

# A Non-Interfering<sup>™</sup> (NI) Protein Assay (high throughput 96-well)

#### **G-Biosciences**

## **Abstract**

The Non-Interfering<sup>m</sup> Protein Assay is a colorimetric assay for determining protein concentrations in protein loading buffer (Laemmli buffer), high  $\beta$ -mercaptoethanol concentrations, and in lipid and vesicle preparations.

Citation: G-Biosciences A Non-Interfering™ (NI) Protein Assay (high throughput 96-well). protocols.io

dx.doi.org/10.17504/protocols.io.e9tbh6n

Published: 13 Sep 2016

## **Guidelines**

### **INTRODUCTION**

The Non-Interfering<sup>m</sup> Protein Assay is a highly sensitive colorimetric assay that overcomes interference by common laboratory agents. The assay removes detergents (non-ionic, ionic and zwitterionic), reducing agents ( $\beta$ -mercaptoethanol, DTT), chelating agents (EDTA), amines (Tris), sugars, and is highly tolerant of strong chaotropic buffers.

The NI<sup> $^{\text{M}}$ </sup> Protein Assay is suitable for determining protein concentrations in protein loading buffer (Laemmli buffer)1-4, high  $\beta$ -mercaptoethanol concentrations (<15%)5-7 and in lipid and vesicle preparations8-9. The NI<sup> $^{\text{M}}$ </sup> Protein Assay has a linear response between 0.5-50 $\mu$ g and has a small sample requirement (1-50 $\mu$ l). The kit components are suitable for 500 assays.

#### **NOTE**

For high throughput 96-well assays, we recommend using 2ml deep round or V- bottom well titer plates. These are available from multiple sources, including VWR, Fisher and USA Scientific. The high throughput protocol requires centrifugation of the 96-well plate at 2-5,000xg and this may require a special centrifuge adaptor.

## ITEM(S) SUPPLIED

Description	786-005	Cat. # 786-896		
UPPA™ I	250ml	250ml		
UPPA™ II	250ml	250ml		
Copper Solution (Reagent I)	50ml	50ml		
Color Agent A	2 x 250ml	2 x 250ml		
Color Agent B	5ml	5ml		
BSA Protein Standard [2mg/ml]	5ml	-		
Non-Animal Protein Standard [2mg/ml]	-	5ml		

Cat #

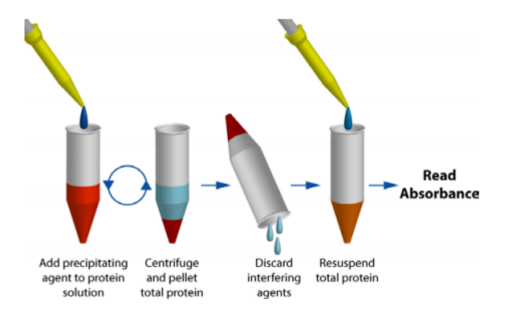
## **STORAGE CONDITIONS**

The kit is shipped at ambient temperature. Upon arrival, store UPPA-I and UPPA-II at room temperature. The remaining kit components should be stored in the dark and refrigerated in its original box. When stored properly, the kit is stable for 1 year.

#### **OVERVIEW**

The Non-Interfering™ Protein Assay is composed of two simple steps (see figure):

- 1. Universal Protein Precipitating Agent (UPPA $^{\text{\tiny M}}$ ) is added to the protein solutions to rapidly precipitate total protein. Protein is immobilized by centrifugation and interfering agents in the supernatant are discarded.
- 2. Protein concentration is assayed by mixing with an alkaline solution containing a known concentration of copper salt; the copper ions bind to the peptide backbone and the assay measures the unbound copper ions. The assay is independent of protein side chains minimizing protein-to-protein variation. The color density is inversely proportional to the amount of protein.



