Microparticle conferred microRNA profiles - implications in the transfer and dominance of cancer traits

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Abstract

Background: Microparticles (MPs) are membrane vesicles which are released from normal and malignant cells following a process of budding and detachment from donor cells. MPs contain surface antigens, proteins and genetic material and serve as vectors of intercellular communication. MPs comprise the major source of systemic RNA including microRNA (miRNA), the aberrant expression of which appears to be associated with stage, progression and spread of many cancers. Our previous study showed that MPs carry both transcripts and miRNAs associated with the acquisition of multidrug resistance in cancer.

Results: Herein, we expand on our previous finding and demonstrate that MPs carry the transcripts of the membrane vesiculation machinery (floppase and scramblase) as well as nucleic acids encoding the enzymes essential for microRNA biogenesis (Drosha, Dicer and Argonaute). We also demonstrate using microarray miRNA profiling analysis, the selective packaging of miRNAs (miR-1228*, miR-1246, miR-1308, miR-149*, miR-455-3p, miR-638 and miR-923) within the MP cargo upon release from the donor cells.

Conclusions: These miRNAs are present in both haematological and non-haematological cancer cells and are involved in pathways implicated in cancer pathogenesis, membrane vesiculation and cascades regulated by ABC transporters. Our recent findings reinforce our earlier reports that MP transfer ‘re-templates’ recipient cells so as to reflect donor cell traits. We now demonstrate that this process is likely to occur via a process of selective packaging of nucleic acid species, including regulatory nucleic acids upon MP vesiculation. These findings have significant implications in understanding the cellular basis governing the intercellular acquisition and dominance of deleterious traits in cancers.

Keywords: Cancer, Microarray, Microparticles, MicroRNA, Multidrug resistance, Selective packaging

Background

Extracellular membrane vesicles are important vehicles of intercellular communication across numerous biological processes. MPs are typically defined by their size (0.1–1 μm in diameter) [1], exposure of phosphatidylserine (PS) and the expression of surface antigens originating from their donor cells [1-3].

MP vesiculation occurs as a cellular response to various physiological conditions including: apoptosis, senescence, cellular activation [4]; shearing stress and biochemical triggers (such as cytokines and chemotherapeutics) [5]. In the steady state the cell membrane is asymmetric in its composition with phosphatidylcholine and sphingomyelin located in the outer layer whereas phosphatidylserine (PS) and phosphatidylethanolamine (PE) present in the inner layer. This asymmetric distribution in the membrane is maintained by a group of two ATP-dependent enzymes namely flippase, floppase as well as a bidirectional ATP-independent scramblase [6-8]. Flippase specially translocates PS and PE from the outside to the inside of the bilayer membrane. Floppase transports phospholipids and cholesterol from the inner to the outer leaflet.
Floppase does not specifically act on transport of amino-phospholipids and probably works together with flippase. Scramblase whose role is thought to be the transportation of phospholipids between the two monolayers of the cell membrane, is inactive in steady state [6-8]. Following stress or under physiological conditions, an increase in intracellular calcium, a subsequent loss of phospholipid asymmetry following the inactivation of flippase and activation of floppase and scramblase, and disruption of the cytoskeletal apparatus occurs leading to MP vesculation [6,7,9]. The released MPs are enriched in PS and PE exposed on their outer surface. Consequently, MPs carry also cellular proteins, second messengers, growth factors and genetic material from their cells of origin [1,10] and comprise the major source of RNA (ribosomal RNA (rRNA), messenger RNA, (mRNA) and microRNA (miRNA) in systemic circulation [11,12].

miRNAs are highly conserved, single-stranded non-coding regulatory nucleic acids, typically 19–25 nucleotides in length. These RNAs modulate the activity of specific mRNA targets and serve as important regulators of a wide range of pathophysiological processes [13]. miRNA synthesis begins in the nucleus by RNA polymerase II to form primary miRNA (pri-miRNA). Primary miRNA is processed by the ribonuclease, Drosha and Dicer to generate mature miRNA. The single stranded miRNA, in association with Argonaute 2, binds to complementary sequences in the 3’ untranslated region (UTR) of target transcripts to regulate gene expression either by translational repression, activation or degradation of the mRNA transcript [1,14]. By targeting several genes, miRNAs play important roles in normal biological processes including cell proliferation, differentiation, apoptotic cell death, stress resistance and physiological metabolism [15,16]. Consequently, aberrant expression of miRNAs has been associated with malignancy, including; cancer stage, disease progression and metastatic spread [17-19]. Furthermore, some miRNAs have been shown to have oncogenic (such as mir-21, the cluster mir-17–92, mir-155, mir-221 and mir-222) [20] and tumour suppressive (such as let-7 in lung cancer and mir-15/16 in leukaemia and prostate cancer) properties [21-23].

