Urinary Hepcidin-25 Is Elevated in Patients That Avoid Acute Kidney Injury Following Cardiac Surgery

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Abstract
Background: Acute kidney injury (AKI) following cardiac surgery leads to increased morbidity and mortality. Characterization and validation of early biomarkers of AKI may ultimately facilitate early therapeutic intervention. We have previously identified that elevated urinary hepcidin-25 is inversely and independently associated with the development of AKI in adult cardiac surgery patients. Hepcidin-25 is an antimicrobial peptide that sequesters iron intracellularly, and its elevation following human ischemia reperfusion injury may represent a renoprotective response to minimize renal injury.

Objective: Our goal was to validate urinary hepcidin-25 as a non-invasive biomarker in an independent cardiac surgery cohort, within the context of clinical AKI predictors.

Design: Prospective observational cohort study.

Setting: Adult cardiac surgery program at St. Boniface Hospital, Winnipeg, Manitoba, Canada.

Patients: Adult cardiac surgery patients undergoing cardiopulmonary bypass (CPB), n = 306.

Measurements: Urine hepcidin-25, measured on post-operative day (POD) 1.

Methods: A prospective, observational cohort of adult CPB patients (n = 306) was collected with serial perioperative urine samples. Urine hepcidin-25 at POD 1 was measured by competitive ELISA. Its diagnostic performance was evaluated in conjunction with clinical parameters and the Thakar clinical prediction score, using multivariate logistic regression.

Results: Urinary hepcidin-25 is elevated following cardiac surgery in AKI and non-AKI patients. Elevated urinary hepcidin-25 concentration was inversely associated with AKI on both univariate (odds ratio [OR]: 0.61, 95% confidence interval [CI]: 0.45-0.83, P = .002) and multivariate analysis (OR: 0.67, 95% CI: 0.50-0.95, P = .02). A combined model with clinical risk factors demonstrated that baseline estimated glomerular filtration rate (eGFR), diabetes mellitus, and urinary hepcidin-25 concentration had an overall area under the curve (AUC) of 0.82 (0.75-0.88) for predicting subsequent AKI development, which was superior to clinical prediction alone as determined by the Thakar score.

Limitations: (1) A single-center observational study. (2) Polyclonal antibody–based competitive ELISA.

Conclusion: Hepcidin-25 is inversely associated with AKI in a multivariate model when combined with eGFR and diabetes mellitus, with an overall AUC of 0.82. Notably, urinary hepcidin-25 improves on clinical AKI prediction compared to the Thakar score alone.

Contexte : L'insuffisance rénale aiguë (IRA) qui survient à la suite d’une chirurgie cardiaque est associée à une augmentation du taux de morbidité et de mortalité. La caractérisation et la validation de biomarqueurs précoces d’une IRA permettraient éventuellement une intervention thérapeutique plus opportune. Nous avions antérieurement déterminé qu’un taux élevé d’hépcidine-25 est inversement et indépendamment associé au développement d’une IRA à la suite d’une chirurgie cardiaque chez les patients adultes. L’hépcidine-25 est un peptide antimicrobien qui capte le fer intracellulaire. L’élévation de sa concentration à la suite d’une lésion d’ischémie-reperfusion pourrait être une réponse rénoprotectrice permettant de limiter les lésions inflammatoires.

Objectifs de l’étude : Nous avions pour objectif de valider la qualité de biomarqueur non invasif de l’hépcidine-25 urinaire dans une cohorte indépendante de patients subissant une chirurgie cardiaque, dans le contexte des prédicteurs cliniques de l’IRA.

Type d’étude : Il s’agit d’une étude de cohorte observationnelle prospective.
Cadre : L’étude a été réalisée au Canada, dans le cadre du programme de chirurgie cardiaque du St. Boniface Hospital de Winnipeg, au Manitoba.

Patients : La cohorte était constituée de 306 patients adultes subissant un pontage cardiopulmonaire.

Mesures : Le taux d’hépcidine-25 urinaire a été mesuré le jour suivant l’intervention.

Méthodologie : Une série d’échantillons d’urine périopératoire a été collectée chez une cohorte prospective et observationnelle constituée de 306 patients adultes subissant un pontage cardiopulmonaire. Le taux d’hépcidine-25 a été mesuré à l’aide d’un test ELISA compétitif, et sa performance diagnostique évaluée conjointement avec les paramètres cliniques et le score prédictif de Thakar en utilisant une régression logistique multivariée.

