Systematic review of prognostic prediction models for acute kidney injury (AKI) in general hospital populations

Luke Eliot Hodgson,1 Alexander Sarnowski,2 Paul J Roderick,1 Borislav D Dimitrov,1 Richard M Venn,3 Lui G Forni2,4

**ABSTRACT**

**Objective** Critically appraise prediction models for hospital-acquired acute kidney injury (HA-AKI) in general populations.

**Design** Systematic review.

**Data sources** Medline, Embase and Web of Science until November 2016.

**Eligibility** Studies describing development of a multivariable model for predicting HA-AKI in non-specialised adult hospital populations. Published guidance followed for data extraction reporting and appraisal.

**Results** 14 046 references were screened. Of 53 HA-AKI prediction models, 11 met inclusion criteria (general medicine and/or surgery populations, 474478 patient episodes) and five externally validated. The most common predictors were age (n=9 models), diabetes (5), admission serum creatinine (SCr) (5), chronic kidney disease (CKD) (4), drugs (diuretics (4) and/or ACE inhibitors/angiotensin-receptor blockers (3)), bicarbonate and heart failure (4 models each). Heterogeneity was identified for outcome definition. Deficiencies in reporting included handling of predictors, missing data and sample size. Admission SCr was frequently taken to represent baseline renal function. Most models were considered at high risk of bias. Area under the receiver operating characteristic curves to predict HA-AKI ranged 0.71–0.80 in derivation (reported in 8/11 studies), 0.66–0.80 for internal validation studies (n=7) and 0.65–0.71 in five external validations. For calibration, the Hosmer-Lemeshow test or a calibration plot was provided in 4/11 derivations, 3/11 internal and 3/5 external validations. A minority of the models allow easy bedside calculation and potential electronic automation. No impact analysis studies were found.

**Conclusions** AKI prediction models may help address shortcomings in risk assessment; however, in general hospital populations, few have external validation. Similar predictors reflect an elderly demographic with chronic comorbidities. Reporting deficiencies mirrors prediction research more broadly, with handling of SCr (baseline function and use as a predictor) a concern. Future research should focus on validation, exploration of electronic linkage and impact analysis. The latter could combine a prediction model with AKI alerting to address prevention and early recognition of evolving AKI.

**INTRODUCTION**

Acute kidney injury (AKI) is defined as an acute increase in serum creatinine (SCr) or reduction in urine volume.1 The incidence of AKI is increasing, affecting up to one in five hospitalised adults worldwide.1 A continuum of injury exists long before sufficient loss of excretory kidney function can be measured with standard laboratory tests (ie, SCr) and is associated with mortality. Associated mortality remains high, in part reflecting the severity of the underlying disease, but may also be due to the limitations of conventional markers to detect early injury.2

Deficits in recognition and management of patients with AKI have led to practice guideline calling for improved risk assessment, at which point interventions could be most beneficial.7 One suggested strategy to achieve...
this aim is through the implementation of clinical prediction models. Though development and validation of AKI prediction models is desirable, clinical application in this and other fields has been hampered for a number of reasons:

- potential predictors and models continuously increase with new studies often finding conflicting results.
- substandard reporting of methodology and results make conclusions problematic.
- few general hospital population studies exist; specialist fields (cardiac and transplant surgery and contrast-induced (CI-AKI)) account for the majority of CI-AKI models and all systematic reviews but are unlikely to be generalisable and
- models rarely enable electronic automation as part of clinical workflow, known to influence uptake.

High-quality systematic reviews of prediction models have been called for. Following recent reporting guidance (CHecklist for critical Appraisal and data extraction for systematic Reviews of prediction Modelling Studies (CHARMS) and Transparent Reporting of a multivariable prediction model for Individual Prognosis or Diagnosis (TRIPOD)), this review appraises hospital-acquired AKI (HA-AKI) prediction models in general populations who, in the UK, account for the majority of hospital episodes and AKI cases.

