

# AssayMax™ Rat BNP-32 ELISA Kit

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For any questions regarding troubleshooting or performing the assay, please contact our support team at support@assaypro.com.

Thank you for choosing Assaypro.

## **Assay Summary**

**Step 1**. Add 50 μl of Standard or Sample per well. Incubate 2 hours.

**Step 2.** Wash, then add 50  $\mu$ l of Biotinylated Antibody per well. Incubate 2 hours.

**Step 3**. Wash, then add 50  $\mu$ l of SP Conjugate per well. Incubate 30 minutes.

**Step 4.** Wash, then add 50  $\mu$ l of Chromogen Substrate per well. Incubate 25 minutes.

Step 5. Add 50  $\mu$ l of Stop Solution per well. Read at 450 nm immediately.

## **Symbol Key**



Consult instructions for use.

# **Assay Template**

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# AssayMax™ Rat Brain Natriuretic Peptide 32 (BNP-32) ELISA Kit

Catalog No. ERB1201-1
Sample insert for reference use only

#### Introduction

Natriuretic peptides (ANP, BNP, and CNP) comprise a family of structurally related peptides, which are derived from three different genes and share a 17-amino acid internal ring (1). A high level of plasma BNP may have a strong, independent association with increased mortality rates in patients with primary pulmonary hypertension (PPH), congestive heart failure and/or after acute myocardial infarction (2-4).

#### Principle of the Assay

The AssayMax™ Rat BNP-32 ELISA (Enzyme-Linked Immunosorbent Assay) Kit is designed for detection of BNP-32 in rat plasma, serum, cell culture, and tissue samples. This assay employs a quantitative sandwich enzyme immunoassay technique that measures rat BNP-32 in approximately 5 hours. A polyclonal antibody specific for rat BNP-32 has been pre-coated onto a 96-well microplate with removable strips. BNP-32 in standards and samples is sandwiched by the immobilized antibody and a biotinylated polyclonal antibody specific for rat BNP-32, which is recognized by a streptavidin-peroxidase (SP) conjugate. All unbound material is washed away and a peroxidase enzyme substrate is added. The color development is stopped and the intensity of the color is measured.

## **Caution and Warning**

- This product is for Research Use Only and is not intended for use in diagnostic procedures.
- Prepare all reagents (diluent buffer, wash buffer, standard, biotinylated antibody, and SP conjugate), as instructed, prior to running the assay.
- Prepare all samples prior to running the assay. The dilution factors for the samples are suggested in this insert. However, the user should determine the optimal dilution factor.
- Spin down the SP conjugate vial and the biotinylated antibody vial before opening and using contents.
- The Stop Solution is an acidic solution.
- The kit should not be used beyond the expiration date.

#### Reagents

- Rat BNP-32 Microplate: A 96-well polystyrene microplate (12 strips of 8 wells) coated with a polyclonal antibody against rat BNP-32.
- Sealing Tapes: Each kit contains 3 precut, pressure sensitive sealing tapes that can be cut to fit the format of the individual assay.
- Rat BNP-32 Standard: Rat BNP-32 in a buffered protein base (10 ng, lyophilized).
- **Biotinylated Rat BNP-32 Antibody (50x):** A 50-fold concentrated biotinylated polyclonal antibody against rat BNP-32 (120 μl).
- MIX Diluent Concentrate (10x): A 10-fold concentrated buffered protein base (30 ml).
- Wash Buffer Concentrate (20x): A 20-fold concentrated buffered surfactant (30 ml, 2 bottles).
- SP Conjugate (100x): A 100-fold concentrate (80 μl).
- Chromogen Substrate (1x): A stabilized peroxidase chromogen substrate tetramethylbenzidine (7 ml).
- **Stop Solution (1x):** A 0.5 N hydrochloric acid solution to stop the chromogen substrate reaction (11 ml).

