BNP FRAGMENT ELISA ASSAY KIT

ENZYME IMMUNOASSAY FOR THE QUANTITATIVE DETERMINATION OF BNP FRAGMENT IN HUMAN SERUM, CITRATE PLASMA, EDTA PLASMA OR HEPARIN PLASMA

CAT. NO. BI-20852W 12 X 8 TESTS

FOR RESEARCH USE ONLY
NOT FOR USE IN DIAGNOSTIC PROCEDURES
Additional information on our products is available on our website.

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1) INTRODUCTION

The natriuretic peptides are members of a family of structurally similar but genetically distinct peptide hormones, consisting of atrial-, brain-, and C-type (ANP, BNP, and CNP, respectively). ANP and BNP preferentially bind to a membrane-bound guanylyl cyclase (GC) receptor called GC-A or NPR1, whereas CNP is the physiological ligand for GC-B (NPR2). The natriuretic peptides play an important role in the regulation of cardiovascular and renal homeostasis and in the regulation of fatty acid metabolism and body weight.

BNP is mainly expressed by ventricular myocardium in response to volume overload and increased filling pressure. BNP has a cleavable signal sequence. Mature BNP consists of 108 amino acids (proBNP or BNP-108), and undergoes cleavage resulting in physiologically active BNP-32 and additional C-terminal fragments (cf. http://www.uniprot.org/uniprot/P16860#PRO_0000001532), along with a physiologically inactive N-terminal peptide comprising amino acids 1-76, which is further degraded proteolytically. BNP fragments in the circulation are therefore very heterogenous.

BNP has a key role in cardiovascular homeostasis with biological actions including natriuresis, diuresis, vasorelaxation, and inhibition of renin and aldosterone secretion. A high concentration of BNP in the bloodstream is indicative of heart failure.

The discovery of natriuretic peptides identified an endocrine system that contributes to diuresis and vascular tone. The biology, biochemistry and the pathophysiological role of natriuretic peptides are described in several reviews.

Areas of Interest
- Cardiac impairment, acute myocardial infarction, (left ventricular dysfunction)
- Obesity and diabetes
- Renal failure
- Various forms of secondary hypertension

2) CONTENTS OF THE KIT

<table>
<thead>
<tr>
<th>CONT</th>
<th>KIT COMPONENTS</th>
<th>QUANTITY</th>
</tr>
</thead>
<tbody>
<tr>
<td>PLATE</td>
<td>polyclonal anti BNP fragment (8-29) antibody coated microtiter strips in strip holder packed in aluminium bag with desiccant</td>
<td>12 x 8 tests</td>
</tr>
<tr>
<td>WASHBUF</td>
<td>Wash buffer, 20x concentrated, natural cap</td>
<td>1 x 50 ml</td>
</tr>
<tr>
<td>ASYBUF</td>
<td>Assay buffer, red cap, ready to use</td>
<td>1 x 20 ml</td>
</tr>
<tr>
<td>STD</td>
<td>Standard 1-7, synthetic human BNP fragment (0; 200; 400; 800; 1,600; 3,200; 6,400 pmol/l), white cap, lyophilised</td>
<td>7 vials lyophilised</td>
</tr>
<tr>
<td>CTRL</td>
<td>Control, yellow cap, synthetic human BNP fragment, lyophilised exact concentration after reconstitution see label</td>
<td>1 vial lyophilised</td>
</tr>
<tr>
<td>CONJ</td>
<td>Conjugate, (synthetic BNP fragment -HRPO), red dye, amber cap, ready to use</td>
<td>1 x 6 ml</td>
</tr>
<tr>
<td>SUB</td>
<td>Substrate (TMB solution), blue cap, ready to use</td>
<td>1 x 22 ml</td>
</tr>
<tr>
<td>STOP</td>
<td>Stop solution, sulphuric acid, white cap, ready to use</td>
<td>1 x 7 ml</td>
</tr>
</tbody>
</table>

