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Identification of diagnostic upper gastrointestinal cancer tissue type-specific urinary biomarkers

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Abstract. Several potential urinary biomarkers exhibiting an association with upper gastrointestinal tumour growth have been previously identified, of which S100A6, S100A9, rabenosyn-5 and programmed cell death 6-interacting protein (PDCD6IP) were further validated and found to be upregulated in malignant tumours. The cancer cohort from our previous study was subclassified to assess whether distinct molecular markers can be identified for each individual cancer type using a similar approach. Urine samples from patients with cancers of the stomach, oesophagus, oesophago gastric junction or pancreas were analysed by surface-enhanced laser desorption/ionization-time-of-flight mass spectrometry using both CM10 and IMAC30 (Cu2+-complexed) chip types and LC-MS/MS-based mass spectrometry after chromatographic enrichment. This was followed by protein identification, pattern matching and validation by western blotting. We found 8 m/z peaks with statistical significance for the four cancer types investigated, of which m/z 2447 and 2577 were identified by pattern matching as fragments of cathepsin-B (CTSB) and cystatin-B (CSTB); both molecules are indicative of pancreatic cancer. Additionally, we observed a potential association of cystatin-B (CSTB); both molecules are indicative of pancreatic cancer. Furthermore, the potential pancreatic cancer biomarkers CSTB and CTSB were validated independently by western blotting. Therefore, the present study identified two new potential urinary biomarkers that appear to be associated with pancreatic cancer. This may provide a simple, non-invasive screening test for use in the clinical setting.

Introduction

A significant number of cancer-related deaths worldwide are associated with malignant tumours of the upper gastrointestinal (GI) tract, such as the stomach, oesophagus, oesophago gastric junction (OGJ) (1) and pancreas. The Global Burden of Disease study undertaken in 2015 identified these types of cancer to have a very poor prognosis, with gastric and oesophageal cancers contributing to 10.3 and 5.4%, respectively, of all cancer deaths globally, with both exhibiting a male preponderance (2). Pancreatic cancer is associated with a dismal 5-year survival rate of 3% and its incidence appears to be increasing annually (3,4). This has led to the development of various therapeutic strategies to prolong survival, which have had limited success, including improved surgical techniques, anti-angiogenesis therapies and adjuvant/neoadjuvant chemoradiotherapy (5,6). Pancreatic cancer cases in particular are usually diagnosed at an advanced stage and, therefore, the possibility of a non-invasive intervention to halt tumour progression is greatly reduced. Therefore, it is crucial to devise methodologies to not only pharmacologically treat, but also to diagnose this type of cancer at an early stage.

The most successful and widely used cancer assays to date are based on the detection and quantification of glycans in the serum using antibodies against CA19.9 (7) or CA125 (8), a pan-cancer marker that targets carbohydrate-associated epitopes of immunoglobulin heavy chains (9). CA19.9, with a reported sensitivity of 90-100% and specificity of 70-98% for pancreatic cancer detection (10), is the best biomarker assay currently in clinical use. However, the positive predictive value of CA19.9 for detecting pancreatic cancer is only 0.9% in the asymptomatic population due to a dependency of blood-group markers where CA19.9 can be used (11). It is also associated with biliary obstruction (12) and has been proven unable to distinguish pancreatic cancer from matched controls in larger studies, highlighting its poor clinical utility as a tumour marker (13).
Other potential cancer protein biomarkers have been identified by antibody arrays, such as H2B histone using the IPO-38 antibody to define gastric cancer (14), M2 pyruvate kinase in faeces as a GI cancer marker (15) and tumour-associated trypsin inhibitor as a marker of liver metastasis and colorectal cancer (16), and by mass spectrometry, such as S100A9 for upper GI cancer (17), as well as S100A6 for a wide array of tissue cancers, including thyroid (18), gastric (19), ovarian (20), hepatocellular (21), bowel (22), breast (23) and upper GI (17) cancers.

Numerous potential biomarkers were also identified by gene array screens of diseased tissue (24-27) and by microRNA array screens of circulating biofluids (28), in the most frequent cancers affecting the GI tract (29,30). The latter is considered as a promising source of clinically relevant biomarkers, when compared with tissue gene array screens, since the use of biopsies as a predictive diagnostic tool remains unrealistic for use in the clinical setting. Additionally, prediction of several cancer types based on serum metabolic profiles is also feasible (31). Concomitantly, collaborative efforts have been made to promote the detection of plasma-derived metabolite markers for pancreatic cancer diagnosis (32,33).

