

MicroBC Assay : protein quantitation kit

Product Description

High quality reagents for the determination of protein concentration by the bicinchoninic acid method

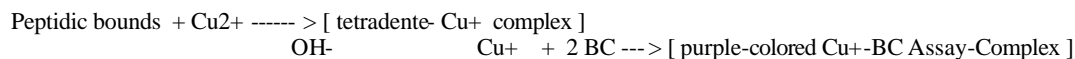
Catalog Number	Name
UP75860A	MicroBC Assay protein Quantitation Kit, complete kit qsp 500 (tube) or 3400 (microplate) determinations Contains: UP67251A Reagent A, 250ml UP67252A Reagent B, 250ml UP67253A Reagent C, 12ml UP36859A BSA standard, 10x1ml
UP75860C	MicroBC Assay protein Quantitation Kit, complete kit qsp 50 (tube) or 340 (microplate) determinations Contains: UP67251C Reagent A, 25ml UP67252C Reagent B, 25ml UP67253C Reagent C, 1.2ml UP36859A BSA standard, 1ml

Storage: 1 year from receipt, at room temperature (long term storage at + 4°C) (.)

For laboratory use only, not for drug, household or medical use

Principle:

The MicroBC Assay is a colorimetric assay: it involves the reduction of Cu²⁺ to Cu⁺ by peptidic bounds of proteins. The BC (BicinChoninic acid) chelates Cu⁺ ions with very high specificity to form a water soluble purple coloured complex. The reaction is increased by high temperatures. As it continues over time, the reaction should be read at a defined time and temperature.



This reaction is measured by the high optical absorbance of the final Cu⁺ complex at 562 nm. Absorbance is directly proportionnal to the protein concentration, with a broad linear range between 1 µg/ml to 100 mg/ml. The protein concentration can be calculated with a reference curve obtained for a standard protein.

Assay Procedure

Labware must be carefully cleaned and rinsed with distilled water to avoid traces of proteins and metals.

Preparation of samples:

The protein concentration must fall in the range of standard curve. Therefore it may be useful to prepare several dilutions to meet this requirement: dilute samples if necessary with their respective buffer (alternatively with water). Each buffer used in samples should be assayed alone to control eventual interference.

Label the tubes and record dilution factor. The dilution factor should be taken in account for the right protein concentration after interpolation from OD_{562nm}. For example :

Sample (name)	Volume sample	Buffer (or water)	Dilution	OD@562 nm	Assayed Protein Concentration (1)	Protein Concentration in Sample (2)
#1: sample1	200	0	1	0.811	67 µg/ml	67 µg/ml
#2: sample2	20 µl	180 µl	1/10	0.592	44 µg/ml	440 µg/ml
#2: sample3						

(1) Protein concentration = calculated from OD@562nm with the standard curve

(2) Protein concentration in sample = assayed protein concentration X dilution factor

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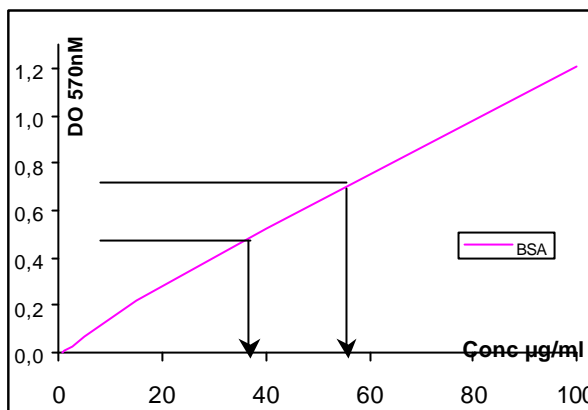
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Preparation of standards :

Uptima recommends to use the protein standard #UP36859A (BSA at 2 mg/ml) for most applications. Prepare a fresh set of protein standards at 100 µg/ml to 0.5 µg/ml, diluted from the stock solution in the same buffer as the samples (alternatively, water may be used; check the sample buffer by analysing it versus water).

