

ELISA PRODUCT INFORMATION & MANUAL

Human Azurocidin/CAP37/HBP ELISA Kit (Colorimetric) NBP2-80317

Sample Insert for Reference Only

Enzyme-linked Immunosorbent Assay for quantitative detection. For research use only.

Not for diagnostic or therapeutic procedures.

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BACKGROUND

Azurocidin (AZU1), also known as heparin-binding protein (HBP) or cationic antimicrobial protein 37 (CAP37), is an azurophil granule antibiotic protein, with monocyte chemotactic and antibacterial activity. The Azurophil granules, specialized lysosomes of the neutrophil, contain at least 10 proteins implicated in the killing of microorganisms. Azurocidin is a member of the serine protease family that includes Cathepsin G, neutrophil elastase (NE), and proteinase 3 (PR3), however, Azurocidin is not a serine proteinase since the active site serine and histidine residues are replaced. Neutrophils arriving first at sites of inflammation release Azurocidin, which acts in a paracrine fashion on endothelial cells causing the development of intercellular gaps and allowing leukocyte extravasation. It thus be regarded as a reasonable therapeutic target for a variety of inflammatory disease conditions.

INTENDED USE

For the quantitative determination of Human Azurocidin/CAP37/HBP concentration in serum, cell culture supernates and plasma.

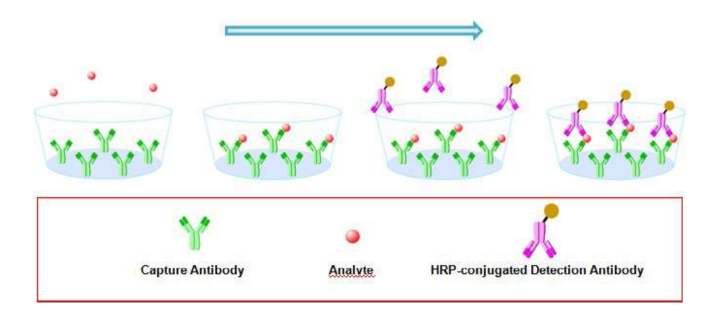
The use of this kit for other sample types need be validated by the end user due to the complexity of natural targets and unpredictable interference.

PRINCIPLE OF THE ASSAY

The principle of this ELISA kit is based on the solid phase sandwich enzyme immunoassay technique. A monoclonal antibody specific for Human Azurocidin/CAP37/HBP has been pre-coated onto well plate strips. Standards and samples are added to the wells and Human Azurocidin/CAP37/HBP present in the sample is bound by the immobilized antibody.

After incubation the wells are washed and a horseradish peroxidase conjugated anti-Human Azurocidin/CAP37/HBP antibody is added, producing an antibody-antigen-antibody "sandwich complex". Following a wash to remove any unbound antibody a TMB substrate solution is loaded and color develops in proportion to the amount of

Human Azurocidin/CAP37/HBP bound. The reaction is stopped by the addition of a stop solution and the intensity of the color can be measured at 450 nm (See schematics below).



MATERIALS PROVIDED

Human Azurocidin/CAP37/HBP Microplate - 96 well polystyrene microplate (12 strips of 8 wells) coated with rabbit mAb antibody against Human Azurocidin/CAP37/HBP.

Human Azurocidin/CAP37/HBP Detection Antibody - 0.2 mg/mL of rabbit pAb antibody against Human Azurocidin/CAP37/HBP conjugated to horseradish peroxidase (HRP) with preservatives.

Human Azurocidin/CAP37/HBP Standard - Recombinant Human Azurocidin/CAP37/HBP in a buffer with preservatives, lyophilized. The amount of standard is lot specific and indicated on the label of standard vial.

Wash Buffer Concentrate - 25 mL of a 20-fold concentrated solution of buffered surfactant with preservatives.

Dilution Buffer Concentrate - 8 mL of a 20-fold concentrated dilution buffer with preservatives.

Color Reagent A - 13 mL of stabilized hydrogen peroxide.

Color Reagent B - 13 mL of stabilized chromogen (tetramethylbenzidine).

Stop Solution - 8 mL of 2 N sulfuric acid.

