Urinary TIMP2·IGFBP7 for the prediction of platinum-induced acute renal injury

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Introduction: Platinum-based chemotherapy (PBC) is a potent antineoplastic treatment, but cisplatin nephrotoxicity is often the limiting factor. Identifying the patients who are at risk for developing platinum-induced renal injury is an important issue. We tested urinary TIMP2·IGFBP7, a new US Food and Drug Administration (FDA)-cleared test to assess the risk of acute kidney injury (AKI), in a cohort of patients with malignant neoplastic disease receiving PBC.

Patients and methods: A total of 58 patients with malignant neoplastic disease were enrolled in this study, of whom 32 patients had both urine samples and subsequent serum creatinine values available for detecting AKI within 72 hours. Urine samples were collected within 6 hours prior to PBC application and within 12 hours after the end of chemotherapy administration. We examined the predictive ability of TIMP2·IGFBP7 for the development of AKI as defined by KDIGO (Kidney Disease: Improving Global Outcomes) criteria within 72 hours after the administration of chemotherapy. Operating characteristics were determined for the previously validated TIMP2·IGFBP7 cutoff of 0.3 (ng/mL)²/1000.

Results: Four (12.5%) patients developed AKI within 72 hours. Primary disease was lymphoma in 13 patients (40.6%) and solid tumors in 19 patients (59.4%). Eight patients (25.0%) received carboplatin and 24 (75.0%) cisplatin. TIMP2·IGFBP7 after PBC administration discriminated for the risk of AKI with an area under the receiver operating characteristic curve (AUC; 95% confidence interval) of 0.92 (0.80–1.00). At the cutoff of 0.3 for TIMP2·IGFBP7, sensitivity was 50%, specificity was 87%, negative predictive value was 95% and positive predictive value was 25% for the prediction of AKI within 72 hours.

Conclusion: Urinary TIMP2·IGFBP7 measured in specimens gathered after PBC may be a useful tool to early identify patients who are at risk for developing platinum-induced AKI.

Keywords: acute kidney injury, biomarker, cisplatin, nephrotoxicity, insulin like-growth factor binding protein-7, tissue inhibitor of metalloproteinases-2

Introduction

Cisplatin and platinum-based chemotherapy (PBC) are broadly used potent antineoplastic treatments. The cytotoxic substance, cisplatin, was first discovered in *Escherichia coli* as an anti-infective agent, and its antitumor activity was reported several years later.¹² PBC is widely applied to treat oncologic diseases, such as lymphoma, testicular, ovarian, cervical and head- and neck non-small cell lung cancers, partially in combination with radiation therapy.³⁴

As with most antitumor therapies, the major reason for PBC discontinuation is the noxious side effects. These side effects are well studied in cisplatin toxicity: the most severe, dose-limiting side effect is nephrotoxicity, which was reinforced by reports of renal failure after accidental overdose of cisplatin.³ Approximately one-third of the...
patients receiving cisplatin chemotherapy develop AKI at doses of 50–100 mg/m². AKI is known to increase mortality and complicate hospital stay. Cisplatin nephrotoxicity is mostly mediated through inflammation and direct cytotoxicity through reactive oxygen species. Renal proximal tubules show vulnerability to cisplatin toxicity that leads to a decline in GFR.

To alleviate the nephrotoxic effects, prevention strategies such as hydration were established which allow the application of higher cisplatin doses. Development of carboplatin and other platinum-containing compounds led to less severe nephrotoxicity but did not eliminate nephrotoxic side effects. Despite hydration and new developments, nephrotoxicity remains a major reason for the discontinuation of therapy and complication of antitumor therapy.

Over the past several years, new potential biomarkers for the early detection of AKI have been identified. TIMP2 and IGFBP7 are the most promising markers. Many studies have shown that the product of the concentration of these proteins TIMP2·IGFBP7 in urine (Astute Medical, San Diego, CA, USA) is an early indicator of risk for AKI. As the first available AKI biomarker test in the USA, urinary TIMP2·IGFBP7 was cleared in 2014 by the US FDA.