## **TOLERANCE GUIDE**

- 2-Mercaptoethanol, 15%
- Ammonium Sulfate, 40%
- Brij® 35, 1%
- CHAPS, 4%
- CHAPSO, 1%
- Digitonin, 0.3%
- DTT, 0.35M
- EDTA, 0.1M
- Glycerol, 30%
- Guanidine Thiocynate, 4M
- Guanidine.HCl, 6M
- HEPES, 0.1M
- Hydrochloric acid, 0.1N
- Imidazole, 0.5M
- · Iodoacetamide, 15mM
- N-Octyl Glucosidase, 0.5%

- Phosphate buffer, 0.2M
- Sarcosyl, 1%
- SDS, 2%
- Sodium azide, 0.1M
- Sodium Chloride, 0.5M
- Sodium hydroxide, 2.5mM
- Sucrose, 30%
- TCEP, 15mM
- Thesit, 2%
- Thiourea, 2M
- Tris.HCl, 0.5M
- Triton® X-100, 3%
- Triton® X-114, 3%
- Tween® 20, 2%
- Urea, 8M
- Zwittergent® 3-12, 1.5%

## **PREPARATION BEFORE USE**

**Prepare Reagent-II** - Prior to use, prepare an appropriate volume of Reagent II by mixing 100 parts of Color Agent A with 1 part of Color Agent B. (e.g. For 10ml of Reagent II, add 0.1ml Color Agent B to 10ml Color Agent A).

Reagent II can be stored refrigerated for one month or as long as the optical density of the solution at 475-490nm is less than 0.025 O.D.

## **REFERENCES:**

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## **Before start**

**Prepare Reagent-II** - Prior to use, prepare an appropriate volume of Reagent II by mixing 100 parts of Color Agent A with 1 part of Color Agent B. (e.g. For 10ml of Reagent II, add 0.1ml Color Agent B to 10ml Color Agent A).

Reagent II can be stored refrigerated for one month or as long as the optical density of the solution at 475-490nm is less than 0.025 O.D.

# **Materials**

NI™ (Non-Interfering™) Protein Assay 786-005 by G-Biosciences

## **Protocol**

## Step 1.

Perform assays at room temperature. Use 2ml tubes for assay.

#### NOTES

# Colin Heath 29 Aug 2016

For high throughput 96-well assays, we recommend using 2ml deep round or V- bottom well titer plates. These are available from multiple sources, including VWR, Fisher and USA Scientific. The high throughput protocol requires centrifugation of the 96-well plate at 2-5,000xg and this may require a special centrifuge adaptor.

## Step 2.

Prepare a set of protein standards using the supplied BSA or Non-Animal Protein Standard as indicated in the table below:

Tube #	1	2	3	4	5	6
Protein Standard [2mg/ml] (μl	0 (	4	8	12	20	25
Protein (µg)	0	8	16	24	40	50

## Step 3.

Add 1-50µl of the protein samples to be assayed to 2ml tubes.

#### NOTES

Colin Heath 06 Jul 2016

**NOTE**: It is recommended that duplicates are used. The total amount of protein should not exceed 50µg and we recommend various protein dilutions are used to ensure samples are below 50µg.

**NOTE**: For determination of protein concentrations in buffers free of interfering agents skip steps 4-7.

## Step 4.

Add 0.5ml UPPA™ I to each tube and vortex.

## Step 5.

Incubate for 2-3 minutes at room temperature.

© DURATION 00:02:00

# Step 6.

Add 0.5ml UPPA™ II to the tubes and vortex.

## Step 7.

Centrifuge the titer plate at 5,000xg for 7 minutes to pellet the precipitate.

© DURATION 00:07:00

## Step 8.

Invert the titer plate to remove the supernatant and shake to remove all excess supernatant.

## Step 9.

Add 100µl Copper Solution (Reagent I) and 400µl deionized water to the tubes and vortex until the protein precipitate pellet dissolves.

## Step 10.

Using 1ml pipette, **rapidly shoot** 1ml Reagent II directly into each tube containing Reagent I plus DI Water and **immediately mix** it by inverting the tubes.

## **Step 11.**

Incubate at room temperature for 15-20 minutes and then immediately read absorbances at 480nm against DI water.

**O DURATION** 

00:15:00

## **Step 12.**

After incubation, transfer 200µl assay reaction to a flat bottom 96 well micro titer plate and measure the absorbances at 480nm against DI water.

# Step 13.

Plot absorbance against protein concentration and determine protein concentrations of unknowns.

## NOTES

Colin Heath 06 Jul 2016

**NOTE**: Do not subtract blank reading from the sample reading as absorbance will decrease as protein concentration increases.