STORAGE

Unopened Kit	Store at 2 - 8°C and the kit is stable for 6 months upon receipt.			
	Diluted Wash Buffer Diluted Dilution Buffer	Stored for up to 1 week at 2 - 8°C		
Opened/ Reconstituted Reagents	Conjugate Stop Solution Unmixed Color Reagent A Unmixed Color Reagent B	Stored for up to 1 month at $2 - 8^{\circ}$ C		
	Standard	After reconstitution, store for up to 1 month at -80°C. The reconstituted standards should be aliquoted and avoid repeated freeze-thaw cycles.		
	Microplate Wells	Return unused strips to the foil pouch containing the desiccant pack and reseal along entire edge of zip-seal. Stored for up to 1 month at 2 - 8°C		

OTHER SUPPLIES REQUIRED

- ·Microplate reader capable of measuring absorbance at 450 nm
- ·Pipettes and pipette tips
- ·Deionized or distilled water
- ·Multi -channel pipette, squirt bottle, manifold dispenser, or automated microplate washer
- ·500 mL graduated cylinder
- ·Tubes for standard dilution
- ·Well plate cover or seals

PRECAUTIONS

- 1. This kit is **for research use only** and is not for use in diagnostic or therapeutic procedures.
- 2. The kit should not be used beyond the expiration date.
- 3. Do not mix reagents from different lots.
- 4. The kit is designed and tested to detect the specific targets and samples shown in the manual. The use of this kit for other purpose should be verified carefully by the end user.

SAFETY INSTRUCTIONS

- 5. The Stop Solution provided with this kit is an acid solution. Take care when using the reagent to avoid the risk.
- 6. All biological materials should be handled and discarded as potentially hazardous following local laws and regulations.
- 7. Personal protective equipments such as lab coats, gloves, surgical masks and goggles are necessary in experiments for safety reasons.

TECHINICAL TIPS

- 8. Bring all reagents and samples to room temperature before use.
- 9. Samples should be thawed completely and mixed well prior to analysis. Avoid repeated freeze-thaw cycles of frozen samples.
- 10. A standard curve should be generated for each set of sample assayed. DO NOT USE the standard curves from other plates or other days.
- 11. Use a new disposable reagent reservoir and new disposable pipette tips for each transfer to avoid cross-contamination.
- 12. Read the absorbance of each well within 20 minutes after adding the stop solution.

SAMPLE COLLECTION AND STORAGE

Serum - Use a serum separator tube and allow samples to clot for 30 minutes before centrifugation for 15 minutes at 1000 x g. Remove serum and assay immediately or aliquot and store samples at $-20 \,^{\circ}\text{C}$ or lower temperature. **Avoid repeated freeze** -thaw cycles.

Cell Culture Supernates - Remove particulates by centrifugation and assay immediately or aliquot and store samples at -20°C or lower temperature. Avoid repeated freeze-thaw cycles. If the use of original supernate sample or low dilution (<5 fold) are necessary due to the expected low concentration of antigen supernates need be adjust to neutral pH condition before assay.

Plasma - Collect plasma using heparin or EDTA as an anticoagulant. Centrifuge for 15 minutes at 1000 x g within 30 minutes of collection. Assay immediately or aliquot and storesamples at -20°C. **Avoid repeated freeze-thaw cycles.**

Note:

The sample should be diluted to within the working range of the assay in $1 \times$ dilution buffer. The exact dilution must be determined based on the concentration of specific target in individual samples.

REAGENT PREPARATION

Bring all reagents to room temperature before use. If crystals have formed in buffer solution, warm to room temperature and mix gently until the crystals have completely dissolved.

Wash Buffer - Prepare 1× wash buffer by adding 20 mL of Wash Buffer Concentrate to deionized or distilled water to prepare 400 mL of Wash Buffer.

Dilution Buffer - Prepare 1× dilution buffer by adding 5 mL of Dilution Buffer Concentrate to deionized or distilled water to prepare 100 mL of Dilution Buffer.

Detection Antibody - Centrifuge at 10,000 x g for 20 seconds. Dilute to **work** concentration of 0.25 µg/mL in Dilution Buffer before use.

Substrate Solution - Color Reagents A and B should be mixed together in equal

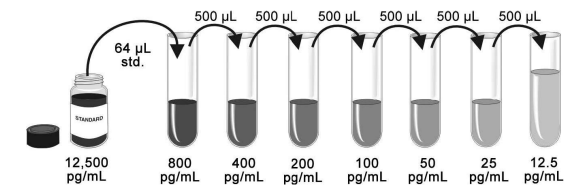
volumes within 15 minutes of use. Protect from light. 200 μ L of the resultant mixture is required per well. Take care not to contaminate the Color Reagent. If the mixed color reagent is blue. DO NOT USE.