Résultats : Le taux d’hépcidine-25 urinaire était élevé en post-opératoire chez tous les patients de la cohorte, qu’ils soient ou non atteints d’insuffisance rénale. Une concentration élevée d’hépcidine-25 urinaire a été inversement associée à la survenue d’une IRA tant selon l’analyse univariée (RC=0,61 ; IC à 95 % 0,45-0,83 ; p=0,002) que selon l’analyse multivariée (RC=0,67 ; IC à 95 % 0,50-0,95 ; p=0,002). Un modèle combinant certains facteurs de risque cliniques a démontré que le DFG initial, le diabète sucré et la concentration d’hépcidine-25 urinaire présentaient une surface sous la courbe (SSC) de 0,82 (0,75-0,88) pour la prédiction du développement d’une IRA à la suite d’une chirurgie cardiaque. Ce niveau d’exactitude s’est avéré supérieur à la prédiction clinique déterminée par le score de Thakar.

Limites de l’étude : Deux principaux facteurs limitent la portée de l’étude : d’abord, le fait qu’il s’agît d’une étude observationnelle menée dans un seul centre, puis, la variabilité inhérente au test ELISA compétitif pour les anticorps polyclonaux.

Conclusion : Dans une analyse multivariée, lorsque combinée au DFG et au diabète sucré, l’hépcidine-25 urinaire est inversement associée à la survenue d’une IRA (SSC globale de 0,82). Ainsi, comparativement au score de Thakar utilisé seul, la mesure du taux d’hépcidine-25 urinaire prédit plus exactement le risque d’IRA.

Keywords
AKI, biomarkers, cardiac surgery, iron, ischemia reperfusion injury, renoprotection

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What was known before
Urinary hepcidin-25 was identified as a novel biomarker of acute kidney injury (AKI) using proteomic techniques, and its diagnostic performance has been evaluated in adult cardiac surgery cohorts. Recently, exogenously administered hepcidin-25 has been shown to be renoprotective in murine models of renal ischemia-reperfusion injury and hemoglobin-mediated AKI that both mimic cardiac surgery–associated AKI.

What this adds
This study validates that urinary hepcidin-25 is a non-invasive early marker for AKI in an independent adult cardiac surgery cohort. This study extends upon the previous work by demonstrating that urinary hepcidin-25 outperforms clinical prediction and serum creatinine.

Introduction
Acute kidney injury (AKI) following cardiac surgery requiring cardiopulmonary bypass (CPB) is a serious complication resulting in increased morbidity and mortality. Mild AKI occurs in 17% of cardiac patients (with CPB), and severe AKI requiring dialysis occurs in 2% of patients with a 60% mortality risk. Mild renal impairment, defined as a serum creatinine (Cr) >25% from baseline, is associated with a doubling in mortality up to 10 years following cardiac surgery.
independent of whether renal function recovers to baseline.\textsuperscript{7,8} Post-operative AKI is also an independent predictor of developing chronic kidney disease.\textsuperscript{9,10} Non-invasive urinary biomarkers may identify early injury in AKI patients and potentially provide a window for therapeutic intervention prior to loss of function.\textsuperscript{11}

Hepcidin-25 is a disulfide-rich antimicrobial peptide that regulates circulating iron by degrading and internalizing the cellular iron transporter, ferroportin, on hepatocytes, macrophages, and enterocytes.\textsuperscript{12} Renal ischemia reperfusion injury (IRI) is known to increase ferroportin expression in the spleen and liver, inducing hepatosplenic iron export.\textsuperscript{13} Tight regulation of ferroportin is necessary to keep circulating iron levels stable. Hepcidin is produced by the liver as an 84-amino acid prepropeptide and is approximately 90\% bound to \( \alpha_2 \)-macroglobulin in the circulation.\textsuperscript{14} Hepcidin is also expressed in the thick ascending limb, connecting tubules and collecting ducts of the kidney.\textsuperscript{15} Cleavage into the bioactive 25-amino acid form\textsuperscript{16} with a specific N-terminus is required for ferroportin interaction and intracellular iron sequestration.\textsuperscript{17} Hepcidin-20 and -22 isoforms have been detected in human serum and urine, but only hepcidin-25 is associated with ferroportin degradation.\textsuperscript{18-20}

Using an unbiased proteomics approach, we identified urinary hepcidin-25 as a marker for AKI avoidance\textsuperscript{21} and demonstrated that it is inversely associated with AKI at post-operative day 1 (POD 1).\textsuperscript{22} The goal of this study was to validate its diagnostic performance in an independent cardiac surgery cohort in the context of clinical AKI predictors.

