Discovery of Urinary Proteomic Signature for Differential Diagnosis of Acute Appendicitis

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Acute appendicitis is one of the most common acute abdomens, but the confident preoperative diagnosis is still a challenge. In order to profile noninvasive urinary biomarkers that could discriminate acute appendicitis from other acute abdomens, we carried out mass spectrometric experiments on urine samples from patients with different acute abdomens and evaluated diagnostic potential of urinary proteins with various machine-learning models. Firstly, outlier protein pools of acute appendicitis and controls were constructed using the discovery dataset (32 acute appendicitis and 41 control acute abdomens) against a reference set of 495 normal urine samples. Ten outlier proteins were then selected by feature selection algorithm and were applied in construction of machine-learning models using naïve Bayes, support vector machine, and random forest algorithms. The models were assessed in the discovery dataset by leave-one-out cross validation and were verified in the validation dataset (16 acute appendicitis and 45 control acute abdomens). Among the three models, random forest model achieved the best performance: the accuracy was 84.9% in the leave-one-out cross validation of discovery dataset and 83.6% (sensitivity: 81.2%, specificity: 84.4%) in the validation dataset. In conclusion, we developed a 10-protein diagnostic panel by the random forest model that was able to distinguish acute appendicitis from confusable acute abdomens with high specificity, which indicated the clinical application potential of noninvasive urinary markers in disease diagnosis.

1. Introduction

Acute appendicitis is one of the most common surgical emergencies in clinic worldwide [1]. However, the precise diagnosis of acute appendicitis is still a challenge, especially in children and childbearing women. The absence of classical clinical signs presents in 30~45% of patients [2]. Also, similar symptoms of other acute abdominal inflammations (such as gastrointestinal disorders, cholelithiasis, and gynecological diseases) with appendicitis lead to misdiagnosis and high negative appendectomy rates, which is reported to a range between 10% and 32% [3–5]. In addition, the missed diagnostic rate of acute appendicitis ranges from 11% to 28% [6, 7], and the delays in diagnosis and treatment would lead to the increased risk of appendix rupture, abscess, and even septicemia [8].

For now, the diagnosis of acute appendicitis mainly depends on patient symptoms and serological results, such
as white blood cell count (WBC), C-reactive protein (CRP),
and neutrophil cells. However, these tests did not present
efficient sensitivity and specificity for appendicitis diagnosis
[9, 10]. Imaging examinations such as ultrasonography and
computed tomography (CT) are also used to assist diagnosis.
Ultrasonography has moderate specificity (81%, 95% CI: 78–
84%) [11] in detecting acute appendicitis, limiting its diag-
nostic ability. Though CT examination is more accurate
[11], the strong radiation risk and high cost make it unsuit-
able for wide application. Additionally, several risk scores
by combing clinical signs have been developed to improve
the predictive ability for appendicitis. The Alvarado score is
one of the most well-known scoring systems; it has good sen-
sitivity (99%, 95% CI: 97–99%) but low specificity (43%, 95%
CI: 36–51%) [12]. Thus, efficient diagnostic methods for
discrimination of appendicitis from other confused acute
abdomens are in urgent need.

Urine is a kind of noninvasive, easily attainable clinical
case. With less complexity than serum, urinary proteo-
ics has become an attractive field in biomarker discov-
y. Current urinary proteomics is mainly applied in
diagnostic or prognostic marker discovery for urogenital
diseases [13–15] and nonurogenital diseases, such as chol-
angiocarcinoma [16], coronary artery disease (CAD) [17],
heart failure [18], and stroke [19]. Also, for acute appendi-
citis, candidate urinary biomarkers have been reported,
including 5-hydroxyindoleacetic acid (5-HIAA) [20, 21]
and leucine-rich α-2-glycoprotein (LRG) [22, 23]. However,
5-HIAA was proved to be unreliable for determining acute
appendicitis in afterward studies [24, 25]. LRG had better
performance in diagnosing pediatric appendicitis with
nonappendicitis, but the diagnostic effects ranged largely in
different studies (area under the curve (AUC): 0.63–0.98)
[22, 23]. So, the improvement of a urinary proteomic system
in acute appendicitis diagnosis is still in great demand.