### METHODS

Published guidance (CHARMS, TRIPOD and Preferred Reporting Items for Systematic Review and Meta-Analysis (PRISMA)) helped frame the review question, data extraction, reporting and appraisal. The research question was: what are the available prognostic prediction models for the development of HA-AKI in adult general populations? Using explicit, systematic methods to minimise bias and provide reliable findings from which conclusions can be drawn and decisions made, the review aimed to collate empirical evidence for AKI prediction models across general hospital settings, fitting prespecified eligibility criteria (online supplementary table 1). Performance was assessed by discrimination and calibration including validation studies. The presence of any impact analysis studies was also investigated. The review aimed to provide recommendations for the most robust, usable models, including the ability to incorporate future electronic data linkage, for example, between the community (primary care) and hospital.

### Data sources, study selection and data extraction

We searched MEDLINE, Embase and Web of Science databases (inception to November 2016) using recommended filters (online supplementary tables 2–4). Titles and abstracts were screened by two reviewers (LEH and AS), and full articles were reviewed if eligible. Disagreements were resolved by iterative screening rounds. Reference lists from retrieved articles, systematic reviews, national and international guidance and our own literature files were also analysed. Data extraction and quality assessment were performed by two investigators (LEH and AS) with disagreements resolved by a third reviewer (LGF). A data extraction form was used based on previous reviews and guidance (summary online supplementary table 5).

### Outcome, model performance and clinical utility

It was anticipated that study outcome, HA-AKI, would vary given the numerous definitions in use prior to Kidney Disease: Improving Global Outcomes (KDIGO) in 2012. Thus, during the search strategy, we included studies with an SCr around admission and repeated during a hospital admission to diagnose HA-AKI. Information was gathered on how a study defined a patients baseline renal function, how community AKI cases were handled, whether SCr was used as a predictor in analysis and finally the magnitude and timeframe used to define the outcome. Discrimination and calibration are the most common methods to assess model performance. Discrimination is usually assessed graphically by the area under the receiver operating characteristic curve (AUROC), representing how well a model separates and ranks patients who experienced the outcome from those who did not. For prediction models, the AUROC, which focuses solely on accuracy, has a number of shortcomings, such as a lack of information on consequences and when used in populations where the outcome prevalence is rare. Calibration describes how well predicted results agree with observed results. The Hosmer-Lemeshow (H-L) test, despite limitations, is the most commonly used calibration statistic. It is also recommended to graphically plot expected and actual outcomes, for example, with a calibration slope. In addition to performance, ease of bedside use and whether the models could be electronically automated—factors known to influence successful uptake—were recorded. A quantitative synthesis of the models was not performed, being beyond the scope of review and formal methods for meta-analysis of prediction models are yet to be fully developed.

### Study quality assessment

A global TRIPOD score for each study was calculated to quantify reporting, consisting of the sum of the scores for each individual item (out of a maximum 37, with a score of 1 for criterion met, score of 0 for each item not met or unclear). As yet there has been no suggested cut-off for what represents a high-quality study, though it would be reasonable to judge that those studies with the most significant gaps in reporting are likely to be
at higher risk of bias. Furthermore, the quality (risk of bias) of each study was assessed by piloting a version of Prediction study Risk Of Bias Assessment Tool (PROBAST), a tool for assessing risk of bias and applicability of prognostic model studies, nearing completion and ready for piloting when this review was undertaken (Wolff R, Whiting P, Mallett S, et al, personal communication, website: http://s371539711.initial-website.co.uk/probast/). Elements were considered in the following domains: study participants, predictors, outcome, sample size and missing data, statistical analysis and overall judgement of bias and applicability.

**Patient involvement**

Patients were not involved in setting the research question, outcome, design and implementation of the study. There are no plans to involve patients in dissemination.

**RESULTS**

From 14,046 articles identified by the search strategy, 254 full articles were reviewed (PRISMA flow chart, figure 1). Specialised fields (predominantly cardiac surgery, transplantation or CI-AKI) accounted for 61 of 74 (82%) of all studies. This review included 11 general model studies (n=474,478 patient episodes), in general surgery,33,34 trauma and orthopaedics (T&O),35 general hospital cohorts (predominantly medicine and surgery)36–40 and heart failure (summarised in table 1 and online supplementary etable 6, with abbreviations in supplementary etable 7).41–43 Two further studies were purely external validations.44,45 HA-AKI incidence was 7% (21,641 events), though this varied from <1% in the general surgery models33,34 to 28% across the heart failure studies, and heterogeneous definitions (timeframe and marker) were employed (see table 1 for definitions used with further information in online supplementary etable 6).

