Studying the molecular mechanism of viral replication in real time using the CLARIOstar^{Plus} with ACU

Dr Marko Noerenberg¹ Dr Vincenzo Ruscica² Dr Manoja Rasamanikkam³ Dr Alfredo Castello^{1,2*} ¹ Centre for Virus Research, MRC-University of Glasgow, 446 Bearsden Road, G61 1QH, Glasgow, Scotland (UK) ² Department of Biochemistry, University of Oxford, South Parks road, OX1 3QU, Oxford, (UK) ³ BMG LABTECH, Aylesbury, (UK) *Correspondence to: alfredo.castello@glasgow.ac.uk

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Introduction

Viruses are obligate intracellular pathogens that can infect all living organisms. They not only assume an outmost medical relevance due to their connection with several human diseases, but they also have a major economic impact given the costs required for medical treatments and for protecting cattle and crops. Viruses encode a limited number of proteins, and they thus interact and hijack host cellular machineries to promote their replication and spread. Therefore, understanding host-virus interactions become crucial to uncover the molecular mechanisms underpinning infection, and for the development of new treatments. With the emergence of 'omics' in the last decade, known host-virus interactions have been massively expanded, calling for novel approaches to test the importance of cellular proteins in virus infection in a systematic way. Moreover, the discovery of antivirals often relies on the screen of large libraries of compounds, which also poses a technical challenge. Hence, to test the importance of cellular proteins in virus infection and to discover new antivirals, robust and efficient highthroughput methods are required to measure viral fitness without using expensive reagents such as luciferin. Here we describe a method to analyse viral gene expression in near real time using fluorescence as a proxy for infection. This assay can discover not only if a given condition inhibits or enhances virus infection, but also provides insights into its effects on the kinetics of infection.

96-well microplate, flat µclear bottom (Greiner Bio-One). This protocol is, also compatible with 384-well microplates.

- Colourless DMEM medium
- SINV-mCherry virus stock (~5x10⁸ pfu/ml)
- BMG LABTECH CLARIOstar Plus microplate reader

every 15 minutes. Addition of drugs or other reagents (e.g. interferon) can be performed before or after starting the programme in the plate reader.

Instrument settings

Materials & Methods

Optical settings	Filters	Excitation 570-15 Dichroic 593.8 Emission 620-20
	Focal height	4.4 mm
	Gain	2900
General settings	Number of flashes	8
	Settling time	0 s
Scan settings	Orbital averaging	
	Scan diameter	4 mm
Kinetic settings	Number of cycles	92
	Cycle time	900 s
Incubation	37 °C, 5% CO ₂	

Assay Principle

with ACU

Experimental Procedure

HEK293 cells were seeded at a density of 5x10⁴ cells in a 96-wells microplate with flat µclear bottom. Infection was performed using SINV-mCherry at 0.1 multiplicity of infection (MOI), and cells were incubated at 37 °C with 5% CO_2 in the CLARIOstar ^{Plus} plate reader for 24 hours. Incubation time should be adapted to the lifecycle duration of each virus (e.g. 72 hours for HIV). The viral genome expression was monitored by the detection of the red fluorescent signal derived from mCherry (excitation: 570 nm, and emission: 620nm), which was measured

Results & Discussion

To assess viral genomic replication, WT (wild type), XRN1+/- (partial knockout), and XRN1-/- (complete knockout) cells were infected with SINV-mCherry and the expression of the red fluorescent protein was monitored every 15 minutes for 24 hours using the CLARIOstar Plus microplate reader. The detection of fluorescent signal every 15 minutes for 24 hours post infection (hpi) was enough to cover both, the early and late phases of SINV on viral gene expression. infection. In HEK293 WT cells, SINV gene expression was detected at around 8 hpi with a subsequent constant increase (figure 2). In cells completely lacking XRN1, viral gene expression drops to background level. These results reflect that XRN1 is a critical host factor for SINV infection and the lack of it causes cells to become refractory to infection¹ (figure 2).

In addition, this assay allows the testing for drug effects on viral fitness. For example, figure 3 shows that cyclosporin A, a known inhibitor of cyclophilin A (PPIA), moderately delays SINV gene expression to a comparable level to that of a PPIA knock out (figure 3, bottom panel). This example goes to highlight the potential of this application for screening the effects of proteins and drugs

A - SINV-mCherry genome



Fig. 1: Schematic representation of (A) SINV-mCherry and (B) the viral fitness assay.

Herein, we illustrate a fluorescent-based method for monitoring viral gene expression using a BMG LABTECH CLARIOstar ^{Plus} microplate reader with Atmospheric Control Unit (ACU). As an illustrative example, Sindbis virus (SINV, Alphavirus genus) with an engineered genome containing the coding sequence for mCherry fluorescent protein was employed. Expression of mCherry can be used as a proxy for viral gene expression and can be measured at regular intervals throughout the infection. Although exemplified with SINV, this approach has been applied successfully to a broad range of viruses including the human immunodeficiency virus (HIV).



Fig. 2: Detection of mCherry fluorescence signal in the indicated HEK293 cells. Adapted from¹

The high sensitivity of the assay allows for the profiling of intermediate phenotypes in a very reproducible manner. For example when XRN1 is partially depleted, SINV infection is delayed but not fully supressed¹ (figure 2).



Fig. 3: Detection of mCherry fluorescence signal in the presence of the PPIA inhibitor (top panel) and in the wild type and PPIA knock out HEK293 cells (bottom panel). Adapted from¹

Conclusion

The viral fitness assay can be successfully performed using the CLARIOstar and CLARIOstar Plus fluorescence microplate readers with ACU, permitting the study of viral gene expression in multi-well plates in real time for up to 72 hours post infection. This method allows for the simultaneous screening of a broad range of experimental conditions, such as drug libraries and knockout cell lines in virus infection and can thus be employed for the study of host-virus interactions.

References

. Garcia-Moreno, *M. et al.* System-wide Profiling of RNA-Binding Proteins Uncovers Key Regulators of Virus Infection. Mol Cell, doi:10.1016/j.molcel.2019.01.017 (2019).