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1. Introduction

EVs are cell-derived membranous vesicles with phospholipid bilayers, ranging from 30 to 1000 nm in diameter.^{1,2} These cell-derived vesicles exist in a variety of body fluids, *e.g.*, serum, urine, saliva, milk, bile, and other fluids.³⁻⁶ EVs share the same topology as cells and contain various biologically important molecules, including proteins, RNA, and DNA.⁷ EVs have play major roles in intercommunication during biological processes, particularly in cancer initiation and progression.⁸⁻¹⁰ It is worth noting that the characterization of EVs has evolved from *in vitro* investigation of cell culture supernatants to more recently *in vivo* studies on directly isolated EVs from animal or human body fluids, *e.g.*, plasma, urine, and saliva.¹¹⁻¹³

A new integrated method for tissue extracellular vesicle enrichment and proteome profiling[†]

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Extracellular vesicles (EVs) are membranous vesicles released by cells that carry a number of biologically important components such as lipids, proteins, and mRNAs. EVs can mediate cancer cell migration, invasion, angiogenesis, and cell survival, greatly contributing to cell-to-cell communication in the tumor microenvironment. Additionally, EVs have been found to have diagnostic and prognostic significance in various cancers. However, the direct isolation of pure EVs remains challenging, especially from tissue samples. Currently available EV isolation approaches, e.g., ultracentrifugation, are time-consuming, instrumental dependent, and have a low recovery rate with limited purity. It is urgent to develop rapid and efficient methods for enriching tissue EVs for biological and clinical studies. Here, we developed a novel isolation approach for tissue EVs using an extraction kit combined with TiO₂ microspheres (kit-TiO₂). The EVs were first precipitated from the tissue fluid using a precipitation agent and then further enriched using microspheres based on the specific interaction between TiO2 and the phosphate groups on the lipid bilayer of the EVs. Kit-TiO₂ approach led to improved purity and enrichment efficiency of the isolated EVs, as demonstrated by western blot and proteomic analysis, compared with previously reported methods. A total of 1966 protein groups were identified from the tissue EVs. We compared the proteomic profiles of the liver tissue EVs from healthy and hepatocellular carcinoma (HCC) bearing-mice. Twenty-five significantly upregulated and 75 downregulated protein groups were found in the HCC EVs. Among the differentially expressed proteins, Atic, Copa, Cont3, Me1, Anxa3, Fth1, Anxa5, Phb1, Acaa2, ATPD, and Glud1 were reported to be highly relevant to HCC. This novel isolation strategy has provided a powerful tool for enriching EVs directly from tissues, and may be applied in biomarker discovery and drug screening of HCC.

While EVs are found in large quantities in the circulatory system, EVs also exist in the interstitial tissues and play a key role in cell-to-cell communication. In tumors, interstitial EVs are essential in modulating the tumor microenvironment for cell seeding and metastatic growth.¹⁴ Compared to EVs obtained from cell cultures, EVs directly isolated from tissues show great advantages in tissue specificity and close relevance to the microenvironment.¹² Therefore, optimized techniques for EV extraction from solid tissue specimens are needed to enable *in vivo* study of tumor EVs and in-depth exploration of their underlying biological mechanisms.

Hepatocellular carcinoma (HCC) is the third most common cancer in the world, accounting for 85–90% of primary hepatic tumors.^{15–17} HCC cells can interfere with the biological behaviors of different types of cells by releasing EVs. For example, secreted EVs regulate the epithelial-mesenchymal transition (EMT) in the microenvironment to transform the surroundings into an inflammatory state, coordinate with neighboring tumor cells to increase invasiveness, and induce the conversion of adjacent fibroblasts and macrophages to CAFs and TAMs.¹⁸ Moreover, HCC-released EVs might interfere with immune cells and endothelial cells to induce immune escape and angiogenesis.^{19,20} HCC-secreted EVs also

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mediate signaling pathways and regulatory factors involved in intercellular interactions.²¹

Several studies have revealed that differentially expressed EV miRNAs might be used as biomarkers for diagnosing HCC. Additionally, the delivery of EV miRNAs and lncRNAs may significantly inhibit cancer development.^{22,23} Therefore, EV components are potential diagnostic markers and therapeutic targets for HCC.