3) ADDITIONAL MATERIAL IN THE KIT
- 2 self-adhesive aluminium films
- Quality control protocol
- Protocol sheet
- Instruction for use

4) MATERIAL AND EQUIPMENT REQUIRED BUT NOT SUPPLIED
- Precision pipettes calibrated to deliver 30 µl, 50 µl, 150 µl, 200 µl, 300 µl and disposable tips
- Distilled or deionised water
- Plate washer is recommended for washing, alternative multichannel pipette or manifold dispenser
- Refrigerator with 4°C (2-8°C)
- ELISA reader capable of measuring absorbance at 450 nm (with correction wavelength at 630 nm)
- Graph paper or software for calculation of results
5) **REAGENTS AND SAMPLE PREPARATION**

All reagents of the BNP Fragment ELISA Assay Kit are stable at 4°C (2-8°C) until the expiry date stated on the label of each reagent.

Sample preparation:
BNP fragments are stable in whole blood, serum or plasma for several hours at room temperature or 4°C (2-8°C). Nevertheless we recommend to separate plasma or serum by centrifugation as soon as possible, e.g. 20 min at 2,000 x g, preferably at 4°C (2-8°C). Aliquot the acquired plasma or serum samples and store them at -25°C or lower. Samples can be subjected to 5 freeze-thaw cycles without any loss of immune reactivity. Lipemic or hemolyzed samples may give erroneous results. Samples should be mixed well before assaying. We recommend duplicates for all values. Samples with values above highest STD could be diluted with STD 1 or BNP fragment negative human serum.

Reconstitution/Handling:
WASHBUF (Wash buffer): Dilute the concentrate 1:20 (1+ 19) e.g. 50 ml WASHBUF + 950 ml distilled water. Crystals in the buffer concentrate will dissolve at room temperature. Buffer is stable at 4°C (2-8°C) until expiry date stated on label. Use only diluted WASHBUF (Wash buffer) for the assay performance.

STD (Standard): Pipette 200 µl of distilled or deionised water into each vial. Leave at room temperature (18-26°C) for 20 min. Swirl gently. The standard concentration is printed on the label. Reconstituted standard is stable at -25°C or lower until expiry date. Avoid freeze-thaw cycles.

CTRL (Control): Pipette 200 µl of distilled or deionised water to the vial. Leave at room temperature (18-26°C) for 20 min. Swirl gently. The final concentration is stated on the label. Reconstituted control is stable at -25°C or lower until expiry date stated on label. Avoid freeze-thaw cycles.

6) **PRINCIPLE OF THE ASSAY**

![Assay principle diagram](image)

7) **ASSAY PROTOCOL**

All reagents and samples must be at room temperature (18-26°C) before use in the BNP ELISA Assay Kit.

Mark position for BLANK/STD/SAMPLE/CTRL (Blank/Standard/Sample/Control) on the protocol sheet.

Take microtiter strips out of the aluminum bag, take a minimum of one well as blank. Store unused strips with desiccant at 4°C (2-8°C) in the aluminum bag. Strips are stable until the expiry date stated on the label.
1. Add 150 µl ASYBUF (Assay buffer) into all wells, except blank.

2. Add 30 µl STD/SAMPLE/CTRL (Standards/Sample/Control) in duplicate into respective well, except blank.

3. Add 50 µl CONJ (Conjugate) into each well, except blank, swirl gently.

4. Cover tightly and incubate overnight (16-25 hours) at 4°C (2-8°C) in the dark.

5. Aspirate and wash wells 5x with 300 µl diluted WASHBUF (Wash buffer), remove remaining WASHBUF by hitting plate against paper towel after the last wash.