In this study, a high throughput urinary proteome
analysis platform was employed to get deep profiling of urine
samples from patients with acute abdomens. The urinary
proteomic data were analyzed using several classification
strategies for diagnosis of acute appendicitis with other
mimic acute abdomen samples. Among these models, a ran-
dom forest diagnosis model with a panel of 10 urine proteins
achieved the best diagnostic result. Without using clinical
signs and imaging examinations, the random forest model
got an accuracy of 83.6% in the validation dataset, and the
sensitivity and specificity were 81.2% and 84.4%, respectively.
Hence, our study provides potential urinary markers and effi-
cient model for acute appendicitis diagnosis.

2. Materials and Methods

2.1. Patient Samples. The study was approved by the Tian-
jin Baodi Hospital Ethical Review Committee. Urine
samples from 134 patients with acute abdomens were collected
between 2013 and 2016 from Tianjin Baodi Hospital,
including 48 cases with acute appendicitis (AA) and 86
cases with control diseases (cholecystitis and cholelithiasis
(CHG), pancreatitis (PAN), gastrointestinal perforation
(GP), intestinal obstruction (IO), and other acute abdom-
enes (OTH)). Patients were diagnosed according to blood
tests, radiological studies (CT and/or ultrasound), or path-
ological results after surgery. Patient information was
shown in Table 1, and the sample metadata were summa-
ized in Table S1.

| Diseases                              | Cases | Gender (M/F) | Age (year) | Cases | Gender (M/F) | Age (year) |
|---------------------------------------|-------|--------------|------------|-------|--------------|------------|
| Acute appendicitis (AA, n = 48)       | 32    | 14/18        | 38.5±17.9  | 16    | 9/7          | 34.2±15.2  |
| Control acute abdomens (CON, n = 86) |       |              |            |       |              |            |
| Cholecystitis and gallstones (CHO)    | 17    | 5/12         | 56.9±17.0  | 15    | 9/6          | 62.7±16.3  |
| Pancreatitis (PAN)                    | 5     | 0/5          | 42.0±13.9  | 13    | 10/3         | 48.0±13.9  |
| Gastrointestinal perforation (GP)     | 6     | 4/2          | 67.2±17.6  | 5     | 5/0          | 66.8±14.1  |
| Intestinal obstruction (IO)           | 9     | 4/5          | 60.3±15.9  | 4     | 1/3          | 58.0±13.0  |
| Other abdomens (OTH)                 | 4     | 0/4          | 34.2±12.3  | 8     | 2/6          | 47.2±25.5  |
| Total                                 | 41    | 13/28        | 55.1±18.2  | 45    | 14/17        | 56.6±17.2  |

2.2. Urine Sample Preparation. Midstream urine was
obtained from all donors before treatment and stored at
-80°C. Urine samples were prepared according to a previ-
ously described method (Figure S1) [26]. Briefly, 8.9 ml of
urine was centrifuged at 200,000g for 70 min. The pellet
was resuspended and heated at 56°C for 30 min to remove
uromodulin. After heating, the solution volume was
adjusted to 8.9 ml and centrifuged at 200,000g for 70 min.

2.3. LC-MS/MS Analysis. The digested peptides were
extracted and loaded onto a homemade trap column and
an analytical column that both packed with C18 (Dr. Maisch
GmbH, Germany). A 75 min gradient was used for online
HPLC-MS. Thermo Fisher Orbitrap mass spectrometers
were used for measuring all urine samples. LC-MS/MS data
were processed using Proteome Discoverer (V1.4, Thermo Fisher) with Mascot algorithm (Mascot V2.3, Matrix Science) against human RefSeq database (2013.07.04). Tryptic peptides of 293T cell lysates were used as quality control samples for evaluation of the instrument reproducibility.

All assigned peptides were filtered with 1% false discovery rate (FDR). Only proteins with ≥2 strict peptides (1% FDR and ion score > 20) were kept for quantification. Intensity-based absolute quantification (iBAQ) algorithm was used for protein quantification. To normalize the differences in loading amounts among samples, the iBAQ value was converted to iFOT (fraction of total, iBAQ value of each protein divided by the sum of all iBAQ values of all proteins in the sample). For better visualization, iFOTs of each protein divided by the sum of all iBAQ values of value was converted to iFOT (fraction of total, iBAQ value of each protein divided by the sum of all iBAQ values of all proteins in the sample). For better visualization, iFOTs of each protein divided by the sum of all iBAQ values of all proteins in the sample). For better visualization, iFOTs were normalized by $10^{5}$ [27]. To eliminate the difference between different groups, the iFOTs were normalized by quantile algorithm [28].