In terms of separating EVs from liver tissues or any other tissue, it is critical to properly disrupt the tissue while minimizing cellular disruption. One or more EV separation steps might be imposed to increase the EV purity with minimal cellular contamination.24 Previously published methods rely on tissue homogenization and filtration, which is likely to disrupt the cellular structure and mix intracellular vesicles with EVs.^{25,26} Current methods for EV isolation include ultracentrifugation, precipitation kits, immunoaffinity approaches, size exclusion chromatography (SEC), and ultrafiltration.27,28 However, ultracentrifugation is time-consuming and repeated centrifugation might lead to EV loss, resulting in limited throughput and recovery rates.29 The kit precipitation method is mainly based on polymer precipitation, but the purity of the extracted EV is relatively low. The introduced polymers are also difficult to remove and may lead to severe interference in downstream analysis.30,31 SEC shows a good recovery rate and reproducibility. However, the size of the purification column limits the quantity of the sample and the throughput of this method. Additionally, the elution step dilutes the sample, resulting in a low concentration of the final product. The immunoaffinity approach isolates EVs through the specific interactions between antibody and EV membrane proteins. The recovery of EVs is unpredictable, depending on the performance of the antibody, and the cost is relatively high.

In this report, we established a novel method to enrich EVs from liver tissue based on a two-step method via a combination of an extraction kit and TiO₂ microspheres. In this method, the aqueous contents and complex components in the tissue are first removed by a kit method, and then the tissue EVs are extracted using TiO₂ microspheres via the specific interaction between TiO₂ and the phosphate groups on the EV lipid bilayers. The isolated EVs were characterized by their size, morphology, and proteomic profiles. The proposed enrichment approach has been demonstrated to have high efficiency and good reproducibility for enriching tissue EVs. Obviously improved purity and quantity of the enriched EVs was achieved as demonstrated by western blot and proteomic analysis compared with previously reported methods. Furthermore, we successfully applied this approach to analyze HCC samples. In total, 25 significantly upregulated proteins were found, 11 of which were associated with liver cancer, demonstrating the potential of the proposed new method for tissue EVs for cancer studies.

2. Materials and methods

2.1 Animal models

All animal procedures were performed in accordance with the Guidelines for Care and Use of Laboratory Animals of Beijing

Proteome Research Center, and approved by the Animal Ethics Committee of Beijing Proteome Research Center (license: IACUC-20211228-42). C57BL/6 mice were purchased from Beijing Weitong Lihua Inc. The HCC model mice (Available from Shanghai Model Organisms Center, Inc) were established as previously described.32 Briefly, 2 week-old C57 male mice (reared at SPF level 5) were injected intraperitoneally with DEN (Sigma N0258, 40 mg kg⁻¹, configured with saline to 4 mg mL⁻¹, 10 μ L g⁻¹) once, and 6 weeks later (8 week-old mice) they were injected with CCL₄ (National Drug 10006464, configured with corn oil (ABCone, C67366) to 10%, 10 μ L g⁻¹) twice a week for 20 weeks. At the 16th week of administration, the liver was dissected and observed to check for primary liver cancer. During the experiment, the mice were observed daily for coat sheen, activity, feces, water, and food administration, and weighed once a week to monitor their health status.

2.2 Liver tissue samples from the mice

Phosphate-buffered saline (PBS) (VIVICUM, VCM3008) was preheated in a 37 °C water bath, and all surgical instruments were sprayed with 70% ethanol and wiped dry. The pump tubing was flushed by running 70% ethanol through and rinsed twice with running water to remove any residue. A sterile 5 cm Petri dish with RPMI-1640 medium (VIVICUM, VCM5140) was prepared to hold the digested liver after perfusion. The mice were injected intraperitoneally with 400 µL of anesthetic (3% sodium pentobarbital) and were checked to ensure complete anesthesia through a skin pinch clamp response or toe stimulation response. All four limbs of the mouse were held down with pins. The abdominal skin was cleaned by spraying with 70% ethanol and wiping with gauze and an alcohol pad, avoiding mouse fur contamination. The skin was cut from the anterior to the pelvic bone using sterile scissors. The portal vein (PV) and inferior vena cava (IVC) were exposed by gently applying a cotton-tipped applicator, and the intestine was moved to the left side of the animal. Then, a lagging needle was inserted carefully from the portal vein. Once the blood returned, the inferior portal cava was perfused with preheated irrigation solution at a flow rate of $3.5-4 \text{ mL min}^{-1}$ for 4 min. During the perfusion, the inferior vena cava was severed and clamped with forceps for 7-10 s until the liver was enlarged. To ensure that the liver was swollen and relaxed, the procedure was repeated at 30 s intervals. After 5-7 perfusions, the liver blood was completely removed through the inferior vena cava as the liver turned white. Then, the liver was cut out with the gallbladder carefully removed and placed in a prechilled Petri dish containing PBS.

