



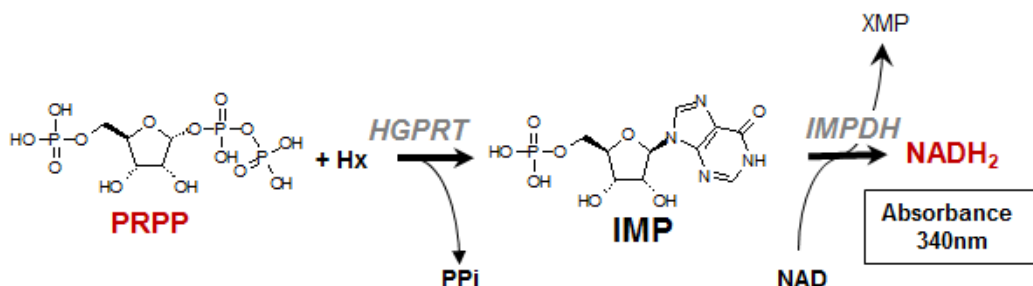
PRECICE® PRPP Assay Kit

For enzymatic measurement of α -D-5-phosphoribosyl-1-pyrophosphate (PRPP)

PRECICE® PRPP Assay Kit is a first non-radioactive assay for *in vitro* measurement of PRPP (α -D-5-phosphoribosyl-1-pyrophosphate) content in cells extracts. This enzymatic assay is based on a coupled reaction involving Hypoxanthine-guanine phosphoribosyltransferase (HGPRT) and Inosine Monophosphate Dehydrogenase (IMPDH).

I. Principle

In the presence of hypoxanthine, HGPRT enzyme converts PRPP to IMP which is further oxidized to XMP by downstream IMPDH enzyme leading to simultaneous NADH_2 formation. The amount of NADH_2 formed is measured spectrophotometrically at 340 nm and is equivalent to the amount of PRPP in the assay.



Short description of procedure:

1. PRPP is extracted from cells with organic solvent.
2. Clarified and filtered extracts are dried in the presence of enzymatic mixtures allowing immediate conversion of PRPP to NADH_2 . For maximal accuracy, the same extracts are dried in parallel in the presence of **HGPRT-IMPDH** enzymes (**Reaction mixture**) or with **IMPDH** enzyme only (**BLANK**).
3. Dried extracts are resuspended in deionized water and clarified by centrifugation.
4. The absorbance of samples is measured at 340nm. The absorbance observed in the absence of HGPRT is used as blank and is subtracted from the absorbance measured in its presence.

Warnings:

1. **Method sensitivity.** PRECICE® PRPP Assay Kit allows PRPP quantification within the range of 8 to 400 μM . Since, according to published data, PRPP concentration in mammalian cells varies from 0.1 to 1 μM , the extracts should be concentrated 10-100 folds to allow spectrophotometric quantification.
2. **PRPP instability.** PRPP is highly unstable cellular metabolite susceptible to degradation by phosphatases and nucleotidases present in all cell types and organisms from prokaryotes to eukaryotes. To avoid PRPP degradation by phosphatases and nucleotidases: all aqueous solutions (deionized water, PBS) should be autoclaved for 30 min; cellular enzymes should be totally precipitated during extraction step.
3. **Control PRPP degradation** using positive control where PRPP (10 μL of freshly prepared 10mM solution) is added to filtered cellular extracts or to cells suspension.

The difficulties of measurement of PRPP in cells due to its low intracellular concentration and high instability can be circumvented by measuring the activity of PRPP-synthetase done in one step.



II. Equipments required

- 1) Plate agitator
- 2) Plate reader fitted with a filter 340nm (ex. Labsystems iEMS Reader MF (Thermo), Epoch (BioTec); PerkinElmer.

III. Kit Contents for 24 analyses (8 samples in triplicate):

Once dissolved, the reagents provided in the kit are not stable and should be used immediately.

A standard PRECICE® PRPP Assay Kit:

- one tube "Cofactor 1" (DTT);
- one tube "Cofactor 2" (NAD);
- one tube "IMPDH enzyme" (1U);
- one tube "HGPRT enzyme" (1U);
- one tube "Reaction buffer 10x";

(1M Tris-HCl; 1MKCl, 130mMMgCL₂ autoclavé, puis ajouté BSA FA-free chauffé 70°C 1h, contains 5mM hypoxanthine et 0.5M KF);

- one transparent 96-well plate (round-bottom 96-well plate Corning, Costar®, ref. 3797)

Not provided:

Autoclaved deionized water

Positive control: PRPP (α -D-5-phosphoribosyl-1-pyrophosphate, available at Sigma-Aldrich, ref. P8296)

Acetone

IV. Recommendations for PRPP extraction / concentration:

Several protocols of PRPP extraction have been described:

- Chemical extraction where proteins are precipitated with perchloric acid (PCA) or trichloroacetic acid (TCA).
- Extraction by boiling where cells or tissue is heated for 85°C.