In this study, we tested the hypothesis that urinary TIMP2·IGFBP7 collected directly prior and after PBC administration can predict the development of AKI. A rapid biomarker test, such as urinary TIMP2·IGFBP7, that can identify the patients who are at risk for developing PBC-induced AKI could lead to a more effective application of these antitumor agents or a further development of prevention strategies.

Patients and methods

This study was conducted at the Department of Internal Medicine at Robert-Bosch Hospital, Stuttgart (Germany), after obtaining approval from the ethics board of the University of Tuebingen (Germany). Urinary biomarkers were obtained in clinical routine, and the analysis was conducted retrospectively. Informed consent was not obtained due to the retrospective study design; patient confidentiality was maintained, and the collected data were anonymized without identifying information. Inclusion criteria were as follows: sCr values within normal range (0.5–1.2 mg/dL) at chemotherapy initiation, known malignant neoplastic disease, therapy with cisplatin or carboplatin and available follow-up sCr values. Exclusion criteria were as follows: already existing AKI or kidney impairment at the start of chemotherapy, no malignant neoplastic disease or no cisplatin/carboplatin chemotherapy. In this study, 26 patients were excluded due to missing subsequent sCr values (Figure 1). Six patients with baseline urinary TIMP2·IGFBP7 had missing urinary TIMP2·IGFBP7 samples after PBC administration. To identify eligible patients, the program for the preparation of cytotoxics was screened for PBC regimens (cisplatin and carboplatin). Urinary TIMP2·IGFBP7 was collected within 6 hours prior to PBC application and within 12 hours after the end of chemotherapy administration as a routine measurement. We examined the predictive ability of TIMP2·IGFBP7 for the development of AKI within 72 hours after the administration of chemotherapy. AKI was defined as at least stage 1 by sCr criteria of the KDIGO guideline. eGFR was calculated using the CKD-EPI equation. Test characteristics (sensitivity, specificity, NPV, PPV) were determined for the previously validated TIMP2·IGFBP7 cutoff of 0.3.

Biomarker assays

sCr and urinary TIMP2·IGFBP7 were measured at Robert-Bosch Hospital. Urinary TIMP2 and IGFBP7 concentrations were measured using the NephroCheck™ test and Astute140 Meter (Astute Medical, San Diego, CA, USA). TIMP2·IGFBP7 levels were reported in units of (ng/mL)²/1000.

Statistical analysis

Predictive performance was assessed using ROC curve analysis, including the determination of the operating characteristics at the previously validated cutoff of 0.3. 95% CIs for AUCs were calculated with the DeLong method. 95% CIs for operating characteristics were constructed with the Clopper–Pearson method. For comparisons of baseline

Figure 1 Study flow diagram.

Abbreviations: AKI, acute kidney injury; sCr, serum creatinine.
characteristics in end point negative and positive patients, categorical variables were analyzed using the Fisher’s exact or chi-square test and continuous variables using the unpaired t-test and Wilcoxon rank-sum test, respectively, for normally and non-normally distributed variables. For comparison of paired samples, Wilcoxon matched-pairs signed-rank test was used. For all analyses, two-sided p-values of <0.05 and one-sided p-values of <0.025 were considered statistically significant. Statistical analysis was performed using Prism (GraphPad Software, Inc., La Jolla, CA, USA).

