Antibody-coupled monolithic silica microtips for high-throughput molecular profiling of circulating exosomes

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Exosome-mediated signal transportation plays a variety of critical roles in cancer progression and metastasis. From the aspect of cancer diagnosis, circulating exosomes are ideal resources of biomarkers because molecular features of tumor cells are transcribed on them. However, isolating pure exosomes from body fluids is time-consuming and still major challenge to be addressed for comprehensive profiling of exosomal proteins and miRNAs. Here we constructed anti-CD9 antibody-coupled highly porous monolithic silica microtips which allowed automated rapid and reproducible exosome extraction from multiple clinical samples. We applied these tips to explore lung cancer biomarker proteins on exosomes by analyzing 46 serum samples. The mass spectrometric quantification of 1,369 exosomal proteins identified CD91 as a lung adenocarcinoma specific antigen on exosomes, which was further validated with CD9-CD91 exosome sandwich ELISA measuring 212 samples. Our simple device can promote not only biomarker discovery studies but also wide range of omics researches about exosomes.

Lung cancer is the leading cause of cancer-related mortality worldwide, accounting for 1,475,117 deaths in 2011 (Global Health Observatory Data Repository, World Health Organization). The high mortality is mainly attributable to a late-stage diagnosis and the lack of effective treatments. Indeed, by means of current cancer screening tests, only 30% of patients are diagnosed at an early disease stage and present surgically resectable tumors. Therefore development of novel biomarkers and establishment of blood-based early detection system for lung cancer is crucial in order to improve clinical outcome and overall survival rate.

Recently biological significance and clinical utility of exosomes have been extensively discussed. Particularly contribution of tumor-derived exosomes to the formation of metastatic microenvironments is one of the most fundamental functions of them, which would provide a better understanding for cancer metastasis and even new therapeutic strategies to prevent metastasis. Exosome-mediated delivery of therapeutic RNAs has been already in a pioneering stage for cancer treatment. In the field of cancer diagnosis, exosomes are also fascinating targets for biomarker discovery due to their molecular characteristics. In principle, a set of molecules expressed in original solid tumor cells would be detectable as exosomal components in blood circulation. Despite the theoretical feasibility of exosomal biomarkers, difficulties in exosome isolation from biological fluids have significantly hindered effective discovery of biomarker candidates. In fact, although ultracentrifugation-based methods are the most common strategies to isolate exosomes from serum samples, the reproducibility, processing time, and purity are not appropriate for biomarker screening studies dealing with a lot of clinical samples quantitatively.

In the present study, we established an antibody-assisted exosome purification tips by immobilizing anti-CD9 antibody to Mass Spectrometric Immunoassay (MSIA) monolith pipette tips. This multi-channeled platform effectively streamlined proteome-wide mass spectrometric profiling of serum exosomes and allowed accurate statistical identification of lung cancer-specific exosomal proteins. We further constructed exosome sandwich ELISA assays for large-scaled replication study to validate screening reliability for an identified exosome surface antigen CD91.

Results

Isolation of serum exosomes by anti-CD9-MSIA tips. To perform reproducible and high-purity separation of exosomes from serum, we employed the antibody-immobilized low back pressure monolithic tips on automated
12-channel pipette system (Figure 1a), which allowed 30 minutes isolation of exosomes from 12 serum samples simultaneously. Here we selected a tetraspanin molecule CD9 as a target of exosome-capturing antibody due to its strong expression on the surface of exosomes secreted from diverse cell types. In order to evaluate the reproducibility of anti-CD9-MSIA tips, exosomes were purified from a pooled serum sample using 6 independent anti-CD9-MSIA tips and analyzed by LC/MS/MS in triplicated measurements. The coefficient of variation (CV) of peak intensities corresponding to CD9 155-170 peptide (GLAGGVEQFISDICPK, m/z = 845.9266) or CD81 149-171 peptide (TFHETFIDCGSTLTALTTSVLK, m/z = 848.0733) was shown. Exosomes were isolated from 46 serum samples by anti-CD9 antibody-coupled monolith tips (anti-CD9-MSIA tips) on 12-channel automatic pipetting platform. The enriched exosome fractions were analyzed by LC/MS/MS and subjected to label-free quantification analysis by RefinerMS software on the Expressionist proteome server system. The quantified peptides underwent 2-step statistical analysis, composed of ANOVA and feature elimination method, and finally extracted biomarker candidate peptides were identified with Sequest database search. The identification threshold was set at false discovery rate (FDR) < 1%.