**Figure 1** PRISMA study flow chart.24 AKI, acute kidney injury; PRISMA, Preferred Reporting Items for Systematic Review and Meta-Analysis.
Table 1  Summary of HA-AKI prediction models

| Population | General surgery | T&O | General (medical and surgical) | Heart failure |
|------------|----------------|-----|--------------------------------|---------------|
| Author, year (n=derivation) | Kheterpal et al, 2007 (n=14066) | Kheterpal et al, 2009 (n=57080) | Bell et al, 2015 (n=6220) | Drawz et al, 2008 (n=360) | Matheny et al, 2010 (n=26107) | Koyner et al, 2016 (n=202961) | Bedford et al, 2016 (n=7556) | Fomie et al, 2013 (n=1867) | Forman et al, 2004 (n=1004) | Breidhardt et al, 2011 (n=657) | Wang et al, 2013 (n=1010) |
| Centres, design | 1, R | 121, R | 3, R | 3, CC | 1, R | 5, R | 3, R | 1, P | 11, R | 1, P | 1, R |
| Age (with outcome) | 59 | 65 (±15) | 77 (±11) | 67 | – | 70 (±16) | – | 80 (70-86) | 68.7 | 79 (72-85) | 73 (67-78) |
| Age (no outcome) | 47 | 54 (±17) | 70 (±16) | 63 | – | 63 (±19) | – | 73 (61-81) | 66.8 | 79 (70-85) | 71 (63-75) |
| Outcome predicted | eGFR <50 (<7 days) | ≥TSCr >177 μmol/L, RRT (30 day) | KDIGO TSCr | TSCr* | RIFLE TSCr | KDIGO TSCr (24 hours) | KDIGO TSCr (72 hours) | KDIGO TSCr (<7 days) | ↑ TSCr >26.5 μmol/l† | ↑ TSCr >26.5 μmol/l† | AKIN TSCr (<48 hours) |
| Events | 121 | 561 | 672 | 120 | 1352 | 17541 | 222 | 95 | 271 | 136 | 341 |
| Mortality with outcome (%) | 15 | 42 | – | – | – | 6 | – | 20 | 27 | 17 | 17 |
| Mortality no outcome (%) | 3‡ | 8‡ | – | – | – | 1 | – | 4 | – | 6 | 2 |
| Predictors tested | 30 | 19 | 11 | 19 | 23 | 29 | 45 | 25 | 29 | 48 | 35 |
| Predictors included | 7 | 9 | 7 | 7 | 27 | 29 | 12 | 7 | 4 | 3 | 8 |
| EPP | 4 | 30 | 61 | 6 | 59 | 605 | 5 | 4 | 9 | 3 | 10 |
| Inappropriate handling of SCr | X | X | X | X | X | X | X | X |
| Derivation AUROC | 0.77 | 0.80 | 0.74 | 0.73 | 0.75 | – | – | 0.72 | – | 0.71 | 0.76 |
| IV AUROC | – | 0.80 | 0.73 | 0.66 | – | 0.74 | 0.67 | 0.76 | – | – | 0.76 |
| EV AUROC | 0.67 | X | 0.71 | X | X | X | 0.71 | 0.65-0.71‡ | 0.65, 0.65 | X | X |
| Derivation calibration | RR | RR | Plot | – | H-L p=0.29 | – | – | H-L p=0.96 | – | – | H-L p=0.98 |
| IV calibration | – | RR | Plot | RR | Plot | – | H-L p=0.04 | – | – | H-L p=0.13 |
| EV calibration | RR | – | Plot | – | – | – | H-L p=0.12 | H-L p=0.06-0.09, plot | RR | – | – |
| TRIPOD items reported | 25 | 28 | 34 | 26 | 28 | 24 | 29 | 29 | 26 | 23 | 30 |
| Bedside calculation | – | – | – | – | – | – | Yes | Yes | Yes | Yes | Yes |
| Electronic automation | – | – | Yes† | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes |

* Increase sCr ≥44 μmol/L if baseline Scr of ≤168 μmol/L, ≥88 μmol/L baseline 177–433 μmol/L and ≥133 μmol/L baseline >442 μmol/L.
† During admission.
‡Propensity matched.
§Propensity matched.
††Used linked community and hospital data.

