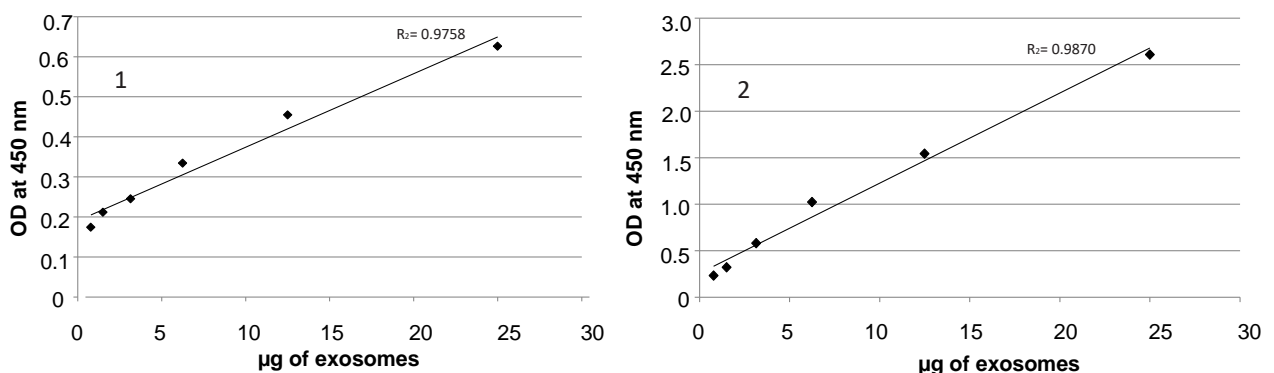






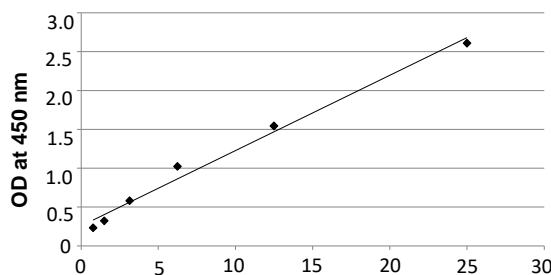


9. Wash the plate as indicated above. Add 100  $\mu$ l of rabbit HRP-conjugated secondary antibody solution to each well (diluted to 1:2000 dilutions in 1X Sample Buffer respectively). Seal the plate with parafilm and incubate at room temperature while shaking for 15 min (2-3 rotations per sec). Then incubate for 1 hr at 4°C (for human plasma and serum samples) or incubate for 1 hr at 37°C in humid chamber (for human urine and cell culture medium samples). Wash the plate as indicated above.
10. **For Colorimetric detection:** Add 100  $\mu$ l of Substrate Chromogenic Solution to each well and incubate at room temperature in the dark for 5-10 min. Be careful not to immerse metallic components of a pipette into substrate solution. Also avoid making bubbles and, if formed, remove them gently with a pipette tip. Do not seal the plate and monitor till a blue color is visible. Intensity of color is proportional to the exosome concentration only within a certain dynamic range. Many plate readers do not deliver accurate results when the OD is above 3. Stop the reaction by adding 100  $\mu$ l of Stopping Solution to each well. The color will change from blue to yellow. Read the absorbance at 450 nm within 10 minutes. If possible, the absorbance should also be read at 570 nm and the measurement should be subtracted from the measurement at absorbance 450 nm.
11. **Calculation:** Exosome standards are provided as assay calibrators and also as the positive control. It is important to note that the origin of purified standard exosomes may change the proportion of common exosomal proteins such as CD9. The amount of proteins on their membrane might differ slightly from the amount on the sample exosomes. The standard curve is used to determine the amount of exosomes in an unknown sample. Figure 3, is an example of standard curves obtained. The curve is obtained by plotting the average readings for different standard concentrations against the corresponding amounts of exosomes. Calculate the mean absorbance for each set of duplicate standards, controls and samples. The values of the negative controls (blanks) must be subtracted from all OD values before the results can be interpreted. The regression curve coefficient should be above 0.95. The estimated sample concentration is reliable if within the linear range of the curve, otherwise the samples must be diluted and the test repeated. For diluted samples, multiply the concentrations with the appropriate dilution factors.

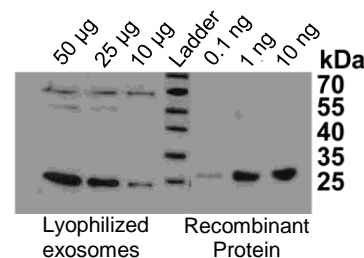


**Figure 3. CD9 Titration of COLO1 (1) and plasma (2) exosomes.** Example of standard curves obtained. Exosome standard preparation derived from (1) COLO1 Cell Culture Supernatant and (2) Plasma of healthy donors. This standard curve is for demonstration only. For quantification purposes, a standard curve must be obtained with every assay.

12. **Sensitivity:** The sensitivity of the ExoQuant™ was compared to Western blot. The data reported in Figure 4 and 5 demonstrate that the sensitivity of the ExoQuant™ is higher than that of Western blotting. Figure 5 shows that 10  $\mu$ g of lyophilized exosomes is equivalent to 0.1 ng of recombinant exosomal protein (CD9). Since the standard curve's lower concentration is 0.39  $\mu$ g of lyophilized exosomes (Figure 4), the sensitivity of our test is around 39  $\mu$ g of protein equivalent.



