Ensure sufficient	
control results are unstable	Incomplete washing	Ensure the washing time and number of washes	
	Incubation time too short	Ensure adequate	
	The experimental	Use recommended	
Very weak or colorless	Insufficient reagent	Check the liquid	
	Incorrect dilution	aspiration and addition	
	Enzyma lahal	Mix enzyme conjugate	
	Enzyme label	and substrate and check	
		在酶标仪上检查波长及	
Reading value is low	Microplate reader	滤 光片设置	
	settings are incorrect	提前打开酶标仪预热	
变异系数大	加液不正确	检查加液情况	
	检测抗体的工作浓度过	使用推荐的稀释倍数	
背景值高	酶标板洗涤不完全	保证每步清洗完全; 如果用自动洗板机, 请检查所有的出口是 否有堵塞;是否使用	
	洗液有污染	配制新的洗液	
灵敏度低	ELISA 试剂盒保存不当	按说明书要求保存相关 试 剂	
火蚁及瓜	读数前未终止	OD 读数前应在每孔中 加 入终止液	

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第15页共20页

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。

【声明】

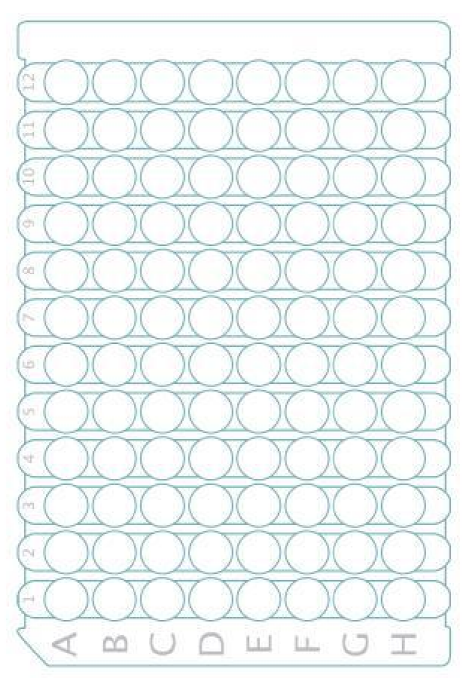
- 1、限于现有条件及科学技术水平,尚不能对所有原料进行全面的鉴定分析,本产品可能存在一定的质量技术风险。
- 2、本试剂盒在研发过程中去除/降低了生物学样本中的一些内源性干扰因素,并非所有可能影响的因素均已去除。
- 3、最终的实验结果与试剂的有效性、实验者的相关操作以及当时的 实验环境等因素密切相关,本公司只对试剂盒本身负责,不对因使用试 剂盒所造成的样本消耗负责,请使用者使用前充分考虑到样本可能的使 用量,预留充足的样本。
- 4、为了达到好的实验结果,请只使用本公司试剂盒内提供的试剂, 不要混用其他制造商的产品,严格按照说明书操作。
- 5、由于操作过程中试剂制备以及酶标仪参数设置不正确,可能导致结果异常,实验前请仔细阅读说明书并调整好仪器。

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- 6、即使是相同人员操作也可能在两次独立实验中得到不同的结果, 为保证结果的重现性,需要控制实验过程中每一步的操作。
- 7、试剂盒发货前会经过严格的质检,然而,因为运输条件、实验设备差异等等因素影响,用户检测结果可能跟出厂数据不一致。
- 8、本试剂盒未与其他厂家同类试剂盒或不同方法检测同一目的物的 产品进行对比,所以不排除检测结果不一致的情况。
- 9、试剂盒仅供研究使用,如将其用于临床诊断或任何其他用途,我 公司将不对因此产生的问题负责,亦不承担任何法律责任。



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第18页共20页

【实验心得】

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