Results

Baseline characteristics

Four (12.5%) patients developed AKI stage 1 within 72 hours. In each patient, urinary TIMP2·IGFBP7 samples after PBC administration were gathered before an AKI-defining rise in sCr was evident. No subject developed AKI stage 2 or 3. SCr increase was significantly greater (p<0.0001) in the AKI group (median: 0.5 mg/dL; IQR: 0.3–0.8) than in the non-AKI group (median: 0.0 mg/dL; IQR: 0.0–0.1). eGFR decrease was significantly (p=0.0004) greater in patients with AKI (median: -28.3 mL/min/1.73 m²; IQR: -34.4 to -18.6) than those without AKI (median: 0.0 mL/min/1.73 m²; IQR: -9.6 to 0.0). AKI was diagnosed in all patients between 24- and 72-hour time frame after PBC administration: in two patients, AKI was diagnosed on the second day (24-48 hours) and in another two patients AKI was manifest on the third day (48-72 hours) after chemotherapy. Maximum creatinine was evident in all AKI patients at 72-hour time point. Oliguric AKI was recorded in none of the patients.

Eight patients (25.0%) received carboplatin and 24 patients (75.0%) cisplatin. No significant differences in age, BMI, cisplatin dose and cisplatin cycle as well as in baseline sCr and baseline eGFR between the groups “no AKI” and “AKI” were observed (Table 1). Primary disease was lymphoma in 13 patients (40.6%) and solid tumors in 19 patients (59.4%). The lymphoma group comprised 12 non-Hodgkin lymphoma and one Hodgkin lymphoma.

### Table 1 Baseline characteristics

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>No AKI</th>
<th>AKI</th>
<th>p-value</th>
<th>Total, n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n (%)</td>
<td>Median</td>
<td>IQR</td>
<td>n (%)</td>
</tr>
<tr>
<td>Total</td>
<td>28 (87.5)</td>
<td>62.5</td>
<td>51.5–73.3</td>
<td>4 (12.5)</td>
</tr>
<tr>
<td>Age (years)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male gender</td>
<td>18 (64.3)</td>
<td>4 (100)</td>
<td>27.7</td>
<td>22 (68.8)</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>24.8</td>
<td>22.0–28.5</td>
<td>4 (16.7)</td>
<td></td>
</tr>
<tr>
<td>Cisplatin</td>
<td>20 (83.3)</td>
<td>73.5</td>
<td>30.0–100.0</td>
<td></td>
</tr>
<tr>
<td>Cycle</td>
<td>2.0</td>
<td>1.0–3.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Carboptatin</td>
<td>8 (100)</td>
<td>257.0</td>
<td>189.5–458.0</td>
<td></td>
</tr>
<tr>
<td>Dose (mg/m²)</td>
<td>2.0</td>
<td>2.0–5.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline sCr (mg/dL)</td>
<td>0.8</td>
<td>0.6–0.9</td>
<td>4 (100)</td>
<td>1.0</td>
</tr>
<tr>
<td>Baseline eGFR (mL/min/1.73 m²)</td>
<td>94.1</td>
<td>85.5–103.7</td>
<td>0.9</td>
<td>0.6–1.1</td>
</tr>
<tr>
<td>Maximum sCr within 72 hours (mg/dL)</td>
<td>90.3</td>
<td>77.6–99.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Minimum eGFR within 72 hours (mL/min/1.73 m²)</td>
<td>90.3</td>
<td>77.6–99.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>sCr changes from baseline within 72 hours (mg/dL)</td>
<td>0.0</td>
<td>0.0–0.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>sCr changes from baseline within 72 hours (%)</td>
<td>0.0</td>
<td>0.0–12.5</td>
<td>0.0</td>
<td>0.0–12.5</td>
</tr>
<tr>
<td>eGFR changes from baseline within 72 hours (mL/min/1.73 m²)</td>
<td>0.0</td>
<td>-9.6 to 0.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>eGFR changes from baseline within 72 hours (%)</td>
<td>0.0</td>
<td>-15.4 to 0.0</td>
<td>0.0</td>
<td>-15.4 to 0.0</td>
</tr>
<tr>
<td>TIMP2·IGFBP7 prior to PBC ((ng/mL)²/1000)</td>
<td>0.12</td>
<td>0.04–0.20</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TIMP2·IGFBP7 after PBC ((ng/mL)²/1000)</td>
<td>0.07</td>
<td>0.03–0.17</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Primary disease</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Solid tumor</td>
<td>17 (89.5)</td>
<td>2 (10.5)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lymphoma</td>
<td>11 (84.6)</td>
<td>2 (15.4)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note: All subjects developed AKI stage 1 defined by KDIGO *p<0.05.