**Figure 4. CD9 titration of plasma healthy donor exosome standards.**



**Figure 5. CD9 marker detection by Western Blotting on lyophilized exosomes from human plasma and the recombinant CD9 protein.**

13. **Reproducibility:** Intra Assay (within run) CV (%) < 10 Inter Assay (interplate run) CV (%) < 13. Coefficient of variation (CV) is expressed as a percentage of variance to the mean calculated for intra assay reproducibility evaluation by assessing at least 4 replicates of three different standard preparations containing different exosome concentrations. Subsequently, assay to-assay reproducibility was assessed by comparing mean absorbance for the same samples on independent plates tested by different operators.







	<ul style="list-style-type: none"> <li>. Stop solution not added</li> </ul>	<p>optimization of results</p> <ul style="list-style-type: none"> <li>. Prepare the substrate solutions immediately before use. Ensure that the stock solutions are in date and have been stored correctly, and are being used at the correct concentration. Ensure the reagents are used as directed at the correct concentration</li> <li>. Ensure reagents have been prepared correctly and are within the expiry date</li> <li>. Ensure that you are incubating the antibody for the recommended amount of time, if an incubation time is suggested. The incubation time may need to be increased for optimization of results. Longer incubation time may be required</li> <li>. Antibodies have optimum binding activity at the correct temperature. Ensure the incubations are carried out at the correct temperature and that incubators are set at the correct temperature and is working. Incubation temperature may require some optimization. Ensure that all reagents are at room temperature before proceeding</li> <li>. Addition of stop solution increases the intensity of the color reaction and stabilizes the final color reaction</li> </ul>
<p><b>Inconsistent absorbance across the plate</b></p>	<ul style="list-style-type: none"> <li>. Plates stacked during incubations</li> <li>. Pipetting inconsistent</li> <li>. Antibody dilutions/Reagents not well mixed</li> <li>. Wells allowed to dry out</li> <li>. Inadequate washing</li> <li>. Bottom of the plate is dirty affecting absorbance readings</li> </ul>	<ul style="list-style-type: none"> <li>. Stacking of plates does not allow distribution of temperature across the wells of the plates. Avoid stacking</li> <li>. Ensure pipettes are working correctly and are calibrated. Ensure pipette tips are pushed on far enough to create a good seal. Take particular care when diluting down the plate and watch to make sure the pipette tips are all picking up and releasing the correct amount of the liquid. This will greatly affect the consistency of results between duplicate wells</li> <li>. To ensure a correct dilution of samples/standards across wells, ensure that all the reagents and samples are mixed before pipetting onto the plate</li> <li>. Ensure that plates are well sealed with film when incubating. Place a humidifying water tray (bottled clean/sterile water) on the bottom of the incubator</li> <li>. This will lead to some wells not being washed well as others, leaving inside different amounts of unbound antibody, which will give inconsistent results</li> <li>. Clean the bottom of the plate carefully before re-reading the plate</li> </ul>
<p><b>Color developing slowly</b></p>	<ul style="list-style-type: none"> <li>. Plates are not at the correct temperature</li> <li>. Secondary Antibody too weak staining</li> <li>. Contamination of solutions</li> </ul>	<ul style="list-style-type: none"> <li>. Ensure plates are at room temperature and that the reagents are at room temperature before use</li> <li>. Prepare the substrate solutions immediately before use. Ensure that the stock solutions are in date and have been stored correctly, and are being used at the correct concentration. Ensure the reagents are used as directed</li> </ul>



		<ul style="list-style-type: none"><li>. Presence of contaminants, such as sodium azide and peroxides can affect the substrate reaction. Avoid using reagents containing these preservatives</li></ul>
<b>Poor Standard Curve</b>	<ul style="list-style-type: none"><li>. Inaccurate pipetting</li><li>. Improper standards and samples preparation</li><li>. Improper washings</li></ul>	<ul style="list-style-type: none"><li>. Check pipettes and increase attention</li><li>. Ensure to reconstitute standards in a proper buffer and mixing thoroughly by vortexing and gentle pipetting.</li><li>. Wash thoroughly as recommended in the assay procedure. Wash for longer time after incubation with the HRP-conjugated secondary antibody</li></ul>
<b>Low signals</b>	<ul style="list-style-type: none"><li>. Low signals in standards due to improper storage or preparation</li><li>. Low signals in samples due to low exosome concentration</li></ul>	<ul style="list-style-type: none"><li>. Ensure to store reconstituted standards at -20°C and strictly avoid thaw and freeze cycles. Assure that standards are properly reconstituted and standard dilutions are prepared as suggested in the data sheet</li><li>. Check out the sample preparation protocols</li></ul>
<b>Low reproducibility of duplicates</b>	<ul style="list-style-type: none"><li>. Inaccurate pipetting</li><li>. Improper standards and samples preparation</li></ul>	<ul style="list-style-type: none"><li>. Check the pipettes and be careful with pipetting</li><li>. Ensure to mix thoroughly by vortexing and gentle pipetting before loading onto wells</li></ul>

*FOR RESEARCH USE ONLY! Not to be used on humans.*