Tumour microvesicles contain retrotransposon elements and amplified oncogene sequences

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Tumour cells release an abundance of microvesicles containing a selected set of proteins and RNAs. Here, we show that tumour microvesicles also carry DNA, which reflects the genetic status of the tumour, including amplification of the oncogene \textit{c-Myc}. We also find amplified \textit{c-Myc} in serum microvesicles from tumour-bearing mice. Further, we find remarkably high levels of retrotransposon RNA transcripts, especially for some human endogenous retroviruses, such as LINE-1 and Alu retrotransposon elements, in tumour microvesicles and these transposable elements could be transferred to normal cells. These findings expand the nucleic acid content of tumour microvesicles to include: elevated levels of specific coding and non-coding RNA and DNA, mutated and amplified oncogene sequences and transposable elements. Thus, tumour microvesicles contain a repertoire of genetic information available for horizontal gene transfer and potential use as blood biomarkers for cancer.
Increasing knowledge of the genetic and epigenetic changes occurring in cancer cells provides an opportunity to detect, characterize and monitor tumours by analysing tumour-related nucleic acid sequences and profiles. Cancer-related changes include specific mutations in gene sequences, up- and downregulation of messenger RNA (mRNA) and microRNA (miRNA) expression, mRNA splicing variations and changes in DNA methylation patterns, as well as amplification and deletion of genomic regions. Brain tumours comprise a variety of phenotypic and genetic subtypes and knowing the expression/mutational profile of individual cancers is critical for personalized medicine as many drugs target specific pathways affected by the genetic status of the tumours. Detection of genetic biomarkers in tumour patient blood samples is challenging because of the need for high sensitivity against a background of normal cellular DNA/RNA found circulating in blood. Microvesicles released by tumour cells into the circulation can provide a window into the genetic status of individual tumours.5,9

Many types of cancer cells release an abundance of small membrane-bound vesicles, which have been observed on their surface in culture12,16. These microvesicles are generated and released through several processes and vary in size (30 nm to 1 μm in diameter) and content.11 Microvesicles can bud off the plasma membrane of cells, much like retrovirus particles,8,21 be released by fusion of endosomal-derived multivesicular bodies with the plasma membrane,13 or be formed as apoptotic bodies during programmed cell death.4,16 In addition, defective retrovirus particles derived from human endogenous retroviral (HERV) elements may be found within microvesicle populations.42 Different microvesicle types often co-purify and given the evolving nomenclature for these various types, our study has focused on collective microvesicle populations of <0.22 μm in diameter.

Microvesicles from various cell sources have been extensively studied with respect to protein and lipid content.6,47 They also contain a select set of cellular RNAs and mitochondrial DNA8,18,19,20 and may facilitate the transfer of genetic information between cells and/or act as a ‘release hatch’ for DNA/RNA/proteins that the cell is trying to eliminate. Both mRNA and miRNA in microvesicles can be functional following uptake by recipient cells8,19,21–24 and it has also been shown that apoptotic bodies can mediate horizontal gene transfer between cells.12

Increased transcription of retrotransposon elements in the human genome has been noted in a number of cancer cell types. These repetitive elements constitute almost 50% of the human genome and include: half a million LINE-1 (L1) elements, of which about 100 are transcriptionally active and encode proteins involved in retrotransposition, including reverse transcriptase (RT) and integrase; a million Alu elements, which depend on L1 functions for integration; and thousands of provirus HERV sequences, some of which contain near-to-full-length coding sequences.45,46 Increased expression of retrotransposon elements in cancer seems to result in part from overall hypomethylation of the genome, which is also associated with genomic instability.27,28 and tumour progression.28,30 Interestingly, increased expression of L1 and HERV RNA and proteins, as well as formation of retrovirus-like particles, have been reported in tumour tissue from breast cancer, melanoma and germ cell carcinoma.31–33 Retrotransposon RNA/proteins, as well as antibodies against HERV proteins and virus-like particles, are also found in blood of some cancer patients.34–36