Abbreviations: AKI, acute kidney injury; BMI, body mass index; eGFR, estimated glomerular filtration rate; IQR, interquartile range; KDIGO, Kidney Disease: Improving Global Outcomes; NV, no value; PBC, platinum-based chemotherapy; sCr, serum creatinine.
solid tumors, the individual malignancies were four testicular tumors, three bronchial carcinomas, two esophageal carcinomas, two breast carcinomas, two urothelial carcinomas, two ovarian cancers, one neuroendocrine carcinoma, one osteosarcoma, one peritoneal carcinoma and one extragonadal germ cell tumor.

**Biomarker results**

Urinary TIMP2·IGFBP7 values after the administration of PBC were significantly \((p=0.04)\) higher in patients with AKI than those without AKI. The median urinary TIMP2·IGFBP7 level was 0.07 (IQR: 0.03–0.17) in the non-AKI group and 0.69 (IQR: 0.30–1.08) in the AKI group (Figure 2).

Similarly, urinary TIMP2·IGFBP7 values prior to the application of PBC differed significantly in patients developing AKI within 72 hours compared to patients without AKI \((p=0.019)\). The median value in the non-AKI group was 0.12 (IQR: 0.04–0.20), whereas in the AKI cohort the median urinary TIMP2·IGFBP7 level was 0.35 (IQR: 0.27–0.89) (Figure 3).

The TIMP2·IGFBP7 test characteristics for the prediction of AKI within 72 hours using the cutoff of 0.3 and urine samples after chemotherapy administration were as follows: sensitivity of 0.50 (95% CI: 0.01–0.99), specificity of 0.87 (95% CI: 0.66–0.97), NPV of 0.95 (95% CI: 0.76–1.00) and PPV of 0.25 (95% CI: 0.01–0.81). Most discriminatory cutoff was 0.28 with the sensitivity of 1.00 (95% CI: 0.16–1.00) and specificity of 0.87 (95% CI: 0.66–0.97). The AUC was 0.92 (95% CI: 0.80–1.00) (Table 2), but in this analysis, there were only \(n=2\) subjects in the AKI group.

![Figure 2](https://www.dovepress.com)  
**Figure 2** Urinary TIMP2·IGFBP7 after the administration of PBC in patients with malignant disease.  
**Notes:** Patients developing AKI within 72 hours had significant higher urinary TIMP2·IGFBP7 values than patients without AKI \((p=0.04)\). No AKI: median: 0.07 (IQR: 0.03–0.17); AKI: median: 0.69 (IQR: 0.30–1.08) \((n=26)\). Box and whiskers show the IQR and total observed range, respectively. Horizontal line within the box shows the median.  
**Abbreviations:** AKI, acute kidney injury; IQR, interquartile range; PBC, platinum-based chemotherapy.

Urinary TIMP2·IGFBP7 measured in samples prior to PBC application showed a sensitivity of 0.75 (95% CI: 0.19–0.99), specificity of 0.82 (95% CI: 0.63–0.94), NPV of 0.96 (95% CI: 0.79–1.00) and PPV of 0.38 (95% CI: 0.09–0.76). Most discriminatory cutoff was 0.24 with the sensitivity of 1.00 (95% CI: 0.40–1.00) and specificity of 0.82 (95% CI: 0.63–0.94). The AUC was 0.86 (95% CI: 0.73–0.98) (Table 3).