#### **Storage Condition**

- Upon arrival, immediately store components of the kit at recommended temperatures up to the expiration date.
- Store SP Conjugate and Biotinylated Antibody at -20°C.
- Store Microplate, Diluent Concentrate (10x), Wash Buffer, Stop Solution, and Chromogen Substrate at 2-8°C.
- Unused microplate wells may be returned to the foil pouch with the desiccant packs and resealed. May be stored for up to 30 days in a vacuum desiccator.
- Store Standard at 2-8°C before reconstituting with Diluent and at -20°C after reconstituting with Diluent.

## Other Supplies Required

- Microplate reader capable of measuring absorbance at 450 nm
- Pipettes (1-20 μl, 20-200 μl, 200-1000 μl, and multiple channel)
- Deionized or distilled reagent grade water

#### Sample Collection, Preparation, and Storage

 Plasma: Collect plasma using a final concentration of 0.1 M sodium citrate as an anticoagulant. Centrifuge samples at 3000 x g for 10 minutes and collect plasma. The sample is suggested for use at 1x for

- medium-high levels of BNP-32; however, user should determine optimal dilution factor depending on application needs. The undiluted samples can be stored at -20°C or below for up to 3 months. Avoid repeated freeze-thaw cycles.
- Serum: Samples should be collected into a serum separator tube. After clot formation, centrifuge samples at 3000 x g for 10 minutes and remove serum. The sample is suggested for use at 1x for medium-high levels of BNP-32; however, user should determine optimal dilution factor depending on application needs. The undiluted samples can be stored at -20°C or below for up to 3 months. Avoid repeated freeze-thaw cycles.
- Cell Culture Supernatant: Centrifuge cell culture media at 1500 rpm for 10 minutes at 4°C to remove debris and collect supernatant. If necessary, dilute samples into MIX Diluent; user should determine optimal dilution factor depending on application needs. The undiluted samples can be stored at -80°C. Avoid repeated freeze-thaw cycles.
- **Tissue:** Extract tissue samples with 0.1 M phosphate-buffered saline (pH 7.4) containing 1% Triton X-100 and centrifuge at 14000 x g for 20 minutes. Collect the supernatant and measure the protein concentration. If necessary, dilute samples into MIX Diluent; user should determine optimal dilution factor depending on application needs. Store remaining extract at -80°C. Avoid repeated freeze-thaw cycles.

Applicable samples may also include biofluids, cell culture, and tissue homogenates. If necessary, user should determine optimal dilution factor depending on application needs.

#### Refer to Dilution Guidelines for further instruction.

	Guidelines for Dilutions of 100-fold or Greater				
	(for reference only; please follow the insert for specific dilution suggested)				
	100x		10000x		
A)	4 μl sample : 396 μl buffer (100x)	A)	4 μl sample : 396 μl buffer (100x)		
	= 100-fold dilution	B)	4 μl of A : 396 μl buffer (100x)		
			= 10000-fold dilution		
	Assuming the needed volume is less than		Assuming the needed volume is less than		
	or equal to 400 μl.		or equal to 400 μl.		
	1000x		100000x		
A)	4 μl sample : 396 μl buffer (100x)	A)	4 μl sample : 396 μl buffer (100x)		
B)	24 μl of A : 216 μl buffer (10x)	B)	4 μl of A : 396 μl buffer (100x)		
	= 1000-fold dilution	C)	24 μl of B : 216 μl buffer (10x)		
			= 100000-fold dilution		
	Assuming the needed volume is less than		Assuming the needed volume is less than		
	or equal to 240 μl.		or equal to 240 μl.		

