

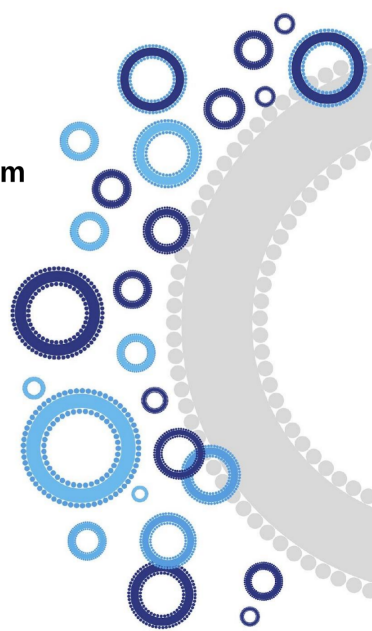
# User Guide

# **CD9 ExoLISA**

# **Exosome Detection System**

Europium time-resolved immunofluorescence assay for detection of exosome antigens

Cat EX501



ISFR	$\sim$ 1	
1SER	(-1	III ) <b>–</b>

# **Contents**

Storage	4
Product components	
Equipment and materials required but not supplied with this kit	4
Product information	5
Reagent preparation	6
Technical positive control preparation	6
Protocol	7
Technical positive control example	9
Related products	9
Purchaser Notification	11

# CD9 ExoLISA Exosome Detection System

### **Storage**

- Store all components at 4°C.
- The kit has a shelf life of at least three months from receipt.

## **Product components**

- 1 x Streptavidin-coated 96-well plate (8-well strip format)
- 1 x Technical positive control, lyophilized, 25 μg.
- 1 x Eu-labelled CD9 mAb (in 40 μl TSA buffer), 2.75 μg
- 1 x Biotinylated CD9 mAb (in 22 μl PBS 7.4 pH, 15 mM NaN<sub>3</sub>), 22 μg

#### The CD9 antibody used in the kit is human specific.

- 1 x Assay buffer, 22 ml
- 1 x Europium fluorescence intensifier (EFI Solution), 11 ml
- 1 x 25x wash buffer, 20 ml
- 1 x User Guide

# Equipment and materials required but not supplied with this kit

- Time-resolved fluorescence microplate reader
- Automatic plate washer
- Plate shaker
- Pipettes for dispensing reagents
- Multichannel micropipette reservoir
- Distilled / Milli-Q water
- Phosphate buffered saline (PBS)

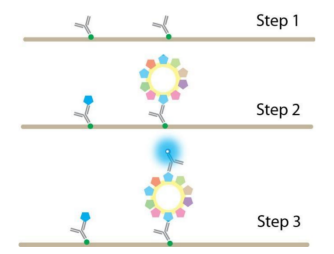
#### **Product information**

#### Introduction and assay principle

In the ExoLISA exosome assay, the same antibody is used for binding of target to the assay plate and for detection. The assay consists of a monoclonal antibody (labelled with biotin) bound to a streptavidin-coated plate which captures protein present on the surface of exosomes (Figure 1). Subsequently, an identical monoclonal antibody (labelled with europium) is used for detection. Because the capture and detection antibody are identical, they require two linked copies of the same epitope for a signal to be detected. Exosomes provide an ideal structure to link CD9 molecules and allow detection of CD9 in this assay. Exosomes typically have multiple copies of CD9 facing towards the attachment surface and additional CD9 molecules available for detection. Any non-specific binding of capture and detection antibodies is unlikely to generate a signal. Using a europium fluorophore (see below) provides high levels of sensitivity for the assay, which is able to detect small changes in the abundance of the target CD9 protein even within unpurified complex biological samples, such as blood plasma and cerebrospinal fluid.

**Fluorophores** are chemical substances that emit light following excitation by light or other electromagnetic radiation. The emission of light from a fluorophore is maximal immediately following excitation and decays over a period of time. Time-resolved fluorimetry uses fluorophores which have long decay periods. For such fluorophores, measurement of emitted light can be performed when the excitation light is no longer present, thus increasing sensitivity.

**Europium** is a fluorophore which produces an extended emission decay and has a wide Stokes shift with maximal excitation at 340 nm and peak emission at 615 nm. ExoLISA assays are time-resolved immunofluorescence assays which utilize europium and have been developed to measure the abundance of CD9 protein specifically associated with exosomes in biological fluids including urine, saliva, cell culture medium, cerebral spinal fluid and blood plasma.



- **Step 1.** Biotinylated antibody is bound to streptavidin-coated assay plates.
- **Step 2.** Biological samples are added. Exosomes and any free antigen are captured by the antibody.
- **Step 3.** Europium-labeled antibody is added and binds specifically to exosome antigen. The epitopes of bound monomers are already occupied and not detected. Samples are read in a time-resolved fluorescence plate reader.

