ATP Kit SL

Monitoring of ATP in e.g. assays of enzymes and metabolites, cell lysis or platelet aggregation

- Fast assay: Results within minutes
- Stable light: Half-life >1 h
- Detection limit: $10^{-15}$ mol or $10^{-12}$ mol/L ATP
- Reliable: ATP Standard in liquid form
- Flexible: Choice of µ-plate & cuvette methods

Leader in luminescent ATP-assays
ATP Kit SL

Intended use
ATP Kit SL is intended for monitoring of adenosine triphosphate (ATP) in the range $10^{-12} - 10^{-6}$ mol/L. The low decay rate of the light emission (around 0.9 % per min) is due to a low consumption of ATP in the firefly luciferase reaction, and a luciferase activity remaining unchanged during the measurement. This allows enzymatic formation or degradation of ATP and cellular release of ATP to be monitored by measuring the light intensity).

Applications
1. Assays of enzymes and metabolites participating in ATP converting reactions.
2. Monitoring of oxidative phosphorylation and photophosphorylation.
3. ATP release during cell lysis.
4. ATP release during platelet aggregation.

Assay principles
Firefly luciferase catalyses the following reaction:

$$\text{luciferase} \quad \text{ATP} + \text{D-luciferin} + \text{O}_2 \rightarrow \text{AMP} + \text{PPi} + \text{oxy} \text{luciferin} + \text{CO}_2 + \text{light}$$

The assay has been optimised to give a stable light at all ATP levels up to $10^{-6}$ mol/L.

Instruments
The stable light makes it possible to use manual single tube luminometers, automatic tube luminometers or microplate luminometers. The detection limit depends on luminometer sensitivity.

Kit contents
1. The kit is intended for 200-1000 assays depending on the final assay volume (0.2-1 mL).
2. ATP Reagent SL. 4 vials of lyophilised reagent containing D-luciferin and luciferase.
3. Diluent C 10 mL. 4 vials for reconstitution of ATP Reagent SL.
4. ATP Standard 5 mL. 4 vials containing 10^{-5} moles/L of ATP.
5. Tris-EDTA Buffer 2x100 mL. 2 bottles containing 0.1 mol/L Tris(hydroxymethyl) aminomethane, 2 mmol/L EDTA and adjusted to pH 7.75 with acetic acid.

Assay procedure using internal ATP Standard
The light emission is measured before and after the addition of a known amount of ATP. This strongly increases the reliability of the assay and makes it possible to express ATP results in moles rather than rlu or other non-chemical units. The procedure below is for tube luminometers. The assay can also be automatically performed in microplate luminometers.

1. Add sample and Tris-EDTA Buffer to give a total volume of 0.8 mL in the cuvette.
2. Add 0.2 mL ATP Reagent SL.
3. Monitor the light emission corresponding to sample ATP, I_{smp}.
5. Measure the light emission corresponding to sample plus standard ATP, I_{smp+std}.

Calculations:
Calculate the sample ATP concentration in the cuvette by the following equation:

$$\text{ATP}_{smp} = 10^{-7} \times \frac{I_{smp}}{I_{smp+std} - I_{smp}}$$

The factor $10^{-7}$ is the concentration of ATP Standard in the cuvette.


Product characteristics
Detection limit: $10^{-12}$ mol/L or $10^{-15}$ mol ATP
No. of determinations (cuvettes): 200
No. of determinations (microplate): 1000

Ordering info
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