The **ARC(CK2)-Fluo Protein Kinase Assay Kit** is used for a homogenous, high-throughput binding/displacement assay for screening/characterization of inhibitors of the protein kinase CK2.

### Table 1. ARC(CK2)-Fluo Protein Kinase CK2 Assay Kit (CK2-FLUO590)

<table>
<thead>
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<th>Application</th>
<th>Determination of $K_d$ values for both ATP- and substrate-competitive protein kinase CK2 inhibitors</th>
</tr>
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<tbody>
<tr>
<td>Targets</td>
<td>CK2 (both the free CK2α catalytic subunit and CK2 holoenzyme)</td>
</tr>
<tr>
<td>Detection principle</td>
<td>Fluorescence anisotropy or polarization Ex: 540 nm/ Em: 590 nm [1]</td>
</tr>
<tr>
<td>Assay setup</td>
<td>Homogeneous, 1-step, mix &amp; read; 384-well low-volume format</td>
</tr>
<tr>
<td>Assay volume</td>
<td>10 - 25 µl</td>
</tr>
<tr>
<td>Probe</td>
<td>CK2-FLUO590 probe</td>
</tr>
<tr>
<td>Cat. No.</td>
<td>Amount</td>
</tr>
<tr>
<td>---------------</td>
<td>-----------------------------</td>
</tr>
<tr>
<td>CK2-FLUO590</td>
<td>1 Kit (384 assay points)</td>
</tr>
</tbody>
</table>

For in vitro use only

Storage: Upon arrival open the package and store the reagent pouch at -20°C, the rest of the package can be stored at ambient temperature.

Quality guaranteed for 6 months since the date of purchase

Kit contents

2x1.1 mL 10x Assay Buffer
30 μL CK2-FLUO590 probe (1 μM in DMSO)
2x solid DTT (1x for 5000 μl 1x assay buffer)
1 Film-covered 384-well low-volume microtiter plate
1 Transparent lid

To be provided by the user

Protein kinase
Cofactors if necessary
Shaking thermostat

A Fluorescence Polarization Plate-Reader possessing high sensitivity (< 5 mP standard deviation at 1 nM Fluorescein), e.g., PHERAStar FS or comparable, with excitation at 540 nm and emission at 590 nm. Tested on PHERAstar plus (BMG Labtech), Synergy Neo (BioTech) and Cytation 5 (BioTech) platereaders.
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1. Description

The ARC(CK2)-Fluo590 Protein Kinase Assay Kit is for a homogenous, high-throughput binding/displacement assay designed for determination of the concentration of active form of the protein kinase and screening/characterization of inhibitors of protein kinase CK2 (Tab. 1).

The principle of the assay relies on the competitive displacement of the fluorescent probe [CK2-FLUO590] from its complex with the protein kinase by the inhibitor (Fig. 1) [1]. CK2-FLUO590 probe possesses high affinity towards CK2α (K_D = 0.4 nM) [2], it is binding with high affinity to both the free CK2α catalytic subunit and CK2 holoenzyme and enables the **characterization of both ATP- and substrate competitive inhibitors** due to its unique bisubstrate character (simultaneous association with both binding sites of the protein kinase).

![Figure 1. Principle of ARC(CK2)-Fluo Protein Kinase Assay Kit](image)

The fluorescent probe CK2-FLUO590 probe has low molecular weight (MW < 2000) and possesses low value of fluorescence anisotropy (polarization) in solution. When CK2-FLUO590 probe binds to the protein kinase CK2, a high molecular weight complex is formed (MW > 40000) which leads to increase of anisotropy of the solution. An inhibitory compound competes with CK2-FLUO590 for binding to the kinase and displaces the probe from the complex resulting in decrease of fluorescence anisotropy.
2. Calculation of $K_d$ values

When $K_i > K_D$, where

- $K_D$ is the dissociation constant of the CK2-FLUO590:kinase complex, $K_D = \frac{[\text{CK2-FLUO590]\text{[kinase]}}}{[\text{CK2-FLUO590:kinase}]}$
- $K_i$ is the dissociation constant of the inhibitor:kinase complex, $K_i = \frac{[\text{inhibitor][kinase]}}{[\text{inhibitor:kinase}]}$