2.4. Feature Selection and Construction of Machine-Learning Models. In the discovery dataset, normalized iFOTs were log2 transformed, and the missing values were imputed with a minimal value. For the selection of feature, firstly, AA and CON outliers were screened with the normal reference intervals (RIs) built with 495 normal urine samples [26]. The criterion of outlier is that the iFOT value of a protein is more than 95% CI of normal RI plus 2 folds of interquartile range (IQR). Outlier proteins in each sample were identified and then were further screened depending on the iFOT cutoff value and frequency in the disease group to build disease-related outlier pool. Outlier pool selection standards: (1) the iFOT of the outlier is more than 75% percentile of all the outlier iFOT cutoff values, and it should be outlier in at least 90% percentile of all the outlier frequencies in the AA group or the CON group; (2) if the protein is not expressed in healthy urinary samples, it should be outlier in at least 95% percentile of all the outlier frequencies in the AA group or the CON group. AA outlier pool and CON outlier pool were established, respectively. Then, the nonparametric Wilcoxon test was performed between proteins in the two outlier pools; proteins with $p$ value $< 0.05$ were applied in feature selection algorithm for further screening. Ten proteins were selected and applied in the construction of classification models with different machine-learning algorithms, including random forest (RF), support vector machine (SVM), and naïve Bayes (NB). Leave-one-out cross validation was used in the evaluation of model robustness. To evaluate the prediction effect, the classification models were verified in the independent validation dataset.

2.5. Statistical Analysis. R software (version 3.5.3) and RStudio (version 0.99.489) were used in the construction outlier models and machine-learning models, statistical analysis, receiver operating characteristic (ROC) analysis, and plotting. Gene ontology (GO) analysis was performed with DAVID (https://david.ncifcrf.gov/).

3. Results and Discussion

3.1. Proteomic Profiling of Acute Appendicitis (AA) and Control Acute Abdomen (CON) Samples. A total of 134 samples passed quality control in protein identification (more than 800 proteins in each sample) were used for data analysis, including 48 AA samples and 86 CON samples (Table 1). High instrument quantification reproducibility was achieved with an average Pearson correlation coefficient of 0.88 between quality control samples. Totally, 5335 proteins were identified and quantified (Table S2), and an average of 1361 ± 432 proteins was quantified in each sample. Among them, 3834 proteins were identified in both groups. There were 138 unique proteins in the AA group and 1363 unique proteins in the CON group (Figure S2). There were 198 (5.0%) proteins identified in all AA samples, 183 (3.5%) in the CON group, and 142 proteins identified in all the samples (Figure 1(b)). The distribution of proteins showed that the CON group had much more low frequency (<10% frequency) proteins (43.7%) than the AA group (31.4%) (Figure S2). What is more, the highly expressed proteins with more than 90% frequency (AA: 449 proteins; CON: 542 proteins) had a greater proportion than the other groups with more than 20% frequency, suggesting that there were a considerable percentage of “core” urinary proteins widely expressed in different urine samples.

3.2. Reliability Demonstration of Profiling Data in Acute Pancreatitis. To assure whether the urinary proteomic method could be applied in the diagnostic biomarker discovery for nonurogenital disease, we checked the classification ability of our data for acute pancreatitis (PAN) from other acute abdomens because there were commonly accepted reliable urinary markers for pancreatitis. The diagnostic capabilities of the urinary level of amylase (the expression products
of AMY2A and AMY2B) and trypsinogen-2 (the expression product of PRSS2) had been proved in many researches [29, 30]. We found that AMY2A, AMY2B, and PRSS2 were significantly highly expressed in urine samples from pancreatitis patients compared with those from other acute abdomen patients (Figures 2(a)–2(c), Kruskal-Wallis test, ** means $p < 0.01$, * means $p < 0.05$). The AUCs of the three proteins between pancreatitis and the rest of the acute abdomen samples were all above 0.75 (Figure 2(d), Table 2). The AUC of AMY2A was 0.83, which was the best one among these three proteins, showing its outstanding ability in diagnosing pancreatitis. This result demonstrated that the urinary proteome system could produce reliable datasets, and proteins derived from nonurogenital tissues could be detected and beneficial in disease diagnosis.

