A Novel Truncated Form of Nephronectin Is Present in Small Extracellular Vesicles Isolated from 66cl4 Cells

Jimita Toraskar,†,§ Synnøve N. Magnussen,⊥ Lars Hagen,∥,¶ Animesh Sharma,†,¶ Linh Hoang,†,§ Geir Bjørkøy,†,‡ Gunbjørg Svineng,⊥,¶ and Tonje S. Steigedal†

†Department of Clinical and Molecular Medicine and ‡Centre of Molecular Inflammation Research, Faculty of Medicine and Health Sciences, and ¶CMIC, Cellular and Molecular Imaging Core Facility, Norwegian University of Science and Technology, N-7491 Trondheim, Norway
‖Cancer Clinic, St. Olav’s Hospital, Trondheim University Hospital, Trondheim, Norway
∥Department of Medical Biology, Faculty of Health Sciences, UiT—The Arctic University of Norway, N-9037 Tromsø, Norway
⊥PROMEC, Proteomics and Modomics Experimental Core Facility, Norwegian University of Science and Technology, N-7030 Trondheim, Norway
§Arctic University of Norway, N-7490 Narvik, Norway

Supporting Information

ABSTRACT: Extracellular vesicles are emerging as biomarkers in breast cancer. Our recent report suggested that an intracellular granular staining pattern of the extracellular matrix protein nephronectin (NPNT) in breast tumor sections correlated with a poor prognosis. Furthermore, the results showed that NPNT is localized in extracellular vesicles derived from mouse breast cancer cells. In this study, we performed proteomic analysis that revealed that several proteins, including tumor-promoting molecules, are differentially expressed in the cargo of small extracellular vesicles (sEVs) derived from NPNT-expressing mouse breast cancer cells. We also identified three different forms of NPNT at 80, 60, and 20 kDa. We report that the native form of NPNT at 60 kDa becomes further glycosylated and is detected as the 80 kDa NPNT, which may be processed by matrix metalloproteinases to a shorter form of around 20 kDa, which has not previously been described. Although both 80 and 20 kDa NPNT are detected in sEVs derived from breast cancer cells, the 20 kDa form of NPNT is concentrated in sEVs. In summary, we show that a novel truncated form of NPNT is found in sEVs derived from breast cancer cells.

KEYWORDS: breast cancer, nephronectin, small extracellular vesicles

1. INTRODUCTION

Extracellular vesicles can be classified according to their size: exosomes (30–100 nm), microvesicles (100–1000 nm), apoptotic bodies (50 nm to 2 μm), and oncosomes (1–10 μm).1 Microvesicles are bilipid-membrane vesicles originating from the plasma membrane,2 initially disregarded as cellular debris but now recognized as biologically significant.3,4 Extracellular proteins may enter into the intraluminal vesicles (ILVs) within multivesicular bodies (MVBs)2 on inward budding of the plasma membrane. The invaginations of the limiting membrane of the MVBs further allow several intracellular proteins to enter into the ILVs.5,6 The content of MVBs is either released as exosomes into the extracellular milieu or into the lysosomes for degradation.7,8 The secreted exosomes may release their contents into a recipient cell by fusion or interact with the target cells via cell surface proteins.9,10 Ultracentrifugation is regarded as the gold standard for exosome isolation.11,12 However, it has been found that the pellet of vesicles obtained after spinning the supernatant of cells at 100 000g for 70 min contains a heterogeneous population of membrane vesicles, in addition to enriched exosomes.13–16 Therefore, the preferred terminology for the isolated fraction is small extracellular vesicles (sEVs).17

The content of sEVs is not random;18 rather, the cargo is cell and disease-type specific and may deliver discrete molecular messages that inflict a biological response.19 sEVs are found to carry proteins,20 lipids,21 transposable genetic elements,22 double-stranded DNA,23 mitochondrial DNA,24 and several types of RNAs.25 Several extracellular matrix (ECM) proteins, such as collagens, fibronectin, vitronectin, and nephronectin (NPNT), are found in sEVs.26 NPNT has previously
been reported in isolated sEVs from human ductal saliva, colorectal cancer cells, and mouse breast cancer cells.