2.3 Preparation of mouse liver tissue fluid

The liver tissue was gently sliced into small sections (2–4 mm³) and incubated for 20 min at 37 °C in the RPMI-1640 medium supplemented with collagenase D (Sigma, c5138-1G, 2 mg mL⁻¹) and DNase I (Solarbio, no. D8071, 40 U mL⁻¹). The incubated tissue blocks were gently pressed through a cell filter on top of a conical tube to remove any undigested tissue. The dissociated tissues were then centrifuged at $300 \times g$ for 10 min at 4 °C. The

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generated pellet ("liver homogenate with collagenase, LHC") was stored at -80 °C prior to protein extraction. The supernatant was transferred to a fresh tube and centrifuged at $2000 \times g$ for 20 min at 4 °C to remove cell debris and aggregates. The separated supernatant was further centrifuged at $10\,000 \times g$ for 30 min at 4 °C to remove any large or apoptotic vesicles. The obtained tissue fluid was stored at -80 °C prior to EV isolation.

2.4 Isolation of tissue EVs by kit combined with TiO₂ strategy

The tissue fluid sample (1 mL) was added to 250 μ L of EV precipitant (Umibio EV extraction and purification kit) and incubated at 4 °C for 2 h. The solution was centrifuged at 10 000 \times *g* for 1 h and the supernatant was removed. The EV precipitate was resuspended and mixed with phenol red-free DMEM (VIV-ICUM, VCM50520). The supernatant was then mixed with 20 mg TiO₂ microspheres (I.D. = 5 μ m, GL Sciences) and incubated for 10 min at 4 °C on a shaker (THZ-C, Thermo Fisher Scientific) for sufficient enrichment. The sample was then centrifuged at 1000 \times *g* for 5 min at 4 °C. The precipitate was washed with 200 μ L 150 mM sodium chloride and centrifuged at 1000 \times *g* for 5 min at 4 °C. After washing three times, the obtained precipitate was TiO₂ microspheres enriched with tissue EVs.

2.5 Isolation of tissue EVs by TiO₂

The above tissue fluid sample (1 mL) was directly mixed with 20 mg TiO₂ microspheres and incubated for 10 min at 4 °C on a constant temperature shaker for adequate enrichment. After incubation, the samples were centrifuged at 1000 × g for 5 min at 4 °C. Next, the separated precipitate was washed three times with 200 µL 150 mM sodium chloride and centrifuged at 1000 × g for 5 min at 4 °C. Finally, the precipitated TiO₂ microspheres with tissue EVs attached were obtained.

2.6 Isolation of tissue EVs by precipitation reagent

Using the Umibio EV extraction and purification kit, the tissue fluid sample (1 mL) was added to 250 μ L EV precipitant and incubated at 4 °C for 2 h. The sample was centrifuged at 10 000 \times *g* for 1 h and the supernatant was removed. The enriched EV pellet was suspended in PBS for downstream experiments.

2.7 Isolation of tissue EVs by ultracentrifugation

According to a previously described method,^{33,34} the tissue fluid sample was ultracentrifuged at 110 000 × g for 70 min at 4 °C. The precipitated pellet was washed twice with PBS by ultracentrifuging the mixture at 110 000 × g for 70 min at 4 °C. The obtained EV pellet was stored at -80 °C until further analysis.

2.8 Liver homogenate preparation

For proteomic analysis, liver tissue was ground in cold PBS for 60 s using a cryogenic tissue grinder (Thermo Fisher Scientific) to obtain collagenase-treated (LHC) liver homogenates. The ground tissue was added to 4% SDS lysis buffer and sonicated on ice for 20 min. The homogenate was then centrifuged at 14 000 \times *g* for

10 min at 4 °C. The resulting protein supernatant was stored at -80 °C until analysis.