Given that MPs are emerging as an important source of miRNA in the circulation in cancer patients [24-26] it is feasible to propose a role for MP in the aberrant miRNA levels displayed in oncogenesis and spread. This reinforces the role that MPs play in cancer biology including cell survival, invasion, metastasis and angiogenesis [27-31]. We recently discovered that MPs serve an important function as mediators in the dissemination and acquisition of multidrug resistance in cancer [32]. Specifically, we have demonstrated that this occurs via the MP-mediated transfer of functional resistance proteins, and nucleic acids including regulatory nucleic acids. In addition, we also showed that the MP transfer ensured the acquisition of the donor cell trait on to the recipient cells [33].

We now expand on these findings and demonstrate that MPs carry the transcripts encoding the membrane vesculation machinery (floppase and scramblase) and the enzymes essential for microRNA biogenesis (Drosha, Dicer and Argonaute). We also demonstrate the selective packaging of miRNAs within MP cargo upon release from the donor cells and propose that this process contributes to the dissemination and acquisition of the donor cell trait.

Results

Microparticles incorporate transcripts encoding the vesiculation machinery and microRNA biogenesis enzymes

qRT-PCR analysis of leukaemic cells and their MPs show that both the drug sensitive and resistant parental cells as well as their MPs, carry the transcripts for the vesiculation enzymes, floppase and scramblase (Figure 1A and 1B). Floppase is present at significantly higher levels in the MPs relative to the donor cells (Figure 1A), whereas scramblase, though present in all samples is present at significantly lower levels in the resistant cells and their MPs relative to the sensitive parental cells (Figure 1B).

MPs originating from VLB100 and CEM cells carry the transcripts encoding the enzymes Drosha, Dicer and Argonaute (Figure 1C, D and E), required for miRNA biogenesis. Both the drug sensitive and the resistant cells have significantly higher levels of the transcripts for Drosha and Dicer relative to their MPs (Figure 1C and D). Argonaute is also present in both the cells and their MPs but with no significant differences in their levels (Figure 1E).

Presence of miRNAs and modulation of the recipient cell miRNA profile following microparticle transfer

The quality of isolated RNAs was confirmed before subjecting the samples to miRNA microarray analysis (Figure 2). After normalization and transformation of the microarray data, the box-whisker plot of probe signal intensity was used to assess and confirm the quality of the microarray data (Figure 3A). Among the 7,815 probe sets in the miRNA microarray (http://www.affymetrix.com/support/technical/datasheets/miRNA_dataset.pdf), 847 probes were annotated as human miRNAs. The scatter plot of the signal intensities of these 847 human miRNAs displayed a correlation between MPs and their donor cells as well as between the acquired cells and the donor cells (Figure 3B). The miRNA microarray data was validated by qRT-PCR using the following selected miRNAs namely miR-150, miR-210, miR-107 and miR-125b (Figure 4).
To explore those miRNAs that were involved in the transfer of drug resistance by MPs to recipient cells, the miRNA expression profiles of MPs, drug sensitive recipient cells, acquired cells and donor cells were compared. The hierarchical clustering analysis of the 847 human miRNA uncovered selectively packaged miRNAs in the MPs relative to the donor cells (Figure 5). Furthermore, the acquired cells displayed a miRNA profile consistent with the donor following MP transfer. The sensitive cells were differential in their miRNA expression with respect to their drug resistant counterparts. In total, 209 miRNAs in leukaemia and 215 in the breast cancer cells were differentially expressed between the resistant donor cells and their MPs (Figure 5). Also, 222 and 155 miRNAs were differentially expressed between the resistant donor cells and their MPs (Figure 5). 208 and 200 miRNAs were also found to be differentially expressed between the drug sensitive and the resistant cells, in leukaemia and breast cancer, respectively (Figure 5). Of these, 195 miRNAs in leukaemia and 140 miRNAs in breast cancer were commonly identified between these two comparisons. The high level of similarity indicated the strong relationship between the MPs, the acquired cells and the donor cells. In addition, hierarchical clustering analysis of the 847 human miRNA expression profiles between all samples displayed common trends across the two cancer cell lines and provides further evidence of the tight correlation between the MPs, acquired cells following MP transfer and the donor cells (Figure 5).