Figure 1. Protocol overview

#### General note

The technical positive control provided with the kit can be used to verify that adequate technical and handling procedures have been conducted and a signal can be detected at the end of the protocol. The provided technical positive control cannot be used to generate a quantification curve for directly quantifying exosomes. Extracellular vesicles of different origin have different CD9 prevalence on their surface.

If absolute quantification is required, an internal quantification curve can be obtained for each population of exosomes studied by correlating CD9 ExoLISA signal intensity with Nanoparticle Tracking Analysis (NTA) measurements at different concentrations of purified exosomes.

## **Reagent preparation**

- Preparation of the wash buffer: Dilute the wash buffer concentrate 25x in Milli-Q water (20 ml concentrate in 500 ml Milli-Q water). The diluted solution may then be stored at room temperature.
- **Preparation of the biotin CD9 in assay buffer**: [Dilutions should be made on the day the assay is performed]. Briefly centrifuge the vial containing the antibody to gather all the liquid at the base of the vial. Prepare a ~2 ng/µl working solution of biotinylated CD9 Ab by diluting 22 µl (22 µg) in 11 ml of assay buffer.
- **Preparation of the europium CD9 in assay buffer**: [Dilutions should be made on the day the assay is performed]. Briefly centrifuge the vial containing the antibody to gather all the liquid at the base of the vial. Prepare a ~0.25 ng/µl working solution of Eu-labeled CD9 mAb, by diluting 40 µl (2.75 µg) in 11 ml of assay buffer.

# **Technical positive control preparation**

- 1. 25  $\mu g$  of exosomes have been purified from cell culture and their total protein concentration determined by BCA assay. The 25  $\mu g$  of exosomes are lyophilized and need to be reconstituted with 250  $\mu l$  of Milli-Q water to generate a final concentration of 100  $\mu g/ml$ .
- 2. Prepare 7 microcentrifuge tubes, each with 125 µl of PBS.
- 3. Prepare 8 serial dilutions as shown in the table below. Ensure that samples are properly mixed in each tube.
  - We recommend beginning with a concentration of 100  $\mu$ g/ml, as this will ensure that the most diluted sample is within the limit of detection of the assay.

Table 1. Positive control preparation

Technical positive control #	Exosomes	Diluted in PBS (μl)	Exosome concentration (125 µl solution)
1	25 μg (resuspended in 250 μL Milli-Q water)	0	100 μg/ml
2	125 µI of Tube #1	125	50 μg/ml
3	125 µI of Tube #2	125	25 μg/ml
4	125 µI of Tube #3	125	12.5 μg/ml
5	125 µI of Tube #4	125	6.3 μg/ml
6	125 µI of Tube #5	125	3.1 μg/ml
7	125 µI of Tube #6	125	1.6 μg/ml
8	125 µI of Tube #7	125	0.8 μg/ml

See example assay reading on page 9.

# **Protocol**

#### Coat the wells with CD9 capture antibody

- 1. Add 100 µl of the freshly prepared dilute solution (2 ng/µl) of biotin-CD9 antibody (prepared as described above) to each well.
- 2. Incubate the plate for 1 hour at room temperature on a plate shaker at 750 RPM.
- 3. Wash the plate using an automatic plate washer. Wash each well three times using 250 µl wash buffer for each cycle.
- 4. Remove the remaining wash buffer.

#### Add the sample

- 5. Clear cells and cellular debris from test samples by centrifuging at 3,000 x g for 20 minutes.
- 6. Transfer 100 μl of the test sample supernatant to each well. Use 100 μl of PBS instead of sample in order to generate a blank reading.
- 7. Incubate the plate for 1 hour at room temperature on the plate shaker at 750 RPM.
- 8. Wash the plate using an automatic plate washer. Wash each well three times using 250 µl wash buffer each time.
- 9. Carefully aspirate the remaining wash buffer.

#### Add the europium-labeled CD9 detection antibody

- 10. Add 100 µl per well of the freshly prepared Eu-labeled CD9 antibody dilution (prepared as described above).
- 11. Incubate the plate for 1 hour at room temperature on the plate shaker at 750 RPM.
- 12. Wash the plate using an automatic plate washer. Wash each well three times using 250 µl wash buffer each time.
- 13. Carefully aspirate the remaining wash buffer.

#### Signal enhancement and reading

- 14. Add 100 µl of EFI solution to each well.
- 15. Incubate the plate for 15 minutes at room temperature on the plate shaker at 750 RPM.
- 16. Measure fluorescence on a time-resolved fluorescence microplate reader as shown on the table below. Measurements should be performed in triplicate. Before taking the readings, make sure that the plate reader is set to read the fluorescence at the bottom of the plate. \*Please check plate reader manual for optimum value of this variable for reading Europium fluorescence.