In this study, we examined the nucleic acid content of microvesicles released by cells in culture and tumours in vivo in: glioblastoma (GBM), the most common and malignant brain tumour in adults; medulloblastoma, the most common and malignant tumour in children with frequent amplification of c-Myc37 and atypical teratoid rhabdoid tumour (AT/RT), a high-grade malignant tumour in children.38 We also included a peripheral tumour: malignant melanoma, one of the most common cancers, which can metastasize to the brain.19 Epidermoid carcinoma tumour cells were used as a control for the in vivo study, as they have amplified epidermal growth factor receptor (EGFR), but not c-Myc genes.39

We examined the nucleic acid content of microvesicles released by cells in culture and tumours in vivo. In addition to RNA, tumour microvesicles contain single-stranded DNA (ssDNA; exoDNA, excreted out of cells in microvesicles), including both genomic and cDNA, as well as high levels of transposable elements. Medulloblastoma cells with the amplified oncogene c-Myc had higher DNA/RNA levels of this oncosine in the microvesicles compared with cells without c-Myc amplification. Elevated human c-Myc RNA was also found in microvesicles isolated from mice-bearing tumours with amplified c-Myc. Further, an abundance of retrotransposon RNA, including HERV, L1 and Alu sequences was found in tumour-derived microvesicles. Tumour microvesicles contain amplified genomic DNA (gDNA), cDNA and retrotransposon elements that may have a role in genetic communication between cells and provide a potential source of tumour biomarkers.

Results

Cultured cells release an abundance of microvesicles. We characterized the size distribution and amount of microvesicles released from tumour cells and normal fibroblasts in culture using Nanosight LM10 nanoparticle tracking analysis (Fig. 1). Medulloblastoma cells were found to release more microvesicles per cell than the other cell types analysed (13,400–25,300 per cell per 48 h for medulloblastomas and 7,000–13,000 per cell per 48 h for the GBM and melanoma cells). Normal human fibroblasts released 3,800–6,200 per cell per 48 h, were of low passage and grew with similar rates as the tumour lines in culture, but of larger size and hence greater surface area per cell. Levels of RNA in microvesicles (exoRNA) from tumour cells as compared with normal fibroblasts were 120- to 310-fold higher for medulloblastoma cells and 2.8- to 6.5-fold higher for GBM cells, with melanoma cells having similar levels of exoRNA compared with fibroblasts despite shedding more than twice as many microvesicles. Thus, medulloblastoma tumour cells, in particular, release abundant microvesicles with a high content of exoRNA.

Characterization of RNA and DNA in microvesicles. Isolated microvesicles were treated extensively with DNase before nucleic acid extraction to reduce the chance of external DNA contamination (external RNase treatment did not affect the RNA yield, indicating no external RNA). After microvesicle lysis and nucleic acid purification, the exoRNA fraction was DNase treated and the exoDNA fraction was RNAse treated. The RNA profile varied among cell types and culture conditions, but in general, RNA with intact 18S and 28S ribosomal peaks was isolated from microvesicles (Supplementary Fig. S1). ExoDNA was more abundant in microvesicles from tumour as compared with normal cells, and appeared to be primarily single stranded. When exoDNA from medulloblastoma tumour cells (D384) was analysed using a double-stranded DNA detection chip, no DNA was detected (Supplementary Fig. S2a). However, when this exoDNA was subjected to second strand synthesis, this same chip detected abundant double-stranded DNA (Supplementary Fig. S2b). The same experiment was performed on GBM cells (GBM 20/3) with similar results. The predominance of ssDNA in exoDNA was also confirmed by its complete sensitivity to S1 exonuclease digestion (Supplementary Fig. S3).