2.9 Characterization of the isolated tissue EVs

2.9.1 Scanning electron microscopy (SEM). Prior to SEM analysis, the TiO_2 microspheres were first fixed in 1 mL 2.5% (v/v) glutaraldehyde (prepared in 150 mM sodium chloride) for 2 h and were washed three times with 150 mM sodium chloride after fixation. Then, the fixed samples were dehydrated with ethanol at different concentrations (30%, 50%, 75%, 90%, 95%, 100%, and 100%) for 15 min. The microspheres were vacuum-dried at 60 °C overnight. For SEM, the obtained samples were spread on double-sided conductive carbon tape, gently blown with nitrogen gas, and then sputtered with gold using a sputter device for 60 s. Finally, images of the TiO_2 microspheres were collected using a SU8010 scanning electron microscope (Hitachi, Japan).

2.9.2 Transmission electron microscopy (TEM). The EV samples diluted in 20 μ L 150 mM sodium chloride were adsorbed on 200-mesh copper grids coated with formvar-carbon and were left aside for 1 min to allow adequate absorption. The solution residual was removed with filter paper. Subsequently, the absorbed samples were negatively stained with saturated uranyl acetate and air-dried at room temperature (RT) for 1 min. Then the samples were observed under a Hitachi HT-7700 transmission electron microscope at 100 kV.

2.9.3 Zeta nanoparticle tracking analysis (NTA). Measurement of the EV size and concentration was conducted by tracking the Brownian motion of individual nanoparticles using ZetaView (Particle Metrix, Germany). The isolated EV samples were diluted in 150 mM sodium chloride until individual nanoparticles could be traced. Based on the automatic analysis and elimination of any outliers from the defined positions, the mean size and concentration of the samples were recorded and computed using the instrument software ZetaView (version 8.05.14.SP7).

2.9.4 Western blot. EVs adhered to TiO₂ microspheres were lysed in 4% SDS. The protein concentration was measured using a BCA protein assay kit (Thermo Fisher Scientific). Equivalent total protein amounts of LHC and isolated EVs were separated using 10% sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) under alkaline conditions and then transferred to PVDF membranes (Immobilon-P). A total of 20 μg protein was applied for the western blot. The same amount of 20 μg were used for both LHC and EV. 20 μg was a typical loading amount for western blot experiments. The membranes were blocked using TBST containing 5% BSA for 1 h at RT and then were incubated overnight at 4 °C with anti-CD63 rabbit polyclonal antibody (Abcam, EPR21151), anti-syntenin-1 mouse polyclonal antibody (Santa Cruz, sc-515538), anti-TSG101 rabbit polyclonal antibody (Abcam, EPR7131), anti-Calnexin (Proteintech, 66903-1-lg), anti-GM130 (Abclonal, A11408), and anti-VDAC (CST, #4866). The PVDF membranes were rinsed three times and further incubated with a second anti-rabbit antibody (Abcam, 6721) or anti-mouse antibody labeled with horseradish peroxidase (Proteintech, SA00001-1) for 1 h at RT. Then the immunoblots were analyzed using an imaging system (Image-Quant LAS 500).

2.10 Proteomic analysis

The isolated EVs attached to TiO₂ microspheres were dissolved in 40 µL of 4% SDS buffer containing 1% protease inhibitor. The resulting suspension was sonicated on ice for 20 min and then centrifuged at 10 000 g at 4 °C for 5 min. The separated supernatant was processed immediately for the filter-aided sample preparation (FASP) digestion. The supernatant was diluted with 8 M urea to a final concentration of 0.1% SDS, and the diluted solution was transferred into a 30 kDa ultrafiltration tube to remove the SDS. Then, the proteins were reduced using 10 mM dithiothreitol for 4 h at 37 °C and alkylated with 20 mM iodoacetamide in the dark at RT for 40 min. The protein concentrations were measured with a Nanodrop 2000 (Thermo Scientific). Trypsin was added at a 1:50 (w/w) ratio and incubated overnight at 37 °C. The final obtained peptide solution was thermally dried using a vacuum rotary drier. Subsequent stable isotope dimethylation labeling the of peptides was conducted. Liver tissue samples from healthy and HCC mice were used for EV protein identification and relative quantification. The peptides were first redissolved in 200 μL of 100 mM TEAB buffer solution. Then, 4 μL of 4% formaldehyde-D2 solution was added to the HCC group and 4 µL of 4% formaldehyde solution was added to the healthy group. Additionally, 4 µL of 0.6 M sodium cyanoborohydride was added to each sample, vortexed and mixed. The reaction was carried out at room temperature for 1 h. Next, 16 µL of 1% ammonia solution acidified with formic acid was added to terminate the reaction. Finally, the light and heavy isotopelabeled groups were mixed and desalted using StageTip C₁₈. The desalted peptides were concentrated and dried using a vacuum spin dryer, and then stored at -80 °C until analysis. The obtained peptides (500 ng) were used for LC-MS analysis.