**Methods**

The study protocol was approved by the University of Manitoba institutional review board (ethics approval HS15221(H2012:097)), and all patients provided written informed consent. All elective coronary artery bypass graft (CABG) and/or valve replacement patients from June 2012-October 2014 were considered for inclusion. Exclusion criteria were off-pump surgeries, pre-existing dialysis dependence or renal transplant, deceased prior to surgery, canceled/delayed surgeries, and missed urine sample bio-banking. Patient demographics and comorbidities were obtained. Intra-operative and post-operative data collection included duration of the operation, CPB and cross-clamp time, urine output, blood pressure, inotropic support, blood products, and potential nephrotoxic exposures (aminoglycoside, non-steroidal anti-inflammatories, angiotensin-converting enzyme inhibitors, angiotensin receptor blockers, and contrast).

**Urine Biobanking**

Samples were obtained at baseline (start of the operation), during early IRI (start of CPB, 1 hour into CPB), on arrival to intensive care unit (ICU), and POD 1 and 3 to 5 (Figure 1). Urine (20 mL) was collected via a Foley catheter placed after anesthetic induction until ICU discharge; ward patients provided a midstream catch. All samples were immediately placed on ice or stored at 4\(^\circ\)C and centrifuged at 870g for 6 minutes. The supernatant was collected and stored at −80\(^\circ\)C.
AKI KDIGO Criteria

Baseline serum creatinine was established using an average of two preoperative serum creatinine values for all patients. AKI was defined using the 2012 Kidney Disease Improving Global Outcomes (KDIGO) criteria: serum creatinine (Cr) rise >50% from baseline or a ≥265 μmol/L serum creatinine increase within any 48 hour period through till POD 4. The KDIGO urine output criterion was not used to determine AKI.

Urinary Hepcidin-25 ELISA

Urinary hepcidin-25 concentration was measured with a competitive ELISA (Penlab S-1337) (BioTek Synergy 4 microplate reader, Gen5 software, Fisher Scientific), as previously described. As urine hepcidin-25 has previously been demonstrated to peak at POD 1, these experiments were performed on POD 1 samples. Briefly, concentrations outside the standard curve range were serially diluted until detection. All samples were run in duplicate with an intra-assay coefficient of variation (CV) of 3.2%. An internal control was included on each ELISA plate using normal urine control as a continuous variable on the final multivariate analysis to control for plate-to-plate variation. Individual ORs (95% CI) were calculated for baseline estimated glomerular filtration rate (eGFR), diabetes mellitus, the internal control, and urine hepcidin-25 concentration. The combined area under the curve (AUC) was determined for receiver operating characteristic (ROC) curves in the final multivariate models. The goodness-of-fit was evaluated using the Hosmer-Lemeshow test. Finally, a supplemental analysis was performed excluding patients that developed AKI on or before POD 1 (n = 6).

Results

Prospective Observational Cohort

Three hundred eighty adult cardiac surgery patients were enrolled in the prospective observational cohort. Twenty-one patients were excluded for off-pump surgeries (n = 3), preexisting dialysis dependence (n = 3), deceased prior to surgery (n = 1), and canceled/delayed surgeries (n = 14). Fifty-three patients were excluded for missed urine sample bio-banking. Therefore, the final prospective observational cohort (n = 306) consisted of 41 AKI and 265 non-AKI patients (Figure 2). The baseline characteristics of patients who were excluded for missing urine samples (n = 53) were compared with the final study population (n = 306), and no significant differences were found between these groups (Supplemental Table 1).

Predictors of AKI

The patient characteristics of the prospective observational cohort were evaluated by univariate logistic regression in AKI versus non-AKI patients (Table 1). Increased urinary hepcidin-25 [AKI 628 (197-1749) vs. non-AKI 1599 (680-2753)] and urine hepcidin-25:Cr [AKI 67 (29-134) vs. non-AKI 121 (65-208)] was associated with a lower likelihood of developing AKI. Urinary hepcidin-25 (OR 0.61, 95%CI 0.45-0.83, p = 0.002) and urine hepcidin-25:Cr (OR 0.65, 95%CI 0.46-0.94, p = 0.02) at POD 1 were both univariate predictors for the subsequent development of AKI Additional univariate predictors for AKI development were baseline renal function as determined by eGFR [OR 0.95, 95%CI 0.94-0.97, p < 0.0001] and serum creatinine [OR 1.02, 95%CI 1.01-1.03, p = 0.0001]; Thakar score [OR 1.63, 95%CI 1.35-1.97, p < 0.0001]; diabetes [OR 3.82, 95%CI 1.93-7.54, p < 0.0001] and a previous history of hospitalized congestive heart failure [OR 3.82, 95%CI 1.72-8.47, p = 0.0005].