#### **Reagent Preparation**

- Freshly dilute all reagents and bring all reagents to room temperature before use.
- MIX Diluent Concentrate (10x): Dilute the MIX Diluent Concentrate 10fold with reagent grade water to produce a 1x solution. When diluting
  the concentrate, make sure to rinse the bottle thoroughly to extract any
  precipitates left in the bottle. Mix the 1x solution gently until the crystals
  have completely dissolved. Store for up to 30 days at 2-8°C.
- Rat BNP-32 Standard: Reconstitute the Rat BNP-32 Standard (10 ng) with 2.5 ml of MIX Diluent to generate a 4 ng/ml standard stock solution. Allow the vial to sit for 10 minutes with gentle agitation prior to making dilutions. Prepare duplicate or triplicate standard points by serially diluting from the standard stock solution (4 ng/ml) 2-fold with equal volume of MIX Diluent to produce 2, 1, 0.5, 0.25, 0.125, 0.063, and 0.031 ng/ml solutions. MIX Diluent serves as the zero standard (0 ng/ml). Aliquot remaining stock solution to limit repeated freeze-thaw cycles. This solution should be stored at -20°C and used within 30 days.

Standard Point	Dilution	[BNP-32] (ng/ml)
P1	1 part Standard (4 ng/ml) + 1 part MIX Diluent	2.0
P2	1 part P1 + 1 part MIX Diluent	1.0
Р3	1 part P2 + 1 part MIX Diluent	0.5
P4	1 part P3 + 1 part MIX Diluent	0.25
P5	1 part P4 + 1 part MIX Diluent	0.125
P6	1 part P5 + 1 part MIX Diluent	0.063
P7	1 part P6 + 1 part MIX Diluent	0.031
P8	MIX Diluent	0.0

- Biotinylated Rat BNP-32 Antibody (50x): Spin down the antibody briefly and dilute the desired amount of the antibody 50-fold with MIX Diluent to produce a 1x solution. The undiluted antibody should be stored at -20°C.
- Wash Buffer Concentrate (20x): Dilute the Wash Buffer Concentrate 20fold with reagent grade water to produce a 1x solution. When diluting
  the concentrate, make sure to rinse the bottle thoroughly to extract any
  precipitates left in the bottle. Mix the 1x solution gently until the crystals
  have completely dissolved.
- SP Conjugate (100x): Spin down the SP Conjugate briefly and dilute the desired amount of the conjugate 100-fold with MIX Diluent to produce a 1x solution. The undiluted conjugate should be stored at -20°C.

#### **Assay Procedure**

- Prepare all reagents, standard solutions, and samples as instructed. Bring all reagents to room temperature before use. The assay is performed at room temperature (20-25°C).
- Remove excess microplate strips from the plate frame and return them
  immediately to the foil pouch with desiccants inside. Reseal the pouch
  securely to minimize exposure to water vapor and store in a vacuum
  desiccator.
- Add 50 µl of Rat BNP-32 Standard or sample to each well. Gently tap
  plate to thoroughly coat the wells. Break any bubbles that may have
  formed. Cover wells with a sealing tape and incubate for 2 hours. Start
  the timer after the last addition.
- Wash the microplate manually or automatically using a microplate washer. Invert the plate and decant the contents; hit 4-5 times on absorbent material to completely remove the liquid. If washing manually, wash five times with 200 μl of Wash Buffer per well. Invert the plate each time and decant the contents; hit 4-5 times on absorbent material to completely remove the liquid. If using a microplate washer, wash six times with 300 μl of Wash Buffer per well; invert the plate and hit 4-5 times on absorbent material to completely remove the liquid.
- Add 50 µl of Biotinylated Rat BNP-32 Antibody to each well. Gently tap
  plate to thoroughly coat the wells. Break any bubbles that may have
  formed. Cover wells with a sealing tape and incubate for 2 hours.
- Wash the microplate as described above.
- Add 50 µl of SP Conjugate to each well. Gently tap plate to thoroughly coat the wells. Break any bubbles that may have formed. Cover wells with a sealing tape and incubate for 30 minutes. Turn on the microplate reader and set up the program in advance.
- Wash the microplate as described above.
- Add 50 µl of Chromogen Substrate to each well. Gently tap plate to thoroughly coat the wells. Break any bubbles that may have formed. Incubate in ambient light for 25 minutes or until the optimal blue color density develops.
- Add 50 µl of Stop Solution to each well. The color will change from blue to yellow. Gently tap plate to ensure thorough mixing. Break any bubbles that may have formed.
- Read the absorbance on a microplate reader at a wavelength of 450 nm immediately. If wavelength correction is available, subtract readings at 570 nm from those at 450 nm to correct optical imperfections.
   Otherwise, read the plate at 450 nm only. Please note that some unstable black particles may be generated at high concentration points after stopping the reaction for about 10 minutes, which will reduce the readings.