Figure 1 | Schematic view of exosomal biomarker discovery workflow. (a) Magnified picture of anti-CD9 MSIA tips (left) and a dedicated holding fixture (right). Pictures were taken by authors. (b) Exosome fractions were purified from a pooled serum sample using 6 independent anti-CD9-MSIA tips and analyzed by LC/MS/MS in triplicated measurements. The coefficient of variation (CV) of peak intensities corresponding to CD9 155-170 peptide (GLAGGVEQFISDICPK, m/z = 845.9266) or CD81 149-171 peptide (TFHETFIDCGSTLTALTTSVLK, m/z = 848.0733) was shown. (c) Exosomes were isolated from 46 serum samples by anti-CD9 antibody-coupled monolith tips (anti-CD9-MSIA tips) on 12-channel automatic pipetting platform. The enriched exosome fractions were analyzed by LC/MS/MS and subjected to label-free quantification analysis by RefinerMS software on the Expressionist proteome server system. The quantified peptides underwent 2-step statistical analysis, composed of ANOVA and feature elimination method, and finally extracted biomarker candidate peptides were identified with Sequest database search. The identification threshold was set at false discovery rate (FDR) < 1%.
identification. Then we next isolated serum exosomes from 10 normal controls (NC), 10 interstitial pneumonia patients (IP), 14 lung adenocarcinoma patients (ADC), and 12 lung squamous cell carcinoma patients (SCC) using anti-CD9-MSIA tips. Purified exosomes were individually analyzed by LC/MS/MS system and subjected to statistical analysis as shown in Figure 1c.

**Proteome-wide overview of human serum exosomes.** The LC/MS/MS analysis of 46 serum samples (Table 1) and subsequent Sequest database search identified 1,369 non-redundant proteins (FDR < 1%, Supplementary Table 1). To assess the purity of anti-CD9-MSIA tip eluates, identified proteins were classified according to subcellular localizations (Figure 2a). The Cellular Component distribution by DAVID GO analysis illustrated highly-enriched 701 intracellular proteins (51.2%) and 290 plasma membrane proteins (21.2%), whereas only 135 extracellular (secreted) proteins (9.8%) were identified. These values clearly represented efficient enrichment of exosomes bearing original cell-derived cellular components. Importantly, most of serum abundant proteins such as albumin and IgG were effectively washed out during MSIA purification steps, which often hindered the sensitive detection of minor exosomal proteins. Moreover to elucidate physiological

| Table 1 | Clinical Information of serum samples |
|---------|---------------------------------------|
|         | Screening set (n = 46) | Validation set (n = 212) |
|         | NC | IP | ADC | SCC | NC | IP | ADC | SCC |
| No. of samples | 10 | 10 | 14 | 12 | 54 | 19 | 105 | 34 |
| Age (average) | 44.7 | 71.6 | 61.5 | 65.3 | 40.2 | 65.7 | 64.4 | 68.8 |
| Gender [female/male] | 4/6 | 3/7 | 6/8 | 2/10 | 29/25 | 5/14 | 35/70 | 7/27 |
| TNM stage | | | | | | | | |
| I | 0 | 0 | 12 | 5 | | | | |
| II | 0 | 0 | 10 | 0 | | | | |
| III | 4 | 5 | 27 | 18 | | | | |
| IV | 10 | 7 | 36 | 11 | | | | |

NOTE: NC, normal control; IP, interstitial pneumonia; ADC, adenocarcinoma; SCC, squamous cell carcinoma.