Urinary hepcidin-25 was inversely associated with AKI avoidance (OR: 0.67, 95% CI: 0.50-0.95, P = 0.02) on multivariate analysis, with higher concentrations associated with a lower likelihood of developing AKI (Table 2). A combined multivariate model was developed using baseline eGFR, diabetes mellitus, and urinary hepcidin-25. This model demonstrated AKI prediction with an AUC of 0.82 (95% CI: 0.75-0.88) and reasonable goodness-of-fit (Hosmer-Lemeshow, P = .90) (Table 2).
Figure 2. Prospective observational cohort of adult cardiac surgery patients (n = 306).
Note. KDIGO = Kidney Disease Improving Global Outcomes; AKI = acute kidney injury; Cr = creatinine; CKD = chronic kidney disease; eGFR = estimated glomerular filtration rate.

Table 1. Study Population Characteristics.

| Characteristic                  | Non-AKI (n = 265) | AKI (n = 41) | Odds ratio | 95% CI     | P value |
|--------------------------------|------------------|-------------|------------|------------|---------|
| Preoperative                   |                  |             |            |            |         |
| Age, y                         | 66 (59-71)       | 70 (64-75)  | 1.03a      | 0.99-1.07  | .09     |
| Male                           | 204 (77%)        | 27 (66%)    | 0.58       | 0.28-1.17  | .13     |
| Baseline eGFR, mL/min/1.73 m²   | 82 (66-97)       | 61 (43-76)  | 0.95       | 0.94-0.97  | <.0001  |
| Baseline creatinine, mg/dL      | 84 (72-100)      | 102 (85-141)| 1.02       | 1.01-1.03  | <.0001  |
| Thakar score                   | 2 (1-2)          | 3 (2-5)     | 1.63       | 1.35-1.97  | <.0001  |
| Diabetes mellitus              | 77 (29%)         | 25 (61%)    | 3.82       | 1.93-7.54  | <.0001  |
| Chronic obstructive pulmonary disease | 22 (8%)          | 2 (5%)      | 0.57       | 0.13-2.51  | .45     |
| Hospitalized congestive heart failure | 20 (8%)          | 11 (27%)    | 3.82       | 1.72-8.47  | .001    |
| Previous myocardial infarction | 99 (37%)         | 12 (29%)    | 0.69       | 0.34-1.42  | .32     |
| Previous CABG                  | 8 (3%)           | 2 (5%)      | 1.65       | 0.34-8.04  | .54     |
| Peripheral arterial disease    | 18 (7%)          | 6 (15%)     | 2.22       | 0.83-5.94  | .11     |
| Amputation or peripheral arterial disease bypass | 2 (1%)         | 0 (0%)      | —          | —         | —       |
| Previous cerebrovascular accident | 12 (5%)          | 4 (10%)     | 2.28       | 0.70-7.44  | .17     |
| Previous transient ischemic attack | 10 (4%)          | 1 (2%)      | 0.64       | 0.08-5.12  | .67     |
| European system for cardiac operation risk, % | 1.4% (0.8%-2.5%)  | 3.0% (1.5%-5.6) | 1.06 | 0.99-1.13 | .07     |
| Type of surgery (isolated CABG) | 169 (64%)        | 23 (56%)    | 0.73       | 0.37-1.41  | .35     |
| Intraoperative                 |                  |             |            |            |         |
| Pump time, min                 | 95 (72-131)      | 97 (82-144) | 1.34b      | 0.93-1.92  | .11     |
| Cross-clamp time, min          | 64 (45-92)       | 61 (45-95)  | 1.10b      | 0.77-1.58  | .60     |
| Operating room duration, min   | 262 (220-339)    | 261 (203-352)| 1.03b    | 0.85-1.24  | .77     |
| Intraoperative urine output, mL | 630 (455-970)    | 628 (197-1749)| 0.57c   | 0.25-1.29  | .18     |
| Postoperative                  |                  |             |            |            |         |
| Urine hepcidin-25, ng/mL       | 1599 (680-2753)  | 628 (197-1749)| 0.61d  | 0.45-0.83  | .002    |
| Urine hepcidin-25:Cr, μg/mmol  | 121 (65-208)     | 67 (29-134) | 0.65a      | 0.46-0.94  | .02     |