There is a clear need to simplify the medium to be screened due to genetic variations and population heterogeneity in order to define reliable disease markers. A substantially less complex system, such as the urine, which contains ~5,000 proteins (34), would be a preferred medium to screen for protein or peptide biomarkers. This has a number of advantages, including non-invasive sampling for patients, ease of sampling, and unrestricted availability under normal conditions (35). Urine itself is also relatively stable in terms of protein and peptide composition and fragmentation state compared with other body fluids, such as the serum, where proteolytic degradation by endogenous proteases has been shown to occur during or after sample collection (36).

In the present study, surface-enhanced laser desorption/ionization-time-of-flight mass spectrometry (SELDI-TOF-MS) was used to screen human urine samples from patients with upper GI tumours to establish biomarker patterns using the CM10 and IMAC30 chip types, as well as independent LC-MS/MS mass spectrometry screening and a combined bioinformatics data analysis, in order to present a potentially useful approach to diagnosing upper GI tissue type-specific cancers in humans using novel potential biomarkers.

Materials and methods

Materials. All buffers, gels and SELDI chips were purchased from Bio-Rad Laboratories Ltd. (Hemel Hempstead, UK), and all other chemicals were obtained from Sigma-Aldrich; Merck KGaA (Gillingham, UK), unless stated otherwise in the text.

Sample collection. Urine samples were obtained from 83 patients with upper GI cancers undergoing potentially curative resection. The participant demographics are summarised in Table I and provided in detail in Table SI. The participants’ age ranged between 43 and 83 years. Fasting urine samples were obtained at induction of anaesthesia. One-third of the patients had pancreatic tumours, one-third had oesophageal cancer, approximately one-sixth had malignancies of the OGJ and one-sixth suffered from gastric cancer. All procedures were approved by the local research ethics committee and written informed consent was obtained from the patients. The study conformed to the standards set by the Declaration of Helsinki. All urine samples were stored at -40°C. Long-term storage of samples (>1 month) was at -80°C.

SELDI-TOF-MS. SELDI chips (CM10 and IMAC30) were prepared for sample application according to the manufacturer's recommendations. Briefly, IMAC30 chips were loaded with 0.1 M CuSO4, washed with water, neutralised with 0.1 M NaHAc (pH 4.0) and again washed with water, followed by two washes with 0.1 M NaHPO4 and 0.5 M NaCl. CM10 chips were washed twice with 0.1 M NaHPO4 (pH 4.0). All chips were processed in a bioprocessor assembly by incubating 0.1 ml urine and 0.1 ml binding buffer [IMAC30: 0.1 M NaHPO4, 0.5 M NaCl; CM10: 0.1 M NaHPO4 (pH4.0)] for 1 h at room temperature with vigorous shaking, followed by three washes with 0.2 ml binding buffer for 5 min each at room temperature with vigorous shaking and two washes with 0.2 ml water at room temperature with vigorous shaking. All chips were removed from the bioprocessor assembly, air-dried and 1 µl energy-absorbing matrix [a saturated solution of sinapinic acid in 50% acetonitrile (ACN) and 0.5% trifluoroacetic acid] was added twice. Air-dried chips were analysed in a PCS4000 SELDI-TOF instrument (Bio-Rad Laboratories, Ltd.) by measuring the 1,000-25,000 Da range with a low laser setting of 2.5 µJ, and spectra were exported as ‘.xml’ files. The SELDI instrument was calibrated using the ProteinChip All-In-one peptide standard (Bio-Rad Laboratories, Ltd.). The source voltage was 25,000 V and the detector voltage was 2,946 V. Quality control and consistency were ensured by using one random pool of urine samples on one spot per chip each. Spectra of the full analysis were recorded in two large batches to minimize instrument variability and drift. Spectral alignments of all quality controls ensured consistency of all spectra.