Figure 2 | Proteome-wide overview of 1,369 identified exosomal proteins. (a) Distribution of protein subcellular localization was shown in a pie chart. (b) The Fisher Exact Statistics in DAVID system was used for functional annotation clustering analysis. The 10 enriched functions detected in 1,369 exosomal proteins were shown with Expression Analysis Systematic Explorer (EASE) scores.
functions of serum exosomes, Expression Analysis Systematic Explorer (EASE) scores were calculated\(^\text{13,14}\) (Figure 2b). This functional estimation suggested the possible association of serum exosomes with immune regulations, cell-to-cell interactions, and stimulation responses, in addition to vesicle transport. These data would contribute to new revelations about the biological functions of not only tumor-derived exosomes but also normal exosomes.

Statistical identification of exosomal biomarkers for lung cancer. The label-free quantification analysis on the Expressionist RefinerMS module (Figure 1c and Supplementary Figure 1) quantified 113,582 non-redundant peptides from 46 serum samples. In the first statistical selection, 3-group ANOVA was used to roughly extract signature peptides specific to ADC patients (230 peptides, \(p < 0.001\)) or SCC patients (316 peptides, \(p < 0.001\)) as shown in Figure 3a or 3b, respectively. Here, to identify lung cancer specific exosomal biomarkers which don’t react with inflammatory lung diseases such as interstitial pneumonia, IP patients were also considered as a non-cancer control group. For the second stage, cross validation-based feature elimination method was employed to compute the minimum biomarker sets which provided the least misclassification rates. Here support vector machine recursive feature elimination (SVM-RFE) method or SVM-SVM method defined 181 or 32 peptides as final candidate biomarker sets demonstrating 90.9% or 100% true prediction rate for ADC or SCC patients group (Figure 3c or 3d, respectively). By referring to protein identification data, 20 peptides derived from 18 proteins were identified (Figure 4 and Table 2). Among them, CD91, Integrin alpha-IIb, and CD317 were selected as favorable exosomal biomarker candidates because their expected localization was on the surface of exosomes and could be measured by exosome sandwich ELISA system (Figure 5a).

Validation experiment for CD91 by exosome sandwich ELISA. To assess the quantitative reproducibility of the label-free quantification results in our single-run screening analysis, as well as the clinical usefulness of a candidate biomarkers, we conducted further validation study by exosome sandwich ELISA using 212 independent serum samples (Table 1). Among three candidate biomarker proteins, we eventually succeeded to obtain good antibody and construct ELISA assay only for CD91. In this assay, we utilized anti-CD9 antibody as an exosome-capture antibody and
biotinylated anti-CD9 or biotinylated anti-CD91 antibody as a detection antibody (Figure 5a). Since serum exosome concentrations, measured by CD9-CD9 sandwich ELISA, had drastic individual variability (Figure 5b), the measurements of CD9-CD91 sandwich ELISA were normalized by exosome concentrations (denoted by U/exosome in Figure 5c). We also tested a classical clinical biomarker CEA in the same sample set (Figure 5d) and compared the diagnostic efficacy to that of exosomal CD91. When we set the cut-off value at 2.04 U/exosome for exosomal CD91 and 5.0 ng/ml for CEA, exosomal CD91 showed significantly higher sensitivity for detecting stage-I, II ADC patients (54.5%) compared with CEA (22.7%), while the detection power of exosomal CD91 in stage-III, IV ADC patients (61.4%) was similar with that of CEA (66.3%). The false positive rate of exosomal CD91 in the control group (NC and IP, n = 73) were 11.0%, while that of CEA was 8.2%. These results indicated that exosomal CD91 possessed better potential for the early detection of lung ADC compared with CEA.

Discussion
In recent years, physiological roles of secreted microvesicles, including exosomes, were actively studied mainly using cell culture supernatants (CCS). However studies on body fluid exosomes have been still at the stage of seeking appropriate technologies which allow high-purity isolation of exosomes. In fact, ultracentrifugal sedimentation methods, the most common techniques to purify exosomes15, can not remove large size proteins (e.g. α2-macroglobulin, IgM, complement factors, and so on), aggregated proteins, circulating proteasomes16,17, and vault ribonucleoprotein particles18 in serum because they have similar diameters (40–100 nm) and sedimentation coefficient to exosomes. Owing to these major contaminants, serum-derived proteins often occupy a great part of identified proteins by LC/MS/MS analysis of ultracentrifugal sedimentation samples (data not shown). In addition to the purity issue, throughput and reproducibility are also critical factors for the development of exosome isolation methods because biomarker screening studies or therapeutic target discovery studies usually deal with lots of clinical