C-Myc oncogene amplification is reflected in exoRNA and exoDNA. The levels of c-Myc amplification were measured at the genomic level (gDNA) by quantitative PCR (qPCR; Fig. 2a). All three medulloblastoma cell lines had a significant amplification of
c-Myc sequences (16- to 34-fold) compared with normal fibroblasts and other tumour cell types. RNA and DNA were extracted from microvesicles shed by these cell lines and measured by quantitative reverse transcription PCR (qRT-PCR) and qPCR, respectively, using primers in exon 3 (Supplementary Table S1). The values for c-Myc sequences were normalized to glyceraldehyde 3-phosphate dehydrogenase (GAPDH), a housekeeping gene constitutively expressed in cells and found in exoRNA\(^3\) and here in exoDNA. Microvesicles from all medulloblastoma cell lines showed elevated levels of c-Myc sequences, both for exoRNA (8- to 45-fold) and exoDNA (10- to 25-fold), compared with microvesicles from fibroblasts and tumour cells with diploid c-Myc copy numbers (Fig. 2b,c). Also, using primers that span a full intron, we successfully detected a 1.6 kbp fragment corresponding to the unspliced c-Myc gDNA (verified by sequencing) in exoDNA from all three medulloblastoma cell lines, but not in any of the other cell lines. Furthermore, to establish that this genomic fragment of c-Myc in microvesicles was derived from a genomic amplicon, we verified the presence of elevated levels of a flanking gene, POU5F1B gene\(^4\) (PCR product also verified by sequencing) at levels matching those of c-Myc (Supplementary Fig. S4). Levels of n-Myc sequences in cellular gDNA or exoRNA were also measured by qPCR and qRT-PCR and none of the tumour types showed genomic amplification of n-Myc sequences or elevated levels of n-Myc exoRNA (Supplementary Fig. S5a,b).

The levels of c-Myc DNA quantified for gDNA and exoDNA/RNA in these medulloblastoma lines were also compared with levels estimated by 250K single-nucleotide polymorphism (SNP) analysis (Table 1; see Supplementary Fig. S5c for a representative heat map). The copy number of c-Myc was increased in medulloblastoma lines and was normal in the AT/RT tumour line. The increased levels of c-Myc exoDNA and exoRNA corresponded well to the genomic copy number estimated by 250k SNP and qPCR in the medulloblastoma cell lines. The normal diploid cell lines showed no amplification of c-Myc on exoRNA or exoDNA. To assess the potential diagnostic utility of using exoRNA to detect c-Myc amplification in tumours, human medulloblastoma cells (c-Myc amplified) and epidermoid carcinoma tumour cells (non-amplified; Fig. 3a) were grown as xenograft tumours in nude mice. Microvesicles were isolated from serum samples in tumour-bearing mice and human c-Myc exoRNA was detected in 2/5 (40%) of the medulloblastoma-bearing mice and in 0/5 of the epidermoid carcinoma-bearing mice (Fig. 3b).

Tumour microvesicles are enriched for retrotransposon elements. Microarray analysis of cellular RNA and exoRNA sequences from a low-passage GBM line indicated high

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**Figure 1 | Analysis of microvesicle profiles and RNA yields from different human cell lines.** Microvesicles were isolated from three medulloblastoma cell lines (a, D384, b, D425 and c, D458), one melanoma (d, Yumel 0106), two GBMs (e, 20/3 and f, 11/5) and two normal fibroblasts (g, HF19 and h, HF27) and measured with Nanoparticle Tracking Analysis (NanoSight). The number of particles per cell per 48 h is shown on the y axis, and the size distribution (particle diameter) on the x axis. The sum refers to the total number of particles released per cell over 48 h and the exoRNA refers to the total microvesicle RNA yield per 1×10⁶ cells per 48 h. The results are presented as the mean±s.e.m. (n=3).
transcription levels of several retrotransposon sequences as evaluated using a whole-genome array. This data is represented on MA plots as the cumulative abundance (in cells and microvesicles) of specific RNAs (x axis) and the relative ratio of these RNAs in microvesicles versus cells (y axis; Fig. 4a). The axis scale is log 2, so RNAs above 4 or below −4 on the y axis have at least 16-fold different levels in the microvesicles versus donor cells. Although RNA from DNA transposons was similar in content in cells and microvesicles (Fig. 4b), RNA from retrotransposons, for example, HERV, Alu and L1, was frequently higher in microvesicles than cells (Fig. 4c–e). This was particularly notable for the HERV sequences.

HERV-H was the most abundant and microvesicle enriched in these GBM cells, followed by HERV-C, HERV-K6 and HERV-W (Fig. 4f). As only a selected subset of transposon/retrotransposon probes are represented on the Agilent arrays, this should be viewed as a partial screen. However, this array analysis supports selective packaging of retrotransposon RNA sequences, especially of HERV, in tumour microvesicles.

