REVIEW

Extracellular vesicles for liquid biopsy in prostate cancer: where are we and where are we headed?

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BACKGROUND: Extracellular vesicles (EVs) are a heterogeneous class of lipid bound particles shed by any cell in the body in physiological and pathological conditions. EVs play critical functions in intercellular communication. EVs can actively travel in intercellular matrices and eventually reach the circulation. They can also be released directly in biological fluids where they appear to be stable. Because the molecular content of EVs reflects the composition of the cell of origin, they have recently emerged as a promising source of biomarkers in a number of diseases. EV analysis is particularly attractive in cancer patients that frequently present with increased numbers of circulating EVs.

METHODS: We sought to review the current literature on the molecular profile of prostate cancer-derived EVs in model systems and patient biological fluids in an attempt to draw some practical and universal conclusions on the use of EVs as a tool for liquid biopsy in clinical specimens.

RESULTS: We discuss advantages and limitations of EV-based liquid biopsy approaches summarizing salient studies on protein, DNA and RNA. Several candidate biomarkers have been identified so far but these results are difficult to apply to the clinic. However, the field is rapidly moving toward the implementation of novel tools to isolate cancer-specific EVs that are free of benign EVs and extra-vesicular contaminants. This can be achieved by identifying markers that are exquisitely present in tumor cell-derived EVs. An important contribution might also derive from a better understanding of EV types that may play specific functions in tumor progression and that may be a source of cancer-specific markers.

CONCLUSIONS: EV analysis holds strong promises for the development of non-invasive biomarkers in patients with prostate cancer. Implementation of modern methods for EV isolation and characterization will enable to interrogate circulating EVs in vivo.

INTRODUCTION

Prostate cancer (PC) is the most common non-skin cancer in males. Thanks to significant advances in the diagnosis and treatment, the 10-year cancer-specific survival of PC patients exceeds 98%1 enabling many men to live with, rather than die of, their disease. In spite of this, more than 180 000 new cases are estimated in 2016 and over 26 000 men are expected to die of PC in the same year.2 Consequently, PC remains the second leading cause of cancer deaths in the United States. Among the key clinical challenges are the inability to accurately distinguish low-risk (indolent) from high-risk (aggressive) disease at the time of diagnosis. In addition, the highly variable clinical course has limited the capacity to predict the biological progression of the disease. Within the existing risk stratification systems for PC, histological grading using the Gleason score system3 remains one of the best independent predictors of clinical outcome. This system was recently modified based on new 2016 World Health Organization PC reporting guidelines.4,5 The five proposed risk groups (PGG1–PGG5) correspond to genomic groups with increasing number of somatic copy number aberrations.6 Metastatic PC exhibits a distinct signature with specific driving genomic alterations that might provide personalized therapeutic targets.6

Indeed, different genomic signatures might characterize diverse facets of the same lethal disease, and this molecular heterogeneity, along with potentially actionable molecular defects has been identified not only in castration resistance (CR) disease but also in naïve PC.7 Identifying alterations that predict high-risk disease, metastasis and/or CR would improve the management of the patients and their long-term outcome.

Although fine-needle biopsies are the standard for PC diagnosis and prognosis, their ability to identify high-risk disease is limited8 Moreover, their invasive nature causes significant morbidity. Analysis of PSA in blood has long been used for early diagnosis and for monitoring of biochemical recurrence, but is flawed with a significant number of false positives.9 These limitations drive ongoing attempts at developing minimally invasive procedures to interrogate cancer-derived macromolecules in circulating tumor cells (CTCs), cell-free DNA (cfDNA) and extracellular vesicles (EVs).

