The Detection of Tear Biomarkers for Future Prostate Cancer Diagnosis

Yong Li* a,b

*Cancer Care Centre, St George Hospital, Gray St, Kogarah, NSW 2217, Australia; bSt George Clinical School, Faculty of Medicine, University of New South Wales, Kensington, NSW 2086, Australia

Abstract: Prostate cancer (CaP) continues to be the second leading cause of cancer-specific death in men in Western countries. The marker currently used for CaP detection is an increase in serum prostate specific antigen (PSA). However, the PSA test may give false positive or negative information and does not allow the differentiation of benign prostate hyperplasia (BPH), non-aggressive CaP and aggressive CaP. Tears are a unique source of body fluid and contain proteins, peptides, mucins and lipids, which is useful for studying clinical proteomics. Advances in the field of proteomics have greatly enhanced the study of tears, with a greater number of proteins now being identified in tears. Identification of novel biomarkers in tear is a new area of development. Modern advances in the field of proteomic techniques hold the promise of providing the clinical oncologists with new tools to find novel CaP biomarkers for early diagnosis and prognosis.

Keywords: Prostate cancer, tear, biomarker, proteomics, diagnosis.

INTRODUCTION

Prostate cancer (CaP) is the most common malignancy and is the second leading cause of cancer death in males [1]. In 2009 alone, it was estimated that CaP would be diagnosed in 192,000 men, and 27,300 would die of the disease in the United State [2]. The marker currently used for CaP detection is an increase in serum prostate specific antigen (PSA) [3]. However, the PSA test may give false positive or negative information and does not allow the differentiation of benign prostate hyperplasia (BPH), non-aggressive CaP and aggressive CaP [3]. The classical value of a PSA cut-off, 4.0 ng/mL, misses more than 75% and a PSA cut-off of 2.0 ng/mL would miss more than 45% of biopsy detectable cancers [4]. Using PSA ranges below 3.0 ng/mL entails a very high risk of overdiagnosis and overtreatment as well as unnecessary biopsies [4]. These observations have also been confirmed by a recent report that found that serum PSA was related to the volume of the prostate; however, with population-based screening, it has become more inaccurate as a marker of cancer and may only relate to BPH [5, 6]. Furthermore, there are emerging data from the Prostate Cancer Prevention Trial and other sources showing that a significant number of men with PSA scores <4 ng/mL may have undetected CaP [7] while as many as 75% of men who have a positive PSA test (> 4.0 ng/mL) do not have CaP [8]. Such reports imply that a single marker will accurately reflect the complex phenotypic changes associated with the development of cancer. Clearly, there is an urgent need for novel biomarkers or protein profiles in body fluids of CaP subjects to improve the early detection and accuracy of diagnosis, determine the aggressiveness of CaP and monitor the efficacy of treatment.

TEARS AS A SOURCE FOR BIOMARKERS

Tears are a unique source of body fluid and contain mucins, glycoproteins, unglycosylated proteins, peptides, and lipids [9]. The major tear proteins are lysozyme, lactoferrin, secretory immunoglobulin A (sIgA), lipocalin, and lipophillin. The diverse composition of the tear fluid may be associated with variations in response to pathophysiological conditions which affect other areas of the organism, making the tear fluid useful for biomarker research. The change of protein components of tears has been reported in some local eye diseases [10, 11], systemic diseases such as diabetes [12] and cancers [13, 14]. The tear proteome has been proposed to contain around 500 proteins [15], whereas that of serum/plasma is estimated containing up to 10,000 proteins [16]. A much lower range of proteins and proportion of serum albumin and immunoglobulins in tears compared to serum makes proteomic analysis more straightforward. Proteomic analysis of tears has been carried out in our laboratories [13, 14, 17]. These results offer the potential for cancer diagnosis by analysing tear fluid proteins.

The collection of tears is relatively safe (e.g. no puncture), non-invasive, inexpensive, and the tears may be collected repeatedly with minimal discomfort to the patients. There is increasing interest in using tears to diagnose systemic diseases because of its simplicity in collection. In addition, tear contains constituents that are frequently altered in the presence of systemic diseases such as cancers, thereby rendering tears as a very desirable source for cancer diagnosis and monitoring.

However, there are still some challenges using tears as source for biomarker identification. Firstly, analysis of tear fluid samples for protein biomarker discovery is complicated by the differences in composition associated with these different collection methods, highlighting the requirement of a standard collection protocol for biomarker research. Another challenge is the requirement of highly sensitive and reliable techniques for analysis of the small amounts of tears able to
be collected to ensure reproducible and quantifiable results. With the development of very sensitive proteomic techniques such as various mass spectrometries that require only 1-5 microlitres of tear, tears will be considered as a useful source of biomarkers for various diseases.