Figure 4 | Identified 18 exosomal biomarker candidates. The LC/MS/MS signal intensities for 18 biomarker candidates acquired from 46 cases were displayed in box plots. The UniProtKB entry protein names accompanied with amino acid numbers were shown over the box plots. N, normal control; IP, interstitial pneumonia; ADC, adenocarcinoma; SCC, squamous cell carcinoma.
the expression level of CD9 might vary by tissues.

Although CD9 is one of the most well-known exosome markers, the present study, we used anti-CD9 antibody to capture exosomes in detection power for limited number of early stage patients (n = 5).

Isolation method could also promote development of mass spectrometry multiplexed biomarker diagnosis measuring two or more exosomal biomarkers simultaneously by MRM/SRM assay.

To improve the comprehensiveness of exosome profiling studies. From the proteome-wide biomarker screening of 46 serum samples, we eventually identified 18 biomarker candidate proteins in Figure 4 and Table 2. Concerning subcellular localizations of them, some; CD91, integrin alpha-IIb, and CD317 on plasma membrane, copine-1 on vesicle membrane, myeloperoxidase in lysosomes, basement membrane-specific heparan sulfate proteoglycan core protein, and tested in clinical trials. This evidence indicated that the lung ADC cells directly release high level of CD317-positive exosomes in blood circulation.

Thus in addition to identification of biomarker candidates, our anti-CD9 MSIA tip can provide diverse novel knowledge about human exosomes. More comprehensive information will be also obtained by replacing the epitope of immobilized antibody with other exosomal surface antigens such as CD63, CD81, tetraspanin-9, or tetraspanin-14.

### Methods

**Serum samples.** Serum samples from lung cancer patients (n = 165), interstitial pneumonia patients (n = 29), and normal controls (n = 64) were collected in Hiroshima University Hospital within the same period. All samples were collected eight days after the disease diagnosis without any anticancer or anti-inflammatory therapy.

**CD91, this molecule is a type-1 transmembrane receptor which mediates ligand endocytosis in clathrin-coated pits and cargo trafficking to lysosomes.** Another fundamental role of CD91 is reported as a signaling receptor regulating cytokine secretion, phagocytosis and migration of cells in the immune system. Notably, with regard to cancer, no or very low expression of CD91 in lung cancer cells was observed in tissues from ADC patients with poor clinical outcome, while strong staining patterns of CD91 were observed in stromal cells surrounding cancer cells from 94/111 ADC patients. Combined with our results, high level of serum CD91–expressing exosomes would be originally secreted from stromal cells surrounding lung cancer cells. Since construction of tumor microenvironments is one of the most well-established functions of exosomes, inhibiting production or function of CD91–positive exosomes might lead to suppression of lung cancer progression. Therefore inhibitors of CD91, which interfere ligand binding, such as the receptor-associated protein (RAP), suramin, v-2-macroglobulin, and lactoferrin would be useful for investigating the association of exosomal CD91 and lung cancer. In contrast to CD91, another candidate of exosomal biomarker CD317 was already confirmed as a potential therapeutic target for lung cancer. Due to its nature of lung cancer specific expression, chimeric or humanized antibody drugs were developed and tested in clinical trials. This evidence indicated that the lung ADC cells directly release high level of CD317–positive exosomes in blood circulation.

