Lyophilized Exosome Standards (2nd Generation)









| | ExoPC3 | 100 μg | Exos. from PC-3 (human prostate adenocarcinoma cell line) | |
|---|----------|--------|---|-----|
| | ExoHT29 | 100 μg | Exos. from HT-29 (human colon cancer cell line) | |
| | ExoMCF7 | 100 μg | Exos. from MCF-7 (human breast cancer cell line) | |
| E | ExoSERUM | 100 μg | Exos. from human serum | |
| | ExoA375 | 100 μg | Exos. from A-375 (human malignant melanoma cell line) | RUC |
| | ExoRPMI | 100 μg | Exos. from RPMI8226 (human myeloma cell line) | |
| ı | ExoCaCo2 | 100 μg | Exos. from CaCo2 (human colon cancer cell line) | |
| | ExoA549 | 100 μg | Exos. from A-549 (human lung cancer cell line) | |
| ı | ExoPANC1 | 100 μg | Exos. from PANC-1 (human pancreas cancer cell line) | |
| | ExoMSC | 100 μg | Adipose-derived Mesenchymal stem/stromal cells | |
| | | | (MSCs) derived exosomes. | |
| | | | | |

1. INTRODUCTION

Exosomes are small (~40-100 nm) extracellular vesicles (EVs) released from all cell types upon fusion of an intermediate endocytic compartment, the multivesicular body (MVB) (1) with the plasma membrane. Exosomes found in body fluids and cell culture supernatants. They are though to provide a means of intercellular communication(2.3) and of transmission of macromolecules between cells allowing the spread of proteins, lipids, mRNA, miRNA and DNA and as contributing factors in the development of several diseases. Exosomes can also modulate cancer microenvironment(4) and the immune response(5, 6).

PRODUCT DESCRIPTION

Lyophilized exosomes (~1x1012) derived from human cancer cell line (8,9). Exosomes are isolated by differential ultracentrifugation (7).

- Tested application: Flow Cytometry (FMC), Nanoparticles Analysis (NTA Tracking Nanosight), Western Blot (WB), BCAProtein Assay, Mass Spectrometry, RT-qPCR.
- Species reactivity: Human
- Presentation: Lyophilized
- Reconstitution of Exosomes: For reconstitution, we recommended adding sterile, distilled water to achieve a final exosome concentration of $1\mu g/\mu I$ (e.g., for 100 μg standard, add 100 µl of dH2O). After the addition of water, recap vial and briefly vortex making sure that the liquid has been gently distributed and has covered the entire inside of the vial. After vortexing, make sure that the solution is collected at the bottom of the vial, if not, centrifuge shortly the vial solution. Now the standard is ready to use.

3. APPROPIATE STORAGE AND HANDLING CONDITIONS

Lyophilized exosomes can be stored between 2ºC and 8ºC for up to 2 years without functional compromise. Immunostep recommends storing small, single -use aliquots of reconstituted exosomes, at - 20°C for up to one month or at - 80°C for longer periods, preferably in locations in frost-free freezers, without appreciable temperature fluctuation. This will minimize protein denaturation that can occur after multiple freeze/thaw cycles.

Reconstituted exosomes, store properly, are functionally guaranteed for up to six months from date of reconstitution. Any unfrozen and/or unused exosome standard can be stored at 4ºC for short term use (<1 week), and should not be re-frozen.

EVIDENCE OF DETERIORATION

Reagents should not be used if any evidence of deterioration is observed. For more information, please contact our technical service; tech@immunostep.com

5. BIOSAFETY LEVEL 1

Biosafety classification is based on 2000/54/EC Directive from the European Council. Customer has to ensure that their facilities comply with biosafety regulations for their own country.

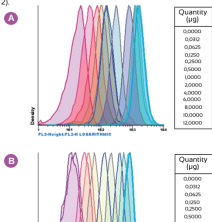
Revision Nº 17I Emission date: 04/10/2021

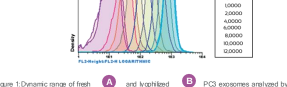
WARRANTY 6.

Warranted only to conform to the quantity and contents stated on the label or in the product labelling at the time of delivery to the customer. Immunostep disclaims hereby other warranties. Immunostep's sole liability is limited to either the replacement of the products or refund of the purchase price.

7. PERFORMANCE DATA

All exosome standard batches has been validated using FCM. WB and NTA Analysis, additionally in order to compare the effects of lyophilization process we have compared all lyophilized batches with respect to fresh exosomes stored at -20°C. Exosome batches are checked and compared for the presence of the CD63 and CD9, a common exosome marker, by FCM (Fig. 1) and WB (Fig. 2).





B PC3 exosomes analyzed by flow cytometry. A and lyophilized Figure 1: Dynamic range of fresh Relationship between background noise and specific signal at different exosome concentrations. Exosomes were captured by CD63+ (Clone TEA3/18) capture beads and subsequently detected by Anti-CD9 PE (Clone VJ 1/20).