Circulating tumor cells

Despite the intrinsic limitation that CTCs are rare,10 analysis of CTCs has allowed a non-invasive, real-time molecular...
characterization of cancer in patients with metastatic disease\textsuperscript{11,12} and has successfully been used to predict response to Docetaxel and Prednisone in patients with metastatic castration resistant PC (mCRPC).\textsuperscript{13} Additionally, nuclear expression of the androgen receptor variant V7 (AR-V7) in CTCs seems to report response to taxanes.\textsuperscript{14} Recently, CTCs have been shown to be heterogeneous and to present characteristics of neuroendocrine differentiation, suggesting that broader criteria for definition and inclusion of CTCs should be used.\textsuperscript{15}

Cell-free DNA
Tumor-derived cfDNA is particularly attractive due to its abundance in plasma. Profiling cfDNA by next-generation sequencing has recently resulted in a strategy for monitoring tumor dynamics and identifying genomic causes of resistance, suggesting a temporal association between cfDNA profiles and clinical progression.\textsuperscript{16} Similar studies have enabled accurate estimations of absolute androgen receptor (AR) copy number\textsuperscript{17} and such an analysis of plasma AR in CRPC patients has identified primary resistance to Abiraterone.\textsuperscript{18} These studies rely on sophisticated computational tools to limit the possibility of missing circulating AR genomic lesions caused by low input cfDNA, which is also contaminated with benign cell-derived DNA.\textsuperscript{19,20} Mounting evidence indicates that cfDNA is not the only source of extracellular DNA in plasma, and that a portion of this DNA along with other cancer-derived molecules is carried and protected in EVs.

Extracellular vesicles
EVs are lipid-enclosed particles released by all cellular organisms and containing a repertoire of macromolecules (cargo) that is reflective of the cell of origin. The EV cargo is stable because it is protected from enzymatic degradation by a lipid membrane envelope. EVs are extremely abundant (\(\sim 1 - 3 \times 10^{12}\) exosomes per ml of plasma).\textsuperscript{21} Direct enumeration of tumor-derived EVs and/or profiling of their molecular cargo in patient body fluids have been shown to provide valuable information about the biology of the tumor.\textsuperscript{19} While EV profiling offers significant advantages because they contain a complex cargo (proteins, lipids, nucleic acids) that could be interrogated simultaneously, discriminating cancer-derived EVs from the non-cancer ones has been challenging.

\textbf{EVs ARE HETEROGENEOUS}

Various populations of EVs have been described.\textsuperscript{22,23} While exosomes and ectosomes are the most frequently studied EVs, biofluids contain highly heterogeneous EV populations, of variable size and originating by diverse mechanisms (Figure 1). Importantly, different EV types exhibit distinct molecular and functional characteristics.\textsuperscript{22,23} Also, benign cells produce various types of EVs.\textsuperscript{24} Thus cancer-derived EVs float in a sea of physiologically normal EVs and the use of EVs in liquid biopsies relies on improving the methodologies to selectively purify single EV populations based on their subcellular origin, size, cargo, and donor cell type and status.

