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A Single Extracellular Vesicle (EV) Flow Cytometry Approach to Reveal EV Heterogeneity

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Abstract: Extracellular vesicles (EVs) actively participate in intercellular communication and pathological processes. Studying the molecular signatures of EVs is key to reveal their biological functions and clinical values, which, however, is greatly hindered by their sub-100 nm dimensions, the low quantities of biomolecules each EV carries, and the large population heterogeneity. Now, single-EV flow cytometry analysis is introduced to realize single EV counting and phenotyping in a conventional flow cytometer for the first time, enabled by target-initiated engineering (TIE) of DNA nanostructures on each EV. By illuminating multiple markers on single EVs, statistically significant differences are revealed among the molecular signatures of EVs originating from several breast cancer cell lines, and the cancer cell-derived EVs among the heterogeneous EV populations are successfully recognized. Thus, our approach holds great potential for various biological and biomedical applications.

Extracellular vesicles (EVs) are membrane-encapsulated vesicles (30–100 nm) secreted by all cell types and present in various biological fluids. They constitute an emerging target for liquid biopsy in cancer diagnosis because they actively participate in cell–cell communication by shuttling signaling molecules between cells and modulate pathological processes such as tumor initiation, progression, and metastasis. Study of the molecular signature of EVs is thus critical for profoundly understanding their biological functions and clinical values. Existing methods for EV study are primarily focused on bulk analysis of a large number of EVs, because the small physical dimensions of EVs limit the total amounts of biomolecules to be carried in each EV, greatly enhancing the difficulty in molecular profiling. However, subtle molecular differences at the single EV level may yield significant variation in EV biological functions, and the highly heterogeneous nature of EV population demands the development of techniques capable of profiling individual EVs. Most recently, single EV counting with a nanochip and imaging single vesicles with advanced fluorescence microscopy have been reported, and showed that EVs from different cell of origin can carry distinct surface markers mimicking their parent cells. Still, they require EV immobilization steps, limiting the down-stream investigations on EV functions and biogenesis.

Flow cytometry has been widely employed to distinguish different cell types in mixed populations based on the expression of cellular markers. Similarly, it could be used to study the heterogeneous EVs. However, the sizes of EVs fall well below the detection limit of conventional flow cytometers, making it impossible to do single-EV analysis without significant instrumentation development. Herein, we report the first single-EV flow cytometry analysis (FCA) in conventional flow cytometers enabled by target-initiated engineering (TIE) of DNA nanostructures on individual EVs. This technique employs a conformation-switchable DNA probe to bind to the EV surface marker, which triggers the engineering of a DNA nanostructure by hybridization chain reaction (HCR). The HCR products not only enlarge the overall size of the single EV to be beyond 500 nm, but also can bind to multiple fluorophores to amplify the signal from the few marker molecules locating on the limited area of EV surface, both enabling visualization of single EVs in a conventional flow cytometer, and greatly simplifying measurement of multiple markers on the same EV.

To test the working principle, we designed the conformation-switchable probe to recognize CD63, a classic tetraspanin marker that is highly abundant in various EVs (Figure 1A). This probe contained a target recognition domain with the sequence of an anti-CD63 aptamer and a trigger domain for initiation of DNA growth via HCR. These two domains were flank by a hinge sequence to achieve conformation change triggered by aptamer–target interaction (Figure 1A; Supporting Information, Table S1): a part of the target recognition domain hybridized with the trigger domain (that is, “deactivated state”) to form a hairpin structure, which could be opened upon target binding to expose the sequence (“activated state”) for hybridization with Hairpin 1 (H1). Then sequential hybridization between H1 and H2 would occur to build the long dsDNA product on EV (Figure 1A).
The design of target-initiated engineering of DNA nanostructure could effectively eliminate non-specific DNA growth without the target. Furthermore, the DNA nanostructures constructed upon recognition of the free, non-EV bound target would not locate on EV surface and should not reach the size range detectible by the conventional flow cytometer. Both features significantly reduce the background in FCA and render our technique ultra-high simplicity: no washing is necessary to remove the unreacted probes.