3.3. Feature Selection for Classification of AA and CON. Based on the type of mass spectrometers (MS), the dataset was divided into discovery dataset and independent validation dataset (Table S2). In the discovery dataset, a total of 73 urine samples including acute appendicitis (AA, $n = 32$) and control acute abdomen (CON, $n = 41$) specimens that processed with multiple MS types were used for feature selection and classification model construction. The independent validation dataset consisted of 16 AA and 45 CON specimens, which were all produced by the same MS type (Orbitrap-QE-Plus).

In order to select the features for classification (Figure 3(a)), AA outliers and CON outliers were identified, respectively, in the discovery dataset against a normal urine database including 495 samples. Outliers were proteins with expression that is higher than 95% confidence interval (CI) of reference interval (RI) plus 2 folds of interquartile range (IQR) of RI. The frequently detected outliers could be considered as markers to reveal changes under pathological conditions. With further screening by frequency (shown in Methods), 287 proteins were selected as AA outliers, and 322 were CON outliers (Table S2).

![Figure 2: Expression and receiver operating characteristic (ROC) analysis of PAN markers. The abundance of AMY2A (a), AMY2B (b), and PRSS2 (c) in different disease groups (Kruskal-Wallis test, ** means $p < 0.01$, * means $p < 0.05$). (d) ROC analysis of AMY2A, AMY2B, and PRSS2, respectively.](image)

Table 2: AUC, sensitivity, and specificity of PAN marker proteins in diagnosing PAN with other acute abdomens.

| Proteins | AUC | Sensitivity | Specificity |
|----------|-----|-------------|-------------|
| AMY2A    | 0.83| 83.3%       | 71.5%       |
| AMY2B    | 0.75| 66.7%       | 83.6%       |
| PRSS2    | 0.82| 83.3%       | 79.3%       |
For the above two outlier pools, Gene ontology (GO) analysis was conducted (Figures 3(b), 3(c), and S3). Among the top 5 processes, three items including platelet degranulation, acute-phase response, and cellular oxidant detoxification were enriched both in AA outliers and CON outliers (Figures 3(b) and 3(c)). For proteins enriched in acute-phase response (ranked second and third in AA outliers and CON outliers, respectively), eight proteins were shared by AA and CON; one protein (ITIH4, inter-alpha-trypsin inhibitor heavy chain family member 4) was CON-specific. Hence, in order to distinguish different diseases, specific proteins in outliers should be further filtered as classification features. The 151 significantly differential proteins between the two groups (nonparametric Wilcoxon test, p value < 0.05, Table S2) were used for feature selection algorithm. Ten proteins (LYVE1: lymphatic vessel endothelial hyaluronan receptor 1; AHCYL1: adenosylhomocysteinase-like 1; APOC1: apolipoprotein C1; SECTM1: secreted and transmembrane 1; SLC31A1: solute carrier family 31 member 1; ITGA6: integrin subunit alpha 6; SLC35F2: solute carrier family 35 member F2; GPX3: glutathione peroxidase 3; TMEM14C: transmembrane protein 14C; SLC47A2: solute carrier family 47 member 2) were further selected for the establishment of classification models, and their expressions were shown in Figure S4.

3.4. Construction and Evaluation of Classification Models. However, AUCs of the ten single-feature-based classification models were all less than 0.75 (Figure 4(a)), indicating the limitation of single feature in classification. Aiming at model optimization, machine-learning algorithms were further adopted to integrate multiple features for model construction, including random forest (RF), support vector machine (SVM), and naïve Bayes (NB) (Figure 4(b)). The robustness of these 3 models was evaluated by leave-one-out cross validation, and the accuracies of the 3 models were 84.9%, 83.6%, and 79.4%, respectively. In the independent validation dataset, 16 AA and 45 CON specimens were included. The RF model could correctly distinguish 83.6% of samples (sensitivity: 81.2%, specificity: 84.4%), SVM model had an accuracy of 78.7% (sensitivity: 25.0%, specificity: 97.8%),
and NB had 70.5% (sensitivity: 68.8%, specificity: 71.1%), respectively. The RF model achieved the best performance among the 3 models and showed advantage in differential classification.