SELDI-TOF-MS data processing. ProteinChip Data Manager Software version 4.1 with integrated Biomarker Wizard cluster analysis (Bio-Rad Laboratories, Ltd.) was used for analysis. SELDI-TOF-MS traces were split into the four cancer type groups. The baseline was subtracted from individual m/z traces and the profiles were normalised using total ion current, followed by identification of peak clusters using the cluster analysis tool. Peaks were selected in the first pass with a signal-to-noise (S/N) ratio of >5 and a valley depth of at least 3, and in the second pass with a S/N of 2 and a valley depth of 2. The cluster mass window was set to 0.2% of the mass. Clustered peaks were only included if they occurred in at least 10% of all spectra. The resulting P-values, mean and median m/z values, and the intensities of the clustered peaks were exported and saved as ‘.csv’ files. A two-sample t-test was used to compare mean normalized intensities between the groups. The P-value was set at 0.05 to indicate statistically significant differences. Clustered peak lists were analysed with the Biomarker Pattern Software (Bio-Rad Laboratories, Ltd.) and m/z vs. intensity matrices were analysed using decision tree analysis, selecting the...
standard error rule of minimum-cost tree regardless of size, and using the Gini method. V-fold testing was set to 1,000. All samples from one cancer type were used to build exploratory networks by using all other cancer samples as ‘negative’ controls. Tree model building was performed using a selected m/z peak as the only deciding factor to obtain sensitivity and specificity values. Sensitivity was defined as the probability of predicting specific cancer cases, and specificity was defined as the probability of predicting other cancers.

Peak isolation and identification by LC-MS/MS and Mascot searching. Peaks observed in the CM10 and IMAC30 chip types (Table SIII) that exhibited marked expression differences between tissue-specific cancer samples, and statistical significance in the cluster analyses with P<0.05, were further investigated. Urine (0.5 ml) from positive or negative samples in relation to specific peaks was added to 30 µl CM10 or IMAC30 (Cu²⁺-complexed) spin column resin (Bio-Rad Laboratories, Ltd.) and 0.75 ml binding buffer [0.1 M NaHPO₄ (pH 4.0) for CM10 resins, and 0.1 M NaHPO₄ (pH 7.0) including 0.5 M NaCl for IMAC30 resins] and incubated for 1 h at room temperature under constant agitation. Unbound material was removed and the resin was washed four times with 0.3 ml binding buffer. Bound material was separated by electrophoresis on a 16.5% Tris-Tricine gel (Bio-Rad Laboratories, Ltd.), and gel bands in the region of 2-10 kDa were excised following Coomassie staining (BioSafe Coomassie; Bio-Rad Laboratories, Ltd.). Positive and negative samples were both selected on the presence and absence of a specific m/z peak to be identified based on SELDI-TOF-MS analysis. Proteins and peptides from gel bands were digested in situ with trypsin, the resulting peptides eluted with ACN, and analysed by LC-MS/MS as described previously (17). Data-dependent acquisition was controlled by Xcalibur software and fragmentation spectra were then processed by Xcalibur and BioWorks software (Thermo Fisher Scientific, Inc., Loughborough, UK) and submitted to the Mascot search engine (Matrix Science, London, UK) using UniProt/SwissProt (release July 2010, Homo sapiens, 18055 sequences) as the reference database. The Mascot search parameters were as follows: Enzyme specificity, trypsin; maximum missed cleavage, 1; fixed modifications, cysteine carbamidomethylation; variable modification, methionine oxidation; precursor mass tolerance, ±3 Da; and fragment ion mass tolerance, ±0.4 Da. Only Mascot hits with a false discovery rate of <0.05 were taken into consideration.

Mascot-SELDI matrix matching. Observed proteins with at least two peptide matches from the LC-MS/MS analysis were then further analysed by pattern matching based on SELDI-TOF-MS measured expression levels of peaks of interest (expected abundance in selected samples). This was performed using software written in-house, which compares observed protein expression patterns in a pre-defined set of samples (LC-MS/MS results) against a matrix of peak patterns (SELDI-TOF clustered peak intensities, where estimated peaks are set to null) in the same set of samples. The scoring is based on sensitivity (percent observed over expected) and specificity (percent not observed over not expected), and the results are presented in descending order of cumulative scores.