As L1 and HERV-K retrotransposons, as well as Alu elements, have been implicated in tumour progression, we further assayed their levels in cellular RNA and exoRNA from tumour and normal cells by qRT-PCR (again with the caveat that the primers do not

**Table 1 | Assessment of c-Myc gene amplification levels in different cell types.**

<table>
<thead>
<tr>
<th>Method</th>
<th>c-Myc genomic copy number</th>
<th>c-Myc amount exoRNA*</th>
<th>c-Myc amount exoDNA*</th>
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<tr>
<td>D425</td>
<td>FISH†</td>
<td>&gt; 25</td>
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<tr>
<td></td>
<td>250K SNP‡</td>
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<td>8±2.0</td>
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<td>250K SNP</td>
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<td></td>
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<tr>
<td></td>
<td>qPCR</td>
<td>12±4.7</td>
<td>42±22</td>
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<tr>
<td>D458</td>
<td>250K SNP</td>
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</tr>
<tr>
<td></td>
<td>qPCR</td>
<td>17±3.0</td>
<td>45±11</td>
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<td>250K SNP</td>
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<td></td>
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<td>qPCR</td>
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<td>2.8±1.4</td>
<td>0.4±0.1</td>
</tr>
<tr>
<td>HF19</td>
<td>qPCR</td>
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</table>

*Reverse transcribed exoRNA (2.5ng) and exoDNA (10ng) were used as template for qPCR. All values were normalized to GAPDH mRNA.
†FISH, Fluorescence in situ hybridization of metaphase chromosome spread‡.
‡See representative heat map Supplementary Figure S5c.

**Figure 2 | Medulloblastomas with amplified c-Myc oncogenes have elevated c-Myc exoRNA and exoDNA in their microvesicles.**

(a) c-Myc amplification levels were quantified in genomic DNA (gDNA) from one normal human fibroblast line (HF19), one GBM line (11/5), one atypical teratoid rhabdoid tumour (AT/RT) line (NS224) and three medulloblastoma (MB) lines (D425, D458 and D384). ExoRNA and exoDNA were also isolated from their corresponding microvesicles. (b) qRT-PCR and (c) qPCR were carried out on nucleic acid from microvesicles from the same cell lines to measure exoRNA and exoDNA, respectively; c-Myc levels were normalized to GAPDH in the same preparation and expressed as fold increase relative to normal fibroblasts. In all cases, values are expressed as mean±s.e.m. (n = 3) and analysed by two-tailed t-test comparing MB lines to HF19 (*P< 0.05, **P< 0.01, ***P<0.001).

**Figure 3 | Detection of amplified c-Myc sequences in serum microvesicles from tumour-bearing mice.** Medulloblastoma (MBT; D425) and epidermoid carcinoma (ECT; A431) cells were used to generate subcutaneous tumours with and without c-Myc amplification, respectively. (a) c-Myc amplification was evaluated on all tumour samples at the RNA level after tumour resection. Values were normalized to GAPDH, presented as fold change compared with epidermoid carcinoma and shown as mean±s.e.m. (n = 3). (b) ExoRNA was extracted from serum samples from five MBT and five ECT (MBT 1–5 and ECT 1–5, respectively). c-Myc PCR product was amplified using human specific primers. Amplified DNA was resolved by electrophoresis in a 2% agarose gel and visualized with ethidium bromide staining. c-Myc is shown as an 89 bp fragment (arrow). MW, molecular weight; NTC, no template control.
Interestingly, the enrichment of the transposable elements at the exoDNA level in the medulloblastoma cell lines corresponded to high levels of endogenous RT activity in exosomes (Fig. 6d), suggesting that a fraction of exoDNA may be cDNA. By comparison, Yumel 0106 and GBM11/5 cells had very little L1 and HERV-K exoDNA in the microvesicles compared with the cells (as shown by the negative values on the bar graph in Fig. 6a,c). As the exoDNA yield in microvesicles was decreased by about 50% following inhibition of DNA replication with mimosine (Supplementary Fig. S7), it seems that some of the exoDNA may also be fragments of gDNA generated during DNA replication and mitosis.