Standard	BSA standard 2 mg/ml #UP36859A	Water or Buffer	Final protein Concentration
Standard A	100 µl of Stock	1900 µl	100 µg/ml
Standard B	666.6 µl of A	1000 µl	40 µg/ml
Standard C	176.5 µl of A	1000 µl	15 µg/ml
Standard D	52.6 µl of A	1000 µl	5 µg/ml
Standard E	25.6 µl of A	1000 µl	2.5 µg/ml
Standard F	10.1 µl of A	1000 µl	1 µg/ml
Standard G	5 µl of A	1000 µl	0.5 µg/ml
Blank	H	1000 µl	0

Figure 1 : Typical standard curve with MicroBC Assay #UP75860A



Preparation of the MicroBC Assay reagent (mix A+B+C, 25:25:1) :

Prepare the MicroBC Assay reagent by adding 25 parts of reagent A to 25 parts of reagent B, then 1 part of reagent C. Mix well (a temporary turbidity may have appeared).

The table below gives the volume of reagent needed for an assay with the recommended 8 points standard curve:

Tube assay (Uniplicates)*

Number of standards	Number of samples	Reagent A	Reagent B	Reagent C
8 points	1 to 12	10 ml	10 ml	400 µl
	13 to 32	20 ml	20 ml	800 µl
	32 to 52	30 ml	30 ml	1,2 ml

Microplate assay (Duplicates)**

Number of standards	Number of samples	Reagent A	Reagent B	Reagent C
8 points	8 to 25	5 ml	5 ml	200 µl
	26 to 59	10 ml	10 ml	400 µl
	60 to 94	15 ml	15 ml	600 µl

A duplicate (*), even a triplicate (**), analysis may be recommended for accurate determination; correct volumes accordingly.

Use the mixed MicroBC Assay reagent in the next hours.

Dispose of any unused reagent because possible contamination or degradation may affect further analysis.

Test Tube assay:

	Standard protocol Working range 1µg/ml to 100µg/ml	Enhanced protocol Working range 0.5µg/ml to 40µg/ml
	Allow the reagents to reach room temperature if needed	
a	Pipet 1 ml of each standard, control, and sample into test tubes. Duplicates are recommended.	
b	Add 1 ml of BC Assay reagent (mix A+B+C 25:25:1) per test tube, and mix	
c	Incubate at +37°C for 60 mn *	Incubate at +60°C for 60 mn
d	Cool all test tubes to room temperature and measure the optical absorbance (OD) at 562 nm against the blank (water, or buffer + MicroBC Assay reagent.	
e	Plot the standard curve, and determine the protein concentration.	

Microplate assay:

	Standard protocol	Enhanced protocol
	Working range 1µg/ml to 100µg/ml	Working range 0.5µg/ml to 40µg/ml
	Allow the reagents to reach room temperature if needed	
a	Pipet 150µl of each standard, control, and sample into microplates wells. Duplicates or triplicates are recommended	
b	Add 150µl of BC Assay reagent (mix A+B+C 25:25:1) per test well, and mix (be careful with cross-contaminations)	
c	Incubate at +37°C for 60 mn *	Incubate at +60°C for 60 mn
d	Right after, cool the microplate at room temperature and read the optical absorbance (OD) at 562 nm against the blank (water, or buffer + MicroBC Assay reagent). Alternatively, wavelengths from 540 to 590 nm have been used.	
e	Plot the standard curve, and determine the protein concentration. An example is given below in figure 1.	

*incubation may be performed for 2-3hours for maximum sensitivity (0.5-5+g/ml)

Scientific and Technical Information

Protocol :

Uptima proposes the above standard protocol, that is suitable and convenient for most applications.

Modified protocols may allow to reach different custom requirements. However, this may have incidence on certain performances.