Western blotting and validation of Mascot search results. Cross-validation of identified peaks was performed by western blotting of raw urine samples for anti-cathepsin B (CTSB) and anti-cystatin B (CSTB) blots (20 µl per sample) using standard protocols (37). The antibodies used were rabbit anti-human CTSB (G-60; 1:1,000; cat. no. 3373; Cell Signaling Technology, Danvers, MA, USA), rabbit anti-human serum albumin (1:1,000; cat. no. A3293; Sigma-Aldrich; Merck KGaA), mouse anti-human CSTB (1:400; cat. no. sc-101510; Santa Cruz Biotechnology, Santa Cruz, CA, USA), and the peroxidase-coupled secondary antibodies were from Upstate (Lake Placid, NY, USA), used at a dilution of 1:5,000. Detection of signals was performed by chemiluminescence using ECL western blotting reagents (Thermo Fisher Scientific, Inc., Cramlington, UK).

Results

Urinary screening via SELDI-TOF-MS analysis. Urine samples from 83 patients diagnosed with various types of upper GI cancer were analysed in the course of this study (Table I). A full demographic with additional clinical data is provided in Table SI. The mean age of the participants was 65 years, 34% of the cohort suffered from pancreatic cancer, 33% from oesophageal cancer, 18% from gastric tumours and 15% from cancer of the OGI. SELDI-TOF-MS analysis was selected to screen the samples mentioned above based on our previous study, where we described global specific markers for upper GI cancer (20). We found that both the metal chelator resin IMAC30 (Cu²⁺-chelated) and the weak cation exchanger CM10 yielded the best and most reproducible results. All samples were measured by SELDI-TOF-MS to obtain a peak pattern of identified molecular constituents, allowing us to stratify tissue type-specific cancers. We selected to compare only samples

| Cancer type       | Oesophagus | Pancreas | OGG | Gastric | Total |
|-------------------|------------|----------|-----|---------|-------|
| No. of patients   | 27         | 28       | 13  | 15      | 83    |
| Mean age (years)  | 66.5 (10.8)| 63.5 (9.1)| 61.5 (8.1)| 70 (7) | 65.3 (9.5) |
| Male              | 22         | 16       | 12  | 9       | 59    |
| Female            | 5          | 12       | 1   | 6       | 24    |

Table I. Demographics of the cohort used in this study (n=83).
from cancer patients against each other to circumvent the issue of accidentally tracking lead markers that may be associated with additional underlying conditions, such as inflammatory responses, which are commonly observed in cancer patients.

SELDI analysis of the CM10 chip type-based screen of the 83 cancer urine samples resulted in 9,379 peaks, and the IMAC30 chip analysis provided a cumulative peak list of 3,346 features (Table SII). Clustering of observed peaks using the thresholds described in Materials and methods resulted in 328 cluster peaks for the CM10 chips, and the IMAC30 chip type gave yield to 92 common peak clusters above the set thresholds (Table SIII). Both analyses were performed by omitting estimated peaks in order to restrict and raise the specificity of potential marker peaks and, therefore, only included well-defined and separated individual peaks from all spectra. Statistical analysis revealed that 8 peak clusters are potentially associated with the various cancer types, namely m/z 2444 and 2557 for pancreatic cancer in the IMAC30 set, and m/z 2447 and 9618 in the CM10 chip-based set for the same cancer type, m/z 5511 and 4908 for OGI cancer and m/z 4639 for gastric cancer in the IMAC30 chips, and m/z 4141 for oesophageal cancer in the CM10 chip set (Fig. 1; Table II). All potential markers have P<0.05 and exhibit upregulation in the associated disease. It is noteworthy to mention that any given peak cluster can comprise more than one molecular entities. Therefore, it is possible that both the 2444 and 2447 m/z peaks share an overlap in their molecular constituents; however, they include other peptides or proteins which are unique in this specific peak cluster based on the sensitivity and specificity distribution. The frequency distribution analysis of those 8 peaks shows that the m/z 9618 cluster from the CM10 chip screen displays the best stratifier for pancreatic cancer (Fig. 1D), with a sensitivity of 61% and a specificity of 82% in upper GI cancer cases. Comparison of all 8 peaks in non-cancer control cases measured in our previous study (17) also demonstrated that the m/z 9618 peak cluster exhibits the same frequency distribution in both non-cancer and non-pancreatic cancer cases of ~30%, whereas this frequency is doubled in pancreatic cancer for this peak cluster. The best predictor, based on a low frequency in non-cancer cases, was the m/z 2577 cluster in pancreatic cancer (Fig. 1B), with a 79% frequency in this cancer type, a 62% frequency in other cancers, and a 10% occurrence in non-cancer patient urine, with a sensitivity of 86% and a specificity of 44%. The elevated amount of the observed m/z 4141 peak cluster associated with oesophageal cancer (Fig. 1H) demonstrates the best specificity of 100%, but exhibits a relatively poor sensitivity of only 19% due to the relatively low frequency of one in four samples where this peak can be measured. Low frequency values were also observed for the OGI cancer-associated elevated peak cluster at m/z 4908 (Fig. 1F), with good sensitivity and specificity values of 92% and 66%, respectively. A more pronounced potential OGI marker in terms of quantitative changes was measured at m/z 5511 (Fig. 1E) with a high specificity (94%), but a relatively low sensitivity (38%).

