

# Quick Manual

## Gaussia-Juice Luciferase Assay

### Package Sizes:

Art.no.	Volume	No. Assays
<b>102541</b>	<b>100 ml</b>	<b>(1.000-2.000 assays)</b>
<b>102540</b>	<b>5 x 10 ml</b>	<b>(500-1.000 assays)</b>
<b>102541-1</b>	<b>1 x 10 ml</b>	<b>(100-200 assays)</b>
<b>102541-0</b>	<b>1 x 4 ml</b>	<b>Sample</b>

### Components included:

#### **Gaussia-Juice**

**Reaction buffer**  
**Store at +4°C.**

#### **Coelenterazine**

**Vial A**  
**Store at -20°C.**

#### **Reconstruction Buffer**

**Vial B**  
**Store at -20°C.**

#### **2x Lysis-Juice**

**dual concentrated Lysis-Buffer**  
**Store at +4°C.**

### Reconstruction:

Please dissolve the substrate in Vial A with the Reconstruction buffer in Vial B and mix gently. The prepared mixture can be stored at -80 °C. After 30 days the activity of the dissolved substrate will start linear to decrease.

Just before measuring the Gaussia Luciferase please dilute the substrate mixture in the Reaction Buffer 1:50 (2µl Substrate in 100 µl reaction buffer).

Before measuring the buffer should be tempered 37 °C - at least room temperature.

Remainders of the mixed Gaussia-Juice should not be frozen again because they will loose noticeable activity.

### Preparation of Cell Lysates:

Gaussia-Juice Luciferase Assay includes a Lysis-Juice. Lysis-Juice is dual concentrated and suitable for mammalian cells which were transfected with Gaussia Luciferase. Please dilute the dual concentrated lysis buffer with water or within your cell culture.

### **Standard Protocol for Cells Cultured in Multiwell Plates**

Required volumes of Lysis-Juice

Culture Plate	Vol. Lysis-Juice
6-well	500µl
12-well	250µl
24-well	100µl
48-well	65µl
96-well	20µl

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### **Standard Protocol for Cells Cultured in Multiwell Plates**

- Remove the growth medium from your adherent cells
- Wash the monolayer two or three times with phosphate buffered saline (PBS)
- Add the required volume of Lysis-Juice to each well (see Table.1)
- Place the plate on a shaker for 15 minutes at room temperature, additional steps of up-down pipetting of cell-buffer mix will increase the lysis (or freeze and defrosting steps could be added)
- Ready to use cell-lysate can placed in storage tubes or measured in the plate by adding Gaussia-Juice.

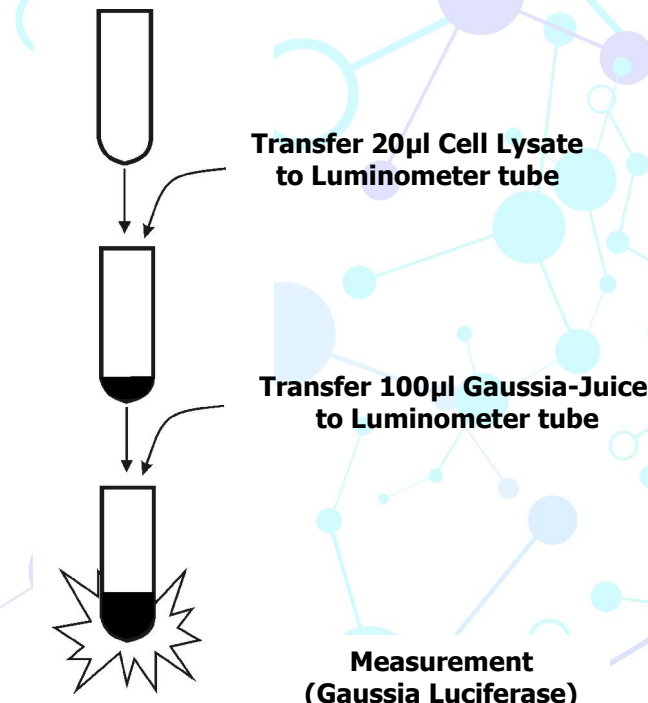
### **Standard Protocol:**

Program Luminometer suggestion: 2 sec. Delay, 5 sec. measurement time (can be customized)

- 1.) Transfer 20µl cell lysate to a luminometer tube / microplate well.
- 2.) Add the prepared **Gaussia-Juice**.
- 3.) Measurement starts after 2 sec. delay for 5 sec. duration.

### **Standard procedure:**

**Program Luminometer**  
(delay 2sec./measurement duration 5sec)



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