#### **Data Analysis**

- Calculate the mean value of the duplicate or triplicate readings for each standard and sample.
- To generate a standard curve, plot the graph using the standard concentrations on the x-axis and the corresponding mean 450 nm absorbance (OD) on the y-axis. The best fit line can be determined by regression analysis using log-log or four-parameter logistic curve fit.
- Determine the unknown sample concentration from the Standard Curve and multiply the value by the dilution factor.

#### **Typical Data**

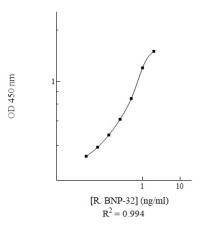
The typical data is provided for reference only. Individual laboratory
means may vary from the values listed. Variations between laboratories
may be caused by technique differences.

Standard Point	ng/ml	OD	Average OD	
P1	2.0	1.912	1.831	
1.2	2.0	1.750	1.031	
P2	1.0	1.312	1.318	
, <u>-</u>	2.0	1.324	1.010	
Р3	0.5	0.710	0.714	
13	0.5	0.718	0.714	
P4	0.25	0.469	0.472	
1 7	0.23	0.475	0.472	
P5	0.125	0.341	0.345	
FJ	0.123	0.349		
P6	0.063	0.266	0.271	
FU	0.063 0.276	0.276	0.271	
P7	0.031	0.219	0.226	
F /	0.031	0.233	0.220	
P8	0.0	0.132	0.132	
го	P6 0.0		0.132	
Sample: Poo	oled Normal	0.692	0.690	
Sodium Citrat	e Plasma (1x)	0.686	0.689	
Sample: Poo	oled Normal	0.689	0.674	
Serun	n (1x)	0.659	0.674	

#### Standard Curve

 The curve is provided for illustration only. A standard curve should be generated each time the assay is performed.

Rat BNP-32 Standard Curve



#### **Performance Characteristics**

- The minimum detectable dose of rat BNP-32 as calculated by 2SD from the mean of a zero standard was established to be 18 pg/ml.
- Intra-assay precision was determined by testing three plasma samples twenty times in one assay.
- Inter-assay precision was determined by testing three plasma samples in twenty assays.

	Intra	Intra-Assay Precision			Inter-Assay Precision		
Sample	1	2	3	1	2	3	
n	20	20	20	20	20	20	
CV (%)	3.8%	5.7%	5.3%	10.1%	11.2%	10.7%	
Average CV (%)		4.9%			10.7%		

## **Spiking Recovery**

 Recovery was determined by spiking one plasma and one serum sample with different BNP-32 concentrations.

Sample	Unspiked Sample (ng/ml)	Spiking Value (ng/ml)	Expected	Observed	Recovery (%)
		0.970	1.079	1.193	111%
Plasma	0.109	0.172	0.281	0.306	109%
		0.065	0.174	0.172	99%
		0.970	1.082	0.891	82%
Serum	0.112	0.172	0.284	0.299	105%
		0.065	0.177	0.202	114%
Average Recovery (%)					103%

## Linearity

• Plasma and serum samples were serially diluted to test for linearity.

Average Percentage of Expected Value (%)				
Sample Dilution Plasma Serum				
1x	115%	89%		
2x	105%	98%		
4x	89%	112%		

## **Cross-Reactivity**

Species	Cross-Reactivity (%)
Canine	40%
Bovine	None
Equine	20%
Monkey	100%
Mouse	10%
Human	30%
Swine	100%
Rabbit	None

• 10% FBS in culture media will not affect the assay.