We then endeavoured to identify the molecular identity of the m/z 2447 peak, since it showed the highest sensitivity of 100% in the detection of pancreatic cancer. This was performed by selecting 8 samples, of which 4 either contained the cluster peak at m/z 2447, or did not show the peak

| Table II. Summary of potential lead candidates for specific cancer types. |
|-----------------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|
| Chip | Cancer | m/z | P-value | Fold-change | Sensitivity | Specificity |
| IMAC | 2444 | Pancreatic | 0.04 | 1.5 | 54 | 76 |
| IMAC | 2577 | Pancreatic | 0.0007 | 1.9 | 44 | 44 |
| CM | 2447 | Pancreatic | 0.01 | 2.1 | 15 | 30 |
| CM | 9618 | OGI | 0.01 | 1.5 | 61 | 31 |
| IMAC | 5511 | OGI | 0.029 | 2.5 | 82 | 94 |
| IMAC | 4908 | Gastric | 0.027 | 2.5 | 38 | 94 |
| IMAC | 4639 | Oesophageal | 0.002 | 1.5 | 43 | 66 |
| IMAC | 4141 | Oesophageal | 0.002 | 1.8 | 57 | 29 |

Fold changes are in comparison to next highest mean peak in other cancer types. Sensitivity and specificity values were calculated by best decision tree analysis using the marker as sole decider, and the SELDI normalised cut-off value are the associated mass spectrometric intensity values where the associated sensitivity and specificity values are met. The number of non-cancer controls was 94. OGI, oesophagogastric junction; SELDI, surface enhanced laser desorption/ionization.
The samples were individually batch-absorbed on CM resin; bands in the 2-4 kDa range were excised after peptide gel electrophoresis and processed as described previously (17). The molecular cluster of interest was shown to consist of fragments from CTSB, $\alpha$-1-antichymotrypsin precursor and immunoglobulin $\gamma$ and $\kappa$ chains (Table III). The same result was obtained using an independent approach, as described below.

We then analysed the molecular constituents of human cancer patient urine in the 2-10 kDa molecular weight range using chromatographic protein and peptide enrichment on CM10 and IMAC30 resins, followed by gel separation, trypsin digestion and LC-MS/MS fragmentation, as before. Samples were selected based on the SELDI-TOF analysis results to either contain the 8 m/z peaks of interest (50% of samples per subset) or not in subsets of groups of 16 samples each. This resulted in the analysis of 145 urine samples, including repeats, from 62 upper GI cancer patients, of which 42 unique patient urines were enriched on CM10, and 40 unique patient samples on IMAC30 chromatography resins. 950 non-redundant proteins were identified in the CM10 resin-based approach by Mascot searching, and 600 unique proteins could be observed in the IMAC30-based enrichment, totalling 1,228 unique and non-redundant proteins in the combined datasets (Table SIV).

Protein expression pattern matching (i.e., molecules identified by LC-MS/MS and Mascot searching, and matching to peak expression patterns observed by SELDI-MS) was performed by using an automated computer program written in-house based on the filtered Mascot dataset, where each individual identification was based on Mascot scores >16 and consisting of at least 2 peptides each. All proteins were identified in at least three independent samples. Table III lists all found by this approach, including a list of publications describing the relevance of these molecules in tumour growth. A fully detailed list of these molecules, including peptide sequences, is supplied in Table SV.