Discussion

Previous studies have shown that microvesicles released from tumour cells into the blood stream of cancer patients contain a representation of the tumour transcriptome, including characteristically high levels of some miRNAs and miRNAs, and mutant/splice variant mRNAs, as well as tumour-related proteins. We now find that microvesicles derived from cultured tumour cells also contain high levels of exoDNA—predominantly ssDNA fragments. Interestingly, exoDNA was virtually undetectable in normal skin fibroblasts used in this study. Of the tumour lines evaluated, both exoDNA and exoRNA were highest in microvesicles from medulloblastoma cells, which had genomic amplification and high-expression levels of the c-Myc oncogene. In addition, tumour microvesicles were found to be highly enriched in certain retrotransposon RNA sequences, especially for some of the HERVs. Interestingly, some of the tumour cell types also had enriched levels of exoDNA for transposable elements (primarily L1 and HERV). We also found that exosomes have RT activity, especially from the medulloblastomas, indicating that some of the exoDNA transposable elements may represent cDNAs. It has previously been shown that reverse transcribed transposable element cDNA is normally degraded by the Trex1 protein, a 3’-exonuclease that functions as an intrinsic cell protection mechanism. If Trex1 protein is knocked out, ssDNA accumulates in the cytoplasm, especially L1, HERV and Alu ssDNA.

Because microvesicles are sticky and can bind free-floating gDNA on their surface, intact microvesicles were treated extensively with DNase before analysis of exoDNA contained within them. Apoptotic vesicles, which are typically larger than the microvesicle fraction we analysed (<0.2 μm), could also potentially contribute gDNA into the exoDNA fraction. To evaluate this contribution we incubated cells with an apoptosis-inducing agent, doxorubicin. Surprisingly, this led to a reduction in exoDNA yield, indicating that cells undergoing apoptosis shed fewer microvesicles than healthy cells (data not shown). In addition, the bar graphs of transposable element DNA (Fig. 6a–c) showed differential levels in microvesicles versus the cells from which they were derived (y≠0), indicating a different composition of exoDNA compared with nuclear DNA.

The elevated c-Myc exoDNA levels in microvesicles from medulloblastoma cells with amplified c-Myc could have several potential sources. They may arise from extra-genomic copies of the amplified c-Myc gene region, which are typically 0.1–1 Mbp and contain one or more origins of DNA replication. These amplified sequences can appear cytogenetically as small ‘double minute’ chromosomes, which can end up in the cytoplasm during mitosis or be eliminated by extrusion out of cells within micronuclei. Fragments of these double-minute chromosomes may end up in microvesicles as supported by the presence of c-Myc intronic and flanking gene sequences. In addition, ssDNA in the cytoplasm and microvesicles may arise during abnormal replication of DNA in cancer cells through faulty re-replication and accumulation of short Okazaki fragments of linear ssDNA. This potential source is supported by results from treatment with mimosine, which blocks DNA replication in late G1 and reduced exoDNA levels in a dose-dependent manner. Further,
some of the exoDNA may be derived from reverse transcription of cellular RNAs, as supported by the RT activity in microvesicles.

Retrotransposable elements can create a dynamic aspect to the genome, with new insertions of retrotransposon and cDNA sequences potentially leading to mutations, deletions, rearrangements and changes in gene expression. Genomic stability is maintained in normal differentiated cells by suppressed transcription of these retrotransposon elements, with tumour cells frequently having elevated retrotransposon expression\(^2\). The specific HERV type found in tumour microvesicles may depend on the cell of origin of the tumour. The abundance of HERV-H in microvesicles from GBM cells is consistent with it being the most active HERV in fetal brain cells with these tumours thought to derive from dedifferentiated glia cells or neuroprecursor cells\(^6,31\). The finding of high HERV-K RNAs in several tumour lines is intriguing as this is the most intact of the HERVs in the human genome\(^2\).

Other studies have noted an association between increased retrotransposon activity and tumourigenesis. In some cases, retrotransposon insertion into the genome acts as a driver mutation in tumourigenesis\(^3\). For example, L1 insertions in oncogenes have occurred in the APC gene in colon cancer\(^4\) and the MYC gene in breast cancer\(^5\). A potential role for increased retrotransposon activity in tumourigenesis is also suggested by the reduced growth of cancer cells when RT activity is inhibited\(^6\) or when translation of HERV RNAs is blocked by RNA interference\(^7\). An important question is whether retrotransposon-derived RNA and proteins in tumour microvesicles can be delivered to other cells and contribute to genomic instability. This transfer is supported by our finding of elevated HERV RNA in endothelial cells exposed to tumour microvesicles. Although new retrotransposon insertion events can prove deleterious to cells in some cases\(^9\), other events could generate cells with increased proliferative or invasive potential. For example, in murine tumours, ongoing endogenous murine leukemia virus integrations lead to genetic changes implicated in increased cell mobility\(^9\).