Breast cancer is a heterogeneous disease and intercellular vesicular communication via sEVs may add to the complexity of the disease. The sEVs released by breast cancer cells can survive in acidic and hypoxic environments and deliver procancerous proteins and transcripts to their target cells. This induces a range of cellular responses within their target cells, to promote breast cancer development, progression, metastasis, and resistance toward therapy. We have previously reported a correlation between intracellular granular NPNT staining pattern and decreased survival of breast cancer patients. This granular staining pattern could represent NPNT-containing MVBs in tumor cells. Similar NPNT-positive granules have been observed in MMTV-PyMT tumor tissues and lung metastases from NPNT-expressing 66Cl4 mouse breast cancer cells. Furthermore, we have shown that NPNT is localized in sEVs isolated from 66Cl4 cells overexpressing NPNT and that the localization of NPNT in sEVs is not dependent on the interaction of NPNT with the integrins. Here we examine the sEVs isolated from the supernatant of the 66Cl4 cells overexpressing either wild-type NPNT (66Cl4-NPNT) or NPNT mutated in the integrin binding sites (66Cl4-NPNT-RGE and 66Cl4-NPNT-RGEAIA). As a control, 66Cl4 cells containing an empty vector (EV) were used (66Cl4-EV). We report several proteins that are differentially packed in sEVs derived from 66Cl4-NPNT cells compared to sEVs derived from 66Cl4-EV cells. We have identified NPNT of 80, 60, and 20 kDa when analyzing whole-cell lysates of 66Cl4-NPNT cells. The 80 kDa NPNT is the highly glycosylated form of the less glycosylated 60 kDa NPNT. We show here for the first time that a 20 kDa truncated form of NPNT is highly concentrated in sEVs.

2. MATERIALS AND METHODS

2.1. Cell Culture

The gene for NPNT was cloned and V5-tagged in the C-terminal, and the integrin-binding motifs were mutated and expressed in the mouse breast cancer cell line 66Cl4, as previously described.

Four different variants of 66Cl4 cell lines were created that harbored either an empty vector (66Cl4-EV), expressed wild-type NPNT (66Cl4-NPNT), NPNT where the RGD integrin-binding motif was mutated (66Cl4-RGE), or where both the RGD and EIE-integrin binding motifs were mutated (66Cl4-RGE-AIA). All 66Cl4 variants were cultured in (1x) minimum essential medium α (Thermo Fisher Scientific, Cat. No. 22561021), supplemented with 10% fetal calf serum, 1% penicillin-streptomycin, and 1 M HEPES buffer (Thermo Fisher Scientific, Cat. No. 15630080).

2.2. Isolation of Extracellular Vesicles

Fetal calf serum (FCS) was depleted of extracellular vesicles using serial centrifugation and standard protocols. The 66Cl4 cell variants were grown in vesicular-free media for 3 days to reach approximately 80% confluency. The collected supernatant was centrifuged at 500 g for 10 min to remove cellular debris. The supernatant was carefully transferred and ultra centrifuged at 12 000 g for 20 min using a Beckman 70Ti rotor to isolate microvesicles (MVs) (pellet fraction 1). The supernatant was transferred to clean tubes and repelleted by ultracentrifugation at 100 000 g for 70 min, to isolate small extracellular vesicles (sEVs) (pellet fraction 2). PBS was used to wash both fractions of MVs and sEVs.

2.3. Preparation for Scanning Electron Microscopy (SEM)

Cultured cells grown on a Thermonox coverslip (Chemtectnik AS, Cat. No. 72280) were fixed with a solution of 2.5% glutaraldehyde (GA) (EM grade distillation purified, Chemtectnik AS, Cat. No. 16310) with 2% paraformaldehyde (PFA) (EMPROVE, VWR part of Avantor, Cat. No. 104005) in 0.1 M HEPES buffer for 2–4 h at room temperature. Cells were washed in 0.1 M HEPES buffer, subsequently dehydrated using increasing ethanol concentrations (25, 50, 70, 90, 90%) for 5 min each, followed by drying using hexamethydisiloxane (HMDS) (50% and 2 × 100%, diluted in absolute ethanol) for 20 min each, and transferred to a desiccator to keep the samples dry. The dried samples were mounted on aluminum pins with double-sided carbon tape and sputter coated (Polaron) with 30 nm of gold/palladium. Samples were examined using a scanning electron microscope (Teneo SEM, Thermo Fisher Scientific) at a voltage of 5 kV.