LC-MS analysis was performed using a Q-Exactive HF-X mass spectrometer (Thermo Fisher Scientific, USA) equipped with an UltiMate 3000 LC system. A 15 cm-long LC column packed with 1.9 µm C₁₈ packing particles was used for the peptide separation. The mobile-phase solvents were HPLC water acidified by 0.1% FA for phase A and 80% acetonitrile acidified by 0.1% FA for phase B. An 88 min gradient of 6-45% buffer B was used for peptide elution at a flow rate of 600 nL min⁻¹. A high voltage of 2.3 kV was applied to the electrospray. The MS system was operated in positive ion mode at a resolution of 1 20 000 (at m/z200) using data-dependent acquisition (DDA) and full scans were acquired with a scan range of m/z 350–1550 with a target of 3×10^{6} ions for the automated gain control (AGC) and 20 ms maximum injection time. For the MS/MS scans, the normalized collision energy was set to 27%, the resolution was 15 000 at m/z200, accumulated for a maximum of 30 ms or until the AGC target of 2×10^4 ions was reached.

The raw LC-MS/MS data files were analyzed using MaxQuant (version 1.6.17.0) with the spectra searched against the UniProt mouse database (updated on February, 2022). For the identification of peptides, the mass tolerances were set to 20 ppm for the initial precursor ions and 0.5 Da for the fragmented ions. Two missed cleavages in tryptic digests were allowed. Cysteine carbamidomethylation was set a static modification. Oxidation of methionine and N-terminal acylation was accepted as the variable modification. Filtering for the protein and peptide identification was set at a 1% false discovery rate (FDR).

Both healthy mouse and HCC mouse samples were used in this study. Three biological replicates were performed for each group. Protein expression was quantified for each group using a dimethyl labeling quantitative approach. The intensities of the EV proteins from healthy and HCC mice were extracted from the MaxQuant result files to represent the final expression of specific proteins in the samples. Any missing values were filled according to the minimum value filling principle. The protein expression matrix was processed with the egdreR package for normalization and differential expression analysis. An adjusted *P* value of <0.01 and \log_2 fold change > 1.5 were used to indicate proteins that were significantly differentially expressed.³⁵ The obtained data were screened for differential proteins by the following principles: (1) proteins identified in at least two of the three experiments. (2) The protein contained at least 1 unique peptide. All statistical analyses were performed in the R statistical programming language (version 4.1.1). GO analysis using online DAVID Bioinformatics Resources 6.8 software.36

3. Results and discussions

Our approach for purifying EVs is to combine an EV extraction kit with TiO_2 microspheres. First, we used the purification kit for the crude separation of tissue EVs to remove aqueous contents and reduce the complexity of the tissue fluid. Then, the EVs were further enriched by exploiting the binding between TiO_2 and phosphate groups on the phospholipid bilayer of EVs. Due to the flexibility of the lipid bilayer, strong multisite interactions between the TiO_2 microspheres and the EVs were achieved.

It is difficult for TiO_2 microspheres to seize all EVs in crude tissue fluid due to the relatively low concentration of EVs. Additionally, any components with phospholipid bilayer structures in the tissue fluid tend to bind to TiO_2 microspheres, resulting in interference with EV isolation. Therefore, a combination of an extraction kit first followed by TiO_2 -based isolation was developed in this work to enrich EVs with high purity from tissues (Fig. 1).