Serum microvesicles derived from tumour cells can provide a window into the genotype and phenotype of tumours in individual cancer patients. The exoRNA in serum microvesicles from GBM patients can be used to detect the EGFRvIII mutant/variant mRNA in tumours\(^4\) and elevated miRNAs in ovarian tumours\(^11\). Further, in the present study, we found that xenograft tumours of human medulloblastoma cells amplified for c-Myc released microvesicles into the serum with elevated expression of c-Myc.

Our present findings indicate additional tumour-specific genetic properties represented in microvesicles from tumour cells. First, the exoDNA and exoRNA content is increased in tumour microvesicles as compared with microvesicles from normal fibroblasts, especially, in the case of tumour cells with an amplified oncogene. As DNA is intrinsically more stable than RNA, this should make quantification and analysis of genomic mutations more robust and sensitive in clinical biomarker assays. Elevation of exoDNA or exoRNA may serve as biomarkers of oncogene amplification, for example, c-Myc in individuals harbouring medulloblastoma tumours. Second, increased expression of certain retrotransposon RNAs appears...
to be a unique feature of tumour cells as compared with normal adult cells. As these retrotransposon sequences are abundant in tumour microvesicles that can be harvested from body fluids, they could be useful as biomarkers. The levels of specific retrotransposon sequences, such as those for specific HERV family members, may be indicative of the cell origin of the tumour. This work expands the list of genetic elements in tumour microvesicles that can potentially be used in blood-based diagnostics for cancer and suggests new modalities of intercellular genetic communication.

Methods

Cells. Primary GLM samples were obtained from decommissioned donors (aged 65–85 years). Exosomes were isolated from samples using the ExoQuick kit (System Biosciences) and stored at −80°C. The exosomes were washed and resuspended in PBS to a concentration of 1 mg/mL. To determine the size distribution of the exosomes, samples were analyzed using the NanoSight LM10 nanoparticle characterization system (NanoSight) equipped with a blue laser (405 nm) illumination which was used for real-time characterization of the vesicles. The result is presented as the average ± s.e.m. of three independent experiments.

Xenograft tumour models. Two groups of five adult immunodeficient mice (nu/nu NCI) were injected subcutaneously in both flanks with 5×10⁶ medulloblastoma cells (line D425) or epidermoid carcinoma cells (line A375). Tumours were allowed to grow for three weeks; the mice were then killed and blood was drawn by cardiac puncture. Subcutaneous tumour mass weights at the time of euthanization were as follows: D425: 1–3.4 g; 2–1.7 g; 3–2.4 g; 4–2.9 g; 5–1.7 g and A375: 1–1.7 g; 2–1.9 g; 3–1.9 g; 4–1.9 g; 5–2.2 g. Approximately 1 ml of blood was obtained from each mouse and allowed to clot at room temperature for 15 min and then centrifuged at 1,300×g for 10 min. The sera were then filtered through a 0.22 µm filter and stored at −80°C. Samples were thawed and centrifuged for 1 h at 100,000×g to obtain microvesicles for RNA extraction, as described above. All animal procedures were performed according to guidelines issued by the Committee of Animal Care of Massachusetts General Hospital.

Estimation of gene copy number by SNP array analysis. Genomic DNA was extracted from medulloblastoma cell pellets using the PureGene DNA Extraction Kit (Genta Systems), according to the manufacturer’s instruction. To obtain signal intensities and genotypes, cDNA samples were digested, labelled and hybridized to Affymetrix 250K SNP arrays, according to the manufacturer’s protocol (Affymetrix). Signal intensities were normalized using rank invariant set normalization, and copy numbers for altered genomic regions were inferred from the MA plot (Gain and Loss of DNA) algorithm available in the GenePattern software package (http://www.genepattern.org). C-Myc copy numbers were inferred by analysing the smoothed copy number data at genomic region chr9:412.4.