**Discussion**

Platinum-based chemotherapeutics are widely applied cytotoxic agents, but side effects are dose limiting and can lead to serious complications.6,7 The most important side effect is nephrotoxicity, which often represents a dose-and therapy-restricting factor.21 Acute kidney impairment appears approximately in one-third of the patients receiving cisplatin chemotherapy.6,7 For the prediction of AKI risk or early detection, sCr is not sufficient because of a delayed increase and a creatinine-blind range.32 Often tubular damage is evident without a notable rise in sCr.33 Acute Dialysis Quality Initiative Consensus Conference has proposed the inclusion of tubular damage markers for the definition of AKI.22 Urinary TIMP2·IGFBP7 seems to be one of the most promising tubular damage biomarkers for the early detection of AKI.23

In this study, significantly \((p=0.04)\) higher median urinary TIMP2·IGFBP7 levels, collected from the patients after receiving PBC, were observed in the AKI group (Figure 2). Interestingly, a significant \((p=0.019)\) increase in urinary TIMP2·IGFBP7 prior to PBC administration was evident in patients developing AKI within 72 hours after chemotherapy application (Figure 3). This could imply that those patients developing AKI already had tubular stress, and therefore had probably an increased vulnerability for the toxic effects of the PBC agents.

Both findings suggest that urinary TIMP2·IGFBP7 could be a helpful tool for the identification of patients who are at risk for developing AKI. As a consequence, for patients with urinary TIMP2·IGFBP7 levels >0.3, intensified prevention strategies could be implemented for the reduction in AKI rates after PBC application. Conversely, maybe higher PBC doses could be tolerated by the kidneys when urinary TIMP2·IGFBP7 levels are low.

Previous works described increased biomarker concentrations such as KIM-1, calbindin and TIMP-1 in cisplatin toxicity models in vitro and in vivo, confirming the relevance of urinary biomarkers in cisplatin nephrotoxicity.34 Our findings are comparable to preceding studies examining...
AKI biomarkers in patients receiving cisplatin: Lin et al demonstrated an AUC of 0.865 for NGAL levels and AKI development. Comparable AUCs were described by Tekce et al for urinary KIM-1 levels after PBC application. In our study, urinary TIMP2·IGFBP7 had an AUC of 0.92 (0.80–1.03) when using the urinary sample after PBC application. Surprisingly, we also showed that urinary biomarkers collected prior to PBC application can also predict AKI risk: the AUC of TIMP2·IGFBP7 for samples collected before PBC was 0.86 (0.73–0.98).

We consecutively enrolled all patients getting PBC; for that reason, we also included patients receiving carboplatin. Carboplatin is known to have less serious, but still existing nephrotoxic effects. Comparable AUCs were described by Tekce et al for urinary KIM-1 levels after PBC application. In our study, urinary TIMP2·IGFBP7 had an AUC of 0.92 (0.80–1.03) when using the urinary sample after PBC application. Surprisingly, we also showed that urinary biomarkers collected prior to PBC application can also predict AKI risk: the AUC of TIMP2·IGFBP7 for samples collected before PBC was 0.86 (0.73–0.98).

We consecutively enrolled all patients getting PBC; for that reason, we also included patients receiving carboplatin. Carboplatin is known to have less serious, but still existing nephrotoxic effects. Previous studies showed that quiescent renal tubular cells reenter the cell cycle during AKI. Similarly, one pathophysiologic mechanism in cisplatin nephrotoxicity is dysregulation of cell cycle proteins. Yu et al demonstrated that cell cycle inhibition can prevent renal proximal tubule cells from cisplatin-induced cell death. Therefore, TIMP2 and IGFBP7 molecules, known to be involved in G2 cell cycle arrest, could serve as a potential target for cell cycle inhibition and could reduce the incidence of PBC-related AKI. Under inhibition, maybe higher chemotherapy doses could be achieved, improving effectiveness of the therapy.

Our main limitation is the small sample size – the results provide a first clue for the usefulness of urinary TIMP2·IGFBP7, but further and larger studies are needed to confirm our findings.