To identify the most prominent miRNAs, linked with the MP-mediated transfer of drug resistance trait to drug sensitive cancer cells, selectively packaged and acquired miRNAs having \( p \)-value less than 0.06 (\( p < 0.06 \)) and fold change more than 1.5 (FC > 1.5) were selected (Figure 6). This comparison between leukaemia and breast cancer cells showed that 17 miRNAs were identified as the important miRNAs selectively packaged into MPs (Figure 7A). Likewise, across both cancers, 18 miRNAs were identified as significantly expressed miRNAs in the acquired cells (Figure 7B).

The MPs have a higher expression of these identified miRNAs relative to its donor cells, thereby being selectively packaged (Figure 7A), which are then transferred to the recipient cells upon coculture (Figure 7B).
acquired recipient cells display higher levels of miRNAs, relative to their parental recipient cells (CEM and MCF-7), following MP transfer. The expression levels in the MP acquired recipient cells with respect to their parental cells, is reflective of the donor cell trends (Figure 7B). Finally, 7 common miRNAs were identified as the most essential (being co-detected across both malignancies), which may be important for the transfer of donor traits via MPs (Figure 6). The 7 miRNAs, which were identified to be selectively packaged and acquired by recipient cells across both cancers, include miR-1228*, miR-1246, miR-1308, miR-149*, miR-455-3p, miR-638 and miR-923 (Table 1).

Microparticles selectively package miRNAs implicating traits specific to membrane vesiculation, cancer etiology and multidrug resistance on the target cell
The 1,571 unique predicted gene targets of the 7 miRNAs were identified from miRBase (prediction score > 60) and EMBL (p < 0.01). The gene target list was uploaded to DAVID Bioinformatics Resources 6.7 web-based program for functional annotation analysis. Significant biological pathways (*EASE score < 0.05) were selected as the important pathways that may be involved in MP formation and MDR trait transfer to recipient cells. The top 9 significant correlated pathways (p < 0.05) include “melanogenesis”, “calcium signalling pathway”, “ABC transporters”, “vascular smooth muscle contraction”, “hypertrophic cardiomyopathy” “steroid biosynthesis”, “maturity onset diabetes of the young”, “regulation of actin cytoskeleton” and “pathways in cancer” (Figure 8). Of these significant pathways identified for the miRNAs in this study, two were related to MP vesiculation (“calcium signalling pathway”, and “regulation of actin cytoskeleton”) and one to MDR (“ABC transporters”). Of all the target genes identified for the miRNAs, the highest percentages (~2.5 %) of these were observed to be related to the “pathways in cancer”. In addition, of all the predicted pathways identified for the miRNAs in this study, eight of them were related to malignancies alone.

Discussion
This study demonstrates that MPs serve as vehicles for intercellular communication and potentially as cancer biomarkers through their discrete miRNA signatures. RT-PCR analysis showed that MPs carry the transcripts of their vesiculation machinery (floppase and scramblase) (Figure 1A and B) together with transcripts encoding miRNA biogenesis enzymes (Drosha, Dicer and Argonaute) (Figure 1C, D and E). The presence of these phospholipid enzymes (floppase and scramblase) may help in intracellular vesicle trafficking either by inducing membrane vesiculation or by providing an environment favorable for binding of vesicle coat proteins [34]. This finding suggests that MPs are self-sufficient and possess the capacity to potentially induce vesiculation in the recipient cell. Although, the presence of scramblase 1 and 3 in exosomes has been previously reported [20,35], to our knowledge, this is the first demonstration of the presence of transcripts encoding the vesiculation machinery in MPs. The presence of Drosha, Dicer and Argonaute transcripts within the shed cargo potentially implicates MPs as key intercellular regulators of miRNA biogenesis in recipient cancer cell populations.