Microarray comparison of transposable elements in microvesicles versus cells. RNA was extracted from microvesicles, as described above. The microarray experiments were performed by Miltenyi Biotec using the Agilent whole human genome microarray (Agilent), 4×44K (44,000 probes), two-color array following the manufacturer’s protocol. The array was performed on two different RNA preparations from primary GBM cells and their microvesicles. The microarray data has been deposited in NCBI’s Gene Expression Omnibus (GSE13470; GEO, http://www.ncbi.nlm.nih.gov/geo).

MA plots. The MA plots for the array data was generated as previously described. The log ratio of the intensities of microvesicle/cell is plotted on the y axis (M = log(Exo)−log(Cel) and the mean log expression of the two on the x axis (A = 0.5×(log(Exo)+log(Cel))).

RT activity assay. Microvesicles were lysed in RIPA buffer (50 mM Tris-HCl (pH 8); 150 mM NaCl, 2.5% sodium dodecyl sulphate, 2.5% deoxycholic acid, 2.5% N-octyl-P-4) for 20 min at 4°C. Microvesicle debris was removed by centrifugation at 14,000×g for 15 min. Proteins were quantified by Bradford assay and diluted 1:6 for each RT reaction. The RT assay was performed using the InvivoGen RT kit according to the manufacturer’s instructions. The RT reactions were performed on a 25 µl reaction, as described by the manufacturer. Fluorescence signal of the samples was measured before and after the RT incubation. The difference between the two values indicates newly synthesized DNA. Serial dilutions of SuperScript III reverse transcriptase (Invitrogen) were used to generate an RT units standard curve. The results are presented as the average ± s.e.m. of three independent experiments.

Microvesicle transfer of HERV-K. HUV-ECs were seeded in 12-well plates at a density of 1.5×10⁵ cells per well. Microvesicles were isolated from 1.2×10⁶ D384 cells, cultured in dFBS–DMEM, over a 48 h period and added to each well in a total volume of 400 µl DMEM. Mock-treated cells were incubated in 400 µl extreme-fibronectin–DMEM. The cells were then replenished with 1.5 ml DMEM (with 5% dFBS). Cells were collected and processed at the indicated time points after the microvesicle exposure and cell RNA was extracted for qRT-PCR analysis. The results are presented as the average ± s.e.m. of three independent experiments.

Statistics. Analysis of variance and Student’s t-test (two tailed).

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LINE-1 and Alu repeats in prostate adenocarcinoma and their relationship to

Rev.

search for novel human retroviruses in chronic disease.

separate packaging of DNA and RNA in apoptotic bodies during apoptosis.


Voisset, J.-L. et al. Tumour-released exosomes and their implications in cancer


Acknowledgments

We thank Ms Suzanne McDavit for skilled editorial assistance, Dr Mikkel Noerholm, Jens Magnusson and Tobias Limperg for editing of the manuscript, and Drs Tom Würdinger and David Noske in the Neuro-Oncology Research Group (NRG), Cancer Center Amsterdam, Amsterdam NL for continuous support. We also thank Dr Liliatele Russo for insightful discussions, Dr Kristan van der Vos for valuable assistance with
microvesicle quantification and Dr Kathleen Burns and Dr Jef Boeke for advice on the RT assay. This work was kindly supported by the Wenner-Gren Foundation (J.S.), Hyugens Scholarship NL (L.B.), Stiftelsen Olle Engkvist Byggmästare (J.S.), NCI CA86355 (X.O.B.), NCI CA69246 (X.O.B./J.S.), and CA141226 NCI (X.O.B.), CA141150 (X.O.B./J.S.), CA109467 (S.L.P., J.C.).

Author contributions
L.B. generated and analysed data, and was involved in experimental strategy and design, writing/editing of manuscript. R.L. generated and analysed data. L.D. was involved in the experimental design. Y.-J. C. generated and analysed amplification data and edited the manuscript. S.L.P. analysed amplification data and edited the manuscript. X.O.B. was involved in experimental strategy and design, analysis of data and writing/editing of manuscript. J.S. generated and analysed data, was involved in the experimental strategy and design and writing/editing of manuscript.

Additional information
Accession codes: The microarray data has been deposited in the Gene Expression omnibus under accession code GSE13470.

Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications

Competing financial interests: Dr Johan Skog has equity options in Exosome Diagnostics. The remaining authors declare no competing financial interest.

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