4. Discussion

Acute appendicitis is one of the most significant acute abdomen diseases in clinic, but the diagnosis of acute appendicitis...
is still difficult. In our study, a urinary proteomic system was applied to discover urinary markers for differential diagnosis of AA with other acute abdomens. A panel of 10 proteins integrated by the RF model achieves 84.9% of accuracy in leave-one-out cross validation and 83.6% in the independent validation dataset, which was better than the reported results of WBC (sensitivity 62%, specificity 75%) and CRP (sensitivity 57%, specificity 87%) [10]. Notably, without relying on any clinical sign, this RF model using urine markers showed high specificity in AA diagnosis, which was helpful in reducing negative appendectomy.

Among the 10 features in this diagnostic model, two features had AUCs higher than 0.7, including LYVE1 (lymphatic vessel endothelial hyaluronan receptor 1) and AHCYL1 (adenosylhomocysteinase-like 1). LYVE1 is an immunity-related factor that may play a role in the migration of immune cells, and the loss of LYVE1 is a sign of activation of inflammation [31, 32]. The expression of LYVE1 is lower in the AA group than in the CON group. Since the appendix is also an immune organ, this may indicate that inflammation is more easily activated in AA. AHCYL1 is upregulated in the CON group, maybe because it acts as a regulator of fluid secretion [33], which is important for the secretory organs in the CON group. APOC1 (apolipoprotein C1) is a feature that upregulated in the AA group. APOC1 can bind to lipopolysaccharide (LPS) on the outer-membrane of gram-negative bacteria to present LPS to macrophages, thereby stimulating the inflammatory response [34]. The upregulation of APOC1 might reflect the existence of bacterial infection in AA than the diseases in the CON group, such as cholecystitis and pancreatitis.

In this study, we found that the urinary protein panel and the random forest model were powerful in the diagnosis of acute appendicitis with the other acute abdomens. The advantages of random forest algorithm, such as nonoverfitting, robustness to noise, make it a robust tool in classification. Our results also demonstrated the potential of urinary protein in the diagnosis of nonurogenital diseases, as the change of acute pancreatitis biomarkers could be well detected in different disease groups. With the relatively high specificity, our result might provide beneficial supplement to the clinical auxiliary diagnosis of acute appendicitis. However, there are still areas that need further improvement. Firstly, this urinary diagnostic model should be further validated in a larger number of samples. Secondly, for facilitation in potential clinical application, the feature proteins need to be absolutely quantified. Except for the immunological methods, the MS-based method for quantification is becoming popular, with the increasingly high-throughput detection of target proteins independent on antibodies. Lastly, since urine is rich of small metabolites, the combination of urinary protein makers with metabolites might contribute to improving diagnostic efficiency.

5. Conclusions

In summary, a urinary proteomic system was applied to discover urinary markers for differential diagnosis of AA with other acute abdomens. A panel of 10 proteins integrated by the RF model achieves 84.9% of accuracy in leave-one-out cross validation and 83.6% in the independent validation dataset. Without relying on any clinical sign, this RF model using urine markers showed high specificity in AA diagnosis, which was helpful in reducing negative appendectomy, indicating the potential of noninvasive urinary markers in clinical application.

Data Availability

The data used to support the findings of this study are available from the corresponding authors upon request.

Conflicts of Interest

All the authors declare no conflict of interest.

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Supplementary Materials

Supplementary 1. Figure S1: the overview of experimental workflow for the urinary proteomic method.

Supplementary 2. Figure S2: the distribution of proteins with different frequencies in the AA and CON groups.

Supplementary 3. Figure S3: GO enrichment analysis of outliers in the AA and CON pools, respectively. (A) Cellular component analysis of the AA outlier pool. (B) Cellular component analysis of the CON outlier pool. (C) Molecular function analysis of the AA outlier pool. (D) Molecular function analysis of the CON outlier pool.

Supplementary 4. Figure S4: relative abundance of 10 feature proteins in the AA and CON groups. The Wilcoxon test was performed between the AA and CON groups, ** means $p < 0.01$, * means $p < 0.05$.

Supplementary 5. Table S1: metadata of urine samples that analyzed in this study, including the sample ID, group label, dataset type, gender age, disease status, sampled date, sampled hospital, and MS instrument.

Supplementary 6. Table S2: sheet 1: relative abundances (iFOT) of discovery dataset; sheet 2: relative abundances (iFOT) of validation dataset; sheet 3: list of 287 outlier proteins in the AA group; sheet 4: list of 322 outlier proteins in the CON groups; sheet 5: list of 151 outlier proteins that are significantly different between AA and CON groups.
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