Since PRPP is stable neither at acidic pH nor at high temperature, NovoCIB has developed an alternative method of PRPP extract based on protein precipitation with organic solvent. The protocol of acetone extraction is described below. This protocol should be validated to every cell type and optimized for cell number to be extracted. For additional information contact us at contact@novocib.com

Important: Use only polypropylene tubes (ex. - Corning, ref. 430829) resistant to acetone. Never use polystyrene tubes.

For 5 ml of cell pellet (to be prepared in duplicate):

1. Chill 45 mL of acetone at -70°C.
2. Prepare cells pellet (4-5mL):
 - a) Transfer 10mL of blood to 50mL tube, add 40mL of autoclaved PBS solution;
 - b) Recover cells by centrifugation. Carefully remove supernatant, wash cell pellet with autoclaved PBS. Repeat 3 times to remove nucleotidases and phosphatases present in medium. Carefully remove all PBS.
3. Chill cell pellet on ice for 5 min.
4. Add 45mL of cold acetone to cell pellet (expected volume 5mL), put immediately at -70°C.
5. Incubate for 3h at -70°C (or overnight at -20°C) to precipitate all proteins.



6. Centrifuge the extract for 30 min at 4°C, carefully recover supernatant (acetone extract) that should be totally clear. While recovering supernatant, avoid to touch and to resuspend protein pellet.
7. Filter acetone extract using PTFE syringe filters (0.2µm) to remove all proteins not precipitated by centrifugation. **The presence of remaining proteins in acetone extract would lead to PRPP degradation during drying of the extracts.**

The efficacy of protein precipitation by acetone can vary for different cell types and can be improved by using the solvents other than acetone or by adding carrier protein.

V. Preparation of enzymatic reaction mixtures

1. Label 15mL tube as **"BLANK"** and fill it with 9mL of autoclaved deionized water. Transfer 1mL of **"Reaction buffer 10x"** to **"BLANK"** tube.
2. Add 100µL of autoclaved deionized water to the tube with **"IMPDH enzyme"**. Agitate until complete dissolution of the powder.
3. Quantitatively transfer the content of 3 tubes with **"Cofactor 1"**, **"Cofactor 2"** and **"IMPDH enzyme"** to a tube **"BLANK"**

To do so:

- pipet 1ml of buffer from **"BLANK"** to each tube and mix them by inverting or pipeting up and down until the powder dissolved.
- transfer the content of two tubes back into a vial **"Reaction mixture 1x"** by pipeting.
- repeat to be sure that all reagent and enzymes of the small tubes and vial are recovered. Mix by gently inverting until complete dissolution. Avoid bubbles.

6. Transfer 5ml of thus prepared **"BLANK"** to orange 15ml tube pre-filled with lyophilized HGPRT enzyme.

You have prepared: 5ml of **"BLANK"**
 5ml of **"Reaction mixture"**

VI. Drying extracts

1. Split filtered acetone extract into 2 glass vials.
2. Add 200µL of **"BLANK"** to one half of acetone extract and 200µL of **"Reaction mixture"** to another.
3. Leave the extract to dry. The faster drying provides better results.

VI. Positive controls:

1. Prepare fresh solution of 10mM PRPP in 10mM NaF solution in water.
2. Add 10µL of 10mM PRPP to 40mL filtered extracts. Split the extract into 2 glass vials, add 200µL of **"BLANK"** and 200µL of **"Reaction mixture"** and proceed with drying.



This positive control allows checking the absence of phosphatases/nucleotidases in cellular extracts and PRPP degradation during drying. The expected optical density (340nm) of positive control after "BLANK" subtraction is 1.225.

To exclude PRPP degradation during extraction, you can add 10µL of 10mM PRPP to cells suspension before or immediately after acetone addition.

VII. Measuring absorbance

1. Resuspend dried extract in 250µL of autoclaved deionized water. Close the vials and incubate for 30min at room temperature.
2. Carefully recover the extracts, centrifuge at 13 000g for 10min. The extracts should be totally clear. If the extracts are not clear, add 50µL of ethanol and centrifuge again.
3. Program plate reader for absorbance reading (absorbance at 340nm), insert the plate into the reader and read absorbance data.

VIII. Calculating PRPP concentration

$$[\text{PRPP}] \text{ in the sample (in } \mu\text{M}) = \frac{\text{Absorbance}_{\text{Reaction}} - \text{Absorbance}_{\text{Blank}}}{0.0049}$$

Where: 0.0049 is extinction coefficient of 1µM NADH at 340nm corrected for l (path-length of round-bottom 200µL-well 96-well plate Corning, Costar®, ref. 3797) as measured with iEMS Labsystems plate reader (Thermo).