Note. Values expressed as median (interquartile range) or n (%). Continuous variables compared using Mann-Whitney test, categorical variables compared using chi-square or Fisher’s exact test. Missing covariate values were estimated via multiple imputation. eGFR, estimated glomerular filtration rate; CABG, coronary artery bypass graft; Cr, creatinine.
Odds ratio: aAge (per year); bTime (per hour); cIntraoperative urine output (per L); dUrine hepcidin-25 (per 1000 ng/mL); eUrine hepcidin-25:Cr (per 100 μg/mmol).
In order to control for the observed plate-to-plate variability, the multivariate analysis was recapitulated including the internal control as a variable. Urinary hepcidin-25 remained a predictor of AKI (OR: 0.70, 95% CI: 0.51-0.96, \( P = .03 \)). The combined AKI prediction model using baseline eGFR, diabetes mellitus, and urinary hepcidin-25 demonstrated an AUC of 0.83 (95% CI: 0.76-0.89) and reasonable goodness-of-fit (Hosmer-Lemeshow, \( P = .72 \)) (Table 3). The Thakar score was not significant in either analysis.

**Discussion**

The principle finding of this study was that urinary hepcidin-25 is inversely associated with AKI avoidance in an independent, prospective observational adult cardiac surgery cohort. Importantly, a combined clinical and biomarker model with baseline eGFR, diabetes and hepcidin-25 demonstrated an AUC of 0.82 for the subsequent development of AKI. These findings are consistent with our previous study\(^2\) and extend them by demonstrating that they outperformed clinical AKI prediction with the Thakar score.
All cardiac surgery patients are exposed to IRI with varying degrees of tubular stress or injury,25 resulting in activation of different renoprotective mechanisms. As urinary hepcidin-25 is increased in non-AKI versus AKI patients following IRI, we previously postulated that it is a renoprotective response which may be mediated via the intracellular sequestration of iron to limit oxidative stress, free radical damage, and renal injury.21 Since then, Scindia et al have found hepcidin to be highly protective in a murine model of renal IRI. Notably, the administration of hepcidin resulted in a significant reduction in tubular injury, apoptosis, renal oxidative stress, and neutrophil infiltration, as well as improved renal function.13 Furthermore, these findings are highly consistent with van Swelm et al who demonstrated that the administration of hepcidin-25 decreased markers of kidney injury in a murine model of hemoglobin-mediated AKI.26

Labile ferrous iron release is an inevitable consequence of cardiac surgery. The mechanical forces exerted during extracorporeal circulation induces hemolysis and free hemoglobin release,27 with increasing CPB duration correlated with increased free hemoglobin release.28 Elevated free hemoglobin levels are associated with AKI following on-pump repair of aortic aneurysms.25 Red blood cell (RBC) storage can result in loss of structural integrity and increased hemolysis following blood transfusion,25,30 while myoglobin release may also contribute to the circulating labile iron pool.31 Finally, renal IRI alone can induce increased serum iron levels, with associated kidney iron accumulation.13 The intersection of these findings with the observed renoprotective effect of hepcidin in murine models of renal IRI and hemoglobin-mediated AKI, all suggest that iron handling is a crucial and potentially modifiable factor of cardiac surgery–associated AKI.

While hepcidin-25 is renoprotective in animal models, its underlying mechanisms in cardiac surgery–associated AKI remains unknown as there are conflicting observations regarding the role of apoptosis and renal ferroptosis.13,26 Notably, these observations suggest that hepcidin-25 may act via alternative pathways to prevent renal tubular epithelial cell death.13,26 Ferroptosis is a novel iron-dependent form of regulated cell death that has recently been demonstrated to play a key role in AKI-induced cell death.32,33 Indeed, inhibition of ferroptosis has added protective effects beyond necroptosis and necrosis inhibition alone in a murine model of severe renal IRI.32 Furthermore, ferroptosis was the primary mediator of cell death and renal injury in folic acid–induced AKI. In this model, inhibition of ferroptosis protected renal function, histological injury, and cell death, whereas inhibition of apoptosis and necroptosis were ineffective.33 These findings are intriguing in that they delineate an alternative iron-dependent pathway to renal injury that is shared between different types of AKI. It is tempting to speculate that the lack of consistently observed inhibition of apoptosis by hepcidin-25 administration13,26 may be due to activation of alternative cell death pathways, such as ferroptosis, but this cannot be clarified in our human model of IRI-AKI. These findings are consistent with data on another iron-binding protein, neutrophil gelatinase-associated lipocalin (NGAL), which has been demonstrated to be renoprotective as part of an NGAL:iron:siderophore complex that upregulates heme-oxygenase.34 Taken together, these findings suggest that hepcidin-25 may be part of a coordinate response involving different iron-binding proteins that act to mitigate the response to human renal IRI.