3.1 Characterization of the isolated EVs

To characterize the enriched EVs, we compared kit-TiO₂ enrichment strategy with conventional enrichment approaches using SEM. As shown in Fig. 2A, the TiO₂ microspheres show a clean and smooth surface, with a uniform diameter of 5 μ m before incubation with the tissue fluid samples. After being incubated with the kit extracted tissue fluid samples, the surface of the TiO₂ microspheres was covered with a large number of round vesicles (Fig. 2B). The attached vesicles on the microsphere surface were further characterized by TEM. As shown in Fig. 2C, numerous vesicles were present on the surface of the TiO₂ microspheres. Under alkaline conditions (pH 10–12), the coordinated binding between the phosphate group and TiO₂ microspheres is destroyed. We introduced 10% ammonia to elute the enriched EVs for TEM and NTA analysis. After elution,

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intact, round vesicle structures were observed (Fig. 2D). This result suggested that the lipid bilayer was preserved during separation and elution.

We performed NTA particle size, number analysis and protein quantification (BCA) on the eluted EV. As shown in Fig. 2E, the size distribution of EVs isolated by the three methods were analyzed by NTA. The average particle size was 175.9 nm, 192.6 nm, and 239.2 nm for EVs extracted using kit-TiO₂, kit only, and TiO₂ only approaches, respectively. EVs extracted using the kit only and TiO₂ only method displayed a broad distribution of vesicles with several peaks. However, the EVs extracted using the kit-TiO₂ approach only had one major peak, indicating its narrower size distribution. As shown in Fig. 2F and G. The number of particles and protein abundance of EVs were measured by NTA and BCA protein assay kit. The results indicated that the concentration of EVs particles extracted using kit-TiO₂ was significantly higher than that of the other two methods. Interestingly, we found the protein of EVs extracted using the kit method was higher than that using kit-TiO₂. A possible explanation is that the EVs enriched by the kit method might contain more high abundant contaminate proteins.37-39

We also conducted western blotting experiments to confirm the above results. EVs were isolated by the kit-TiO₂ approach, TiO₂ alone and the kit alone to simultaneously validate the EV markers TSG101, CD63 and syntenin-1. As shown in Fig. 2H, the intensity of the maker protein syntenin-1 and CD63 were significantly higher in EVs isolated by the kit-TiO₂ approach than that obtained by the other two methods. Next, western blot analysis was performed to assess the purity of the extracted EVs. Typical cellular proteins, *e.g.*, Golgi (GM130), endoplasmic reticulum (Calnexin), and mitochondrial proteins (VDAC), were detected in high abundance in liver tissue homogenates, but were absent in the EV samples extracted by the combined method (Fig. 2I), suggesting that cell debris was completely removed and that the isolated EVs were of high purity.

The EVs enriched by the combined method were examined by SDS-PAGE and Coomassie Blue staining. The supernatant, washing buffers (first wash, second wash, and third wash solution), and tissue EVs eluted from the TiO₂ microspheres were all collected and examined. Protein signals were negligible in the washing buffer. Significant differences were detected between the tissue proteins (LHC) and EV proteins (Fig. 3A and B), indicating the heterogeneity between tissue cells and EVs. These results demonstrated that the EVs were effectively isolated through the binding to TiO₂.

To assess the reproducibility of this kit-TiO₂ isolation strategy, three biological replicates were tested. EVs were extracted from three identical tissue fluid samples using the proposed strategy. The obtained EVs were lysed with 4% SDS, and EV proteins were collected. Then, the isolated EV proteins were characterized by SDS-PAGE. As shown in Fig. S1,† similar proteomic distribution patterns were observed for the three replicates in lanes 1, 2 and 3, suggesting that good reproducibility was achieved.

3.2 Optimized enrichment strategy

To further improve the enrichment of the EVs, we evaluated the amount of TiO_2 microspheres used and the incubation time for EV isolation. Using 200 µL tissue fluid as the sample, the abundance of the EV protein marker TSG101 was traced before and after EV enrichment by western blotting. The protein intensity was relatively quantified based on the grayscale and normalized to the maximum value.⁴⁰ As illustrated in Fig. 4A, the abundance of the protein marker was extremely low when using 1 mg TiO₂. As the amount of TiO₂ microspheres increased, the marker intensity was enhanced and then saturated at 20 mg. This might be due to insufficient dispersion of