Successful DNA hybridization cascade initiated by CD63 or the CD63-containing EVs was verified by native agarose gel electrophoresis. We can see from the gel image (Figure 1B) that, long DNA products were formed in the presence of CD63 protein (Lane 3), accompanied with significant consumption of H1 and H2, similar to the reaction between the hairpins and a simple initiator (Lane 2), a sequence of the trigger domain that can directly hybridize with H1 without target recognition and initiate DNA growth. With the CD63-containing EVs (referred to as “standard EVs” hereafter), most of the long DNA products were trapped inside the loading well probably by binding to the EVs (Lane 4). We employed transmission electron microscopy (TEM) to confirm the size enlargement in EVs produced by TIE. As clearly illustrated in Figure 1C, after TIE, each standard EV exhibited the “hairy” morphology on its surface, with the overall size increasing from the original tens of nanometers to hundreds of nanometers and maintaining the spherical shape (Supporting Information, Figure S1A,B). In contrast, the DNA hybridization products triggered by simple initiator were in random shapes without the densely stained core (Supporting Information, Figure S1C). The growth effect was also viewed by atomic force microscopy (AFM) (Supporting Information, Figure S2) and nanoparticle tracking analysis (NTA) Supporting Information, (Figure S1D): after TIE, DNA strands up to 250 nm long were seen surrounding the individual EVs under AFM; and the size distribution profile in NTA shifted to the larger size region with a new peak at 500 nm appeared.

We examined the enlarged EVs by a confocal fluorescence microscope. Labeled the hairpins with biotin, the long DNA products can be tagged with the streptavidin-conjugated fluorescent probes, like the Qdot 525 streptavidin conjugate (QDs, Ex 488 nm/Em 525 nm) and the streptavidin/rhodamine-conjugated iron oxide nanoparticles (RhNPs, Ex 547 nm/Em 572 nm). These two tags were used to produce two distinct EV populations in separate reaction tubes, which were mixed and inspected under microscope. Indeed, the engineered EVs labeled with either the QDs or the RhNPs were clearly discernible in the confocal mode with an objective of 40× (Figure 1D). We even found particles that were overlapped (highlighted by a triangle), located close to each other (highlighted by a square), or completely separated (indicated by circles), showing that each fluorescent particle represented one individual vesicle. The negative controls, that is, the QD-labeled DNA nanostructure established by the simple initiator, and the RhNPs or QDs by themselves, were not observable even under ahiger magnification of 60× or 100× (Supporting Information, Figure S3), which was due to the few numbers of QDs on each DNA nanostructure. On contrary, each EV could carry multiple surface markers, and be labeled by multiple dsDNAs (Supporting Information, Figure S2) that not only enlarge the overall size, but also

![Figure 1. A) The single extracellular vesicle flow cytometry analysis technique enabled by target-initiated engineering of DNA nanostructures. B) Analysis of the long DNA products by gel electrophoresis: Lane 1: reaction probes only; Lane 2, 3, and 4: reaction triggered by the simple initiator, CD63 protein, and CD63 + EVs, respectively. C) TEM image of the Engineered EV (Engr. EV). D) Fluorescence microscopy image of the Engr. EV tagged with rhodamine-nanoparticles (Engr. EV-RhNPs; shown in red) or QDs-525 (Engr. EV-QDs shown in green). □ Engr. EV-RhNPs or Engr. EV-QDs, well separated from each other; □ one Engr. EV-QDs located nearby another Engr. EV-RhNPs; △ Engr. EV labeled with both QDs and Rh-NPs.](image-url)
amplify the total fluorescence signal, making the single EV visible under the same imaging setting. This result confirms that the EVs enlarged by TIE were visible by conventional optical imaging tools.

Compared to microscopic imaging, flow cytometry provides fast and automatic particle counting at the rate of thousands particles per second; and can sort pure particle populations defined by fluorescence patterns, much more ideal for interrogating single biological particles. Thus, we explored whether the engineered EVs could be detected in a conventional flow cytometer FACSCanto that is widely used in research and clinical settings. Analysis of the standard beads confirmed that this instrument could not see particles smaller than 500 nm. In this experiment, the DNA nanostucture carried by the engineered EVs were tagged by either Alexa Fluor 488 streptavidin conjugates (that is, Engr. EV-Alexa488) or the green QDs-525 (Engr. EV-QDs). Interestingly, both EV samples revealed a significant particle cluster in the flow cytometry plots of FSC (forward scatter) vs. SSC (side scatter) and FL1 (fluorescence channel, $\lambda_{ex} = 488$ nm) vs. SSC (Figure 2A). These particles exhibited larger FSC and higher fluorescence (FL) than the background particles, confirming the significant EV size enlargement induced by the CD63-initiated EV engineering; and the signal amplification from the multiple fluorescent tags bound to the long DNA chains on each EV. Gating the particle cluster by R1 (for the QD label) or R2 (for the Alexa488 label) illustrated that the enlarged EVs represented about 50\% of the total particles detected for each sample (Supporting Information, Table S2). In contrast, no distinct particle cluster was detected on the flow plots obtained from the negative control samples, including the EVs labeled by the anti-CD63-conjugated QDs without any size enlargement (that is, EV-QDs) (Figure 2A), and the EVs mixed with just the hairpins and QDs (Supporting Information, Figure S4). The background particles present in Figure 2A and the Supporting Information, Figure S4 with low forward scatter and fluorescence signals might be produced from non-specific adsorption of the hairpins and QDs on the EVs, as well as random aggregation of QDs in solutions. Furthermore, TIE on free CD63 did not produce observable particles, ensuring the counting was from intact EVs instead of free markers released by cells or from vesicle breakage.