6. Add 200 µl SUB (Substrate) into each well.

7. Incubate for 20 min at room temperature (18-26°C) in the dark.

8. Add 50 µl STOP (Stop solution) into each well, shake well.

9. Measure absorbance immediately at 450 nm with reference 630 nm, if available.

8) CALCULATION OF RESULTS

Read the optical density (OD) of all wells on a plate reader using 450 nm wavelength (correction wavelength 630 nm). Subtract the blank OD from the values of STD, CTRL and sample. Construct the standard curve from the OD values of the STD. Use software or graph paper. Obtain sample concentration from this standard curve. The assay was evaluated with a 4PL algorithm. Different curve fitting methods need to be evaluated by the user. Respective dilution factors have to be considered.

Example typical STD-curve:

[Graph showing typical STD-curve]

The quality control (QC) protocol supplied with the kit shows the results of the final release QC for each lot at production date. Data for OD obtained by customers may differ due to various influences and/or due to the normal decrease of signal intensity during shelf life. However, this does not affect validity of results as long as an OD of 1.5 or more is obtained for STD 0 and the value of the CTRL is in range (target range see label).

9) ASSAY CHARACTERISTICS

| Values from apparently healthy individuals (n=76): | In a panel of blood donors the median was 392 pmol/l. |
| Standard range: | 0 to 6,400 pmol/l |
| Sample volume: | 30 µl human serum or plasma (Citrate, EDTA or Heparin) |
| Detection Limit: | 171 pmol/l at 95% B/Bo |
| Incubation time: | 16-25 hrs / 20 min |

For further information on assay characteristics please visit our website www.bmgrp.com technical file or contact our customer service by e-mail export@bmgrp.com or by phone +43/1/29107-45.
10) PRECISION

Intra-Assay: 2 samples of known concentrations were tested 3 times in 1 assay.
Inter-Assay: 2 samples of known concentrations were tested in 8 times in 2 assays by different operators.

<table>
<thead>
<tr>
<th>Intra-Assay (n=3)</th>
<th>Sample 1</th>
<th>Sample 2</th>
<th>Inter-Assay (n=8)</th>
<th>Sample 1</th>
<th>Sample 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean (pmol/l)</td>
<td>763</td>
<td>3,236</td>
<td>Mean (pmol/l)</td>
<td>781</td>
<td>3,199</td>
</tr>
<tr>
<td>SD (pmol/l)</td>
<td>43</td>
<td>251</td>
<td>SD (pmol/l)</td>
<td>45</td>
<td>236</td>
</tr>
<tr>
<td>CV%</td>
<td>6%</td>
<td>8%</td>
<td>CV%</td>
<td>6%</td>
<td>7%</td>
</tr>
</tbody>
</table>

11) TECHNICAL HINTS

- Do not mix or substitute reagents with those from other lots or sources.
- Do not mix stoppers and caps from different reagents or use reagents between lots.
- Do not use reagents beyond expiration date.
- Protect reagents from direct sunlight.
- Substrate solution should remain colourless until added to the plate.
- To ensure accurate results, proper adhesion of plate sealers during incubation steps is necessary.
- Avoid foaming when mixing reagents.

12) PRECAUTIONS

All test components of human source were tested with 3rd generation tests against HIV-Ab and HBsAg; and were found negative. Nevertheless, they should be handled and disposed as if they were infectious. Liquid reagents contain ≤0.1% Proclin 300 as preservative. Avoid contact with skin and mucous membrane. Proclin 300 is not toxic in concentrations used in this kit. It may cause allergic skin reactions – avoid contact with skin or eyes.

- Do not pipette by mouth.
- Do not eat, drink, smoke or apply cosmetics where reagents are used.
- Wear gloves, glasses and lab jacket while performing this assay.
- Sulfuric acid is irritating to eyes and skin. Avoid contact with skin and mucous. Irritations are possible. – Flush with water if contact occurs!!