#### Table 2. Instrument settings

	Time resolved fluorescence (TRF)	
Optic settings	Filters	Excitation: 340 nm Emission: 615 nm
	Number of flashes	200 with flash lamp
Conoral cottings	Settling time	0.1 - 0.3 s*
General settings	Integration start/lag time	60 - 200 μs*
	Integration time	400 μs

# **Technical positive control example**

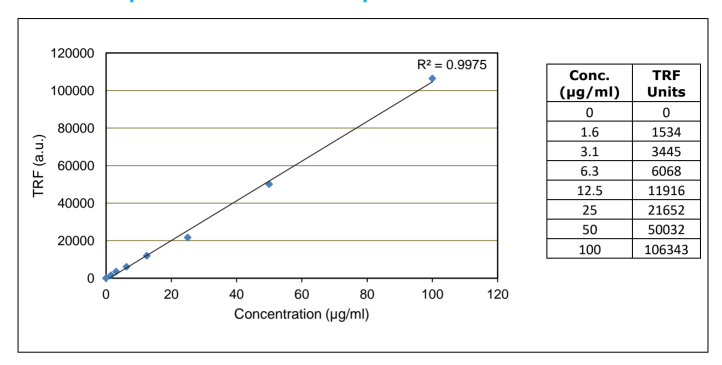


Figure 2. Example readings obtained using the ExoLISA exosome assay with the provided technical control sample (exosomes purified from cell culture). Note the linearity of response over a wide range of concentrations.

# **Related products**

Related products	Product description	Product code
Exosome purification	Exo-spin™ purification kit	EX01, EX02, EX03, EX04, EX05, EX07
Nanoparticle Tracking Analysis (NTA) size profiling service	ZetaView NTA Particle Analysis Service	ZV-1 and ZV-12

#### Exo-spin<sup>™</sup> purification kit

The Exo-spin<sup>™</sup> technology combines precipitation and size exclusion chromatography techniques, making it a superior method for exosome separation and concentration, allowing for high specificity and high recovery of exosomes. Exo-spin<sup>™</sup> is available in 5 different configurations represented with catalogue codes EX01, EX02, EX03, EX04, EX05, and EX07; specifically designed and optimized for different sample types and downstream applications.

#### NTA size profiling service

Exosome characterization service for analysis of particle size and particle concentration using the ZetaView instrument from Particle Metrix.

1	ISED	$\alpha$	IIDE

#### **Purchaser Notification**

Limited warranty Cell Guidance Systems and/or its affiliate(s) warrant their products as set forth in the Terms of Sale found on the Cell Guidance Systems web site at www.cellgs.com/Pages/Terms\_and\_Conditions.html

If you have any questions, please contact Cell Guidance Systems.

This product incorporates licensed technologies. The purchase of this product conveys to the purchaser the limited, non-transferable right to use the purchased amount of the product only to perform internal research for the sole benefit of the purchaser. No right to resell this product or any of its components is conveyed. This product is for internal research purposes only and is not for use in commercial services of any kind, including, without limitation, reporting the results of purchaser's activities for a fee or other form of consideration. For information on obtaining additional rights, please contact info@cellgs.com.

CELL GUIDANCE SYSTEMS AND/OR ITS AFFILIATE(S) DISCLAIM ALL WARRANTIES WITH RESPECT TO THIS DOCUMENT, EXPRESSED OR IMPLIED, INCLUDING BUT NOT LIMITED TO THOSE OF MERCHANTABILITY, FITNESS FOR A PARTICULAR PURPOSE, OR NON-INFRINGEMENT. TO THE EXTENT ALLOWED BY LAW, IN NO EVENT SHALL CELL GUIDANCE SYSTEMS AND/OR ITS AFFILIATE(S) BE LIABLE, WHETHER IN CONTRACT, TORT, WARRANTY, OR UNDER ANY STATUTE OR ON ANY OTHER BASIS FOR SPECIAL, INCIDENTAL, INDIRECT, PUNITIVE, MULTIPLE OR CONSEQUENTIAL DAMAGES IN CONNECTION WITH OR ARISING FROM THIS DOCUMENT, INCLUDING BUT NOT LIMITED TO THE USE THEREOF.

Cell Guidance Systems' reagents and services enable control, manipulation and monitoring of the cell, both *in vitro* and *in vivo*.

#### **Growth Factors**

- Recombinant
- PODS® Sustained Release

#### **Exosomes**

- Purification
- Detection
- Purified Exosomes
- NTA Service

#### Cytogenetics

- Karyotype Analysis
- Array Hybridization

#### Other research products and services

- Matrix Proteins
- Small Molecules
- Cell Counting Reagent
- Lipid Quantification Assay







General info@cellgs.com
Technical Enquiries tech@cellgs.com
Orders order@cellgs.com

www.cellgs.com

#### **EUROPE**

Cell Guidance Systems Ltd
Maia Building
Babraham Bioscience Campus
Cambridge
CB22 3AT
United Kingdom

T +44 (0) 1223 967316 F +44 (0) 1223 750186

#### USA

Cell Guidance Systems LLC Helix Center 1100 Corporate Square Drive St. Louis MO 63132 USA

T 760 450 4304 F 314 485 5424