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Fig. 2 Characterization of the isolated EV (A) SEM image of TiO_2 microsphere; (B) SEM image of TiO_2 microsphere after incubation with kit extracted tissue EVs; (C) TEM image of the tissue EVs adsorbed on the TiO_2 microsphere; (D) TEM images of EVs isolated by kit- TiO_2 method was pointed by red arrows; (E) the size distribution of the isolated EVs separated by different methods and measured by NTA; (F) concentration of the EVs were measured by NTA for the different extraction methods, n = 3; (G) quantification of total protein by BCA protein assay for the EVs isolated by different methods, n = 3; (H) western blot of marker protein TSG101, CD63, and syntenin-1 for EVs extracted using the three different enrichment methods, including extraction kit only, TiO_2 only, and kit combined TiO_2 ; (I) western blot analysis of cellular interfering protein VDAC, calnexin, and GM130 in the extracted liver tissue EV and liver tissue homogenate; (H + I) 20 µg protein was loaded per lane.

 TiO_2 in the solution for amounts higher than 20 mg. As a result, the optimal amount of TiO_2 microspheres was 20 mg for enriching EVs from 200 μ L tissue fluid.

Furthermore, the incubation time of the tissue fluid with microspheres was assessed. As suggested above, the optimal amount of TiO₂ (20 mg) was used for 200 μ L of tissue fluid, and different incubation times of 1 min, 2 min, 5 min, 10 min, and

20 min were tested. Fig. 4B shows that the protein marker intensity increased with increasing incubation time and was saturated at 10 min of enrichment. Prolonged incubation might result in irreversible adsorption of EVs on the surface of TiO_2 microspheres, and lead to a lower recovery rate of EVs. Therefore, 10 min was the optimized enrichment time and was used



Fig. 3 (A) Separation of tissue EV proteins by SDS-PAGE (blue-stained), in which EVs were isolated by the kit-TiO₂ approach; (B) distribution of tissue EVs proteins and liver tissue proteins (LHC: liver homogenates collagenase) by SDS-PAGE (20 µg protein was loaded per lane).

for subsequent experiments for isolating EVs from 200 µL tissue fluid.

3.3 Proteomic analysis of EVs from mouse liver tissue

To apply the proposed EV isolation strategy, mouse liver tissue was selected as the model system. Tissue EVs were enriched by the kit-TiO₂ approach and characterized using mass spectrometry. As shown in Fig. 5A, we compared the proteins identified by different EV isolation methods with the mouse EV proteins in the existing EV database Exocarta.41 The kit-TiO2 method isolated 1317 EV proteins, while the kit alone, TiO₂ alone, and the

gold standard ultracentrifugation method isolated 1135, 1036, and 1166 EV proteins. The greater number of EV proteins isolated by kit-TiO₂ methods indicated its advantages for tissue EV studies. Further comparison in Fig. 5B showed that a total of 1966 protein groups were present in EVs isolated from mouse liver tissue fluids using kit-TiO₂ enrichment strategy, while a total of 1801, 1629, and 1501 protein groups were isolated with ultracentrifugation, extraction kits, and TiO₂-based isolation, respectively. In addition, the majority of the EV protein groups isolated using ultracentrifugation (56.8%), the extraction kit (60.2%), and TiO₂-based enrichment (63%) were also isolated by

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Fig. 4 Optimization of experimental conditions for EVs isolation. The protein marker TSG101 was relatively quantified based on the greyscale of western blot. Relative abundance of TSG101 was a function of the quantity of TiO₂ (A) and incubation time (B). Protein intensity was normalized to the maximum intensity value.



Fig. 5 (A) Ven diagram of EV proteins isolated using ultracentrifugation, kit only, TiO₂ only, and kit-TiO₂ methods; (B) overlap of protein groups identified in the tissue EV isolated using ultracentrifugation, extraction kit, TiO₂ only, and kit-TiO₂ approaches; (C) overlap of proteins identified from biological triplicates using kit-TiO₂ method; (D) reproducibility evaluation of the kit-TiO₂ enrichment strategy.

the kit-TiO₂ approach. Further analysis revealed that the kit-TiO₂ strategy not only efficiently covered 62% proteins obtained by the other three methods, but also provided 321 extra proteins, indicating its higher capability in identifying EV proteins from tissue sample. Further comparison in Fig. S2[†] revealed lower protein abundance of the uniquely identified proteins by the kit-TiO₂ method than the shared proteins with other methods, which indicated the higher sensitivity of the kit-TiO₂ method to identify more EV proteins. Over 84.5% of the protein groups were identified in at least two of the three replicates by the kit-TiO₂ approach (see Fig. 5C). Good reproducibility with Pearson correlation coefficients greater than 0.83 was achieved (see Fig. 5D). This result demonstrates the high reproducibility of the method for extracting EVs from tissues using kit-TiO₂ approach.