The $IC_{50}$ values for the inhibitors are linearly proportional to the $K_i$ values [2]. Comparison of $IC_{50}$ values for a compound under evaluation (X) and a reference compound (Y) with previously established $K_i$-value enables the determination of the dissociation constant $K_i$ for X from the comparison of $IC_{50}$-values for these 2 compounds without knowing the concentration of the kinase in the assay

$$K_i(X) = \frac{IC_{50}(X)}{IC_{50}(Y)} \times K_i(Y) \quad \text{(Eq.1)}$$

- $IC_{50}$ is the concentration of the inhibitor that displaces 50% of CK2-FLUO590 probe from CK2-FLUO590:kinase complex

3. Experimental set-up

3.1 Preparation of 1x Assay Buffer

Prepare 5 mL of fresh 1x Assay Buffer (sufficient for 8 inhibitors) by diluting the 10x Assay Buffer:

- Dissolve one batch of provided DTT in 550 μl 10x Assay Buffer
- Transfer 500 μl of the prepared DTT solution in 10x Assay Buffer to a centrifuge tube with a capacity of at least 5 mL
- Add 4500 μL deionized water (e.g. Milli-Q Quality)*

* reduce the volume of water accordingly if cofactors and/or activators will be added (Tab.2)

Please note: 1x Assay buffer is active for 18 hours and should not be used beyond that time. Always prepare fresh buffer and do not store the 1x Assay Buffer!

3.2 Preparation of the CK2-FLUO590 probe stock solution

Thaw the CK2-FLUO590 probe (1 μM in DMSO) at room temperature and dilute 10-fold in 1x assay buffer.

The resulting 100 nM solution can be stored at 4°C in the dark for one day.
4. Determination of the Optimal Protein Kinase Concentration

This assay is tested for both the free CK2α catalytic subunit and CK2 holoenzyme. The recommended concentrations can be found in Table 2. However, the following procedure is mandatory since protein kinase activity may vary in different preparations.

This is a simplified approach and sufficient for the procedure of choosing the optimal concentration of the kinase for the inhibition assay. For the determination of the $K_D$ value of the protein kinase:CK2-FLUO590 probe complex or the concentration of the active form of the protein kinase, please refer to: www.kinasera.com/assay_support

Table 2. Recommended concentrations of the CK2-FLUO590 probe and protein kinases in the measurement solution

<table>
<thead>
<tr>
<th></th>
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</tr>
</thead>
<tbody>
<tr>
<td>CK2α</td>
<td>2</td>
<td>3</td>
<td>0.4</td>
</tr>
</tbody>
</table>

Please note: Centrifugation of all solutions before dispensing them into the wells is recommended.

a. Prepare 30 µl of 125 nM solution of protein kinase in 1x assay buffer
   - Note that the prepared concentration is 1.25-fold higher than the final concentration in the well (Tab. 3)
b. Dispense 16 µl of 1x assay buffer into wells (#1B to #1N) of a single column of the 384-well plate
c. Dispense 24 µl of 125 nM protein kinase into well #1A
   - Prepare a titration series by transferring 8 µl of protein kinase sample from well #1A into #1B, mix thoroughly by careful pipetting. Transfer 8 µl from #1B to #1C and continue stepwise until #1L. After mixing thoroughly in #1L, discard the tip with 8 µl solution
d. Dispense 20 µL of 1x assay buffer into wells #1O and #1P of the column (for blank correction)
e. Prepare 60 µl of 5x CK2-FLUO590 probe working solution in 1x assay buffer
   - mix 6 µl of 100 nM CK2-FLUO590 probe with 54 µl of 1x assay buffer
f. Add 4 µL of 5x CK2-FLUO590 probe working solution into wells #1A to #1N
g. Incubate the plate on a shaking thermostat for 15 minutes at 30°C and 400 rpm
h. Measure the fluorescence anisotropy values of each well with a fluorescent plate reader
   - Use well #1M or #1N for the adjustment of instrument settings (the polarization value of free CK2-FLUO590 probe should be set to 0.02)
   - Use wells #1O and #1P for blank correction
Plot fluorescence anisotropy values against the concentration of the kinase. Fit the curve to the hyperbolic function
Table 3. Final concentrations of the CK2-FLUO590 probe and protein kinase (final volume: 20 µl per well)