Both CTSB and CSTB were also cross-validated by western blot analysis (Fig. 2), and both molecules exhibited a very good correlation to the expected cluster-peak pattern observed by SELDI analysis, thereby validating both the predictor model and the Mascot identification, as well as the SELDI peak clustering of these molecules, and may represent viable prognostic biomarkers for pancreatic cancer.
Table III. List of potential biomarkers associated with specific cancer types.

| m/z peak | Chip type | Cancer type | Protein ID | Protein name | Mass (kDa) | Mascot score | Mean number of peptides | Mean emPAI | % sequence coverage | Expected % sequence coverage | PMID | Pattern matching | Mascot-SELDI pattern matching score |
|----------|-----------|-------------|------------|--------------|------------|--------------|------------------------|------------|---------------------|-------------------------------|------|-----------------|-------------------------------|
| 2444 IMAC30 | Pancreatic | AACT_HUMAN | α-1-antichymotrypsin precursor | 48 145 8.4 0.26 13.4 | 11.5 | 16212428, 15709178 | A | 56/75 |
| 2577 IMAC30 | Pancreatic | CYTB_HUMAN | Cystatin-B | 11 77 7.4 1.57 | 41.7 | 50.0 | 18754876 | A | 50/100 |
| 2447 CM10 | Pancreatic | CATB_HUMAN | Cathepsin B precursor | 39 44 3.7 0.1 | 6.1 | 14.1 | 15367886, 11185708 | A, M | 50/100 |
| 9618 CM10 | Pancreatic | KV304_HUMAN | Ig κ chain V-III region Ti | 12 117 5.7 0.83 | 38.9 | 45.8 | 18615426, 20571237 | A | 61/63 |
| 5511 IMAC30 | OGJ | PDC6L_HUMAN | Programmed cell death 6-interacting protein | 97 29 31 0.04 | 6 | 5.7 | 11683497 | A | 75/100 |
| 4908 CM10 | OGJ | VMO1_HUMAN | Vitelline membrane outer layer protein 1 homolog precursor | 22 101 5.6 0.39 | 21.4 | 25.0 | A | 71/82 |
| 4639 IMAC30 | Gastric | AACT_HUMAN | α-1-antichymotrypsin precursor | 48 145 8.4 0.26 | 13.4 | 11.5 | 18813785, 17884789 | A | 71/76 |
| 4141 CM10 | Oesophageal | CO4A_HUMAN | Complement C4-A precursor | 194 81 6.3 0.05 | 3.0 | 2.8 | 20116351 | A | 71/84 |
| 3918 IMAC30 | Oesophageal | PPAP_HUMAN | Prostatic acid phosphatase precursor | 45 44 3.3 0.08 | 7.1 | 12.2 | 20645695 | A | 42/100 |
| 2789 CM10 | Oesophageal | CAP7_HUMAN | Azurocidin precursor | 27 43 2.7 0.14 | 10 | 20.4 | 15473694 | A | 42/96 |

Protein ID is the Swiss‑Prot identifier. PMID, PubMed identifier relating to articles where the protein is differentially regulated in, or associated with, cancer. Pattern matching was performed using either an automated approach (A) using in‑house written software, or by manual comparison (M) of the observed presence of proteins in four positive and four negative urine samples based on the measured normalised intensities of m/z peak clusters. Mascot‑SELDI pattern matching scores are based on ‘sensitivity/specificity’ and relate to the observed expression patterns of individual peaks in the samples tested based on SELDI analysis (expected pattern) and pattern of identified molecules by LC‑MS/MS. OGJ, oesophagogastric junction; SELDI, surface‑enhanced laser desorption/ionization; MS, mass spectrometry; emPAI, exponentially modified protein abundance index.
Development of a cancer profiling database. Over the last decade, the emergence of high-throughput screening platforms opened the possibility to mechanistically characterize at the molecular level disease phenotypes, disease subtypes, assess disease progression and monitor response to therapy. However, despite a crescendo in the availability of these large-scale multi-omics data for numerous conditions, no apparent or adequate effort has been made to curate and integrate those data resources.

Therefore, large-scale multi-omics data handling differential expression of several molecular entities, for example, microRNA, gene, protein and metabolite across multiple tissues and biological fluids in several cancer types (with special emphasis on GI neoplasms) were collected from the literature and general scope databases, and subjected to extensive manual curation and annotation. This effort yielded a cancer profiling database: the Multi-Omics Cancer database MoCadb (www.padb.org/mocadb), a database module of the Pan-Omics Analysis Database (PADB) (www.PADB.org) which ensures a long-term lifecycle of the created repository, along with providing the appropriate framework that encloses curated resources, in order to assist in the development of integrative -omics disease models, straightforward translation of findings from experimental animal models using established ortholog maps, creation of cellular regulatory networks based on the association of miRNAs and transcription factors to targets, protein-protein and gene-gene interactions, enzymatic reactions, delineation of pathways and drug interactions.