Affymetrix miRNA microarray was used to explore the miRNA expression profiles of MPs and their donor cells from both leukaemia and breast cancer cells in this study (Figure 5). This data was validated using RT-PCR
studies where we observed almost similar trends for most miRNAs examined (miR-107, miR-125b and miR-210) except for miR-150 (Figure 4). The basis of this difference in currently unknown and may be attributed to the differences in assays. Our analysis showed that several human miRNAs are selectively packaged into MPs. Upon coculture with recipient cells, we observe an increase in select miRNAs, including miR-1246, miR-1308, miR-1228*, miR-149*, miR-638 and miR-923 (Figure 7). These miRNAs displayed a 2-12-fold increase in expression levels in the MPs relative to their donor cells (Figure 7A). These observations are consistent with selective packaging, which we have previously shown for miR-451 and miR-326 in MPs shed from MDR resistant breast and leukaemia cells [33]. In addition, our findings are consistent with previous reports demonstrating that these same miRNAs are also selectively packaged into exosomes [24,36-38].

We also demonstrate that following MP transfer to recipient breast cancer and leukaemia cells, the same miRNAs were significantly increased in the acquired cells, with levels ranging from 2-15-fold increase (Figure 7B). Once again the expression level of the miRNAs in the MP compartment was directly correlated to the levels observed in the acquired population following MP transfer (Figure 5). In saying this however, we cannot exclude the possibility that increased miRNA levels in the recipient cells are caused by either/both direct or indirect MP-mediated effects on the transcription of the miRNA. Interestingly, the acquired population displayed miRNA expression trends reflective of the donor cells (Figure 7B). This demonstrates that the recipient cell reflects the donor

Figure 3 Quality of data and gene expression signal correlation among identified miRNA. After normalization and transformation of the microarray data (A) the box-whisker plot of probes signal intensity assessed and confirmed the microarray data quality. (B) The scatter plot of the signal intensity of the 847 annotated human miRNAs showed that certain level of correlation was identified between the MPs (VLBMP or DXMP), the acquired cells (CEM+ VLBMP or MCF-7+ DXMP), the donor cells (VLB or DX) and the parental recipient cells (CEM and MCF-7).
trait after MP-mediated transfer of cargo. These findings are consistent with our previous report where we have shown the “re-templating” of recipient cells to reflect donor cell traits following MP-mediated transfer of MDR transporter transcripts [33].

The miRNAs identified in our study play crucial roles in cancer cell biology. Using the miRDB target prediction program [39], NKIRAS1, which is a NFKB inhibitor was identified as one of the targets of miR-1308. Nuclear factor kappa B (NFκB) is a family of transcription factors that play important roles in regulating cell differentiation, proliferation, immune response and blocking apoptosis [40,41]. This family of transcription factors have been reported to chemosensitize P-gp overexpressing cancer cells [42]. This miRNA has also been shown to be upregulated in cancerous tissues and also in the more aggressive inflammatory breast cancer (IBC) in comparison to the non-IBC tissues [43,44]. Similarly, miR-1246 targets the NKF3 kinase family member gene, SGK269 (miRDB database). NKF3 or PEAK1 promotes anchorage independent growth and tumour progression in pancreatic cancer cells transplanted in mice [45]. miR-149+ is known to induce apoptosis by the direct inhibition of Akt1 and E2F1 in neuroblastoma cells [46]. Akt is the key kinase of the signal pathway, which mediates the regulation of divergent cellular processes including apoptosis, proliferation, differentiation and metabolism [47].

miR-638 has been found to be consistently, highly expressed in human plasma and its presence in the plasma may be physiologically necessary [48]. As such, the ratio of miR-92a/miR-638 in blood is associated with diagnosis in acute leukaemia patients [48]. Gene targets of miR-638, cyclin G2 and transcription elongation regulator 1-like factor (miRDB database), were involved in p53 and platelet-derived growth factor (PDGF) signalling pathways [49]. miR-638 was one of the downregulated miRNAs in colorectal liver metastases with respect to the adjacent liver tissues that have the potential to serve as a diagnostic and predictive marker of colorectal liver metastases [50]. Likewise, miR-1228+ has been previously shown to be highly expressed in malignant mesothelioma tumour samples compared to normal samples [51].