-increased temperatures (i.e. +60°C) and durations (i.e. 2-3hours) improve the sensitivity. Undesired evaporations may lead to a lower accuracy.

-increasing the volume of sample to reagent can also increase the minimal detecte concentration, but take care the assay is not affected because of pH change, or by interfering substances.

-reading the absorbance can be done between 540 and 590 nm, for example with microplate readers lacking a 562 nm filter. Absorbances, hence the sensitivity, is however not optimal (decreased).

Protein Standard :

Uptima complete kits includes the Bovine Serum Albumin #UP36859A because BSA is a common standard that works for most

applications (see below the standard curve). Each user / application may include in the analysis other purified proteins or even any known sample (for example the extract of a reference strain). Ask Uptima for other available standards.

Protein to protein variations :

As with any other protein assay, but with a far lower extend, protein to protein variations may occur with some degree depending on several parameters :

. amino-acid sequences rich in cysteine, tryptophane, tyrosine may increase the MicroBC Assay reaction

. the coloured response may be affected by the primary structure (sequence order), secondary and tertiary (steric conformation) structures of the protein, isoelectric point (pI), side chains, prosthetic groups...

The MicroBC Assay can quantitate immobilized proteins (e), for example gel-coupled proteins, or cells adhering to plates.

Interfering / compatible substances

Some substances may interfere with the BC reaction. It is however remarkable that the MicroBC Assay procedure should be well known for being compatible with a lot of substances, notably most detergents (b). The following table gives some compatible and incompatible substances / concentration :

<i>Compatible (*) Substances</i>	<i>Incompatible Substances</i>
< 3% SDS	Creatinin, Cystein, Tyrosin, Tryptophan
< 2% CHAPS	> 40µM DTT and mercaptans
< 3% Tween 20	Ascorbic acid, H2O2, hydrazides
< 3% Triton X100	EGTA
< 3 M Urea	Phenol Red
< 1% DMSO, DMF	Iron, Copper salts
< 1% Glycerol	
< 1 mM PMSF	
< 0.5% NaN ₃	
TBS (20 mM Tris, 150 mM NaCl, pH 7.6)	
PBS (0,1 M phosphate, 150 mM NaCl, pH 7.2)	
Carbonate / Bicarbonate 100 mM	

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Bicarbonate 40mM

Inquire for other substances

(*) compatibility is determined if there is less than 10% variation of absorbances for the BSA standard at 40µg/ml

The compatible concentrations may depend on protein nature and concentration.

Labware may bear traces of metals that affect the BC Assay reaction. Use cleaned or disposable vials.

To limit the interference of some substances (b), (c), (d) :

-the samples can be diluted provided the protein concentration remains sufficient

-the interfering substance can be removed (f), for example by prior desalting (dialysis...), precipitation (TCA...), purification...

-overcome the presence of copper chelators by increasing the A:B:C ratio of BC Assay reagent up to 20:30:1 (v/v).

In any case, all standards, blanks and controls must be treated in the same way to preserve the accuracy of the assay.

Other Information

ForR&D in vitro use only

Ask Uptima for any question

Literature

(a) Smith P et al, 1995, Measurement of protein using bicinchoninic acid, Anal. Biochem. 150, 76-85

(b) Kaushal et al, 1986, Effect of Zwitterionic buffers on measurement of small masses of protein with bicinchoninic acid, Anal.Biochem, 157, 291-294

(c) Hill et al, 1988, Protein determination using bicinchoninic acid in the presence of sulfhydryl reagents, Anal.Biochem. 170, 203-208

(d) Kessler R & Fanestil D, 1986, Interference by lipids in the determination of protein using bicinchoninic acid, Anal.biochem.159, 138-142

(e) Stich T, 1990, Determination of protein covalently bound to agarose supports using bicinchoninic acid, Anal.biochem.191. 343-346

(f) Brown et al, 1989, Protein measurement using bicinchoninic acid: elimination of interfering substances. Anal.biochem.180, 136-139

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