2.4. Preparation for Transmission Electron Microscopy (TEM)

MVs and sEVs isolated and purified as described above were resuspended and fixed in cold 2% PFA in PBS. Droplets of MVs and sEVs were put on Formvar-carbon coated copper grids (200 mesh) for 5 min, fixed in 2.5% GA in 0.1 M Sorenson’s phosphate buffer for 10 min, washed in reverse osmosis (RO) water, stained with 2% uranyl acetate (UA) in Milli-Q water for 4 min, and embedded in a solution of 2% UA and 3% poly(vinyl alcohol) (diluted 1:10) for 10 min. The excess liquid on the grids was then removed with filter paper (hardened) and air-dried. Grids were examined using a transmission electron microscope (JSM-1011 TEM, JEOL), at 100.000× magnification and a voltage of 80 kV. Images were captured with a Morada digital camera with TEM software (BoRAS).

2.5. Uptake of Small Extracellular Vesicles

The pellet of sEVs was labeled with PKH26 (Sigma, MINI26), a dye taken up by the lipid membrane of sEVs. The 66Cl4-EV cells supplemented with sEVs from 66Cl4-EV and 66Cl4-NPNT cells were fixed with 4% PFA for 4 h. Images were captured using a multichannel fluorescence imaging system (Thermo Fisher Scientific, EVOS).

2.6. Immunoblotting

Protein lysates were prepared using a mix of RIPA buffer (Thermo Fisher Scientific, Cat. No. 89901) and HALT phosphatase inhibitor single-use cocktail (Thermo Fisher Scientific, Cat. No. 78428). In some experiments, cells were pretreated with a final concentration of 100 nM of bafilomycin A1 (Sigma, Cat. No. SML1661) or 1× eBiocience protein transport inhibitor cocktail (PTI) (Invitrogen, Cat. No. 00-4980-93) for ≤6 h. To test the effect of GM6001 (Abcam, Cat. No. ab120845) on NPNT, cells were exposed to 10 μM of GM6001 for 24 h. To investigate the post-translational modification of NPNT, whole cell lysates were digested with different deglycosylation enzymes as per the manufacturer’s protocol (New England Biolabs, Cat. Nos. P0704, P0733, and P0720). Samples equivalent to 30 μg each were loaded on a 10% Bis-Tris gel (Invitrogen). PVDF membranes were incubated with anti-V5 (CST, 13202) (1:1000) overnight at 4 °C. Unspecific binding was prevented by preincubating membrane in 5% BSA for 1 h at room temperature. Anti-GAPDH (Abcam, Cat. No. ab9484) (1:5000) and Anti-Alix (CST, Cat. No. 2171) were used as markers for whole cell lysate and sEVs, respectively. Pelleted fractions were checked for several other vesicular markers as previously reported. HRP linked secondary antibody anti-mouse.
(Dako, Cat. No. P0447) (1:5000) or HRP-linked anti-rabbit (Dako, Cat. No. P0399) (1:5000) were used. SuperSignal West femto substrate (Pierce, Cat. No. 34096) with an Odyssey FC system (LI-COR Biosciences) was used for image analysis.

2.7. Immunoprecipitation and Gel Electrophoresis

V5-tagged NPNT was pulled down from 7 mg of whole cell lysates using 100 μL of anti-V5 (CST, Cat. No. 13202) (1:50) coated Dynabeads protein G (Thermo Fisher Scientific, Cat. No. 10003D) for 2 h at 4 °C. The beads were washed twice in PBS and proteins were eluted in LDS sample buffer (Invitrogen) for gel electrophoresis. 1D-PAGE of eluted proteins was performed in 10% NuPAGE Noves Bis-Tris gels using MOPS buffer. Proteins were visualized using SimplyBlue gel stain, and three bands (20, 60, and 80 kDa) were excised from the gel.