| m/z | z | Protein Description | UnIProtKB entry | Protein Accessions | Sequence | XCorr |
|-----|---|---------------------|-----------------|-------------------|----------|-------|
| 713.401 | 2 | Brain-specific angiogenesis inhibitor 1-associated protein 2 | BAIP2 | Q9UQ88 | EGDULLVPEAR | 3.22 |
| 579.285 | 2 | Copine-1 | CPNE1 | Q99829 | SDPFLFFI | 2.67 |
| 507.583 | 3 | Fibrinogen alpha chain | FIBA | P02671 | GLDENVQDFTRN | 4.54 |
| 1047.638 | 2 | Complement C5 | CO5 | P10313 | LNVLATLPFTKGPIYPK | 3.79 |
| 1011.136 | 3 | Proliferation lipoprotein | CD91/1/LRP1 | Q07954 | DGHNVGEAVDWMGDNLNYWTDGPK | 4.57 |
| 953.123 | 3 | Mannan-binding lectin serine protease 1 | MAN1B1 | P48740 | ETTOETQPTGQEVVSPLGSFMSTSFP | 5.28 |
| 1027.069 | 2 | Apolipoprotein B-100 | APOB | P01414 | IIGELGFASLDIQLQDG | 5.58 |
| 1100.068 | 2 | Complement component C8 alpha chain | COBA | P07357 | YNPWVIDEMQHPHEVL | 5.22 |
| 619.324 | 2 | Plasminogen | PLMN | P00747 | FVTWIEGVMR | 3.31 |
| 481.584 | 3 | Basement membrane-specific heparan sulfate proteoglycan core protein | PGBM | P98160 | IAHVELADAGYR | 3.52 |

**Table 2** | 18 exosomal lung cancer biomarker candidates

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| m/z | z | Protein Description | UnIProtKB entry | Protein Accessions | Sequence | XCorr |
|-----|---|---------------------|-----------------|-------------------|----------|-------|
| 670.872 | 2 | Myeloperoxidase | PERM | P05164 | IGLDLPALMQR | 3.30 |
| 1441.657 | 2 | IgG-binding protein | FCGBP | Q99687 | VPSYAEAALCGCGNDFPADDLAR | 4.49 |
| 512.332 | 2 | Integrin alpha-IIb | ITA2B | P08514 | IVDLVPR | 3.21 |
| 1020.122 | 3 | Ig kappa chain V-III region HIC | KV313 | P18136 | LEPEDFAVYQCGYSWPFGGHTK | 4.62 |
| 742.352 | 2 | Plasma protease C1 inhibitor | IC1 | P15515 | VTTSDQMLSIMEK | 2.65 |
| 542.811 | 2 | Alpha-2-macroglobulin | A2MG | P01023 | GHFSIPVK | 4.69 |
| 1141.520 | 3 | Bone marrow stromal antigen 2 | BAIP2 | Q10589 | GFQDVEAQATCNHTVMLMASDAEAQGQ | 3.11 |
| 727.377 | 2 | Hyaluronan-binding protein 2 | HABP2 | Q14520 | LGPVDGHcALESK | 2.95 |

**NOTE:** m/z, mass/charge; z, charge; XCorr, cross-correlation score from Sequest database search.
from untreated patients at the initial visit to hospital. The nonanticoagulated blood samples were collected and allowed to clot at room temperature for 1–2 hours. Sera were then separated by centrifugation at 1500 rpm for 15 min and stored frozen at \(-80\)°C. Written informed consents were obtained from all participants. This study was approved by The Ethical Committee of RIKEN (Approval code: Yokohama H20-12) and The Ethical Committee of Hiroshima University Hospital. All experiments were performed in accordance with relevant guidelines and regulations.

**Exosome purification by anti-CD9 MSIA tips.** All procedures were performed on Novus-i 12-channel electronic pipettes and adjustable pipette stand (Thermo Fisher Scientific, Waltham, Massachusetts, USA). The MSIA D.A.R.T.'s, Protein G tips (Thermo Fisher Scientific) were equilibrated with 300 μl PBS 3×10 cycles prior to 25 μl 100 cycles of immobilization of 1 μg anti-CD9 antibody (provided by Shionogi & Co., Ltd., Osaka, Japan) in 50 μl PBS. After crosslinking with 0.25 mM BS3 (Thermo Fisher Scientific) in PBS (100 μl 100 cycles), reaction was quenched with 50 mM ethanolamine-HCl (pH 8.0) (100 μl 100 cycles). Following 300 μl 10 cycles of equilibration in PBS, the anti-CD9-MSIA tips were incubated with 350 μl of seven-fold diluted serum samples by 300 μl 100 cycles pipetting. Tips were washed three times by 300 μl 25 cycles pipetting in PBS. Finally captured exosomes were eluted by 20 μl 100 cycles pipetting in 30 μl of [8M Urea, 50 mM ammonium bicarbonate] solution.