Using R1 and R2, we compared the scatter and FL signals obtained with Alexa488 and QDs (Figure 2B). Both labels gave out comparable FSC readings (the mean FSC intensity ratio of QD/Alexa = 0.92 ± 0.07 ($n = 3$)), indicating the size enlargement effect was mainly resulted from the growth of the dsDNA, not the additional size of the fluorescent tag. Labeling with QDs resulted in $70.5 \pm 6.2$ ($n = 3$) times higher FL than with Alexa488. The higher FL shifted the enlarged EVs further away from the background particles, making the population more distinct, which is a critical characteristic for future sorting of specific EV populations. This is the first time a new population of detectible EVs appearing on the flow cytometric plots to enable clear and unambiguous recognition of pure EV population in a conventional flow cytometer, owing to TIE-enabled transformation of the invisible EVs to the visible particles.

The kinetics of TIE was studied by changing the HCR time from 0 h to 24 h. A gate was applied on the flow plots to identify the particle cluster displaying higher FL and larger FSC than background, which were considered as the detectible events. We found that, a reaction time of 2 h produced only very few numbers of the detectible events, which dramatically increased at 4 and 8 h (Supporting Information, Figure S5). NTA measurement also confirmed that with the reaction time increasing to 4 and 8 h, more engineered EVs larger than 250 nm were produced (Supporting Information, Figure S1D). The mean fluorescence intensity (MFI) gradually increased with the reaction time going from 0 to 8 h (Supporting Information, Figures S5,S6). However, extending the reaction to 12 h did not induce significant changes in the number of detectible events and the MFI value detected in the flow cytometer, neither big change was observed in the size distribution profile obtained with NTA. This result indicates that recognition of the CD63-positive EVs may have reached a steady state and no new engineered EVs could be produced with longer reaction time. Still, the length of the DNA product continued to increase with longer reaction duration, which generated the engineered EVs larger than 500 nm at 24 h observed by NTA (Supporting Information, Figure S1D). However, space hindrance and winding of the long DNA strand may have prevented more QDs from binding to the DNA; and after being kept at 37°C for 24 h, the EVs may no longer be stable, both leading to the decrease in MFI and detectible events measured by flow cytometry (Supporting Information, Figure S6). While detailed study of the kinetics of engineering DNA nanostructures on EVs needs to be conducted in future works, the present work...
employed overnight reaction to obtain stable signals in flow cytometric analysis of single EVs.

Under the optimized TIE conditions, we confirmed that the number of detectible events was linearly ($R^2 = 0.9974$) proportional to the number of EVs in the sample within the range of 20 ng mL$^{-1}$ to 500 ng mL$^{-1}$ (Supporting Information, Figure S7), proving the capability of our method in EV quantification. Furthermore, the robustness of our method was verified by the intraday and interday replicates (Supporting Information, Table S3). The Student’s $t$ test obtained a $p$ value smaller than $10^{-4}$ when evaluating the MFI or mean forward scatter (MFS) among all these replicates, showing that EV engineering is highly reproducible. More importantly, we showed that our method can directly engineer the EVs in biofluids such as cell culture medium (Supporting Information, Figure S8) and serum (Supporting Information, Figure S9) by simply treating the matrix with sodium citrate to inhibit exonuclease activity.$^{[13]}$ Direct EV detection in biological samples with minimum sample pretreatment and without EV enrichment or immobilization is a big advancement compared to existing technologies and makes our method highly suitable for studying the heterogeneity of EV populations and examining their clinical values.