2.8. In Gel Digestion and Mass Spectrometry

SimplyBlue-stained protein bands corresponding to 80, 60, and 20 kDa, respectively, were manually cut out from the gel, and in-gel tryptic digestion was performed as previously described. After desalting, peptides were dried down in a SpeedVac centrifuge and resuspended in 0.1% formic acid. The peptides were analyzed on a LC–MS/MS platform consisting of an Easy-nLC 1000 UHPLC system (Thermo Fisher Scientific) interfaced with an LTQ-Orbitrap Elite hybrid mass spectrometer (Thermo Fisher Scientific) via a nanospray ESI ion source (Proxeon, Odense). Peptides were injected into a C-18 trap column (Acclaim PepMap100, 75 μm i.d. × 2 cm, C18, 3 μm, 100 Å, Thermo Fisher Scientific) and further separated on a C-18 analytical column (Acclaim PepMap100, 75 μm i.d. × 50 cm, C18, 2 μm, 100 Å, Thermo Fisher Scientific) using a multistep gradient with buffer A (0.1% formic acid) and buffer B (CH3CN, 0.1% formic acid): From 0 to 6% B in 5 min, 6–12% B in 39 min, 12–20% B in 80 min, 20–28% B in 9 min, 28–40% B in 4 min, 40–100% B in 1 min, 100% B in 9 min, 100–0% B in 1 min, and 10 min with 100% A. The flow rate was 250 nL/min. Peptides eluted were analyzed on the LTQ-Orbitrap Elite hybrid mass spectrometer operating in positive ion and data-dependent acquisition mode using the following parameters: electrospray voltage 1.9 kV, CID fragmentation with normalized collision energy 35, and automatic gain control parameters: electrospray voltage 1.9 kV, CID fragmentation parameters: electrospray voltage 1.9 kV, CID fragmentation 250 nL/min. Peptides eluted were analyzed on the LTQ-Orbitrap Elite hybrid mass spectrometer operating in positive ion and data-dependent acquisition mode using the following parameters: electrospray voltage 1.9 kV, CID fragmentation with normalized collision energy 35, and automatic gain control parameters: electrospray voltage 1.9 kV, CID fragmentation parameters: electrospray voltage 1.9 kV, CID fragmentation internal calibration target value of 1E6 for Orbitrap MS and 1E3 for MS/MS scans. Each MS scan (m/z 300–1600) was acquired at a resolution of 120 000 fwhm, followed by 20 MS/MS scans triggered for intensities above 500, at a maximum ion injection time of 200 ms for MS and 120 ms for MS/MS scans.

Proteins were quantified by processing MS data using MaxQuant v 1.5.8.3.35 Preview 2.3.5 (Protein Metrics Inc.) was used to inspect the raw data to determine optimal search criteria. The following search parameters were used: enzyme specified as trypsin with a maximum of two missed cleavages allowed; mass tolerance set to 20 ppm; oxidation of methionine and deamidation of asparagine and glutamine as dynamic post-translational modification, and carbamidomethylation of cysteine as a fixed modification. These were imported in MaxQuant, which uses m/z and RT values to align each run against each other sample with 1 min window match-between-run function and 20 min overall sliding window using a clustering-based technique. These are further queried against the UniProtKB/Swiss-Prot database (Release April 2017 Mouse proteome with isoforms; 59684 sequences and MaxQuant’s internal contaminants database) using Andromeda built into MaxQuant. Both protein and peptide identifications FDR were set to 1%; thus, only peptides with high confidence were used for final protein group identification. Peak abundances were extracted by integrating the area under the peak curve. Each protein group abundance was normalized by the total abundance of all identified peptides for each run and protein by calculated median summing all unique and razor peptide ion abundances for each protein using a label-free quantification algorithm with minimum peptides ≥1. Protein group abundances were imported and analyzed using R software. Given the structure of the data, the statistical analysis was performed using a two-way analysis of variance (ANOVA) in order to consider the levels of variance at batch as well as test groups. Data were log 2-transformed before the analysis. Proteins were considered identified if they were quantified in at least 75% of the biological replicates. Noise with a standard deviation of 0.01 was added to compensate for the missing values followed by principal component analysis to find and remove the batch effect.