After reduction with 5 mM TCEP at 37°C for 30 minutes and alkylation with 25 mM iodoacetamide at room temperature for 45 minutes, samples were diluted 7 times with 50 mM ammonium bicarbonate. Proteins were digested by Immobilized Trypsin beads (Thermo Fisher Scientific) in a 96-well filter plate with continuous shaking at 37°C for 6 hours. Tryptic digests were desalted by Oasis HLB 96-well μElution Plate (Waters Corporation, Milford, Massachusetts, USA) and subjected to LC/MS/MS analysis.

**LC/MS/MS analysis.** The dried peptide samples were resuspended in 2% acetonitrile with 0.1% trifluoroacetic acid and analyzed by LTQ-Orbitrap-Velos mass spectrometer (Thermo Fisher Scientific) combined with UltiMate 3000 RSLC nano-flow HPLC system (DIONEX Corporation, Sunnyvale, California, USA). Samples were separated on 75 μm × 150 mm C18 tip-column (Nikkyo Technos, Tokyo, Japan) using solvent A [0.1% formic acid] and solvent B [0.1% formic acid in acetonitrile] with multistep linear gradient of solvent B 6.4 to 30% for 95 minutes and 30 to 95% for 10 minutes at a flow rate 250 nl/min. The eluted peptides were ionized with the spray voltage 2000 V and MS data was acquired in a data-dependent fragmentation method in which the survey scan was acquired between m/z 400 to 1600 at the resolution 60,000 with automatic gain control (AGC) target value of 1.0 × 106 ion counts. The top-20 intense precursor ions in each survey scan were subjected to low resolution MS/MS acquisitions using normal CID scan mode with AGC target value of 5,000 ion counts in the linear ion trap.

The protein identification analysis was performed by SEQUEST database search on Proteome Discoverer 1.3 software (Thermo Fischer Scientific). The MS/MS spectra were searched against human protein database SwissProt 2013_03 (20,255 sequences).
using search parameters as follows: Enzyme Name = Semtrypsin, Precursor Mass Tolerance = 10 ppm, Fragment Mass Tolerance = 0.8 Da, Dynamic Modification = Oxidation (Met), and Static Modification = Carbamidomethyl (Cys). False discovery rate of 0.01 and Peptide Rank of 1 were set for peptide identification filters. The gene ontology analysis was performed on EASE (Expression Analysis Systematic Explorer, version 2.0) (http://www.geneontology.org/) to compute the overrepresented functional categories in "Cellular Component," and "Molecular Function".

**Label-free quantification analysis on Expressionist server.** The LC/MS/MS data set was loaded to Expressionist RefinerMS module (Genedata AG, Basel, Switzerland) for data processing and label-free quantification analysis. The whole workflow of RefinerMS software was shown in Supplementary Figure S1. The Spectrum Grid was set at every 10 data points on 2D MS chromatogram planes (x = m/z and y = RT). The Structure Removal was sequentially performed with RT = 2 scans and m/z = 6 points in the first and second Chemical Noise Subtraction, respectively, followed by the third Subtraction using RT Window = 500 scans and Quantile = 90%. After forth Subtraction with RT = 2 scans, signals with intensity < 2,000 were clipped off by Intensity Thresholding. Finally further Structure Removals with RT = 2 scans and m/z = 5 points were run as the fifth and sixth Chemical Noise Subtractions, respectively.

