### CONSORT-EHEALTH (V 1.6.1) - Submission/Publication Form

The CONSORT-EHEALTH checklist is intended for authors of randomized trials evaluating web-based and Internet-based applications/interventions, including mobile interventions, electronic games (incl multiplayer games), social media, certain telehealth applications, and other interactive and/or networked electronic applications. Some of the items (e.g. all subitems under item 5 - description of the intervention) may also be applicable for other study designs.

The goal of the CONSORT EHEALTH checklist and guideline is to be

- a) a guide for reporting for authors of RCTs,
- b) to form a basis for appraisal of an ehealth trial (in terms of validity)

CONSORT-EHEALTH items/subitems are MANDATORY reporting items for studies published in the Journal of Medical Internet Research and other journals / scientific societies endorsing the checklist.

Items numbered 1., 2., 3., 4a., 4b etc are original CONSORT or CONSORT-NPT (non-pharmacologic treatment) items.

Items with Roman numerals (i., ii, iii, iv etc.) are CONSORT-EHEALTH extensions/clarifications

As the CONSORT-EHEALTH checklist is still considered in a formative stage, we would ask that you also RATE ON A SCALE OF 1-5 how important/useful you feel each item is FOR THE PURPOSE OF THE CHECKLIST and reporting guideline (optional).

Mandatory reporting items are marked with a red \*.

In the textboxes, either copy & paste the relevant sections from your manuscript into this form - please include any quotes from your manuscript in QUOTATION MARKS, or answer directly by providing additional information not in the manuscript, or elaborating on why the item was not relevant for this study.

YOUR ANSWERS WILL BE PUBLISHED AS A SUPPLEMENTARY FILE TO YOUR PUBLICATION IN JMIR AND ARE CONSIDERED PART OF YOUR PUBLICATION (IF ACCEPTED). Please fill in these questions diligently. Information will not be copyedited, so please use proper spelling and grammar, use correct capitalization, and avoid abbreviations.

DO NOT FORGET TO SAVE AS PDF <code>\_AND\_</code> CLICK THE SUBMIT BUTTON SO YOUR ANSWERS ARE IN OUR DATABASE !!!

Citation Suggestion (if you append the pdf as Appendix we suggest to cite this paper in the caption):

Eysenbach G, CONSORT-EHEALTH Group

CONSORT-EHEALTH: Improving and Standardizing Evaluation Reports of Web-based and Mobile Health Interventions

J Med Internet Res 2011;13(4):e126

URL: http://www.jmir.org/2011/4/e126/

doi: 10.2196/jmir.1923 PMID: 22209829

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Your e-mail address \* abc@gmail.com sejoong@snubh.org Title of your manuscript \* Provide the (draft) title of your manuscript. Real-time clinical decision support based on recurrent neural networks for in-hospital acute kidney injury: external validation and model interpretation Name of your App/Software/Intervention \* If there is a short and a long/alternate name, write the short name first and add the long name in clinical decision support based on recurrent ne Evaluated Version (if any) e.g. "V1", "Release 2017-03-01", "Version 2.0.27913" V1 Language(s) \* What language is the intervention/app in? If multiple languages are available, separate by comma (e.g. "English, French") English URL of your Intervention Website or App e.g. a direct link to the mobile app on app in appstore (itunes, Google Play), or URL of the website. If the intervention is a DVD or hardware, you can also link to an Amazon page. 내 답변 URL of an image/screenshot (optional) 내 답변 Accessibility \* Can an enduser access the intervention presently? access is free and open access only for special usergroups, not open access is open to everyone, but requires payment/subscription/in-app purchases app/intervention no longer accessible ○ 기타:

Primary Medical Indication/Disease/Condition * e.g. "Stress", "Diabetes", or define the target group in brackets after the condition, e.g. "Autism (Parents of children with)", "Alzheimers (Informal Caregivers of)"  Acute kidney injury	
Primary Outcomes measured in trial * comma-separated list of primary outcomes reported in the trial  Acute kidney injury occurrence	
Secondary/other outcomes  Are there any other outcomes the intervention is expected to affect?  Severe acute kidney development	
Recommended "Dose" * What do the instructions for users say on how often the app should be used?  Approximately Daily Approximately Weekly Approximately Monthly Approximately Yearly "as needed" 7 Eh:	
Approx. Percentage of Users (starters) still using the app as recommended after 3 months *   unknown / not evaluated  0-10%  11-20%  21-30%  31-40%  41-50%  51-60%  61-70%  71%-80%  81-90%  91-100%  7 E :	