The role of the miRNAs and their targeted pathways in the cells examined in this study are currently unknown. It is feasible that the same miRNAs may serve similar functions in these cells as is the case in the other reported cancers. However, the pathway analysis of the predicted targets of the 7 identified miRNAs in this study showed the maximum percentage of target genes to be significantly related to “pathways in cancer” and at least seven other pathways as well that were cancer related (Figure 8). The malignancy-
related upregulated expression of these miRNAs may serve as potential biomarker in the treatment of cancer.

Chemotherapy comprises the major therapeutic strategy for clinical cancer treatment. However, chemotherapy fails to eliminate all tumour cells because of intrinsic or acquired drug resistance, which is the most common cause of tumour recurrence [52,53]. The role of miRNAs in the regulation of resistance mediated by multidrug transporters has only been examined recently [33]. Interestingly, we found that some of the significantly expressed miRNAs (like miR-455-3p) identified in this study target the multidrug resistant protein, P-glycoprotein (P-gp). For example, miRDB target prediction shows MDR member 1 or (P-gp) and HIF1AN (hypoxia-inducible factor 1, alpha subunit inhibitor) as miR-455-3p targets. HIF-1 alpha has been shown to induce MDR in hepatocellular carcinoma cell line [54]. In this study, the microarray analysis showed that the resistant leukaemia cell line has a lower expression level of this miRNA relative to its drug sensitive cells, which is consistent with its overexpression of P-gp. In the acquired cells we observed a suppression of miR-455-3p implicating potentially increased P-gp levels after MP transfer. Hence, our previous observations of P-gp protein transfer in the drug sensitive recipient cells after MP coculture [32] may be due to the transfer of these regulatory miRNAs together with protein via the MP cargo. In breast cancer, the overexpression of miR-923 was shown to be upregulated in the Taxol resistant cancer cells relative to the normal cells [55].

Other than pathways related to malignancies and MP vesiculation, “ABC transporters” was identified as a significant biological pathway with the highest percentages of the identified miRNA target genes. Previous studies have also reported on the role of miRNAs involved in MDR in cancer. These include miR-27a and miR-451, whose expressions were shown to induce MDR1/P-gp expression in resistant human ovarian cancer cells [56]. The overexpression of MRP1 and miR-326 levels was inversely related in breast cancer tissues and leukaemia [33,57]. Recently, miR-345 and miR-7 have shown to target MRP1 in MDR breast cancer cells relative to parental cells [58]. Apart from MDR, the ABC transporter, ABCA1 has also been shown to possess floppase activity and is related to MP production [59]. In addition, of the significant predicted pathways identified for the miRNAs in this study, two of them namely: “calcium signalling pathway” and “regulation of actin cytoskeleton” were related to MP vesiculation (Figure 8). This suggests that MPs not only carry the enzymes for its release but also carry miRNAs and genes which may be potentially involved in its production and release.

Our study gives an implication of the role miRNAs contained within the MP cargo may play in contributing to the emergence of MDR and in regulating transporter expression in recipient cancer cells. Previous studies have shown miRNA transferred by other vesicular bodies like microvesicles to be functional in the recipient cells [60,61]. However, the functional role of the identified miRNAs in this study needs further exploration to have any clinical relevance.

Conclusions

In conclusion, this study reveals miRNAs not only as oncogenic or tumour suppressive, but also highlights the potential role of these molecules as a potential class
of diagnostic biomarkers across both haematological
and non-haematological malignancies. The miRNAs
transferred by MPs also play an important role in the
regulation of biological processes involved in anticancer
drug resistance. Indeed, the detection of circulating
tumour-derived transcripts from melanoma, breast and
lung cancer patients has identified MPs as potential mar-
kers of diagnostic and prognostic significance [26]. Thus,
miRNA profiling has the potential to serve as a non-
invasive approach to probe for the presence of deleterious
cancer traits clinically.