AKIN, Acute Kidney Injury Network; AUROC, area under the receiver operating characteristic curve; CC, case-control; eGFR, estimated glomerular filtration rate; EPP, events per predictor; EV, external validation; IV, internal validation; HA-AKI, hospital-acquired acute kidney injury; H-L, Hosmer-Lemeshow test; KDIGO, Kidney Disease: Improving Global Outcomes; mortality, in-hospital; P, prospective; plot, calibration plot; R, retrospective; RIFLE, Risk, Injury, failure, loss of kidney function; RR, risk range; RRT, renal replacement therapy; TSCr, serum creatinine; T&O, trauma and orthopaedics; TRIPOD, Transparent Reporting of a multivariable prediction model for Individual Prognosis or Diagnosis.
Table 2  Summary of limitations in methodology and reporting

| Area of concern                                      | Description                                                                                                                                                                                                 |
|------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Missing data                                         | Multiple imputation recommended to avoid bias—rarely described.12 50                                                                                                                                       |
| Definitions of outcome and predictors                | No consistent strategy used to differentiate CA-AKI from HA-AKI. Two studies excluded patients with pre-existing CKD33 37; five studies took admission SCr as baseline; five included SCr as predictor despite it forming the outcome; comorbidities inconsistently defined: including from admission diagnoses or coded history. |
| Blinding of predictors or outcome                    | Not reported.                                                                                                                                                                                              |
| Sample size                                          | Calculations not described, six studies had <10 EPP. Small sample increases risk of overfitting and underfitting.12 49                                                                                       |
| Univariate to select for multivariate analysis       | Technique not recommended, used in 10 of 11 models.12                                                                                                                                                       |
| Bootstrapping                                        | Adjust for optimism, without losing information—rarely described.35 43                                                                                                                                   |
| Calibration plots                                    | Important part of model performance,12 present in only one model and one external validation.36 45                                                                                                         |
| External validation and model updating               | Validation adjusts for optimism, assesses generalisability but was scarce, while model updating is recommended but not described.12                                                                        |
| Newer performance measures                           | Techniques such as decision curve analysis offer insight into clinical consequences—not described.29                                                                                                           |
| Use of data linkage                                  | Only one study used data linkage.35                                                                                                                                                                          |

AKI, acute kidney injury; CA-AKI, community-acquired AKI; CKD, chronic kidney disease; EPP, events per predictor; HA-AKI, hospital-acquired AKI; SCr, serum creatinine.

For example, five studies took admission SCr to represent a patients baseline, potentially confusing CKD, established and emerging AKI.34 38 40 42 43 Of note, one study produced a model to predict AKI admission as well as HA-AKI at 72hours with the former not considered suitable for analysis in this review.39

In seven of the nine studies reporting age, this was significantly higher in the group with the outcome, with eight studies reporting a mean or median age over 65 years in the outcome group (table 1). Mortality was significantly higher in those who developed the outcome in the six studies where data were available (ranging 6%-42%). No impact analyses were retrieved.

Study reporting

A median 28 (IQR 25–30) of 37 recommended items were reported, suggesting significant shortcomings (key shortcomings are summarised in table 2 with TRIPOD reporting summarised in online supplementary etable 8). By design, eight studies were retrospective, two were prospective and one was a case control. Five studies were single centre. USA (n=6) and UK (n=3) accounted for the majority of the models. Only three studies used imputation techniques for missing data.34 35 38 Definitions were heterogeneous (table 1) with five using Risk, Injury, failure, loss of kidney function (RIFLE),37 Acute Kidney Injury Network (AKIN),42 or KDIGO criteria for changes in SCr.35 36 39 One study used KDIGO SCr change within a 24-hour timeframe of predictors being measured.40