Table S1 of the Supporting Information (SI) contains normalized log 2 intensity counts and corresponding values after the removal of the component. Perseus plugins are used to carry out these steps. The latter values are only presented for those which have permutation based FDR < 5% with S0 of 0.1. These are computed over Z-scored values with their medians, namely, the row representing the proteins intensities and the column representing the samples are shifted by the median of
the row and scaled by the standard deviation. Pearson’s correlation coefficients were calculated in both dimensions over these values, which were further used for hierarchical clustering using an average linkage procedure (Figure 2d).

2.9. Ingenuity Pathway Analysis

In order to characterize the enrichment of functional category for selected protein groups (Table S1, SI, list of protein groups with permutation based FDR < 5% and S0 of 0.1), the
two broad clusters were mapped to the canonical pathways database of Ingenuity Pathway Analysis (IPA) (Qiagen). The p-values based on Fisher’s exact test over categorical distribution are corrected using Benjamini Hochberg procedure. They represent the statistical significance of the identified canonical functions, namely, the lower the p-value, the higher the proportion of identified proteins, overlapping with the particular canonical function.
3. RESULTS AND DISCUSSION

3.1. Overexpression of NPNT in 66cI4 Cells Does Not Affect the Secretion or Uptake of sEVs

To visualize the surface morphology and sEVs of 66cI4 cells, we utilized SEM. Several spherical particles in the nanometer range were observed on/around the cell surface (Figure 1a), suggesting the presence of extracellular vesicles. To confirm the presence of extracellular vesicles, we used differential ultracentrifugation and isolated extracellular vesicles from the supernatant of the 66cI4 cells, where pellet fraction 1 was collected upon spinning at 12,000 g for 20 min and pellet fraction 2 was collected upon spinning at 100,000 g for 70 min. The isolated extracellular vesicles in the two pellets were characterized by TEM (Figure 1b), which is considered to be a standard tool for characterizing extracellular vesicles. Double membrane vesicles were identified in both pelleted fractions 1 and 2. Fraction 1 displayed a homogenous population of vesicles within the range 100–1000 nm, characteristic of MVs. Fraction 2 displayed vesicles within the range of 30–100 nm in diameter, characteristic of sEVs (Figure 1b). Overexpression of NPNT in 66cI4 cells did not alter the morphology nor the size of the MVs and sEVs, as observed by TEM analyses. The ability of sEVs to exert a functional effect mainly depends on their internalization and subsequent release of its content in recipient cells. We have previously detected NPNT protein in sEVs derived from 66cI4-NPNT cells. To investigate if the presence or absence of NPNT in the sEVs affects its uptake, we treated sEVs derived from 66cI4-EV and 66cI4-NPNT cells with PKH26, a lipophilic dye, for 4 h prior to fixing and thorough washing of cells with PBS. PKH26-labeled vesicles were observed as red dots on and most likely within 66cI4-EV cells (Figure 1c), indicating that these vesicles are taken up or are bound to the surface of 66cI4-EV cells irrespective of their source and content.

3.2. Overexpression of NPNT in 66cI4 Cells Alters the Content of sEVs

To investigate the protein composition of sEVs upon NPNT overexpression, shotgun MS was performed on lysed sEVs derived from 66cI4-EV and 66cI4-NPNT cells. The Venn diagram (Figure 2a) shows the number of proteins identified in sEVs isolated from 66cI4-EV and 66cI4-NPNT cells. A total of 1754 proteins were identified to be common between the sEVs isolated from both cell lines. As expected, several well-known tumor and vesicular markers were detected in this common fraction between sEVs of both cell lines (Figure S1a, S1). A 62% similarity to the previously reported sEVs derived from parental 66cI4 cells was observed. IPA analyses suggest that the majority of proteins in our data came from the cytoplasmic fraction of the cell (Figure 2b) and were enzymatic in nature (Figure 2c). NPNT was detected as one of the 276 proteins found only in sEVs derived from 66cI4-NPNT cells [Figure 2a and Table S1 (SI)]. Several kinases known for their role in tumor development, such as mTOR and JAK, were detected only in sEVs from 66cI4-NPNT cells (Table S1, SI). Ceruloplasmin, a known biomarker for breast cancer, was only present in sEVs from 66cI4-NPNT cells. Two major clusters were identified when protein IDs passing a false discovery rate of 5% were clustered over Z-scores [Figure 2d and Table S1 (SI)]. NPNT is known to bind to different types of integrins, and IPA analyses showed that several proteins identified in sEVs derived from 66cI4-NPNT cells are involved in integrin signaling pathways and are predicted to increase proliferation, cell survival, and organization of cytoskeleton/cytoplasm (Figure 2e). Several other studies have shown that a change in a single oncprotein/oncopogene can alter the contents of vesicular cargo and further incorporate tumor-promoting proteins in sEVs. Taken together, the proteins enriched in sEVs derived from NPNT-expressing breast cancer cells might represent markers that are involved in mechanisms affecting breast cancer progression and metastasis.