Furthermore, Gene Ontology (GO) analysis was performed to probe the subcellular location of the identified tissue EV proteins. As illustrated in Fig. S3,† liver EV proteins were highly enriched in the cellular component of extracellular exosome, suggesting the vesicular and extracellular nature of the proteins. Importantly, the identified proteins were irrelevant to nuclear, Golgi, endoplasmic reticulum or blood microparticles, indicating negligible contamination from cellular debris or other vesicles. In the biological process category, the identified protein groups were mainly relevant to metabolic, oxidationreduction, and protein transport processes. Additionally, the most abundant molecular functions included poly(A) RNA binding, catalytic activity, and cadherin binding involved in cell-cell adhesion. We further performed gene ontology (GO) analysis of EVs proteins enriched by four methods. In Fig. S4,†

the number of proteins classified as EVs in the cellular component were obviously higher in EVs extracted using kit-TiO₂ than using the other three methods. This result demonstrates the reliability of the kit-TiO₂ method and the higher enrichment efficiency.

3.4 Proteomic analysis of tissue EVs from hepatocellular carcinoma and healthy mice

Characterization of EV proteomic profiles may contribute to finding therapeutic targets for the treatment of hepatocellular cancer. We compared the EV proteins extracted from liver tissue of HCC and healthy mice using kit-TiO2 method. Three HCC mice and three healthy mice were analyzed. The proteomic profile showed that a total of 1176 and 1083 EV protein groups were isolated from healthy and HCC mice, respectively (Fig. 6A). Multivariate statistical analysis was performed on proteins identified more than two times to screen the proteomic difference between the two groups. Proteins with a p value <0.01 and $\log_2 |FC|$ > 1.5 were considered significantly differentiated. The volcano plot generated in Fig. 6B reveals 25 upregulated and 45 downregulated proteins in the HCC group relative to the normal group. The 70 differentially expressed proteins in HCC were further probed using unsupervised hierarchical clustering analysis (Table S1[†]). As shown in Fig. 6C, the HCC and healthy groups were clearly separated, indicating that the differentially expressed EV proteins may be useful as biomarkers or therapeutic targets for HCC.

Furthermore, the biological functions of the significantly changed proteins were investigated. Among the 25 upregulated proteins, Atic, Copa, Cont3, Me1, Anxa3, Fth1, Anxa5, Phb1, Acaa2, ATPD, and Glud1 were found to be highly relevant to



Fig. 6 Proteomic analysis of tissue EV from the healthy and HCC mice. (A) Overlap of EV protein groups identified from the two groups; (B) volcano plot of the tissue EV proteins; (C) unsupervised clustering of the significantly changed proteins.

HCC. For example, Atic aided in HCC growth and migration by functioning in the AMPK and mTOR signaling pathways.⁴² Secretory Anxa3 contributes to tumorigenic, metastatic, and self-renewal processes in HCC, and its blockade has been reported as a therapeutic target for hepatocellular carcinoma.⁴³ In addition, several proteins, *e.g.*, Ndufv1, Pygl, Sardh, and Ft11, were previously associated with other cancers including breast cancer and colorectal cancer. Therefore, the significantly altered proteins in tissue EVs might be considered useful markers for the diagnosis of HCC.

4. Conclusions

In this work, we presented a novel approach to enrich EVs from animal tissue using kit-TiO₂ method. High-quality and highpurity EVs were obtained as demonstrated by western blot and proteomic analysis. Successful application of this method led to the identification of dozens of differentially expressed EV proteins in the HCC samples. This new isolation strategy provides a powerful tool for enriching EVs from animal tissues and may support biomarker discovery and the diagnosis of hepatic tumors.

Conflicts of interest

There are no conflicts to declare.

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