\begin{itemize}
  \item \textbf{Size–nomenclature–biogenesis–origin}
  \begin{itemize}
    \item \textbf{Exosomes}. Exosomes are nanosized vesicles (50 to > 100 nm in diameter)\textsuperscript{25} that originate from the fusion of multivesicular bodies with the plasma membrane (PM),\textsuperscript{26} or bud directly off the PM\textsuperscript{26} as recently reviewed.\textsuperscript{20,21} Any type of cell from any living organism can shed exosomes.\textsuperscript{27} Prostate-specific EVs termed ‘prostasomes’ were originally described as exosome-like structures that are released by normal prostate epithelial cells in the seminal fluid.\textsuperscript{28} Functionally, it is known that prostasomes mediate intercellular communication between epithelial secretory cells and sperm cells. More specifically, secretory cells seem to use this mechanism to nurture sperm cells, thus increasing their motility.\textsuperscript{29} The term prostasomes has often been used interchangeably with exosomes due to their high cholesterol/phospholipids ratio and to a protein pattern that suggests the origin of prostasomes from multi-vesicular bodies. Tavosidana \textit{et al}.\textsuperscript{30} reported that detection of prostasomes in peripheral blood is useful for early diagnosis and prognosis in organ-confined PC. However, recent studies have clarified that, unless these EVs are identified in the seminal fluid, the term exosomes is more appropriate to describe them.\textsuperscript{31}
    \item \textbf{Ectosomes}. The term ectosomes indicates exquisitely EVs originating by direct budding and pinching off of the PM.\textsuperscript{32} These vesicles are generally larger than exosomes (250–1000 nm) and result from PM budding as a consequence of activation of GTPases.\textsuperscript{32,33} Ectosomes are also frequently referred to as ‘microvesicles’ (MV). MV were originally described as platelet-derived EVs with a functional role in coagulation\textsuperscript{34} but several new types of bioactive vesicles derived from a variety of cells, including tumor cells, have been described.
    \item \textbf{Large oncosomes}. Large oncosomes (LO) are atypically large EVs (1–10 \(\mu\)m diameter) that are released by tumor cells transitioning from a mesenchymal to a more rapid, amoeboid motility mode, which results in highly migratory, invasive, and metastatic phenotypes.\textsuperscript{35–38} This process occurs in cells with active Akt1 and EGFR signaling,\textsuperscript{39} and can be induced by silencing of the activin receptor-like kinase 1, 2 and 7 (ALK1, ALK2 and ALK7).\textsuperscript{40} They do not appear to be produced by benign cells.\textsuperscript{38} More recently, the identification of similarly large EVs has been reported\textsuperscript{31,40,41} but it is unclear whether they share similar functional or molecular characteristics to the LO.
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  \item \textbf{EVs AS A SOURCE OF BIOMARKERS IN PC}
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    \item While a discrete portion of molecules is enriched or depleted in EVs in comparison with the cancer cell of origin, suggesting a finely regulated mechanism of export,\textsuperscript{42,43} the EV cargo largely reflects the donor cell cargo. Because these molecules are protected from degradation and can be measured in biological fluids, they represent a valuable, non-invasive source of tumor-derived markers, as demonstrated in a large number of neoplasms (Table 1).\textsuperscript{44–49}
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Table 1. Examples of commercially available kits for EVs isolation

| Cancer type | Biological fluid | Correlation |
|-------------|------------------|-------------|
| GPC-1       | Pancreatic       | Biological fluid | Correlation |
| TYRP2       | Melanoma         | Plasma       | Tumor detection |
| VLA-4       | Melanoma         | Plasma       | Tumor detection |
| HSP70       | Melanoma, NSCLC  | Plasma       | Tumor detection |
| CD9         | NSCLC            | Plasma       | Progression |
| CD81        | NSCLC            | Plasma       | Progression |
| Tsg101      | NSCLC            | Plasma       | Progression |
| EGFRvIII    | NSCLC            | Plasma       | Progression |
| CD151       | NSCLC            | Plasma       | Progression |
| miR-21      | Glioblastoma     | Cerebrospinal fluid | Progression |

The majority of these kits lead to the isolation of a mixed extracellular vesicles (EV) population.

Mass spectrometry

Protein profiling in plasma, serum and other biological fluids are limited by an intrinsic high dynamic range, and by the fact that most of the FDA-approved serum biomarkers are at least five orders of magnitude lower than albumin or other abundant serum proteins. For this reason, most of the preliminary studies on the characterization of the protein cargo of PC-derived EVs have been performed on EVs obtained from PC cell lines. This approach has created an invaluable resource of potential biomarkers and has circumvented the limitations deriving from a direct interrogation of body fluids.

