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Fluorescence assay for MERS-CoV Mpro activity measurement

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Charline Giroud¹, oleg.fedorov¹

¹Center for Medicines Discovery, University of Oxford

ASAP Discovery



Charline Giroud

Center for Medicines Discovery

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Protocol status: Working We use this protocol and it's

working

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Abstract

This protocol details the fluorescence assay for MERS-CoV Mpro protease activity measurement. This method is intended to measure the activity of viral proteases by using a specific labelled peptide that allows the detection of the cleaved product. The substrate contains the cleavage-sequence specific to the tested protease and is labeled in C-terminal by the fluorophore Edans (ex 336 nm; em: 455 nm) and in N-ternimal by the quencher Dabcyl (ex 472 nm). In the case of a non-cleaved substrate, the proximity of Dabcyl to Edans prevents the emission and the detection of the fluorescence at 455 nm. The cleavage of the peptide by the protease allows Edans' fluorescence emission and detection.

Attachments



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Materials

Reagents

Assay buffer:

A	В
Tris pH 7.0	50 mM
NaCl	150 mM
Glycerol	10%
DTT (optional)	0.5 mM

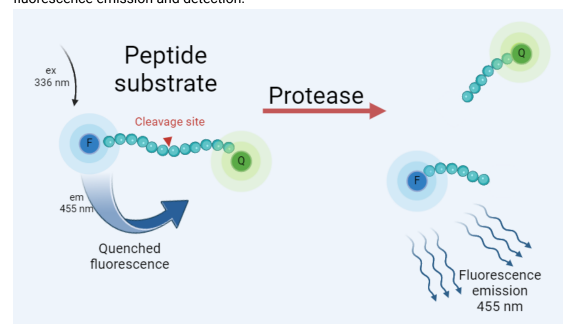
- Incubation: (5) 01:00:00 at

 Room temperature
- EV-D68 3C: Protein stocks were stored at 3 -80 °C and used as 2x solution ([M] 1 micromolar (μM) , [M] 0.5 micromolar (μM) final assay concentration) in assay buffer.
- **Positive control**: GC376 (Pubchem CID 71481119), [M] 50 micromolar (µM) top final assay concentration.
- Plates: ProxiPlate-384 Plus, white, Greiner cat# 6008280.
- Liquid handler: Echo® acoustic liquid handler (Beckman Coulter, USA).
- **Plate reade**r: Pherastar FS, BMG Labtech (Germany), 350-460 FI optic module, the plate is read every 30 s for 2 hours and shacked during 5 s before the first reading.

MERS-CoV Mpro Assay

3h

This method is intended to measure the activity of viral proteases by using a specific labelled-peptide that allows the detection of the cleaved product. The substrate contains the cleavage-sequence specific to the tested protease and is labeled in C-terminal by the fluorophore Edans (ex 336 nm; em: 455 nm) and in N-ternimal by the quencher Dabcyl (ex 472 nm). In the case of a non-cleaved substrate, the proximity of Dabcyl to Edans prevents the emission and the detection of the fluorescence at 455 nm. The cleavage of the peptide by the protease allows Edans' fluorescence emission and detection.



Asset URL:

Reagents and equipment

2 Assay buffer: [M] 50 millimolar (mM) Tris pH 7.0, [M] 150 millimolar (mM) NaCl,

3h 0m 35s

[M] 10 % (V/V) glycerol and [M] 1 millimolar (mM) TCEP (optional).

Incubation: 01:00:00 at room temperature.

MERS-MPro: protein stocks were stored at -80C and used as [M] 2 x solution (

[M] 1.2 micromolar (μ M) , [M] 0.6 micromolar (μ M) final assay concentration) in assay buffer.

Positive control: Ebselen (Pubchem CID 3194), [M] 50 micromolar (μM) top final assay concentration.

Substrate: Edans-GVLQSGLV-LysDabcyl-K (LifeTein, USA) prepared as a stock solution at 10



mM in DMSO and used at 2x solution ([M] 20 micromolar (μ M) , [M] 10 micromolar (μ M) final concentration assay concentration) in assay buffer.

Liquid handler: Echo R acoustic liquid handler (Beckman Coulter, USA).

Plate reader: Pherastar FS, BMG Labtech (Germany), 350-460 FI optic module, the plate is read every 00:00:30 for 02:00:00 and shacked 00:00:05 before each reading.

MERS-MPro IC50 Measurement

3h

Add Δ 50 μ L of 2x protein [M] 1.2 micromolar (μ M) solution to each well containing the compounds to be tested previously dispensed onto the plate.



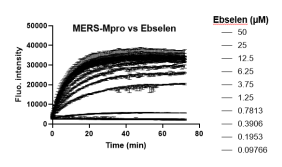
Incubate the mix for \bigcirc 01:00:00 at \bigcirc Room temperature to initiate the enzymatic reaction by the addition of \bigcirc 50 μ L of 2x (\bigcirc 1M] 20 micromolar (μ M)) substrate solution using the plate reader injector.

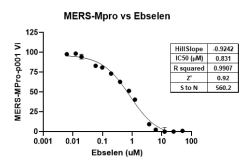
1h

Read the fluorescence intensity at 350/460 nm every 00:00:30 for 00:00:30 in kinetic mode, which includes a shaking step of the plate before each measurement.

1m

6 Calculate the IC50 by plotting the initial velocity against various concentrations of tested inhibitors by using a four-parameter dose-response curve in Prism (v8.0) software.





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7 Calculate the mean (μ) and the standard deviation (σ) of fluorescence intensity and then calculate the signal-to-background ratio and the Z' or Zfactor. (s: signal; c: control).



$$\text{Estimated Z-factor} = 1 - \frac{3(\hat{\sigma}_s + \hat{\sigma}_c)}{|\hat{\mu}_s - \hat{\mu}_c|}$$

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