Differentiation of heterogeneous EVs requires recognition of multiple markers. The TIE system can be easily adjusted to target different surface biomarkers by simply switching the aptamer sequence in the conformation-switchable DNA probe. To demonstrate this, we designed another TIE system to target human epidermal growth factor receptor 2 (HER2), a typical breast cancer marker overexpressed in about 20% of breast cancer patients and present on a subset of EVs derived from breast cancer cell lines.$^{[14]}$ We substituted the anti-CD63 aptamer with an anti-HER2 aptamer$^{[15]}$ in the target-recognition domain, and kept the trigger domain intact. The hinge domain was adjusted slightly to achieve effective conformational switching upon HER2 recognition (Supporting Information, Table S1). Since the HCR triggered by HER2 remains the same as that by CD63, we define it as the “single hybridization cascade system”, and applied it to analyze HER2 and CD63 expression separately on single EVs secreted by three different cell lines: the breast cancer cells of SKBR3 (highly metastatic) and MCF-7 (poorly metastatic) and the non-tumor epithelial cell MCF-10A. We found that the anti-HER2 system produced more detectable events from the SKBR3-derived EVs than those from MCF-7 and MCF-10A (Supporting Information, Figure S11a). Furthermore, two times more events were detected by the anti-HER2 system on the SKBR3 EVs than by the anti-CD63 system. On the other hand, the EVs originating from MCF-10A and MCF-7 resulted in comparable numbers of detectable events (Figure 3C), agreeing with the mixing ratio of the two EV sub-populations (Figure 3A; Supporting Information, Figure S12). Viewing CD63 as the internal standard, we calculated the MFI ratio of FL4/FL1 to represent the relative content of HER2 and CD63 on the same EV. Significant difference in the MFI ratios between EVs from the MCF-10A and SKBR3 cells ($p = 0.009$), or between those from the MCF-7 and SKBR3 cells ($p = 0.02$) was confirmed with Student’s $t$ test (Figure 3C), confirming the effectiveness of dual labeling on the same EV for robust differentiation of the SKBR3-derived EVs from those produced by the MCF cells. The EVs from MCF-7 and MCF-10A were not differentiable using these two markers, because the expression levels of both markers were comparable in these two EVs (Supporting Information, Figure S10).

The significant difference between the MFI values from different EV sub-populations indicates that our technique has the power to reveal the presence of EV sub-populations among heterogeneous mixtures. To confirm this, we tested the mixture of EVs from MCF-10A and SKBR3 prepared at a 1:1 ratio. Using the distribution patterns identified with the pure EV populations, we could clearly see the presence of two EV populations in the mixture (Figure 3D). The particle count ratio for those locating within R1 and R3 was equal to 0.98 ± 0.07 ($n = 3$), agreeing with the mixing ratio of the two EV sub-populations. In contrast, bulk analysis of the same EV mixture using ELISA only reflected a slight increase in the HER2 content, but failed to recognize the presence of two distinct EV sub-populations (Supporting Information, Figure S10). More strikingly, varying the mixing ratios of these two EV sub-populations, the ratio of the particle counts in the gated regions of R1 and R3 showed a strongly linear relationship ($R^2 = 0.9581$) with the ratio of the added EV numbers between two cell lines (Supporting Information, Figure S13). These results demonstrate the great potential of our method in recognizing EV heterogeneity and differentiating EV sub-populations.
In summary, we have developed an single-EV FCA technique to visualize individual EV in a conventional flow cytometer for the first time, which can help gain in-depth insights into the molecular signatures of EV sub-populations under regular lab settings. While our work demonstrates simultaneous recognition of dual surface markers on the same vesicle, more markers can be targeted by simply revising the conformation-switchable probes to improve more effective differentiation of more EV sub-populations. Furthermore, our technique opens the opportunity for EV sorting and collection based on surface marker profiles under typical clinical lab settings. The pure EV sub-populations obtained will definitely benefit clear correlation between EV composition and their biogenesis and functions. Overall, we believe this single-EV FCA technique is a valuable tool for gaining more understanding on the roles of EVs in cell-cell communication and pathological development, and its high simplicity and good adaptivity to clinical labs will be highly beneficial for screening for EV markers for liquid biopsy applications.

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**Conflict of interest**

The authors declare no conflict of interest.

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Figure 3. A) Representative flow cytometry plots of the particle cluster determined by the relative expression levels of HER2 and CD63 for the EVs from different cell lines. B) The dual hybridization cascade system for recognition of two markers on the same EV. C) Box chart of the mean fluorescence intensity (MFI) ratio between FL4 and FL1 of the EVs from three cell lines. *p < 0.05, and **p < 0.01, n = 5. IQR = interquartile range. D) Flow cytometry scatter plots and fluorescence histograms for analysis of EV mixtures. R1 and R3 are defined in Figure 3A using the EVs from the corresponding cell lines. The particle clusters were colored on the scatter plot of FSC vs. SSC and the histograms based on the gates defined in the HER2 vs. CD63 plots.


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