Hepcidin is freely filtered by the glomerulus and reabsorbed at the proximal tubule via megalin-dependent endocytosis26,35; the fractional excretion is negligible with up to 97% reabsorbed in physiological conditions.26 While the source of the observed urinary hepcidin-25 in our human model is unknown, hepcidin is locally synthesized in kidney and released in the distal nephron in AKI.26 These findings are consistent with cardiac surgery models of AKI which demonstrated increased urinary hepcidin-25 and increased fractional excretion of hepcidin-25 in non-AKI patients, in the absence of substantive changes to plasma/serum hepcidin-25, even after excluding chronic kidney disease patients.37,38 Taken together, we speculate that elevation of urinary hepcidin-25 following cardiac surgery is the result of increased distal nephron production in response to IRI, but it remains to be determined whether luminal enzyme activity plays an in vivo regulatory role by inactivating hepcidin-25.

This study had several limitations. First, this is strictly an observational study. Although we speculate that urinary hepcidin-25 is elevated in human IRI as part of a coordinated renoprotective response, causation cannot be determined. Second, this is a single-center study. These findings confirm those of other single-center studies,22,37,38 although urine hepcidin:Cr was not significant after correcting for dilutional factors in this study. It is possible that the urinary hepcidin-25 concentration in the lumen of the distal tubule is predictive of protection, instead of mass excretion rate. An alternative possibility is that urinary hepcidin-25 may be less sensitive for milder phenotypes of AKI, as this cohort consisted of largely Stage 1 AKI (Stage 1, n=38; Stage 2, n=1; and Stage 3, n=2). Therefore, these findings need to be verified in a multi-center setting. The patterns of hepcidin-25 excretion in other human AKI models remains unknown; however, the observed effect of iron-dependent cell death in different mouse AKI models suggests that iron may be a key regulator to cell death/injury beyond IRI or hemoglobin-mediated AKI. Fifty-three patients were excluded for missed bio-banking, although no significant differences were identified between the excluded and final study population.

Hepcidin-25 is limited as an early, non-invasive biomarker by its polyclonal antibody–based ELISA assay. This resulted in significant plate-to-plate variation between lots, which we identified with our internal controls. We accounted for this variability in the analysis and demonstrated that it did not substantially impact the overall findings. While a hepcidin-25 monoclonal antibody sandwich ELISA has been developed,39
these antibodies were not available at the time of our experiments. Clinically applicable biomarkers require a robust, reproducible assay that is readily translatable to clinical laboratories and conform to Good Laboratory Practice guidelines. Finally, hepcidin-25 peaks later at POD 1, similar to KIM-1 (kidney injury molecule-1), and therefore has limited utility for early diagnosis compared with biomarkers that peak earlier. Nevertheless, this cohort provides useful independent observations regarding the consistent response of urinary hepcidin-25 as a marker of AKI avoidance and helps shed insight into the pathophysiology of human renal IRI.

**Conclusions**

Urinary hepcidin-25 is elevated following cardiac surgery and is inversely associated with AKI on an independent level. These findings independently validate the existing literature in a prospective, observational cohort of adult cardiac surgery patients, and extend them to show that a combined clinical and biomarker model with baseline eGFR, diabetes mellitus, and hepcidin-25 has an AUC of 0.82 for the subsequent development of AKI. These findings suggest that iron-dependent pathways are key mediators of renal IRI and are consistent with animal models that demonstrate hepcidin-25 is renoprotective in AKI. It remains to be determined whether iron modulation can mitigate the effects of human AKI, and the results of a clinical trial evaluating the utility of iron chelation in AKI with deferoxamine are pending (NCT00870883).

**Ethics Approval and Consent to Participate**

The study protocol was approved by the University of Manitoba institutional review board, and all patients provided written informed consent (ethics approval HS15221 [H2012:097]).

**Consent for Publication**

We have authors consent for publication.

**Availability of Data and Materials**

Data is available upon request.

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**Declaration of Conflicting Interests**

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