Candidate predictors, model building and sample size

A median of 29 (IQR 19–35) predictors were considered, though frequently studies only reported those significant on univariate or multivariate analysis. Blinding of assessment of predictors and study outcome was not mentioned. Continuous predictors were dichotomised in four studies, and 10 studies used univariate analysis to select for multivariate analysis. No models mentioned shrinkage techniques or sample size calculations. Median number of outcome events was 271 (121–672). For statistical power, all of the studies had more than 10 events per predictor (EPP) included in the model. However, the EPP was <10 in six studies, when accounting for the total number of candidate predictors assessed.35 36 38 39 41 43 Of a total of 56 different predictors, a median of 7 (7–12) was included per model, including demographics, history, procedure information, laboratory parameters, physiological observations and hospital admission diagnoses (most common presented in figure 2, full details online supplementary etables 9–10). Only four studies included physiological parameters in their final model.36 38 40 43 Seven studies included admission SCr as potential predictor with five including this in the final model, thus potentially confusing prediction with a diagnosis of AKI.34 37 40 42 43 Each study’s handling of SCr in terms of when a baseline was calculated (prior or at admission) and whether SCr was used as a predictor are summarised in online supplementary etable 11.

Model performance

Median AUROC (or C-Statistic) was 0.745 (range 0.71–0.80) for derivation (eight studies) and 0.74 (range 0.66–0.80) for internal validations reporting discrimination (seven studies). Excluding the studies using non-consensus-based definitions and those including admission...
Figure 2  Predictors most frequently included in the 11 HA-AKI prediction models. ACEi, ACE inhibitors; ARBs, angiotensin-receptor blockers; Bloods, laboratory parameters; CKD, chronic kidney disease; HA-AKI, hospital-acquired acute kidney injury; ↓HCO₃, reduced serum bicarbonate; SCr, serum creatinine; ↑WCC, raised white cell count.

SCr as predictor and/or baseline, only four studies were left.35 36 39 41 In these studies, AUROCs ranged 0.71–0.74 in derivation (three studies), 0.67–0.76 for internal validation (three studies) and 0.65–0.71 for external validation (three studies). Only one model study presented a calibration plot for derivation and validation.35 The H-L statistic was used in three derivations36 37 42 and two internal validations.39 42

Five models have been externally validated: on separate populations within the same study,35 36 39 other model studies43 or stand alone external validations,35 45 where the AUROCs were moderate, ranging 0.65–0.71. One validation provided a calibration plot,35 one the H-L statistic39 and one reported both.45 In the Bell external validation cohort calibration suggested the model overpredicted the outcome requiring recalibration.35 In the internal validation of the Forni study calibration plots showed agreement at low probability rates, while at higher rates calibration deviated in the medical cohort.45 Two of the three surgical models have been externally validated: the Kheterpal model,35 in a Chinese population (AUROC 0.66),44 and the UK T&O study used a third centre for external validation.35 Two of five mixed general population models have external validation,36 39 the latter having been derived on medical patients and externally validated in medical and surgical cohorts.45

The first of the three heart failure studies was externally validated in the subsequent studies with inferior discrimination (AUROC 0.65 in both validations).41–43 No model updating was reported.

Quality assessment and risk of bias summary

Quality assessment was based on a draft version of the PROBAST tool. This suggested evidence in 9 of the 11 included studies of a high risk of bias (summarised in table 3) with shortcomings across the major domains...
of the assessment. For example, one study used a case-control design, which is inappropriate for developing a prediction model as it does not enable calculation of absolute risks and thus yields incorrect estimates of model intercept or baseline hazard. A wide variety of predictors were considered with use of univariate analysis to select for multivariate in 7/11 of the studies. Six studies were potentially underpowered having less than 10 EPP assessed. Seven of the studies introduced potential bias in handling of renal function and SCr either in failing to establish a reliable baseline renal function, excluding patients with reduced renal function or employing it as a predictor. Finally, outcome definition frequently varied in part owing to a number of the studies preceding consensus definitions.

**DISCUSSION**

**Principal findings**

In this first systematic review of HA-AKI prediction in general hospital settings, the most common predictors were age, diabetes, CKD, drugs, heart failure, SCr and bicarbonate. Modest discrimination performance of all the models is unsurprising when attempting at a single time point to predict a future event reflecting diverse aetiologies, affecting heterogeneous patient groups. Significant shortcomings mirror those described elsewhere.

- multiple similar models, rarely externally validated
- no impact analysis or evidence of clinical implementation
- incomplete reporting and
- little consideration of electronic automation (allowing presentation without additional data input beyond usual clinical care), which influences uptake.