**Methods**

**Cell Lines**

Two cell lines were used for these studies. The first cell
line included the drug-sensitive human acute lympho-
blastic leukaemia cell line CCRF-CEM (designated CEM
for simplicity), and its MDR variant VLB100. The second
included the drug-sensitive human breast adenocarcinoma
cell line MCF-7, and its MDR variant MCF-7/DX (desig-
nated as DX for simplicity). These cells were kind gifts
from Dr Rosanna Supino (Istituto Nazionale per lo Studio
e la Cura dei Tumouri, Milan, Italy) and Dr Suzanne M.
Cutts (La Trobe University, Victoria, Australia). Both of these cell lines have been validated earlier by our group as an appropriate model for the study of P-gp-mediated MDR in vitro [32,62,63]. All cell lines were cultured in RPMI 1640 (Invitrogen Australia, VIC, Australia) containing 10% FCS (Invitrogen, Australia) and maintained under a humidified incubator at 37°C in an atmosphere of 5% CO₂.

MP harvesting and identification
MPs were isolated from confluent CEM, VLB100 and DX cells by differential centrifugation, as previously described [32,33]. The MPs were designated as CEMMP, VLBMP and DXMP for simplicity. Briefly, culture supernatants were collected and centrifuged at 500 g for 5 min to pellet whole cells. The collected supernatant was re-centrifuged at 15,000 g for 1 h at 15°C to pellet the MPs. The final

| miRNA name         | miBase sanger accession number | Sequence                              | Length | Precursor sanger annotations | Chromosomal location |
|--------------------|--------------------------------|----------------------------------------|--------|-----------------------------|----------------------|
| hsa-miR-149-star   | MIMAT0004609                   | 5' - agggaggagggcgccguggc - 3'          | 21     | MI0000478                   | 2q37.3               |
| hsa-miR-455-3p     | MIMAT0004784                   | 5' - gcagucggcggcggcgccg - 3'           | 21     | MI0003513                   | 9q32                 |
| hsa-miR-638        | MIMAT0003308                   | 5' - agggauucgcggcggccggc - 3'          | 25     | MI0003653                   | 9p13.2               |
| hsa-miR-923        | MIMAT0004973                   | 5' - gucagcggacggagaaagac - 3'          | 23     | MI0005715                   |                       |
| hsa-miR-1228-star  | MIMAT0005583                   | 5' - ucacaccucggccgcccc - 3'           | 20     | MI0006318                   | 12                   |
| hsa-miR-1246       | MIMAT0005898                   | 5' - auggauuuuuugggaccg - 3'           | 19     | MI0006381                   | 2q31.1               |
| hsa-miR-1308       | MIMAT0005947                   | 5' - gauggggguucagguug - 3'            | 18     | MI0006441                   | Fragment of a tRNA   |

**Table 1** The list of identified miRNAs with their characteristics

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**Figure 8** Selected miRNAs have common biological pathways. The common biological pathways were identified with the list of predicted targets for the seven co-detected miRNA genes by DAVID Bioinformatics Resources. The percentages of the total identified target genes (y-axis) were plotted against the pathways they regulate (x-axis) with increasing p-value. The pathways in red bars indicate most significant pathways with p-value < 0.05. The green bars indicate the pathways associated with malignancies. Significant biological pathways (p-value < 0.05) were selected as the important pathways. *Ease Score Threshold (Maximum Probability): The threshold of EASE Score, a modified Fisher Exact p-value, for gene-enrichment analysis. p-value <0.05 represents strong enrichment in the annotation categories.
pellet was resuspended in serum free RPMI 1640 media and centrifuged at 2000 g for 1 min to remove debris. The clear MP suspension was further centrifuged at 18,000 g for 30 min at 15°C to pellet MPs. Validation of the isolated MP pellet was performed using flow cytometric analysis (FCM) (Cytomics FC500 MPL, Beckman Coulter) after FITC-annexin V (Beckman Coulter, NSW, Australia) staining as previously described [32]. Total protein content of MPs was determined using the Quant-iT™ protein assay as per the manufacturer’s instructions (Invitrogen Australia).

MP transfer experiment and isolation of mRNA

In a 96-well U bottom culture plates, 180 μg of VL BMP or DXMP was cocultured with 1 x 10^5 CEM or MCF-7 cells, respectively, for 4 h in total of 200 μL complete RPMI culture medium at 37°C and 5% CO₂. Unbound MPs were removed by washing with PBS and centrifuging at 500 g for 5 min at 25°C after 4 h. The cocultured samples were designated as CEM + VL BMP and MCF-7 + DXMP for simplicity and were referred to as the “acquired” cells.