TEAR BIOMARKERS IN CANCER RESEARCH

Using two-dimensional gel electrophoresis (2DGE) technique, our research team found that lactoglobin (mammaglobin, a low molecular weight protein) could be detected in a breast cancer patient with bone metastasis but not in healthy controls, suggesting that lactoglobin in human tears may be a potential marker for breast cancer [13]. Later on, mammaglobin was further confirmed to be a potential marker for metastasis of breast cancer cells to lymph nodes [18].

The surface-enhanced laser desorption/ionization time-of-flight mass spectrometry (SELDI-TOF-MS) ProteinChip has been introduced [19] as an alternative to 2DGE. This technology utilizes affinity surfaces to retain proteins based on their physical or chemical characteristics, and is followed by direct analysis using TOF-MS. It is a rapid and reproducible technique to generate protein expression profiles known as "phenomic fingerprints". Furthermore, SELDI-TOF-MS is more sensitive and requires only small amounts of sample (2-3 μL) compared to 2DGE. This system has enabled detection of critical proteins directly from crude mixtures without labour intensive pre-processing and has been proven to be a very useful tool for identifying biomarkers in various cancers as well as other diseases.

Lebrecht et al. (2009) recently demonstrated a biomarker panel from tear fluids successfully generated by SELDI-TOF-MS allows breast cancer patients to be differentiated from healthy women [20]. The same groups further confirmed that the diagnostic pattern differentiated cancer patients from controls with a specificity and sensitivity of approximately 90% in tear fluid using the same technique [21]. These results indicate that tears may be a potential source of biomarkers for breast cancer. We have analyzed pooled tear proteins from CaP patients (n=8) and normal controls (n=3) using SELDI-TOF-MS. A reversed-phase surface protein chip (H50) was used to analyze the samples. Our results indicate that there are two peptides missing (7110 and 14213 Da) in the tears from CaP patients compared with normal, and have shown the potential of proteomic analysis of tears as a possible new way for finding novel CaP biomarkers by SELDI-TOF [14].

However, whilst this approach has been used in multiple studies, the major disadvantage is that significant further work is necessary to identify and verify biomarker proteins that are detected in the peptide fingerprint.

KEY PROTEOMIC TECHNOLOGIES FOR FUTURE TEAR BIOMARKER RESEARCH

Proteomic studies are the emerging era for searching biomarkers. Key elements of proteomic studies are the ability of separating thousands of peptide/proteins simultaneously, comparing protein profiles from different samples and indentifying proteins. Several technologies are important for this study and they include gel based technologies and non-gel based technologies (Mass spectrometry-MS).

Two-Dimensional Differential in-Gel Electrophoresis (2D-DIGE)

2D-DIGE is a gel-based and fairly recent improvement of the 2DGE technology using fluorescent molecules (CyDyes), or fluors, that are used to pre-label samples prior to separation by 2DGE. The internal standard is a pool of all samples used in the experiment and is used not only to normalize the Cy3- and Cy5-labeled samples, but also to compare across the other 2D gels in the experiment. This significantly reduces the intergel variability that plagues other 2D gel-based analyses. CyDyes also show a linear response to protein concentration up to 5 orders of magnitude [22]. The gel is scanned at different emission wavelengths and multiple images corresponding to the different samples are generated. This methodology significantly improves sample throughput and greatly enhances gel reproducibility [23]. Our preliminary results indicate that the 2D-DIGE technique is promising for CaP tear biomarker identification in the future.

Isobaric Tags for Relative and Absolute Quantitation (iTRAQ)

Due to better sensitivity and reproducibility over 2D-based methods [24], mass spectrometry (MS)-based technology is one of the most powerful tools in analyzing proteome. iTRAQ is a shot-gun based technique developed by Darryl Pappin and colleagues at Applied Biosystems in 2004 [25] which allows the concurrent identification and relative quantification of hundreds of proteins in up to 8 different biological samples in a single experiment [25, 26]. This unique approach labels samples with 8 independent reagents of the same mass that, upon fragmentation in MS/MS, give rise to 8 unique reporter ions (m/z =113–121) that are subsequently used to quantify the 8 different samples, respectively. MS/MS fragmentation, in addition to giving strong reporter ion signals, also yields strong signature y- and b- ions without changing the charge state of any given peptide to allow for more confident protein identification simultaneously with the quantification. iTRAQ based quantitative proteomics is a promising approach for global comparison of protein expression in relatively small amounts of samples such as tear. The iTRAQ technology has many advantages over other proteomic techniques:

- iTRAQ reagents allow parallel proteomic analysis of 8 different samples under the same experimental conditions, resulting in reduced systematic error and increased electrospray ionization efficiency leading to higher sensitivity; reducing the analysis time.
- Protein sequence coverage obtained using iTRAQ reagents is similar to that obtained using other shot-gun proteomic approaches.
- iTRAQ reagents can be used to identify and quantify proteins across diverse molecular weight (MW) and pI ranges, functional categories, cellular locations and abundances.
- The multiplexing capacity of these reagents allows information replication within certain liquid chromatography/ mass spectrometry (LC-MS/MS) experimental regimes, providing additional statistical validation within any given experiment. iTRAQ is relatively high throughput due to sample multiplexing.
Because protein identification and quantification is based on tandem mass spectrometric (MS/MS) evidence, increased selectivity, specificity, and confirmatory power are achieved [25].