3.3. Truncated Form of NPNT Is Concentrated in sEVs Isolated from 66cI4-NPNT Cells

The predicted molecular weight of mouse NPNT (561 amino acids) is 61 kDa. Our immunoblot analysis of 66cI4 cells overexpressing full length NPNT with a C-terminal V5-tag showed three bands of approximately 20, 60, and 80 kDa (Figure 3a). Interestingly, immunoblotting of lysed sEVs revealed only two bands, one at 20 and 80 kDa (Figure 3b). At the same time there was an increase in the relative amount of the 20 kDa NPNT in sEVs compared to the 80 kDa NPNT (Figure 3b). This could indicate that there is a selective sorting process involved in the vesicular packaging of NPNT. This phenomenon of the 20 kDa NPNT being concentrated was seen in both sEVs (Figure 3b) and MVs (Figure S1b, SI). Of note, we have observed that the quality of immunoblot is compromised upon freeze-thaw cycles of sEVs, and this might explain the difference in intensities in the 20 kDa bands in parts b and e of Figure 3.

To investigate whether the different apparent sizes of NPNT were due to post-translational modifications, we pulled-down NPNT from the whole cell lysates using V5 antibody and analyzed it by MS. Six unique NPNT peptide sequences were detected in the 80 kDa NPNT (Table 1), covering almost the full span of the protein (Figure 3c). This verifies that the 80 kDa NPNT is a full-length version of the protein. Four unique peptides matched the 60 kDa NPNT, including the first peptide (amino acids 28–37), indicating that the 60 kDa NPNT is most likely also a full-length version of the protein. One unique NPNT-derived peptide was detected in the 20 kDa NPNT (amino acids 446–459). This indicates that the 20 kDa NPNT is mostly harboring the MAM domain of the C-terminal part of the protein. Further, to identify whether post-translational modifications such as glycosylation contribute to differences between the three NPNT bands, whole cell lysates and sEVs were treated with PNGase F, O-glycosidase, or α-2-3,6,8 neuraminidase. These enzymes remove N- and O-linked glycans with or without a sialic acid cap, thereby reducing the apparent molecular size of NPNT on immunoblots. Upon PNGase F treatment (-N), a reduction in the molecular size was observed for both the 60 and 80 kDa bands in whole cell lysates (Figure 3d) and for the 80 kDa NPNT of sEVs (Figure 3e). There was no shift in any
of the bands upon treatment with O-glycosidase (-O) alone. However, cotreatment with O-glycosidase and α2-3,6,8 neuraminidase (-S) resulted in a major band shift in the 80 kDa NPNT for both cell and sEVs lysates. This suggests that the 80 kDa NPNT has O-glycosylation but with a sialic acid cap. A similar glycosylation pattern of NPNT secreted by osteoblasts has previously been reported.56 A high degree of N- and O-glycosylation is predicted in the region between the EGF repeats and the MAM domains.52 Aberrant glycosylation patterns of secreted glycoproteins are known to contribute in tumor development and progression.57,58 It remains to be investigated whether the glycosylation pattern of NPNT would influence breast cancer progression. The glycosylation patterns are also crucial for sorting of proteins into extracellular vesicles.59−61 Our results indicate that the 60 kDa band is far less glycosylated than the 80 kDa band, while the 20 kDa band is not glycosylated. The heavily glycosylated 80 kDa NPNT, containing both N- and O-glycosylations is recruited into sEVs (Figure 3e), while the less glycosylated 60 kDa NPNT is not detected in either sEVs or MVs. N-Glycosylations are mainly added when proteins transition through the endoplasmic reticulum,62 while O-glycosylations are generally initiated in Golgi.63