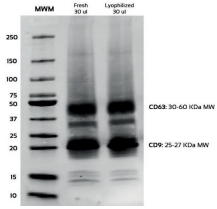
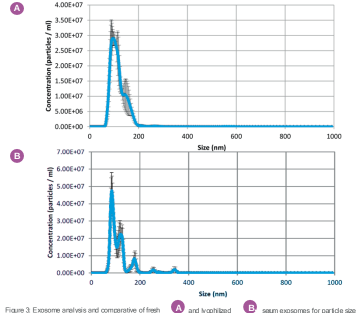


Figure 2: Fresh and Tyophilized MCF-7 exosome batches were analyzed and compared by WB in native conditions for exosomal markers, by anti-CD9 (Clone VJ1/20) and anti-CD63 (Clon TEA3/18) antibodies at a 1:1000

All exosome batches are also subjected to NTA analysis for concentration and particle size estimation (Fig. 3).

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and concentration by NTA. NanoSight LM10HSB. Analysis was carried out with 1ul of purified exosomes diluted

in 999 ul of HEPES buffer (diution 1:1000). The purified exosomes showed a size distribution profiles, with peak diameters from 50 - 150 nm and concentrations about 1x1010 exosomes/ml.

Exosomes have been proposed to provide means for intercellular exchange of macromolecules allowing the transfer of proteins, lipids, mRNA and miRNA, MiRNAs are a class of 17-24 nt small. noncoding RNAs. Exosomal miRNAs play an important role in disease progression. These miRNAs can stimulate angiogenesis or facilitate metastasis in cancers. Therefore, exosomal miRNas present potential for uses as noninvasive biomarkers that can indicate the stage of the disease

The miRNA content of each of our lyophilized exosomes has been analyzed (figure 4) Differential expression analysis was performed with EdgeR software. Besides, sequencing quality was validated with FASTQC software. Below is a series of graphs where the most relevant results of the study are collected.

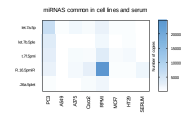


Figure 4: This heatmap shows the common miR-NAs in exosomes from our cell lines and serum. Serum samples are non cancer donors, however other nathologies cannot be ruled out miRNAs selected were those who presented more than 50

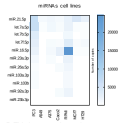
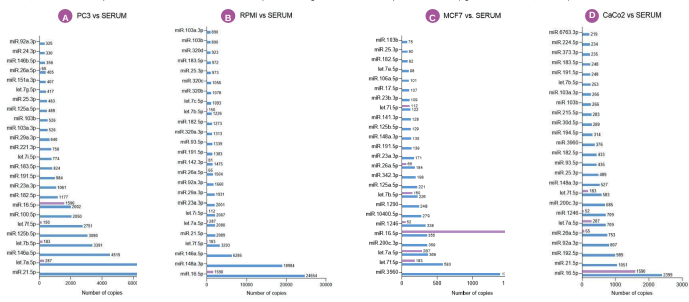


Figure 5: This heatmap shows the common miRNAs in exosomes from our cell lines. miRNAs selected were those who presented more than 50 copies.

In addition, It was made a comparison of the miRNAs with more than 50 copies of readings of each line with respect to the serum (figure 6: A, B, C, D, E, F, and G).



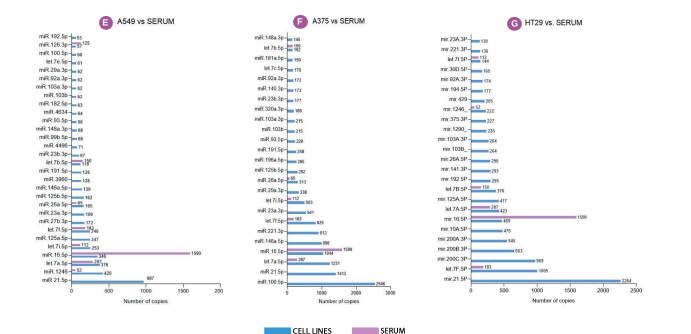


Figure 6 (A, B, C, D, E, F, and G). Comparison of miRNAs of each cell line with respect to serum.



8. ADDITIONAL INFORMATION

For research use only. Not for diagnostic use.

Not for resale. Immunostep will not be responsible of violations that may occur with the use of this product. Any use of this product other than the specified in this document is strictly prohibited.

Unless otherwise indicated by Immunostep by written authorization, this product is intended for research only and is not to be used for any other purpose, including without limitation, for human or animal diagnostic, the

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9. EXPLANATION OF SYMBOLS

| \. | Fluorochrome |
|----------------|------------------------------|
| REF | Product reference |
| \sum | Content for <n> analysis</n> |
| ₽ | Regulatory Status |
| E ₀ | Description |
| RUO | Research Use Only |
| *** | Manufacturer |

10. REFERENCES

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11. MANUFACTURED BY



IMMUNOSTEP SL.

Address: Avda. Universidad de Coimbra, s/n Cancer Research Center (C.J.C) Campus de Unamuno 3700 7 Salamanca (Spain)

7/00 / Salamanca (S)
Telf.fax: (+34) 923 294 827
E-mail: info@immunostep.com
www.immunostep.com