MoCadb aims to incorporate existing molecular information and clinical metadata, ultimately holding the potential to unravel and allow an in-depth understanding of the regulation of key molecules modulating pathophysiology and progression in several cancer types.

Discussion

The identification of potential novel biomarkers for ailments such as pancreatic cancer is of utmost importance, particularly in light of the shortcomings of the markers currently used in the clinical setting for diagnosis and disease progression monitoring. As an example, in a retrospective study, cohort screening using a bead-based antibody array of a multi-parametric signature of three serum biomarkers, namely CA19.9, intercellular adhesion molecules 1 and osteoprotegerin, were able to distinguish patients with pancreatic ductal adenocarcinoma (PDAC) from healthy subjects, with a sensitivity/specificity of 88/90% and an area under the curve (AUC) of 0.93 (38). However, in a follow-up study using a larger prospective cohort, the latter biomarker panel and the respective AUCs associated with the biomarker model were not statistically different, thereby failing to distinguish PDAC cases from matched controls (39). This highlights the current issues and challenges associated with biomarker research in pre-diagnostic risk assessment of pancreatic cancer.

We selected an MS-based proteomic screening of urine from upper GI cancer patients for prospective biomarkers for the four most prominent cancer types associated with the upper GI to establish a proteomic fingerprint pattern, as well as defined molecular markers, which can potentially be used in clinical diagnostics. In the clinical setting, urine is an optimal sample source, as it is easy to obtain, the collection is non-invasive, and is relatively stable in terms of sample...
integrity. A substantial number of previous studies have found the SELDI-TOF-MS technique ideally suited for urine analysis, with a combination of high throughput, speed and relatively low cost (40-42). However, a main drawback of this technique is the comparatively medium resolution of the spectra obtained, but this is adequate to resolve peaks in the 1-25-kDa range from spectra with <500 peaks. We have previously demonstrated that both IMAC30 and CM10 are useful chip types for the analysis of human urine (17), and we were able to generate a decision-tree model for upper GI cancer with an overall sensitivity of 98% and a specificity of 87%. In the present study, we identified 8 peak clusters by SELDI-TOF-MS for the four cancer types studied and, using expression pattern matching, we were able to assign several proteins identified in the urine to our proposed biomarkers.

An elevated amount of SERPINA3 may be associated with both gastric and pancreatic cancer, and has also been reported as a potential urinary biomarker in non-small-cell lung cancer (43). SERPINA3 is a protease inhibitor that can modulate cathepsin G (44), and its glycosylated form is directly associated with colorectal cancer (45). This protein was already known to be associated with pancreatic cancer and could be detected in higher levels in the plasma (46,47). Furthermore, this marker was specifically negatively correlated with survival in patients with advanced pancreatic cancer (48). However, it is also worth noting that several different fragments from the same molecule were observed in our SELDI-TOF experiments, namely m/z 2444, 2447 and 4639, of which the latter is specific for gastric cancer. Additionally, we also identified the same molecule in our upper GI cancer screen (17) as m/z 8803. It is possible that a specific proteolytic fragment of SERPINA3 is associated with and generated in a specific type of cancer, where the m/z 8803 fragment may be generic for upper GI cancer and the small fragments specific to pancreatic cancer. This is further substantiated by our observation that the protease CSTB is also a potential biomarker for pancreatic cancer. Additionally, CSTB was validated as a potential urinary biomarker in pancreatic carcinoma by western blotting. Furthermore, CSTB was reported to be a prognostic marker in pancreatic adenocarcinoma (49), which confirms our findings for this molecule and its potential as a lead marker in pancreatic cancer diagnostics. Another promising candidate as a pancreatic cancer biomarker was identified and verified as CSTB. The cysteine protease inhibitor CSTB was described as a protein that is able to stimulate cancer cell growth in vitro and in vivo (50). Other potential pancreatic cancer markers identified in this study comprise fragments of immunoglobulins. The occurrence of specific fragments of antibodies may be associated with the increased amounts of CSTB, or may be due to a host response to pancreatic tumour growth. Antibodies are also well-described and used in the clinical setting to assess various cancer types (e.g., CA19-9 in pancreatic cancer) (9,10).