Figure 4. Intracellular protein trafficking controls the protein level of the truncated form of NPNT. (a) Immunoblotting for V5-tagged NPNT was used to detect differential expression of NPNT when 66cl4-overexpressing cells were exposed to 100 nM BaF1 or 1X PTI. (b) Immunoblotting was used to detect differential expression of V5-tagged NPNT in 66cl4-RGE cells after exposure to 100 nM BaF1 or 1X PTI for 6 h. (c) Immunoblotting for V5-tagged NPNT using whole cell lysates of 66cl4-RGE-AIA cells after exposure to 100 nM BaF1 or 1X PTI for 6 h. Results here are presented in terms of a fold change after normalizing with GAPDH. Quantification of optical density represents the mean of at least three independent experiments.
3.4. Intracellular Protein Trafficking Controls the Protein Levels of the Truncated NPNT

The intracellular protein trafficking pathways are often manipulated in cancer cells, and a better understanding of this aspect is important for developing therapeutic interventions. To investigate the intracellular NPNT trafficking, we employed a mixture of brefeldin A and monensin, to block transport from the endoplasmic reticulum to the Golgi apparatus, and bafilomycin A1 (BafA1), which inhibits degradation of endocytosed "cargo" by neutralizing the lysosomes (Figure 5c). Hence, any

Figure 5. Matrix metalloproteinases are involved in proteolytic processing of full-length NPNT. (a) Immunoblotting for V5-tagged NPNT after exposure to 10 μM GM6001 for 24 h. Results here are presented in terms of a fold change after normalizing with GAPDH. Quantification of the optical density represents the mean of at least three independent experiments. (b) Illustration showing the structural differences in NPNT at 80, 60, and 20 kDa. The 80 kDa NPNT is the full-length protein having a putative signal peptide, the EGF repeats, glycosylation sites, integrin binding motifs, and the MAM domain. The 60 kDa NPNT is also a full-length protein but with fewer attached glycans, as opposed to the 80 kDa NPNT. The 20 kDa NPNT is the truncated form of NPNT harboring mainly the MAM domain and the C-terminal V5-tag. (c) Illustration showing the probable route of NPNT secretion, endocytosis, degradation, or selective sorting into extracellular vesicles. Upon glycosylation at the endoplasmic reticulum and Golgi, NPNT is secreted via secretory vesicles. The secreted NPNT probably interacts with cell surface receptors such as integrins or is cleaved by matrix metalloproteases (MMPs). The cleaved NPNT adhering to the cell surface could further be taken up and be released as exosomes or degraded by lysosomes, depending on the fate of multivesicular bodies.
endocytosed protein, including NPNT, ends up being accumulated in the cytoplasm. 66cl4-EV and -NPNT cells were exposed to 100 nM BafA1, and whole cell lysates were harvested and analyzed by immunoblotting for the V5-tag. By inhibiting degradation of endocytosed proteins, there was a significant accumulation of the 20 kDa NPNT (Figure 4a) and a slight increase in 80 kDa NPNT. 66cl4-EV and -NPNT cells were exposed to 1× protein transport inhibitor cocktail (1× PTI), and whole cell lysates were harvested and analyzed by immunoblotting for the V5-tag. Inhibiting the secretion resulted in an accumulation of the 60 kDa band of NPNT, with a corresponding reduction of the 80 and 20 kDa bands (Figure 4a). Cells expressing the mutated forms of NPNT showed similar changes when exposed to PTI or BafA1 (Figure 4b,c). Taken together, these results indicate that the 60 kDa NPNT is not a secreted version of NPNT, but rather the “native” protein. This could also explain the limited amount of glycosylations present in the 60 kDa NPNT (Figure 3a) and its absence in the sEVs (Figure 3b). Both 80 and 20 kDa accumulate in the cytoplasm upon BafA1 treatment, indicating that they are both endocytosed from the extracellular milieu. The 80 kDa NPNT is most likely the secreted full-length version of NPNT, having both N- and sialic acid capped O-glycosylations.