One of the first deep mass spectrometry (MS) analyses of EVs was performed by the Llorente group in exosomes derived from PC cell lines to test whether the proteins released in the serum are likely to be packaged in EVs. Strikingly, most of the proteins identified in mouse serum and in PC cell-derived exosomes, including glycerolaldehyde 3-phosphate dehydrogenase (GAPDH), ENO1 and lactate dehydrogenase B (LDHB), were found only in the exosome fraction, and not in the exosome-depleted supernatant, suggesting they are exported in EVs. Identification of TM256 by Øverbey et al. in urine exosomes discriminated PC patients (n = 16) from controls (n = 15) with 94% sensitivity. The sensitivity reached 100% when TM256 was analyzed in combination with the autophagy inhibitor late endosomal/lysosomal adaptor, MAPK and MTOR activator 1 (LAMTOR1). These proteins are candidate EV urinary markers, and the validity of this discovery is supported by the identification, in the same study, of well-known PC markers such as PSA, prostate-specific membrane antigen (FOLH1/PMSA), transglutaminase (TG4M) and transmembrane protease, serine 2 (TMPRSS).

Exosome proteins have also been interrogated for identification of biomarkers of treatment response and particularly for early detection of drug resistance. Kharaziha et al. provided evidence that exosomes report acquisition of resistance to docetaxel. In comparison with parental DU145 PC cells, which are sensitive to docetaxel, DU145 cells that have acquired resistance to the compound (Tax-Res) shed more abundant exosomes as measured by NTA. Additionally, exosomes derived from Tax-Res cells carry a different set of proteins while expressing similar levels of exosome cargo.19,61 It would be interesting to investigate whether EV-bound molecules.

In order to improve reproducibility and sensitivity of protein detection, Webber et al. applied a novel MS platform to exosomes and donor DU145 PC cells. The resulting SOMAscan platform uses a class of protein-binding reagents called SOMA- mers (slow off-rate modified aptamers) that allows the detection of a significant number of proteins and can handle a large number of samples. Milk fat globule-EGF factor 8 (MFG-E8), Notch3, disintegrin and metalloproteinase domain-containing protein (ADAM9) were enriched in exosomes in comparison to the tumor cells. Not only did this study contribute to the identification of novel putative extracellular markers of PC, but it also led to the discovery that proteins that are typically considered to be secreted in a free form, are actually exported in EVs. The authors further demonstrated by sucrose density gradient that the soluble form of transforming growth factor β1 (TGF-β1) does not co-isolate with exosome-bound TGF-β1 and they concluded that this is the case also for granulocyte-colony stimulating factor, vascular endothelial growth factor, and interleukin 8, confirming the importance to analyze soluble molecules separately from EV-bound molecules.

A MS approach was also employed by Hosseini-Beheshti et al. to profile exosome proteins from AR positive and negative PC cell lines. A number of candidate biomarkers, including calysntenin-1 (CLSTN1), growth differentiation factor 15 (GDF15), fatty acid synthase (FASN) and clusterin (CLU) were identified in EVs from these tumor cells but not from non-tumorigenic prostate epithelial cells. Finally, Dujíevs et al. identified higher levels of programmed cell death 6-interacting protein (PDCD6IP/Alix), FASN, CD9 and enolase 1 (ENO1) in exosomes derived from malignant prostate cell versus benign cell lines.

The Jenster group performed MS on sera of mice with androgen-naïve or androgen resistant PC thus providing one of the first evidences for in vivo significance of EVs as biomarkers in this disease. They compared the sera profiles to the profiles of exosomes and exosome-deprived supernatant of PC cells to test whether the proteins released in the serum are likely to be packaged in EVs. Strikingly, most of the proteins identified in mouse sera and in PC cell-derived exosomes, including glycerolaldehyde 3-phosphate dehydrogenase (GAPDH), ENO1 and lactate dehydrogenase B (LDHB), were found only in the exosome fraction, and not in the exosome-depleted supernatant, suggesting they are exported in EVs. Identification of TM256 by Øverbey et al. in urine exosomes discriminated PC patients (n = 16) from controls (n = 15) with 94% sensitivity. The sensitivity reached 100% when TM256 was analyzed in combination with the autophagy inhibitor late endosomal/lysosomal adaptor, MAPK and MTOR activator 1 (LAMTOR1). These proteins are candidate EV urinary markers, and the validity of this discovery is supported by the identification, in the same study, of well-known PC markers such as PSA, prostate-specific membrane antigen (FOLH1/PMSA), transglutaminase (TG4M) and transmembrane protease, serine 2 (TMPRSS).