Table 2 Urinary TIMP2·IGFBP7 sample after chemotherapy administration using the cutoff of 0.3

<table>
<thead>
<tr>
<th>Sensitivity</th>
<th>Specificity</th>
<th>NPV</th>
<th>PPV</th>
<th>AUC</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.50 (0.01–0.99)</td>
<td>0.87 (0.66–0.97)</td>
<td>0.95 (0.76–1.00)</td>
<td>0.25 (0.01–0.81)</td>
<td>0.92 (0.80–1.00)</td>
</tr>
</tbody>
</table>

Note: Operating characteristics (95% CI) for the prediction of AKI within 72 hours after chemotherapy administration in patients receiving PBC (n=26).

Abbreviations: AKI, acute kidney injury; AUC, area under the ROC curve; CI, confidence interval; NPV, negative predictive value; PBC, platinum-based chemotherapy; PPV, positive predictive value; ROC, receiver operating characteristic.

Table 3 Urinary TIMP2·IGFBP7 sample prior to chemotherapy administration using the cutoff of 0.3

<table>
<thead>
<tr>
<th>Sensitivity</th>
<th>Specificity</th>
<th>NPV</th>
<th>PPV</th>
<th>AUC</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.75 (0.19–0.99)</td>
<td>0.82 (0.63–0.94)</td>
<td>0.96 (0.79–1.00)</td>
<td>0.38 (0.09–0.76)</td>
<td>0.86 (0.73–0.98)</td>
</tr>
</tbody>
</table>

Note: Operating characteristics (95% CI) for the prediction of AKI within 72 hours after chemotherapy administration in patients receiving PBC (n=32).

Abbreviations: AKI, acute kidney injury; AUC, area under the ROC curve; CI, confidence interval; NPV, negative predictive value; PBC, platinum-based chemotherapy; PPV, positive predictive value; ROC, receiver operating characteristic.

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Figure 3 Urinary TIMP2·IGFBP7 prior to the administration of PBC in patients with malignant disease.

Notes: Patients developing AKI within 72 hours had significant higher urinary TIMP2·IGFBP7 values than patients without AKI (p=0.019). No AKI: median: 0.12 (IQR: 0.04–0.20). AKI: median: 0.35 (IQR: 0.27–0.89) (n=32). Box and whiskers show the IQR and total observed range, respectively. Horizontal line within the box shows the median.

Abbreviations: AKI, acute kidney injury; IQR, interquartile range; PBC, platinum-based chemotherapy.
Conclusion
Urinary TIMP2·IGFBP7 measured in specimens gathered after PBC application may be a useful tool to identify patients who are at risk for developing platinum-induced AKI. Surprisingly, urinary TIMP2·IGFBP7 measured in samples collected prior to PBC administration could also serve as an indicator of AKI risk and may reflect the individual vulnerability to PBC toxicity. Therefore, TIMP2 and IGFBP7 could be used as a potential target for cell cycle inhibition to prevent the incidence of platinum-induced AKI in future, but larger studies are needed to confirm our results.

Abbreviations
ADQI, Acute Dialysis Quality Initiative; AKI, acute kidney injury; AUC, area under the ROC curves; BMI, body mass index; CI, confidence interval; CKD-EPI, Chronic Kidney Disease Epidemiology Collaboration; FDA, Food and Drug Administration; eGFR, estimated GFR; GFR, glomerular filtration rate; IGFBP7, insulin-like growth factor-binding protein-7; IQR, interquartile range; KDIGO, Kidney Disease: Improving Global Outcomes; NGAL, neutrophil gelatinase-associated lipocalin; NPV, negative predictive value; PBC, platinum-based chemotherapy; PPV, positive predictive value; ROC, receiver operating characteristic; sCr, serum creatinine; TIMP-2, tissue inhibitor of metalloproteinases-2.

Acknowledgments
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Disclosure
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References