## **Troubleshooting**

Issue	Causes	Course of Action
	Use of improper	Check the expiration date listed before use.
	components	<ul> <li>Do not interchange components from different lots.</li> </ul>
		<ul> <li>Check that the correct wash buffer is being used.</li> </ul>
		<ul> <li>Check that all wells are empty after aspiration.</li> </ul>
	Improper wash step	<ul> <li>Check that the microplate washer is dispensing properly.</li> </ul>
		<ul> <li>If washing by pipette, check for proper pipetting</li> </ul>
_		technique.
Low Precision	Splashing of reagents while loading wells	Pipette properly in a controlled and careful manner.
re	Inconsistent volumes	<ul> <li>Pipette properly in a controlled and careful manner.</li> </ul>
> ₽	loaded into wells	Check pipette calibration.
ò	loaded litto Wells	<ul> <li>Check pipette for proper performance.</li> </ul>
_	Insufficient mixing of	<ul> <li>Thoroughly agitate the lyophilized components after</li> </ul>
	reagent dilutions	reconstitution.
	reagent anations	Thoroughly mix dilutions.
		<ul> <li>Check the microplate pouch for proper sealing.</li> </ul>
	Improperly sealed	<ul> <li>Check that the microplate pouch has no punctures.</li> </ul>
	microplate	Check that three desiccants are inside the microplate
		pouch prior to sealing.
_	Microplate was left	Each step of the procedure should be performed
na	unattended between	uninterrupted.
ig	steps	
h	Omission of step	Consult the provided procedure for complete list of steps.
Unexpectedly Low or High Signal Intensity	Steps performed in incorrect order	<ul> <li>Consult the provided procedure for the correct order.</li> </ul>
	Insufficient amount of	Check pipette calibration.
w o	reagents added to	Check pipette for proper performance.
ly Low o	wells	
≥ ⊑	Wash step was skipped	Consult the provided procedure for all wash steps.
ec	Improper wash buffer	<ul> <li>Check that the correct wash buffer is being used.</li> </ul>
ect	Improper reagent	<ul> <li>Consult reagent preparation section for the correct</li> </ul>
άx	preparation	dilutions of all reagents.
ne	Insufficient or	<ul> <li>Consult the provided procedure for correct incubation</li> </ul>
Ō	prolonged incubation	time.
	periods	
		Sandwich ELISA: If samples generate OD values higher
⊭		than the highest standard point (P1), dilute samples
e F	Nan autimal assuals	further and repeat the assay.
≧	Non-optimal sample dilution	<ul> <li>Competitive ELISA: If samples generate OD values lower than the highest standard point (P1), dilute samples</li> </ul>
S	ullution	further and repeat the assay.
5		User should determine the optimal dilution factor for
qa		samples.
Deficient Standard Curve Fit	Contamination of	A new tip must be used for each addition of different
St	reagents	samples or reagents during the assay procedure.
, ut	Contents of wells	Verify that the sealing film is firmly in place before placing
icie	evaporate	the assay in the incubator or at room temperature.
efi	•	Pipette properly in a controlled and careful manner.
	Improper pipetting	Check pipette calibration.
		Check pipette for proper performance.

Insufficient mixing of reagent dilutions	Thoroughly agitate the lyophilized components after reconstitution. Thoroughly mix dilutions.
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#### References

- (1) Wiedemann K et al. (2000) Exp Clin Endocrinol Diabetes. 108(1):5-13.
- (2) Nagaya N et al. (2000) Circulation. 102(8):865-70.
- (3) Cheng V et al. (2001) J Am Coll Cardiol. 37(2):386-91.
- (4) Bettencourt P et al. (2000) Clin Cardiol. 23(12):921-7.

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