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progression66–68 and has been suggested as a circulating marker of metastatic PC by the Thompson group,69 emerged as a promising LO-associated protein detectable in the circulation of patients with metastatic disease but not in patients with organ-confined PC.70 Importantly, Cav-1 expression was also identified in exosomes, confirming the data from other groups.71 However, flow cytometry experiments indicated that Cav-1-positive LO, but not exosomes, are significantly increased in patients with metastasis.70 One putative marker of LO is cytokeratin 18 (CK18), a protein that is almost completely excluded from exosomes, as demonstrated by gradient centrifugation.72 The presence of CK18-positive EVs in human plasma was associated with cancer in a small cohort of PC patients.23 Additional LO markers are HSPA5, HSPD1, GAPDH and other functionally interesting proteins, including membrane proteins and metabolic enzymes that seem to be implicated in the biological functions of LO.23 Interrogating larger cohorts of patients to validate clinical significance of these markers is essential. Because of the tumor-specificity, differentiating LO from other EV types may increase the granularity of cancer-specific signal. Given their large size, LO could also be easily captured and analyzed by NGS, thus informing the genetic composition of the tumor, risk of recurrence, and disease-free or overall survival. Interestingly, the identification of very large objects (sometimes as large as CTCS) positive for epithelial cell adhesion molecule (EpCAM) and CK18 in the circulation of PC patients was reported in 2010 by Coumans et al.72 and was correlated with reduced survival. Unfortunately, the system currently approved by the FDA uses particularly strict parameters to define a CTC, therefore these abnormally large vesicular structures, which could be LO, are discarded and not incorporated in the analysis. Developing or repurposing microfluidic devices to capture and analyze, specifically, tumor-derived LO, might clarify the function and clinical significance of these vesicles.

In summary, several studies on protein cargo have identified potential EV biomarkers. For most of these, it is unclear whether they are specific to a given EV population (Figure 2). Confirming these data in larger cohorts of patients, and determining whether one or more of these markers are associated with specific EV populations will provide significant advantages to the development of EV-based non-invasive tools for personalized medicine.

Extracellular nucleic acids

A few reports on DNA, and several reports on RNA in EVs have been recently published.73,74 The evidence that nucleic acids travel protected in EVs offers generous opportunities to nominate tumor type or disease-stage-specific variants or other genomic alterations, as well as aberrant transcripts or noncoding RNAs in EVs by massive parallel sequencing.

DNA. In 2011, Balaj et al.75 were the first to demonstrate that EVs contain DNA. A few years later, the presence of double-stranded DNA (dsDNA), reflecting the genomic DNA (gDNA) of the originating cell, was reported and low-coverage whole-genome sequencing demonstrated that the DNA contained in EVs spans all chromosomes.73,74 However, single primer PCR was necessary to detect specific DNA aberrations in plasma EVs. Thus, KRAS and PS3 mutations, which typically occur in more than 90% of patients with pancreatic cancer,76,77 were identified in exosomes from pancreatic cancer cells73 and Braf and EfrG mutations were identified in exosomes from melanoma and lung cancer cells.74 These seminal findings suggest that more sensitive methods for discovery will improve detection of EVs in which the tumor-derived DNA is highly concentrated. An interesting report from Lázaro-Ibáñez et al. recently tested whether different populations of EVs (exosomes, MV and apoptotic bodies) contain extracellular gDNA with different profiles, and PC-specific mutations.78 Additional studies are needed to understand the functional distribution of extracellular DNA in different EV populations that might accommodate different amounts ad types of DNA.