Then Chromatogram Grid was set at every 10 scans on noise-subtracted data, followed by Chromatogram RT Alignment using parameters: m/z Window = 11 points, RT Window = 11 scans, Gap Penalty = 1, RT Search Interval = 2 minutes, and Alignment Scheme = pairwise alignment based tree. Next, the Summed Peak Detection Activity detected the peaks on a temporarily-averaged chromatogram with parameters as follows: Summation Window = 20 scans, Overlap = 10, Minimum Peak Size = 6 scans, Maximum Merge Distance = 1 data point, Gap/Peak Ratio = 5, Method = curvature-based peak detection, Peak Refinement Threshold = 5, and Consistency Filter Threshold = 1. Finally Summed Isotope Clustering Activity grouped isotopic peaks derived from single molecule into an isotope cluster. Here parameters were used as follows: Minimum Charge = 1, Maximum Charge = 6, Maximum Missing Peaks = 0, First Allowed Gap Position = 10, Ionization = protonation, RT Tolerance = 0.1 minute, m/z Tolerance = 0.01 Da, and Minimum Cluster Size Ratio = 0.5.

**Statistical analysis on Expressionist Analyst.** Two-step statistical selection was employed for the effective identification of biomarker set. In the first stage, 3-group ANOVA was performed to roughly extract the candidates showing significantly distinct expression level among three clinical groups (p < 0.001). Next the minimum combination of biomarkers which provided the best classification rate was estimated by the Support Vector Machine-Recursive Feature Elimination (SVM-RFE) Ranking method in the Expressionist Analyst module (Genedata AG). SVM-RFE is an iterative algorithm that works backward from an initial set of statistical features. As a classifier, SVM was used for inferring decision rules. RFE was used as a ranking method which iteratively dropped the 10% peptides with the lowest weights in each step. Finally a Line Plot visualizer was returned displaying the average misclassification rate. Each classifier was represented by a line and the optimal classifier and biomarker peptide set size for the chosen ranking method could be read off.

**Exosome sandwich ELISA assay.** The 250 ng/well of anti-CD9 antibody was immobilized to Nunc MaxiSorp flat-bottom 96 well plate (Thermo Fischer Scientific). The blocking solution (150 μl/well of 5% BSA in PBS) was then added and incubated on the plate shaker at ambient temperature for 60 minutes. After 3 times wash with PBS, [5 μl serum + 95 μl PBS] or [30 μl serum + 70 μl PBS] was loaded to the upper 48 wells or the lower 48 wells, respectively. Following 5 hours incubation, plates were washed three times with PBS. The 100 μl/well of biotinylated anti-CD9 antibody (125 ng/ml) or biotinylated anti-CD91 antibody (500 ng/ml, Abcam, Cambridge, UK) in 1% BSA was loaded to the upper 48 wells or the lower 48 wells, respectively. After 60 minutes incubation, plates were washed three times with PBS and then covered with 100 μl/well of 1×HRP-Streptavidin (Abcam) in 1% BSA solution. After 30 minutes incubation, plates were washed three times with PBS and covered with 100 μl/well of TMB Ready Solution (Thermo Fischer Scientific). The reaction was stopped after 10 minutes incubation using 100 μl/well of 2N HCl. The OD at 450 nm was immediately measured. The concentration data from CD9-CD9 and CD9-CD91 ELISA assays were normalized with gradient curves (5, 10, 15, 20, and 25 μl) obtained from a common pleural effusion sample. Finally CD91 concentrations were normalized with exosome concentration calculated with CD9-CD9-CD91 ELISA above.

**Box plot analysis and ROC curve analysis.** The intensities of mass spectrum peaks corresponding to candidate biomarker peptides were displayed by box plot using R algorithm. For each study the box represents the middle half of the distribution of the data points stretching from the 25th percentile to the 75th percentile. The line across the box represents the median. The lengths of the lines above and below the box are defined by the maximum and minimum data point values, respectively, that lie within 1.5 times the spread of the box. ROC curves were also depicted by R. The cut-off value was set at the point whose distance from the (sensitivity, specificity) (1, 1) reached the minimum. The sensitivity (Sens), specificity (Spec), positive predictive value (PV+), negative predictive value (PV−), and area under the curve (AUC) were shown on each graph.

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**Author contributions**

K.U. designed the study and developed the method. K.U., N.I., A.T., N.S. and R.F. performed experiments and analyzed data. K.U. and H.N. interpreted the results and wrote the manuscript.

**Additional information**

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