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Confidential

QL12 QL12-80

Done

Title: QL12-90

Goal: PTO Cy5 uptake, 500 nM, 1 h preincubation + 2 h incubation with different inhibitors
+ activation followed by dialysis

Started on: March 2, 2021

Modified on: June 9, 2021

Closed date

Project: Phosphorothioates

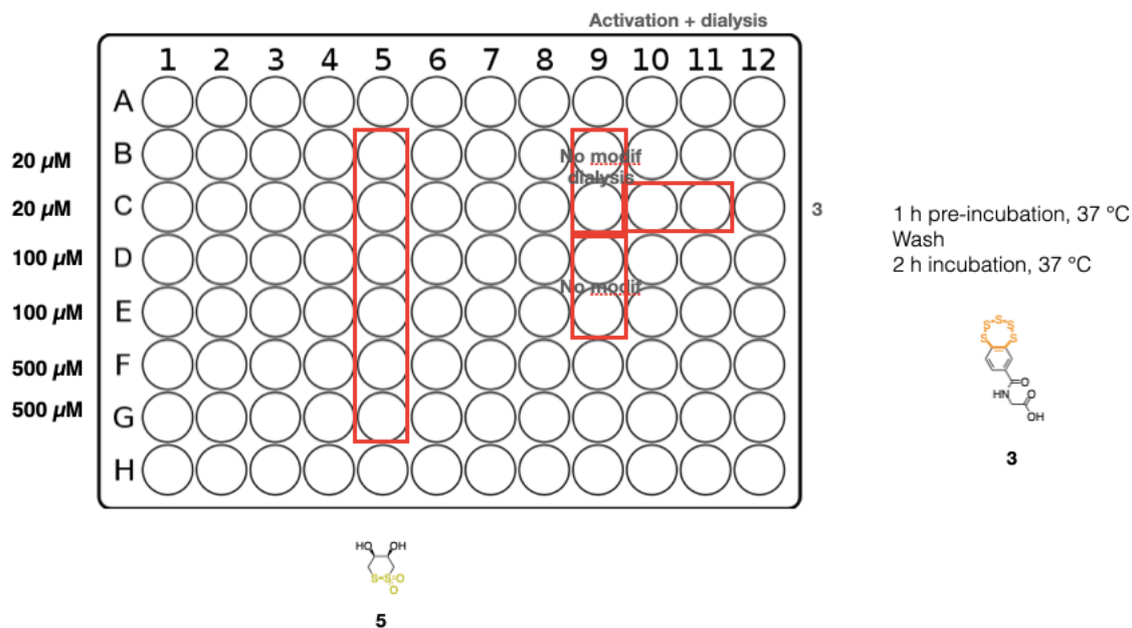
Type: PTO Cy5 Inhibition

HeLa Kyoto cells were seeded at 8×10^4 cells/mL in FluoroBrite DMEM + 10% FBS on μ -Plates 96-well ibiTreat sterile and kept at 37 °C with 5% CO₂ overnight. The following day, cells were washed with FluoroBrite DMEM (3 × 3 mL/well) using a plate washer, keeping a final volume of 135 μ L/well.

Dilutions of Cy5-labelled phosphorothioate and inhibitor in PBS were prepared in a 96-well V-bottom plate and added to the μ -Plate containing the cells at the desired time (15 μ L/well, 10x final concentration in PBS) to reach a final volume of 150 μ L/well, and cells were incubated for 1 h at 37 °C with 5% CO₂ in DMEM. Cells were washed with FluoroBrite DMEM (3 × 3 mL/well) using a plate washer, keeping a final volume of 135 μ L/well, and the corresponding solution of phosphorothioate was added (15 μ L/well, 10x final concentration in PBS) to reach a final volume of 150 μ L/well, and cells were incubated for 2 h at 37 °C with 5% CO₂ in DMEM.

Then, Hoechst 33342 (15 μ L/well, 100 μ g/mL in PBS) was added. After 15 min of incubation at 37 °C with 5% CO₂, the medium was removed and cells were fixed with a PFA solution (3% in PBS, 80 μ L/well) for 20 min. The plate was washed with PBS (3 x 3 mL/well) and imaged using a high-content automated microscope.

Sequence: AGGTCCCCATACACCGAC



Attached files

QL12-80 Plate.pdf
 QL12-80-Treated.xlsx
 QL12-63+80-Activation.pzfx