



Pseudovirus Ebola Zaire (Luciferase)

Lot #230825



Quality control report

1. Summary

The lot number #230825 is a lentivirus-based pseudovirus pseudotyped with the glycoprotein of the Ebola Zaire variant. This quality control report demonstrates that the lot #230825 is efficient for cell transduction and can be effectively neutralized by a standard neutralizing antibody.

2. Transduction efficiency assay

| | |
|------------------------------|--|
| Target cells | HEK293 cells (TIM-1 ⁺) |
| Volume of pseudovirus | 0, 1, 5, 10, 20, 30 or 40 μ L/well |
| Detection signal | Luminescence (firefly luciferase) |
| Detection method | Microplate reader |

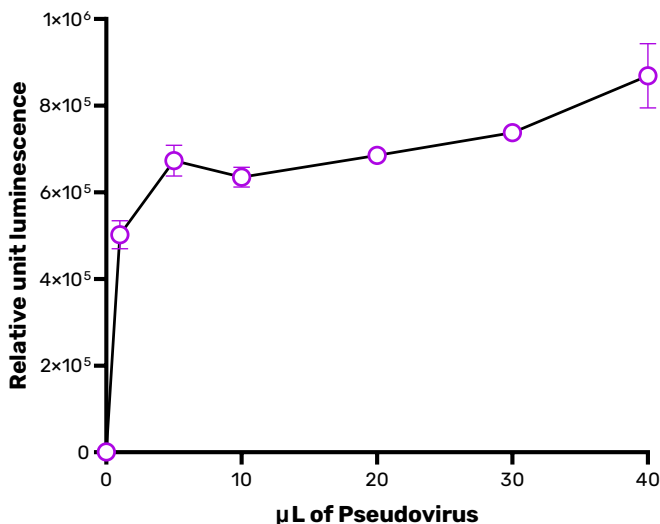


Figure 1: Transduction efficiency curve.

A volume range of pseudovirus was mixed in a final volume of 50 μ L of culture complete medium, in a 96-well plate. Then, an additional 50 μ L containing 20 000 HEK293 cells (TIM-1⁺) was seeded in each well. Luc expression analysis was performed 48 hours post-infection by a luminescence microplate reader.

Conclusion The Luc Ebola Zaire pseudovirus (#230825) can transduce the target cells.

3. Neutralization assay

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|-------------------------------------|--|
| Target cells | HEK293 cells (TIM-1 ⁺) |
| Volume of pseudovirus | 1 μ L/well |
| Neutralizing antibody (Nabs) | Anti-Ebola surface glycoprotein [KZ52], AB00690-10.0 |
| Detection signal | Luminescence (firefly luciferase) |
| Detection method | Microplate reader |

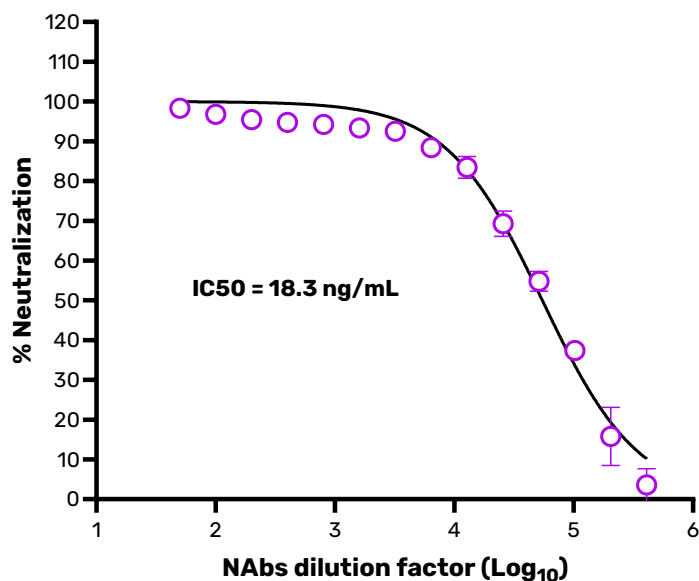


Figure 2: Neutralization curve.

A monoclonal neutralizing antibody ([AB00690-10.0](#)) at 1 μ g/ μ L was serially diluted in a final volume of 50 μ L of complete medium and incubated for 1 hour at 37 °C, with 1 μ L of pseudovirus, in a 96-well plate. Then, an additional 50 μ L containing 20 000 HEK293 cells (TIM-1⁺) was seeded in each well and incubated for 48 hours. Finally, an additional 100 μ L of [Bright-Glo™ Luciferase](#) buffer was added in each well and incubated for 2 minutes. Data in relative unit luminescence (RLU) were obtained from the analysis of 150 μ L of the cell lysate with a microplate reader. Raw data were analyzed using a log(inhibitor) vs normalized-response (variable slope) non-linear regression model in Prism v10 (GraphPad). Percentages of neutralization were normalized considering only cells into wells as 100% neutralization and cells transduced by pseudoviruses without any NABs as 0% neutralization. Data are representative of duplicates.

Conclusion The Luc Ebola Zaire pseudovirus #230825 can be efficiently neutralized by neutralizing antibodies.

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3. Additional information

| | |
|-----------------------------|---|
| Caution | We recommend determining the optimal pseudovirus volume to use according to your specific experimental conditions. |
| Pseudovirus | 3 rd generation, replication incompetent |
| Pseudotyping | Ebola virus glycoprotein from the 2014 outbreak in west Africa (GENBANK : KP096420.1) |
| Glycosylation origin | Human |
| Reporter protein | Firefly luciferase |
| Storage | - 80 °C, avoid freeze/thaw cycles |
| For more information | mathias.mangion@ivanobioscience.com Message object should contain : "#230825" |