Methodological and reporting shortcomings in the studies (summarised in table 2) included six studies having less than 10 EPP potentially leading to overfitting, with only three employing multiple imputation to handle missing data that can increase sample size and power.

Handling of SCr and CKD was of particular concern in a number of areas. First, in part due to a previous lack of a consensus definition, the outcome in question, HA-AKI, had heterogeneous definitions, both in magnitude of SCr rise and timeframe. For example, the Kheterpal study used a rise in SCr ≥177 μmol/L, which has been shown to significantly underestimate rates of AKI when compared with more recent definitions. Koyner et al. used a rolling timeframe of 24 hours while others used SCr elevation at any point during an admission. Indeed one study produced a separate model to predict AKI at admission to hospital. This was further confused in seven studies by the inclusion of admission SCr as a potential predictor (with inclusion in five models), five studies taking admission SCr to represent a patient’s baseline and two studies excluding all patients with a reduced admission estimated glomerular filtration rate from their analysis. This risks confusing prediction and detection of AKI events. Issues with differing definitions have been described before in systematic reviews of prediction models and should be considered when researchers embark on future studies.

A formal risk of bias assessment (PROBAST) suggested the majority of studies had domains placing the studies at high risk of bias. Published after TRIPOD, Bell and colleagues’ model provides researchers with a good template for adherence to reporting guidance, with a low risk of bias and demonstrates the utility of data linkage (eg, between community and hospital), though lack of validation in other populations tempers recommendation for implementation.

**Strengths and limitations of this review**

This review summarises the currently available AKI prediction models in general populations who account for the majority of hospital admissions and AKI cases. The models were selected following an extensive literature search, and the review employed the most recent critical appraisal guidance and risk of bias assessment. The large number of patient episodes provides important insights into AKI prediction complementing other recent reviews in cardiac surgery, CI-AKI, liver transplantation and non-cardiac surgery. In-patient mortality in those who developed the outcome ranged 6%–42% (in the six studies reporting mortality), emphasising this is a crucial group to promptly identify.

The first limitation is the small number of externally validated models, which tempers recommending one model over another. Second, though we aimed to include general populations, caution should be employed, for example, when comparing a model derived on heart failure patients to one from an orthopaedic cohort. However, in many UK hospitals, such populations share similarities (predominantly elderly demographic with comorbidities), and if one aim of a prediction model is generalisability, a model should be tested in these different fields. Third, as study outcome definitions and handling of SCr (baseline and as predictor of outcome) were heterogenous, model comparisons are problematic, though recent studies were more likely to use KDIGO SCr change. Fourth, no studies included urine output, probably reflecting the small number of patients who have this marker closely monitored. Fifth, TRIPOD recommendations were used as a reporting benchmark; however, the relative importance of individual items and what constitutes an acceptable ‘score’ is arguable, though a formal risk of bias assessment was also carried out (PROBAST) providing further insight into respective study strengths and weaknesses. The absence of impact analysis limits the recommendation of one model over another. Finally, a meta-analysis was not performed without access to individual participant data. Expert guidance now exists in this area and offers opportunities to improve the scope of external validation research.
Comparison with previous systematic reviews

Both this study and a review of CI-AKI models found pre-existing predictors: age, CKD, diabetes and heart failure to be the most commonly included. A cardiac surgery review reported specialty-specific predictors in addition to these chronic comorbidities. A non-cardiac surgery review (five of six studies in liver transplantation or resection) reported age, CKD and diabetes in at least two models. Finally, a liver transplantation review highlighted the importance of CKD and (unsurprisingly) liver dysfunction. The present review found drugs or acute laboratory values frequently included, though only four models included acute physiological parameters. Our study and the non-cardiac surgery review included adherence to recommended TRIPOD reporting with similar shortcomings. Across the other reviews, only in the fields of CI-AKI and cardiac surgery were external validations reported. Ease of use (including if necessary a calculator) and potential for electronic automation were rarely considered across the models reviewed. No impact analysis studies have been described.

