GLYCOWORKS RapiFluor-MS QUICK START PROTOCOL

Streamlined Protocol for Disulfide Rich Glycoproteins - 96 sample (4 x 24 Format)

- Set heat blocks to at least 90 °C and 50 °C.
- Protocol is based on 1.5 mg/mL glycoprotein starting concentration.
- DTT is not provided.



STEP 1: Rapid Deglycosylation

- 1. Reconstitute 1 vial of the Intact mAb Mass Check Standard (1 mg/vial) in 670 μ L 18.2 M Ω water to create a 1.5 mg/mL solution.
 - Note: For glycoproteins with a formulation buffer containing nucleophiles or anionic reagents (e.g., His, Gly, Tris, PQ_s^{-3}), if a Glycan C_mAX Column is applied for LC separation, it is highly recommended to desaft the sample with water prior to Sten 1.
- 2. Prepare RapiGest™/DTT denaturing buffer RapiGest 3% (w/v); DTT 2 μg/μL (approx. 15 mM):
 - a) Dissolve 10 mg RapiGest SF Surfactant in 200 μ L of rapid buffer, vortex.
 - b) Dissolve DTT in water to 5 μg/μL. Add 135 μL DTT solution to RapiGest SF solution, vortex.
- 3. Dilute PNGase F enzyme (35 $\mu L)$ with 255 μL water for a total of 290 $\mu L.$
- Add 10 µL of 1.5 mg/mL glycoprotein into the reaction tube or well.
- 5. Add 10 μL of 3% (w/v) RapiGest SF/2 μg/μL DTT denaturing buffer to above tube, aspirate to mix.
- 6. Heat at least to 90 °C for 3 minutes.
- 7. Cool at room temperature for 3 minutes.
- 8. Add 10 μL Rapid PNGase F and aspirate to mix.
- 9. Incubate at 50 °C for 5 minutes.
- 10. Cool at room temperature for 3 minutes.



STEP 2: Rapid Labeling of Glycosylamines

- 1. Aliquot 15 μ L of deglycosylated glycoprotein reaction mixture to a new tube and dilute with 15 μ L of 18.2 M Ω water.
 - Note: For maximum consistency in yield, 2x concentration of RapiFluor-MS reagent is recommended, as compared to the original GlycoWorks protocol (720005343EN). This protocol describes halving the volume of the deglycosylated mixture to achieve the 2x increase in RapiFluor-MS label concentration.
- Add 280 µL of anhydrous DMF directly to one vial of 23 mg of RapiFluor-MS™ Reagent. Mix to solubilize.
- 3. Add 10 µL of the *Rapi*Fluor-MS solution to the deglycosylation mixture and aspirate to mix.
- 4. Allow the labeling to proceed at room temperature for 5 minutes.
- Dilute the reaction with 360 µL of acetonitrile (ACN) and aspirate to mix.





STEP 3: HILIC Cleanup of Labeled Glycosylamines

- Set up a GlycoWorks™ HILIC µElution Plate and add in shims or spacer and waste tray.
- 2. Condition wells by adding 200 µL of water per well.
- 3. Equilibrate wells by adding 200 µL 85% ACN.
- 4. Load ACN-diluted samples (~400 μL).
- Wash wells with two (2) 600 μL volumes of 1% formic acid, 90% ACN.
- Replace waste tray with sample collection tray loaded with 600 µL tubes.
- 7. Elute glycans with three (3) 30 µL volumes of SPE elution buffer into 600 µL tapered bottom inserts.
- Dilute SPE eluate with 310 μL of the GlycoWorks SPE Diluent (DMF/ACN). Aspirate to mix.
 Note: For a Glycan C_a AX separation sample, either skip dilution Step 8 or dilute with 310 μL of water.
- 9. Cap the tubes with pre-slit cap mats.

► For the complete Care and Use Manual, visit waters.com and search 715004793EN.

► For more details on this method, download Application Notes 720005506EN and 720007038EN.