Total RNA was extracted and pooled using Trizol® Reagent (Molecular Research Center, Inc, OH, U.S.A) as per manufacturer’s recommendations from (i) the parental drug sensitive CEM or MCF-7 cells, (ii) the MDR strain VL+B100 or DX cells, (iii) VL BMP or DXMP, and (iv) the cocultured samples CEM + VL BMP or MCF-7 + DXMP from duplicate experiments.

Gene expression analysis of MP vesiculation and miRNA biogenesis enzymes

Quantitative real-time polymerase chain reaction (qRT-PCR) was used to assess the presence of scramblase and floppase RNA transcripts (involved in MP budding), as well as Argonaute, Dicer and Drosha RNA transcripts (involved in miRNA biogenesis) in isolated MPs. Briefly, cDNA was synthesized using the Advantage RT-for-PCR Kit (Clontech Laboratories, Inc., Mountain View, CA). The specific primers against the target genes were used with GAPDH as the housekeeping primer (Sigma-Aldrich, St Louis, MO, USA) (Table 2). Reactions were carried out at the volume of 10 μL using 2 x SYBR Green Premix ExTaq (Takara Bio Inc., Shiga, Japan) with 10 pmole of target specific primer pairs and the amplification were performed on the LightCycler 2.0 (Roche, NSW, Australia). The thermal profile for the qRT-PCR was 95°C for 5 min followed by 45 cycles of 95°C for 5 sec, 55°C for 10 sec, and 72°C for 15 sec. The Ct data of each sample was compared with the housekeeping gene to obtain the ΔCt using the following formula: ΔCt = target gene Ct – housekeeping gene Ct. The relative expression level was calculated using $Δ\Delta Ct = 2^{-ΔΔCt}$ and expressed as fold difference from the experimental control [(ΔΔCt of sample ÷ ΔΔCt of drug-sensitive cells) x 1] as arbitrary units (a.u.).

### Table 2 The list of sequences of primers used for real-time RT-PCR experiments

| Primers   | Sequences                                      |
|-----------|------------------------------------------------|
| Scramblase | Forward: 5'-AAATGGATGTGCTGTCTTCC -3'          |
|           | Reverse: 5'-TCTACTACACACTCTCCTGATT -3'        |
| Flopase    | Forward: 5'-TGGAACTAGGACGATCACG -3'           |
|           | Reverse: 5'-GAAACGTGTCACAGGCCCCAT-3'          |
| Argonaute  | Forward: 5'-TCTATGCTGCTGTAAGTACGA -3'         |
|           | Reverse: 5'-TCAAGCATTCTCGGACACG -3'           |
| Dicer      | Forward: 5'-TGCAAAGCTGTACCCACAGAG -3'         |
|           | Reverse: 5'-ACACTGCTGAAAGCGTTGATT -3'         |
| GAPDH      | Forward: 5'-TGCCAAATATGATGACATCAAGAA -3'      |
|           | Reverse: 5'-GGAGTGGGGTCGTCGTGTTG -3'          |
| miR-U6     | Forward: 5'-CTCGCTTCTGGCCAGACACA -3'          |
|           | Reverse: 5'-AAAGCTTCTCAGGTTGATTCCGT -3'       |
| miR-107    | Forward: 5'-AAGCGACATTGGCTACAGGTTAC -3'       |
| miR-125b   | Forward: 5'-TCCCTGAGACCCTTAACCTTGA -3'        |
| miR-150    | Forward: 5'-TCCCTCCAACCCTTGAACCTGAGT -3'      |
| miR-210    | Forward: 5'-CTGTCGGTGCTGACAGGCGG -3'          |

Analysis of RNA integrity and Affymetrix miRNA Arrays

Total RNA integrity was analysed using the Agilent RNA 6000 Nano kit (Agilent Technologies Inc., Santa Clara, CA) and the result was analysed by Agilent 2100 Bioanalyzer (Agilent Technologies Inc.) as per the manufacturer’s recommendations; the RNA integrity number (RIN) of 10 represents the highest RNA integrity with minimal degradation and score of 1 is the lowest integrity [64]. Nanodrop-1000 spectrophotometer (Nanodrop technologies, DE, USA) was used for the quantification of RNA and 500 ng of RNA from each sample was used for miRNA microarray analysis. RNA labelling, hybridization (Affymetrix™ Fluidics Station 450), scanning (GeneChip™ Scanner 3000 7 G) and raw data acquisition of the Affymetrix GeneChip® miRNA Array (P/N 901326) were performed by Australian Genome Research Facility Ltd, VIC, Australia following a standard procedure from Affymetrix™ (Santa Clara, CA).