Wu et al. compared 2D-DIGE, identified by isotope-coded affinity tag (ICAT), and iTRAQ to show that iTRAQ is the most sensitive proteomics quantitation method among the three techniques evaluated, based on the number of detected peptides [27]. iTRAQ was observed to provide a more consistent quantitation in comparison with 2DGE for proteins stained with lower intensity on 2DE gels [28]. Zhou et al. (2009) used iTRAQ technology to identify 10 potential tear protein biomarkers for dry eye syndrome: α-enolase, α-1-acid glycoprotein1, S100A8 (calgranulin A), S100A9 (calgranulin B), S100 A4, S100 A11 (calgizzarin), prolactin-inducible protein (PIP), lipocalin-1, lactoferrin and lysozyme. In particular, α-enolase and S100 A4 were successfully verified by enzyme-linked immunosorbent assay (ELISA) [29]. This technique is promising for future CaP tear proteomic studies.

**Multiple Reaction Monitoring (MRM)**

MRM-MS has emerged as an alternative and a highly specific, sensitive technology to immunoassays for quantification of target proteins or peptides. It offers superior multiplexing capabilities, allowing for the simultaneous quantification of numerous proteins in parallel. MRM achieves high analytical specificity and sensitivity by selecting predetermined precursor molecular ions for collision-induced dissociation and monitoring the appearance of several diagnostic product ions. When identified in combination, the precursor and product ions confirm the presence of the analyte of interest. Importantly, addition of isotopically labeled internal standards allows absolute quantification. Numerous precursor ions can be monitored in 1 experiment, particularly when using scheduled MRMs, which incorporate prior knowledge of peptide elution times into the LC-MS/MS program [30, 31].

Recently, a large National Cancer Institute–sponsored interlaboratory study showed that an MRM approach can quantify target proteins in a background of unfractionated human plasma with highly robust and reproducible results, supporting the notion that this technology is suitable for biomarker discovery [32].

Under MRM, the mass spectrometer is set up to monitor only specific mass/charge (m/z) values of interest; as a consequence, the probability of detecting even low levels of a peptide in the presence of a complex mixture of peptides is much higher. MRM has been the method of choice for quantification of low levels of small molecules, including drugs or metabolites, as well as peptides [30, 33]. Since a specific antibody does not need to be developed for MRM, this technique can be quickly used to verify candidate biomarkers and shorten the advance to validation stages using MRM or ELISA. An attribute of MRM is its multiplexing capability for many target proteins per run; antibody-based assays measure a limited number of proteins simultaneously. MRM would be the inevitable choice for biomarker verification when there is no appropriate antibody. Use of MRM in biomarker development also can enhance the measure of post-translation modifications (PTMs), which are difficult tasks for antibody-based system [34-36].

Using sequential dyes to analyze phospho-, glyco- and total tear protein profiles (Pro-Q Diamond for phosphoprotein, Pro-Q Emerald for glycoprotein and Sypro Ruby for total protein in 2DGE, our research team have recently identifies a novel tear protein, dermecidin [37]. The identified dermecidin was further validated by MRM. This study provides the groundwork for understanding the PTM of tear proteins and consequently these methods could be useful in the search for CaP biomarkers in tears. Studies of several potential tear biomarkers in CaP patients, BPH patients and control subjects by MRM are currently being investigated in our laboratory.

**CONCLUSIONS**

Proteomic techniques are modern and emerging technologies used to study proteins in biological samples. The techniques can separate, simultaneously, thousands of proteins in samples and the proteins can be sequentially identified. Tear is a unique source of body fluid and contains proteins and peptides, and can be collected as a non-invasive procedure. Identification of novel biomarkers in tear is a new area of development. Identification of tear proteins may lead to the discovery of novel biomarkers for cancer diagnosis or monitoring cancer progression. The biomarkers could be useful to determine whether a patient's cancer is surgically curable and predict the future clinical course of CaP and/or the response to future chemotherapy.

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**ABBREVIATIONS**

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<th>Abbreviation</th>
<th>Definition</th>
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<tr>
<td>BPH</td>
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<td>CaP</td>
<td>Prostate cancer</td>
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<td>2D-DIGE</td>
<td>Two-dimensional differential in-gel electrophoresis</td>
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<tr>
<td>LC-MS/MS</td>
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<td>MRM</td>
<td>Multiple reaction monitoring</td>
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MS = Mass spectrometry
MW = Molecular weight
PSA = Prostate specific antigen
PTM = Post-translation modification
SELDI-TOF-MS = Surface-enhanced laser desorption/ionization time-of-flight mass spectrometry

REFERENCES