3.5. Matrix Metalloproteinases (MMPs) Are Involved in Proteolytic Processing of Full-Length NPNT

Having identified the truncated form of NPNT, we further analyzed whether proteases such as MMPs are involved in this processing by utilizing GM6001, a broad-spectrum MMP inhibitor. 66cl4-NPNT cells were treated with 10 μM GM6001 for 24 h and whole cell lysates were analyzed by immunoblotting for the V5-tag. A significant reduction in the relative amounts of 20 kDa NPNT was observed (Figure 5a). We already know that the extracellularly located 80 kDa NPNT can be endocytosed by the cells. When MMP cleavage is inhibited, the amount of endocytosed 20 kDa NPNT is reduced. This indicates that the 20 kDa NPNT is a cleaved form of NPNT and that this cleavage is at least in part mediated by MMPs. MMPs can cleave proteins intracellularly and extracellularly.66 Several metalloproteinases are also packed inside extracellular vesicles, capable of further altering the vesicular cargo by proteolytic processing.67,68 Cleaved proteins packed in tumor-derived extracellular vesicles are reported to be biologically active.55,69 Further investigation is needed to identify if the cleaved NPNT packed in sEVs has a biological significance. Our study on NPNT trafficking in 66cl4 cells is the first to report a truncated form of NPNT at 20 kDa, mainly harboring the MAM domain part of the protein (Figure 5b). The exact MMP cleavage site and the identity of the MMPs capable of cleaving NPNT remain to be determined.

Though there are several challenges while analyzing the vesicular proteome using MS, the composition of sEVs derived from 66cl4-NPNT cells helps us to picture the molecular fingerprint of exosomes in cells with high NPNT levels. Results from the current study are summarized in Figure 5c and show that the native form of NPNT gets glycosylated and modified at the endoplasmic reticulum and Golgi and that the secreted full-length NPNT (80 kDa) probably gets cleaved by MMPs. Both the highly glycosylated NPNT (80 kDa) and the truncated form of NPNT (20 kDa) are perhaps endocytosed/pinocytosed and get incorporated into the ILVs within a MVB. These ILVs packed with NPNT inside a MVB can either be released as exosomes or be degraded by lysosomes.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jproteome.8b00859.

Table S1 in spreadsheet format (XLSX)

Figure S1, a list of well-known tumor and vesicular markers detected in sEVs isolated from 66cl4-EV and 66cl4-NPNT cells and immunoblotting for V5-tagged NPNT using lysates made from the microvesicles isolated from the supernatant of 66cl4-EV and 66cl4-NPNT cells, and Table S1, a list of proteins identified in sEVs isolated from 66cl4-EV and 66cl4-NPNT cells (PDF)

AUTHOR INFORMATION

Corresponding Author

*E-mail: jimita.toraskar@ntnu.no. Mobile phone:+47-99673377.

ORCID

Jimita Toraskar: 0000-0002-5404-061X
Gunbjørg Svineng: 0000-0002-5947-5647

Notes

The authors declare no competing financial interest.

The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the dataset identifier PXD010999.70,71

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REFERENCES

(5) Palmulli, R.; van Niel, G. To be or not to be... secreted as exosomes, a balance finely tuned by the mechanisms of biogenesis. Essays Biochem. 2018, 62 (2), 177–191.


(13) Bobrie, A.; Colombo, M.; Krumichev, S.; Raposo, G.; Thery, C. Diverse subpopulations of vesicles secreted by different intracellular mechanisms are present in exosome preparations obtained by differential ultracentrifugation. *J. Extracell. Vesicles* 2012, 1, 18397.


(26) Zhao, X.; Wu, Y.; Duan, J.; Ma, Y.; Shen, Z.; Wei, L.; Cui, X.; Zhang, J.; Xie, Y.; Liu, J. Quantitative proteomic analysis of exosome protein content changes induced by hepatitis B virus in Huh-7 cells using SILAC labeling and LC-MS/MS. *J. Proteome Res.* 2014, 13 (12), 5391–402.


(29) www.exocarta.org.


(57) Munkley, J.; Elliott, D. J. Hallmarks of glycosylation in cancer. Oncotarget 2016, 7 (23), 35478−89.


(71) PRIDE. https://www.ebi.ac.uk/pride/archive/projects/PXD010999.