In order to render this effort clinically applicable, it will be essential to test the presence of frequent mutations in specific clinical groups, standardizing the methods and preserving the EV cargo. For example, because genomic alterations of the phosphoinositide 3-kinase (PI3K) axis, which includes PTEN mutations and deletions, have been identified in tumor tissues from 50% of patients with CRPC by whole-exome and RNA seq.79 this group of patients could be an ideal target to measure PTEN copy number as a EV-based circulating DNA assays. Collaborations with other fields will also help standardization of EV profiling. For example, it is still unclear what the best way to preserve EVs is and perhaps some insights can be derived from the CTC community. Similarly, the cfDNA field might inspire future development of EV assays, since they have established the computational pipelines to analyze highly fragmented plasma DNA. It is important to note that the size of DNA fragments in LO is significantly larger than in other types of EVs (unpublished observations), supporting further investigation in those EVs. We speculate that specific selection of these vesicles might improve designing the approach to detect tumor signal in complex biofluids, thus allowing to monitor the tumor genomic make up by liquid biopsy.

RNA. Donovan et al. recently developed an assay that reports a patient score (EXO106) based on mRNA levels of PC antigen 3 (PCA-3), ETS-related gene (ERG) and SAM pointed domain-containing Ets transcription factor (SPDEF) in urine-derived exosomes.80 In receiver operating characteristic (ROC) analysis, the EXO106 score predicted both PC and high-grade disease (Gleason score 7 or higher) with an area under the curve (AUC) of 0.764. Importantly, EXO106 showed its usefulness in the diagnosis of high-grade PC in patients in ‘grey zone’ serum PSA levels (>2 and <10 ng ml−1). In a follow-up study, the gene expression assay score of PCA-3, ERG and SPDEF assayed in urine exosomes (ExoDx Prostate Intelliscore) outperformed standard of care values. This combination resulted to be more predictive in discriminating PC patients with ≥GS7 from GS6 or with negative biopsy results (AUC 0.73) in multiple cohorts obtained from different sites. The addition of the ExoDx Prostate Intelliscore seems also to prove useful in identifying patients who are eligible to biopsy among the ones with elevated PSA levels.81 Nilsson et al. had previously examined PSA, PCA-3 and TMPRSS:ERG mRNA levels in urine-derived exosomes and demonstrated a correlation between these transcripts and high Gleason score.82

Candidate EV biomarkers for PC

| Proteins: | mRNA: | miRNA: |
|---|---|---|
| CD151 | GAPDH | mir-438-5p | PCA-3 |
| CDCP1 | LDHB | mir-34a | ERG |
| MFG-E8 | TM256 | mir-1290 | SPDEF |
| Notch3 | LAMTOR1 | PSAP | PSA |
| ADAM9 | PSA | TGM4 | TMPRSS:ERG |
| CLSTN1 | FOLH1/PSMA | Endophilin-A2 |  |
| GDF15 | TGM4 | MDR-1 and 3 |  |
| FASN | Survivin | δ-catenin |  |
| CLU | Cav-1 | Cav-1 |  |
| CYP17 | PSCD6IP/Alix |  |  |
| PDCD6IP/Alix | Survivin |  |  |
| CD9 | Survivin |  |  |
| ENO1 | Survivin |  |  |
| PSMA | Survivin |  |  |