miRNA Microarray analysis

**Data processing**

Affymetrix™ “CEL” and “CHP” data files of each sample were processed with Affymetrix™ miRNA QCTool software and following the guided workflow as described in the user manual (http://www.affymetrix.com/support/technical/manuals.affx). Briefly, the signal intensities data was extracted from data files and probes level intensity data were obtained using Wilcoxon-Rank Sum test, followed by background adjustment based on the GC content of ‘anti-genomic’ probes, quantile normalization, addition of a small constant (value 16) to avoid negative signal after background-GC correction, and finally applying median summarization to all probe set in each sample. Probe intensities data presented are all log₂.
transformed and p-values are obtained from the software after Wilcoxon-Rank Sum test. All microarray data discussed in this manuscript have been deposited in NCBI's Gene Expression Omnibus and are accessible through GEO Series accession number SEGSE34560 (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?token=dvkxzwnqwgmsrw&acc=GSE34560).

Data mining
Data was filtered to include only annotated as Homo sapiens and miRNA, and probes with the p-value less than 0.06 (p < 0.06) were selected as a significant for further analysis. To identify those miRNAs that were correlated to drug resistance and its transfer from the donor to recipient cells by MPs, expression profiles of (i) MP donor cells (VLB100 or DX), (ii) isolated MPs (VLBMP or DXMP), and (iii) MP cocultured with drug sensitive cells (CEM + VLBMP or MCF-7+DXMP) were compared, and miRNAs with fold change more than 1.5 (FC > 1.5) were identified.

Hierarchical clustering and targeting pathway analysis
‘Cluster 3.0’ program [65] was used for hierarchical clustering analysis, where selected miRNAs were clustered by centroid linkage using Euclidean distance, and depicted result was generated using ‘Java Tree View’ program [66]. Furthermore, a complete list of predicted gene targets on the selected miRNAs was downloaded from miRBase (Release version 16) (http://www.mirbase.org), miRDB (http://www.mirdb.org) and the EMBL Nucleotide Sequence Database (http://www.ebi.ac.uk/emb/l). The target genes for individual miRNAs with a score > 60 (miRBase) or with p-value < 0.01 (EMBL) were selected and uploaded to the online DAVID Bioinformatics Resources 6.7 program (http://david.abcc.ncifcrf.gov/) for their functional annotation clustering analysis. The biological pathways and gene regulation by the selected miRNAs were identified.

Microarray gene expression validation by qRT-PCR
Total RNA from isolated MPs, acquired cells and whole cells were extracted as described above. cDNA for miRNA was synthesized using the Ncode miRNA First Strand cDNA Module kit (Invitrogen Australia) on the GeneAmp PCRsSystem 9700 (Applied Biosystems), mir-150, mir-210, mir-107 and mir-125b miRNA specific primers (10 pmole/reaction) were used for PCR for the detection of miRNAs using mir-U6 as the housekeeping primer (all primers were from Sigma-Aldrich) (Table 2). SYBR Green qRT-PCR amplifications were performed on the Mastercycler® ep realplex (Eppendorf, NY, USA). Reactions were carried out in a 20 μL volume containing 10 μL of 2× SYBR Green Premix ExTaq (Takara). The thermal profile for the qRT-PCR was 91°C for 5 min followed by 45 cycles of 91°C for 15 sec, 60°C for 30 sec, followed by melting curve detection. The Ct data of each sample was collected automatically and data expressed as described above.

Statistical analysis
A one-way analysis of variance (ANOVA) was used for comparison and statistical analysis between the sample populations and the control drug-sensitive cell population using the Graph Pad Prism software. p-values less than 0.05 were accepted as statistically significant.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
RJ conducted all experiments and drafted the manuscript; FL participated in manuscript preparation including microarray and RT-PCR data analysis; JG participated in microarray breast cancer sample preparation; J-MM participated in the design of the microarray study and provided reagents; MB, GERG participated in work planning and manuscript preparation. All authors read and approved the final manuscript.

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