

# PLASMA

Next generation of Exosome isolation for biomarker discovery in liquid biopsy

## **EXOSOME PURIFICATION KIT FOR PLASMA**

# USER GUIDE

#### STORAGE

All components can be stored at room temperature.

### **PRODUCT COMPONENT**

EXOGAG - Plasma<sup>m</sup> isolation kit 50ml (50-25 samples). 1 x User guide. Not supplied: collection tubes.

EXOGAG - PLASMA is a specific, quick and inexpensive method to optimize the process of exosome isolation.

#### **PRODUCT INFORMATION**

The EXOGAG - Plasma precipitation reaction is based on the interaction between the precipitation solution and glycosaminoglycans located on exosome surface. After a simple incubation, the exosomes can be isolated by a short centrifugation.

EXOGAG - Plasma technology allows the isolation of exosomes from plasma based on the affinity of exosomes to the precipitation reagent, which allows their isolation from a complex sample. EXOGAG - Plasma is a patented exosome purification method that allows the isolation of exosomes from a plasma sample with a minimal amount of co-precipitated material, such as protein or genetic material (DNA, RNA or microRNA) which makes it an ideal product for biomarkers research and their transfer to the clinic.

High sensibility and specificity		Inexpensive	Small sample needed	No specific equipment needed
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SAMPLE	SAMPLE, vol.	EXOGAG - PLASMA, vol.
Plasma	500µl	1ml
Plasma	1ml	2ml





#### **EXOGAG - Plasma Exosomes Isolation Protocol.**

**1. Collect plasma sample.** Samples can be frozen until the moment they are used; if the samples have been frozen, thaw and temper them before processing.

- **2. Centrifuge the sample** at 2000 x g for 5' to remove cells and cell debris.
- **3. Transfer the supernatant** to a new tube and discard the pellet of possible cell debris.

4. Add the volume of sample to isolate exosomes to a new tube and add twice the volume of EXOGAG - Plasma precipitation reagent, as shown in the table.

**5. Mix the sample** and EXOGAG - Plasma precipitation reagent by inverting the tube or vortexing to homogenize the final solution (the solution will have a characteristic blue colour).

- **6. Incubate the sample** for 5' at 4°C.
- 7. Centrifuge the sample at  $3000 \times g$ ; 15' at 4°C.
- 8. Remove the supernatant being careful not to remove the pellet containing the exosomes (this pellet will be dark blue).
- 9. Resuspend the exosomes in the appropriate buffer (pipetting repeatedly up and down), depending on the technique.