13) LITERATURE

2. Comparison of Pleural Fluid N-Terminal Pro-Brain Natriuretic Peptide and Brain Natriuretic-32 Peptide Levels. Long AC et al., Chest 2010; 137: 1369-1374
3. N-Terminal Pro-B-Type Natriuretic Peptide as an Indicator of Possible Cardiovascular Disease in Severely Obese Individuals: Comparison with Patients in Different Stages of Heart Failure. Hermann-Arnhof K et al., Clinical Chemistry 2005; 51:138-143
8. Natriuretic peptides/cGMP/cGMP-dependent protein kinase cascades promote muscle mitochondrial biogenesis and prevent obesity. Miyashita K et al., Diabetes 2009; 58: 2880-2892
SYMBOLS

Expiry date / Verfallsdatum / Date de péremption / Data di scadenza / Fecha de caducidad / Data de validade / Uiterste gebruiksdatum / Udløbsdato / Utgångsdatum / Termin Ważności / Lejárati idő / Doba exspiracie / Doba exspirace

Consider instructions for use / Bitte Gebrauchsanweisung beachten / Consultez la notice d'utilisation / Consultare le istruzioni per l'uso / Consulte las instrucciones de utilización / Consulte as instruções para utilização / Raadpleeg de gebruiksaanwijzing / Proszę przeczytać instrukcje wykonania / Vegyük figyelembe a használati utasításban / Postupujte podľa pokynov na použitie / Postupujte dle návodu k použití

Lot-Batch Number / Charge-Chargenummer / Lot-Code du lot / Lotto-Nummer di lotto / Lote-Código de lote / Lote-Código do lote / Lot-Partijnummer / Lot-Batchcode / Lot-Satskod / Numer serii / Lot-Batch szám / Číslo šarže / Číslo šarže

Manufactured by / Hergestellt von / Fabriqué par / Prodotto da / Fabricado por / Fabricado por / Vervaardigd door / Fabrikation af / Tillverkad av / Wyprodukowane pr / Gyártotta / Vyrobené / Vyrobeno

Catalogue Number / Bestellnummer / Numéro de référence / Numero di riferimento / Número de referencia / Número de referência / Referencenummer / Katalognummer / Numer katalogowy / Katalógus szám / Katalógové číslo / Katalogové číslo

Store at between / Lagerung bei zwischen / Conserver à entre / Conservare a tra / Conservar a temp. entre / Armazene a entre / Bewaar bij tussen / Opbevares mellem / Förvaras vid / Przechowywać w / Tároljuk …… között / Skladujte v rozsahu / Skladujte v rozmezí

Contains sufficient for x tests / Inhalt ausreichend für x Tests / Contient suffisant pour x tests / Contenido suficiente para x tests / Contiene suficiente para x pruebas / Contém suficiente para x testes / Bevat voldoende voor x bepalingen / Indeholder tilstrækkeligt til x prøver / Innehållerräcker till x analyser / Zawartość na x testów / Tartalma X teszt elvégzésére elegendő / Obsahuje materiál pre x testov / Obsahuje materiál pro x testů
BI-20852W BNP FRAGMENT EIA
ASSAY PROTOCOL AND CHECKLIST

PREPARATION OF REAGENTS:

- Bring all reagents to room temperature (18-26°C).
- Prepare reagents and samples as instructed.
- Bring unused and prepared components to the storage temperature mentioned in the package insert.
- Take microtiter strips out of the aluminium bag and mark positions on the protocol sheet.

TEST PROCEDURE:

- Add 150 µl ASYBUF (Assay buffer) into all wells, except blank.
- Add 30 µl STD/SAMPLE/CTRL (Standard/Sample/Control) into respective wells, except blank.
- Add 50 µl CONJ (Conjugate) into each well, except blank, swirl gently.
- Cover tightly and incubate overnight (16-25 hours) at 4°C (2-8°C), in the dark.
- Aspirate and wash wells with 300 µl diluted WASHBUF (Wash buffer) five times. Remove remaining buffer by hitting plate against paper towel.
- Add 200 µl SUB (Substrate) into each well.
- Incubate for 20 minutes at room temperature (18-26°C), in the dark.
- Add 50 µl STOP (Stop solution) into each well.
- Read Optical Density at 450 nm with reference 630 nm, if available.
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