Figure 2. Summary of candidate PC biomarkers (protein and nucleic acids) identified in EVs for which it is not clear if they are specific for given EV populations. EV, extracellular vesicles; PC, prostate cancer.
score as well as poor response to androgen deprivation therapy. Even if on a small sample size, the results of this study are remarkable and support investigations of the prognostic role of exosome-enclosed mRNA in larger cohorts of patients. Additional studies from Dijkstra et al. and Hendriks et al. have tested the suitability of using urine-derived exosomes for PC diagnosis. However, while the first study suggests the clinical usefulness of urine-derived exosomes, the second proposes whole urine as substrate for biomarkers analysis. Of note, both studies have attempted to perform extensive, unbiased screenings through array-based approaches to identify novel PC RNA biomarkers (Figure 2). These studies focus particularly on miRNAs, which are stable in human biological fluids because resistant to ribonuclease degradation. Korzeniewski et al. reported three miRNAs (miR-483-5p, miR-1275, miR-1290) as the most abundant miRNAs released from PC cells. When tested in urine samples, miR-483-5p alone as well as in combination with the other two miRNAs, was able to significantly differentiate PC patients with biopsy-proven tumor mass from patients with microscopic tumor. However, PSA was a better disease predictor, and overall, these miRNAs failed to show advantages over conventional methods. Corcoran et al. examined the potential use of exosomal miR-34a as a predictive biomarker of response to docetaxel. Among the miRNAs whose levels correlated with disease progression (miR-598, miR-34a, miR-146a, miR-148a), miR-34a showed the highest clinical relevance based on *in silico* analysis of publicly available datasets. α-synuclein and solute carrier family 7 member 5 (SLC7A5), which are two common targets of these four miRNAs, also correlated with prognosis in CRPC patients. These results collectively suggest that analyses of EV-bound miRNA might be more useful than analyses of whole miRNA fraction in biofluids (obtained without purifying EVs). However, comparative studies are necessary to make this conclusion.

A large-scale screening was performed by Huang et al., who reported that miR-1290 and miR-375 can be used as prognostic markers of CRPC. Rather than using array-based approach, RNA sequencing with short RNA libraries was conducted in 23 patients with CRPC to identify survival-related miRNAs in plasma exosomes. We believe this is the first RNA sequencing screening performed on plasma EVs in CRPC. Candidate miRNAs were further validated for their prognostic values using PCR in 100 patients. Plasma exosomal miR-1290 and miR-375 were chosen as the two most promising prognostic biomarkers. Interestingly, while miR-375 upregulation in tumor tissues correlated with PC progression, miR-1290 was not significantly upregulated in PC tissues. This is an interesting result suggesting that EV analyses could allow identification of cancer-derived molecules in biological fluids even if their levels are not increased in tumor tissue. Last but not least, little is known about the potential for use of exosomal long noncoding RNAs (IncRNA) as circulating diagnostic markers for PC. Isn et al. tested the feasibility of using IncRNA-p21 for the detection of PC. Urine IncRNA-p21 was detected increased levels of EpCAM and PSMA in patients compared to controls starting from a 25 μl of plasma. An aqueous two-phase system was recently proposed for EV isolation from plasma of patients with PC (*n* = 82). Also, in this study, PSMA-positive EVs were associated with high-risk PC and higher risk of biochemical failure. Flow cytometric analysis of EVs has made significant advances. In particular, the use of microflow technologies (Apogeeflow.com) has enabled the direct analysis of EVs of various sizes. The recent inclusion of standardization, particles has been particularly important for achieving consistent detection and characterization of each EV population. The inclusion of fluorescent detection with small and wide-angle scatter enables microflow technologies to interrogate each particle with fluorescent antibodies, thereby bringing the advantages of traditional flow cytometry to the analysis of EVs.

**CONCLUSION**

The EV field is developing rapidly to try to better understand the biology and significance of the cargo with raising opportunities to use EV as biomarkers. Keys to success will be the miniaturization of the assays as well as standardization of EV purification and analysis.
systems. These recent developments hold high potentials for applicability of EV profile to clinical settings if validated in larger cohorts of patients. Indeed, preliminary investigations in different tumors like glioblastoma, colorectal cancer and ovarian cancer demonstrate that these techniques can discriminate patients from controls, and also identify patients that respond to therapy from the ones who do not.

CONFLICT OF INTEREST
The authors declare no conflict of interest.

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