Human Azurocidin/CAP37/HBP Standard - Reconstitute the Human Azurocidin/CAP37/HBP Standard with 1 mL of Dilution Buffer to make stock solution. Shake the vial gently until the lyophilized powder totally dissolved (**Do not turn the vial upside down**). Mix the standard to ensure complete reconstitution prior to making dilutions.

Prepare serially diluted standards as described in the following step:

Pipette 1000 μ L of Dilution Buffer into the 800 pg/mL tube. Pipette 500 μ L of Dilution Buffer into the remaining tubes. Use the stock solution to produce a dilution series as the following figure. Mix each tube thoroughly before the next transfer. The 800 pg/mL standard serves as the high standard. The Dilution Buffer serves as the zero standard (0 pg/mL). **Ensures each assay has a standard curve. DO NOT USE the standard curve on other plates or other days.**

The following graph is only for demonstration purposes. The concentration of stock solution is lot specific and need be calculated with the actual amount of standard labeled on the standard vial.



ASSAY PROCEDURE

Bring all reagents and samples to room temperature before use. It is recommended that all samples and standards be assayed in duplicate.

- 1. Prepare all reagents, working standards, and samples as directed in the previous sections.
- 2. Remove unused microplate strips from the plate frame, return them to the foil pouchcontaining the desiccant pack, and reseal.
- 3. Wash each well three times with Wash Buffer (300 µL/well) using a squirt bottle, multi-channel pipette, manifold dispenser or autowasher. Complete removal of liquid at each step is essential to good performance. Remove any remaining Wash Buffer by aspirating or decanting. Invert the plate and blot it against clean paper towels.
- 4. Add 100 μ L of each serially diluted protein standard or test sample per well including a zero standard. **Ensure reagent addition is uninterrupted and completed within 15 minutes.** Cover/seal the plate and incubate for 2 hours at room temperature.
- 5. Repeat the aspiration/wash as in Step 3.
- 6. Add 100 μ L of Detection Antibody in working concentration to each well. Cover/seal the plate and incubate for 1 hour at room temperature.
- 7. Repeat the aspiration/wash as in Step 3.
- **8.** Add 200 μ L of Substrate Solution to each well. Incubate for 20 minutes at room temperature. **Protect from light.**
- 9. Add 50 μL of Stop Solution to each well. If color change does not appear uniform, gently tap the plate to ensure thorough mixing.
- 10. **Determine the optical density of each well within 20 minutes**, using a microplate reader set to 450 nm.

CALCULATION OF RESULTS

If samples generate values higher than the highest standard, dilute the samples and repeat the assay.

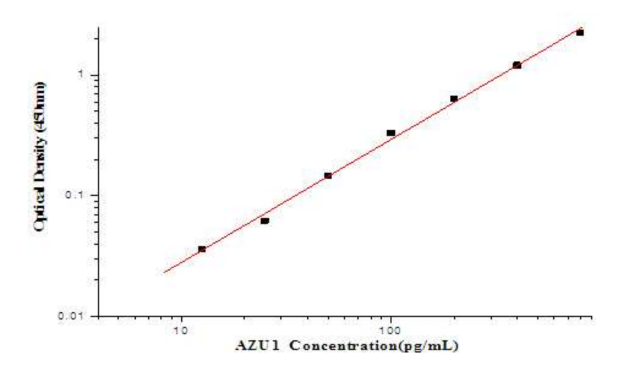
Calculate the mean absorbance for each standard, control and sample and subtract average zero standard optical density (O.D.) .

Construct a standard curve by plotting the mean absorbance for each standard on the y-axis against the concentration on the x-axis and draw a best fit curve through the points on the graph. Most graphing software can help make the curve and a four parameter logistic (4-PL) usually provide the best fit, though other equations (e.g. linear, log/log) can also be tried to see which provides the most accurate. Extrapolate the target protein concentrations for unknown samples from the standard curve plotted.

TYPICAL DATA

This standard curve is only for demonstration purposes. A standard curve should be generated for each assay.

Concentration (pg/mL)	Zero standard subtracted OD
0	0
12.5	0.036
25	0.062
50	0.146
100	0.330
200	0.635
400	1.211
800	2.241



PRECISION

Intra-assay Precision (Precision within an assay)

Three samples of known concentration were tested twenty times on one plate to assess intra-assay precision.