<table>
<thead>
<tr>
<th>Well #</th>
<th>Protein conc.</th>
<th>CK2-FLUO590 probe conc.</th>
</tr>
</thead>
<tbody>
<tr>
<td>1A</td>
<td>100 nM</td>
<td>2 nM</td>
</tr>
<tr>
<td>1B</td>
<td>33.3 nM</td>
<td>2 nM</td>
</tr>
<tr>
<td>1C</td>
<td>11.1 nM</td>
<td>2 nM</td>
</tr>
<tr>
<td>1D</td>
<td>3.7 nM</td>
<td>2 nM</td>
</tr>
<tr>
<td>1E</td>
<td>1.2 nM</td>
<td>2 nM</td>
</tr>
<tr>
<td>1F</td>
<td>412 pM</td>
<td>2 nM</td>
</tr>
<tr>
<td>1G</td>
<td>137 pM</td>
<td>2 nM</td>
</tr>
<tr>
<td>1H</td>
<td>45.7 pM</td>
<td>2 nM</td>
</tr>
<tr>
<td>1I</td>
<td>15.2 pM</td>
<td>2 nM</td>
</tr>
<tr>
<td>1J</td>
<td>5.1 pM</td>
<td>2 nM</td>
</tr>
<tr>
<td>1K</td>
<td>1.7 pM</td>
<td>2 nM</td>
</tr>
<tr>
<td>1L</td>
<td>0.6 pM</td>
<td>2 nM</td>
</tr>
<tr>
<td>1M</td>
<td>-</td>
<td>2 nM</td>
</tr>
<tr>
<td>1N</td>
<td>-</td>
<td>2 nM</td>
</tr>
<tr>
<td>1O</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>1P</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

The optimal kinase concentration for the inhibition assay equals the concentration that gives **80% of the maximal anisotropy signal**.
5. Determination of IC$_{50}$ and $K_d$ values of inhibitors

**Please note:** Centrifugation of all solutions before dispensing them into the wells is recommended.

a. Dispense 16 µl of 1x assay buffer to one column per one inhibitor of the 384-well plate (wells #B to #O)

b. Prepare a dilution of each inhibitor in 1x assay buffer (the starting concentrations should be 1.25-fold higher than the desired final well concentrations as CK2-FLUO590 probe and protein kinase will be added)
   - E.g. if the highest concentration of the inhibitor in the dilution series should be 1 mM, prepare 1,25 mM solution in 1x assay buffer
   - Dispense 24 µl of each inhibitor into #A in separate columns
   - Prepare a serial dilution by transferring 8 µl from #A into #B, mix thoroughly by careful pipetting and continue until row #M for each inhibitor

c. Dispense 20 µL of 1x assay buffer into well #P of the column (for blank correction)

d. Prepare a 5x Master Mix of the kinase and CK2-FLUO590 probe in 1x assay buffer:
   - Use the optimal kinase concentration determined earlier (step 4)
     - Example: For a final kinase concentration of 3 nM and CK2-FLUO590 concentration of 2 nM prepare a solution of 15 nM kinase and 10 nM CK2-FLUO590 probe
   - Volume: 56 µL per inhibitor and extra for pipetting loss

e. Add 4 µl of the 5x Master Mix to each well #A to #N
   - This leads to a final well volume of 20 µl

f. Prepare 5x CK2-FLUO590 probe in 1x assay buffer
   - Volume: 4 µl per inhibitor and extra for pipetting loss

g. Add 4 µl of 5x CK2-FLUO590 probe solution to well #O

h. Incubate the plate on a shaking thermostat for 15 minutes at 30°C and 400 rpm

i. Determine the anisotropy values of each well with a fluorescence plate reader
   - Use well #O for the adjustment of instrument settings
   - Use wells #P for blank correction

j. Plot the obtained anisotropy values against log C (inhibitor) and analyze the curves using sigmoidal dose-response (variable slope) function

k. The $K_d$ values of competitive inhibitors can be calculated using Eq. 1

**Selected references**