Stratification of OGJ cancer cases by SELDI-TOF-MS revealed two potential m/z peak clusters, m/z 4908 and 5511. The latter peak cluster was identified to be a fragment of PDCD6IP (also referred to as AIP1 or ALIX), which has been described to participate in programmed cell death, and it was reported that its overexpression can block apoptosis (51). The m/z 4908 peak cluster consists of fragments from VMO1 and TPI. No role of VMO1 has been implicated in cancer, and this may be a novel target for OGJ cancer, whereas TPI was described in the literature to be upregulated in oesophageal cancer (52), as well as in hepatocellular carcinoma (53).

The oesophageal cancer marker of m/z 4141 appears to contain several molecular constituents, namely C4A, ACPP, AZU1 and fragments from Histone H1. C4A is an important component in the activation of the classical pathway of the complement system and proteolytic breakdown products of C4-A have been suggested as biomarkers in breast cancer (54), although a specific proteolytic product, C4a anaphylatoxin, is a mediator of local inflammatory processes (55). This protein is therefore potentially unsuitable as a diagnostic marker in oesophageal cancer. ACPP, a non-specific tyrosine phosphatase, is well-described to be associated with prostate cancer (56), and is used clinically as a diagnostic marker. AZU1, an antibacterial and monocyte- and fibroblast-specific chemotactic glycoprotein, which acts in conjunction with cathepsin G in host-defense mechanisms (57), was hypothesized to be a potential pancreatic cancer biomarker in the pancreatic juice (58). Histones H1 have been reported to be involved in the survival of breast cancer cells (59), and H1.2 specifically was identified as an apoptogenic factor (60).

In conclusion, the approach of using SELDI-MS to identify potential lead candidates as biomarkers associated with specific upper GI cancers is a useful tool that enabled us to identify potential global upper GI cancer markers, as well as potentially specific markers for individual cancer types, such as gastric, pancreatic, OGJ and oesophageal cancer. CSTB and CTSB, in particular, appear to be promising lead candidates, since these molecules are not commonly found in the urine, and have already been associated with pancreatic cancer in situ in the literature (61-63). Other potential lead markers may require further validation, such as western blotting and, ultimately, exact determination of the sensitivity/specificity values of our novel proposed markers associated with specific disease states and their usefulness in a more general setting. Further studies, including an extended cohort, will help determine the validity of our findings, and specific assays monitoring the expression levels of our proposed biomarkers will help to translate our findings into the clinical setting. One of the main hurdles to overcome is a methodological/technical dependency, such as the use of the SELDI technology, which is not straightforward to translate into a clinical environment. A more appropriate approach would rely on techniques that are commonly in use, such as specific protein detection methods. Additional investigations are needed to explain the biological involvement of the proposed biomarkers in tissue type-specific cancers and their presence in the urine. To aid in this task, and to add a higher level of contextualisation of individual markers or a panel of biomolecules, information held in our MoCadb database, and in other databases within the PADB initiative, can guide the biomarker identification process. Future expansion of this resource, by also incorporating large-scale datasets derived from other cancer tissue type studies, will allow us to not only investigate the biological relevance, importance and usefulness of individual biomolecules, but also to identify potential intervention points or markers of high prognostic/diagnostic value prior to evaluation and validation in patient cohorts. Potential applications of such
findings will most likely include the use of multiplexed platforms, since it was shown that integrating various molecule types, such as proteins and miRs, can exert a beneficial effect on overall sensitivity and specificity of a bioassay for pancreatic cancer (64). However, this also emphasises that further study is required to use novel approaches and may lead to better outcomes in patient stratification and disease treatment.

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Authors’ contributions

HH performed the sample preparations, SELDI measurements, data analysis, Mascot searches, software design and coding, and wrote the manuscript; ADC performed the LC-MS/MS measurements; MF co-wrote the manuscript and developed MoCadb; RJS and JM co-wrote the manuscript; JAR and KCHF supervised the research.

Availability of data and materials

All the datasets generated and analysed in the present study are included in this published manuscript.

Ethics approval and consent to participate

All procedures were approved by the local research ethics committee and written informed consent was obtained from the patients. The study conformed to the standards set by the Declaration of Helsinki.

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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