Inter-assay Precision (Precision between assays)

Three samples of known concentration were tested in five separate assays to assess inter-assay precision.

8	Intra -assay Precision			Inter-assay Precision		
Sample	1	2	3	1	2	3
N	20	20	20	3	3	3
Mean (pg/mL)	129	272	569	180	358	768
SD	3.16	9.53	23.41	12.98	26.15	41.91
CV (%)	2.4%	3.5%	4.1%	7.2%	7.3%	5.5%

RECOVERY

The recovery of Human Azurocidin/CAP37/HBP spiked to different levels throughout the range of the assay in related matrices was evaluated.

Sample	Average % Recovery	Range	
Serum (n=3)	98	93-103%	
EDTA plasma (n=3)	98	93-100%	
Cell culture supernates (n=3)	112	108-113%	

LINEARITY

		Serum	EDTA plasma	Cell culture supernates
1:2	recovery of detected	91%	90%	101%
1:4	recovery of detected	109%	106%	99%
1:8	recovery of detected	93%	94%	103%
1:16	recovery of detected	71%	86%	91%

SENSITIVITY

The minimum detectable dose (MDD) of Human Azurocidin/CAP37/HBP is typically less than 8.76 pg/mL. The MDD was determined by adding three standard deviations to the mean optical density value of twenty zero standard replicates and calculating the corresponding concentration.

CALIBRATION

This immunoassay is calibrated against a highly purified HEK 293-expressed recombinant Human Azurocidin/CAP37/HBP.

SAMPLE VALUES

The average concentration of Human Azurocidin/CAP37/HBP in 10 normal human serum is 11.46 +/-

6.10 ng/mL ranging from 2.44 to 18.88 ng/mL. The average concentration of Human Azurocidin/CAP37/HBP in 10 normal human plasma is 2.50 +/- 0.95 ng/mL ranging from

1.03 to 4.32 ng/mL. Human peripheral blood mononuclear cells (1 x 10E6 cells/mL) were cultured in RPMI supplemented with 10% fetal calf serum, 50 μ M β -mercaptoethanol, 2 mM L-glutamine, 100 U/mL penicillin, and 100 μ g/mL streptomycin sulfate, stimulated with 10 μ g/mL PHA for 5 days. Aliquots of the cell culture supernates were removed and assayed for levels of natural Azurocidin/CAP37/HBP, and measured 9.37 ng/mL.

SPECIFICITY

This assay recognizes both recombinant and natural Human Azurocidin/CAP37/HBP. The factors listed below were prepared at 50 ng/mL in dilution buffer and assayed for cross-reactivity. No cross-reactivity was observed.

Recombinant h	uman			
IL-1	IL-2	IL-33	IL-10	
IL-8	TNF-α	IL-4	IL-6	
IFN-γ	MIP-1β	TIMP-1	TIMP-2	
GM-CSF	SCF	PDGF	VEGF	

TROUBLE SHOOTING

Problems	Possible Sources	Solutions	
	Incorrect or no Detection Antibody was added	Add appropriate Detection Antibody and continue	
No signal	Substrate solution was not added	Add substrate solution and continue	
	Incorrect storage condition	Check if the kit is stored at recommended condition and used before expiration date	
	Standard was incompletely reconstituted or was inappropriately stored	Aliquot reconstituted standard and store at -80 °C. The reconstituted standards should be aliquoted and avoid repeated freeze-thaw cycles.	
Poor Standard Curve	Imprecise / inaccurate pipetting	Check / calibrate pipettes	
	Incubations done at inappropriate temperature, timing or agitation	Follow the general ELISA protocol	
	Background wells were contaminated	Avoid cross contamination by using the sealer appropriately	
Door detection	The concentration of antigen in samples was too low	Enriching samples to increase the concentration of antigen	
Poor detection value	Samples were ineffective	Check if the samples are stored at cold environment. Detect samples in timely manner	
		Use multichannel pipettes without touching the reagents on the plate	
	Insufficient washes	Increase cycles of washes and soaking time between washes	
High Background	Color Reagent should be clear and colorless prior to addition to wells	Color Reagent should be clear and colorless prior to addition to wells	
	Use clean tubes and pipettes tips	Use clean plates, tubes and pipettes tips	
Non av = -: C:	Samples were contaminated	Avoid cross contamination of samples	
Non-specificity	The concentration of samples was too high	Try higher dilution rate of samples	

ASSAY SUMMARY