Future directions

Management of HA-AKI presents a significant challenge that could be helped by robust prediction models to risk stratify populations, encourage prevention and promote prompt recognition. Appraisal and synthesis of prediction studies may enable clinicians and policymakers judge model utility; however, this is problematic when key study details are not reported. Though much of the AKI literature is on (often assumed) HA-AKI, the majority of cases arise from the community (community-acquired AKI). Indeed, a recent study demonstrated a significant proportion of such patients are never hospitalised. This review suggests even in HA-AKI, the strongest predictors are pre-existing patient factors. The two laboratory measures frequently included—SCr and bicarbonate—may also reflect a chronic component. It is likely a proportion of cases classed as HA-AKI represent (evolving) community cases; thus, models using such pre-existing risk factors makes clinical sense. This continuum of harm between community and hospital could suggest that a risk prediction model in place at or even before hospital admission, combined with early flagging of those who have met AKI criteria, may be required to improve outcomes.

Electronic linkage of patient records between community and hospital data is desirable to ensure accurate inclusion of predictors (chronic morbidity, medication, laboratory and physiological parameters). This may also enable bedside automation as part of clinical workflow, where there is evidence that beneficial implementation can be achieved. Acute physiological parameters assessed as predictors in seven studies and subsequently included in only four studies could be an avenue of future research to improve the modest performance of all models at a single time point (admission to hospital) described to date. As hospitals increasingly employ electronic track and trigger observation systems, this may then enable the application of complex statistics (eg, machine learning) to account for the effects of trends and repeated measures. Risk stratification using chronic comorbidity and medication(s) with trends in physiology could be further enhanced by measurement of urine output and/or newer biomarkers. Unfortunately, to date, such research has not been published, with reliance on using retrospective databases often only providing information at a single time point. A future study in this area would thus require prospective collection of rich data, with the aim to achieve accurate prediction modelling demanded by clinicians and patients prior to implementation.

Impact analysis in prediction research is sparse making it difficult to conclude whether a model is worth implementing alongside, or replacing, usual care. This is important as, for example, one study suggested clinical acumen may be superior to prediction models, while another found the combination of a model with clinical acumen was better than either alone. Some impact analyses have suggested benefit, but conclusions are limited due to their rarity and design (mostly before-after without control). There are a number of potential areas for impact analysis and clinical implementation (summarised in Table 4). First, in specific populations, a model could influence location of perioperative care of surgical patients or drug and/or contrast dosing in patients with heart failure. Second, in a wider hospital setting, the effects of highlighting those at highest risk to teams (ward, outreach critical care or nephrology) with an adequate effector arm could be investigated. This has been demonstrated by existing AKI alerts in established AKI where outcome benefit has been limited to patients who had best practice delivered. Third, as healthcare embraces complex technology, the inclusion of physiological (including urine output) or laboratory trends may be the only way to significantly improve model performance.

| Table 4 | Potential areas for future impact analysis of AKI prediction models |
|---------|---------------------------------------------------------------------|
| Population | Impact analysis to inform clinical use |
| General surgery | Perioperative: haemodynamic targets, place of care, drugs, contrast delivery |
| Trauma and orthopaedics | Risk stratification of large populations: for example, influencing intensity of observations, remote monitoring, application of biomarkers in subgroups at high risk |
| General populations | Optimise haemodynamic status: diuretic dosing, use/volume of contrast |
| Heart failure | |
Fourth, a model could identify a high-risk group to be further risk stratified by employing one of the (increasing number of) available renal biomarkers, or response to an intervention such as a frusemide stress test. Finally, one external validation study found those patients high risk on the prediction model who did develop AKI had a higher rate of mortality than the low-risk group who developed HA-AKI, indicating the model predicts disease severity. This could allow early review of such patients to help inform whether escalation of care may be required, or indeed be appropriate in the increasing number of frail elderly patients admitted to hospitals. To conclude, improving the management of patients to prevent AKI, or reduce associated complications, is a global health priority. This systematic review suggests there are few externally validated prediction models to help identify those at risk of AKI across general hospital populations. Future research should concentrate on validation, utility of additional markers, exploration of electronic implementation to enable clinical uptake and impact analysis.

Twitter First systematic review of AKI prediction models in general hospital populations

Contributors LEH, LGF, RMV, BDD and PJR developed the idea for the study. LEH, AS, LGF, BDD and PJR were involved in the study conception, preliminary literature review and design of the search strategy and the study protocol. LEH and LGF were involved in screening and data extraction of papers. All authors reviewed data extraction output. LEH drafted the manuscript, which was critically reviewed and approved by all authors.

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