Overall, was the app/intervention effective? *
yes: all primary outcomes were significantly better in intervention group vs control
O partly: SOME primary outcomes were significantly better in intervention group vs control
ono statistically significant difference between control and intervention
O potentially harmful: control was significantly better than intervention in one or more outcomes
inconclusive: more research is needed
○ 기타:
A title Burnerity Challes (Charles)
Article Preparation Status/Stage *  At which stage in your article preparation are you currently (at the time you fill in this form)
onot submitted yet - in early draft status
onot submitted yet - in late draft status, just before submission
submitted to a journal but not reviewed yet
submitted to a journal and after receiving initial reviewer comments
submitted to a journal and accepted, but not published yet
O published
○ 기타:
Journal *
If you already know where you will submit this paper (or if it is already submitted), please provide the journal name (if it is not JMIR, provide the journal name under "other")
onot submitted yet / unclear where I will submit this
Journal of Medical Internet Research (JMIR)
JMIR mHealth and UHealth
JMIR Serious Games
JMIR Mental Health
JMIR Public Health
JMIR Formative Research
Other JMIR sister journal
○ 기타:
Is this a full powered effectiveness trial or a pilot/feasibility trial? *
Pilot/feasibility
Fully powered

Manuscript tracking number If this is a JMIR submission, please p tracking number can be found in the JMIR. If the paper is already publishe the end of the DOI, to be found at the  ono ms number (yet) / not (y	orovide the submissic ed in JMIR bottom o	on acknowl , then the r f each pub	edgement ns trackin lished arti	t email, or g number icle in JMI	when you I is the four- R)	ogin as author in				
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Identify the mode of delivery. Prefera title. Avoid ambiguous terms like "onl includes non-web-based Internet con offline products are used. Use "virtua only in the context of "online support terms for the class of products (such	1a-i) Identify the mode of delivery in the title  Identify the mode of delivery. Preferably use "web-based" and/or "mobile" and/or "electronic game" in the title. Avoid ambiguous terms like "online", "virtual", "interactive". Use "Internet-based" only if Intervention includes non-web-based Internet components (e.g. email), use "computer-based" or "electronic" only if offline products are used. Use "virtual" only in the context of "virtual reality" (3-D worlds). Use "online" only in the context of "online support groups". Complement or substitute product names with broader terms for the class of products (such as "mobile" or "smart phone" instead of "iphone"), especially if the application runs on different platforms.									
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subitem not at all important	0	0	•	0	0	essential 선택해제				
Does your paper address subitem 1a-i? *  Copy and paste relevant sections from manuscript title (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study  Real-time clinical decision support based on recurrent neural networks in clinical practice such as university hospitals.										
1a-ii) Non-web-based components support").		•								
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1a-iii) Primary condition or ta Mention primary condition or target ( Example: A Web-based and Mobile Ir Randomized Controlled Trial	group in th	e title, if a	ny (e.g., "f			
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## Does your paper address subitem 1b-iv? Copy and paste relevant sections from the manuscript abstract (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study Model 1 predicted any AKI development with an area under the curve (AUC) of 0.88 (internal

Model 1 predicted any AKI development with an area under the curve (AUC) of 0.88 (internal validation) and 0.84 (external validation), and stage 2 or higher AKI development with an AUC of 0.93 (internal validation) and 0.90 (external validation). Model 2 predicted the future creatinine values within 3 days with mean square errors of 0.04-0.09 for patients with higher risks of AKI and 0.03-0.08 for those with lower risks.

## 1b-v) CONCLUSIONS/DISCUSSION in abstract for negative trials Conclusions/Discussions in abstract for negative trials: Discuss the primary outcome - if the trial is negative (primary outcome not changed), and the intervention was not used, discuss whether negative results are attributable to lack of uptake and discuss reasons. (Note: Only report in the abstract what the main paper is reporting. If this information is missing from the main body of text, consider adding it) 1 2 3 4 5

subitem not at all important  $\bigcirc$   $\bigcirc$   $\bigcirc$   $\bigcirc$   $\bigcirc$  essential

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#### Does your paper address subitem 1b-v?

Copy and paste relevant sections from the manuscript abstract (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Not applicable for this study (This is not a negative trial)

#### INTRODUCTION

#### 2a) In INTRODUCTION: Scientific background and explanation of rationale

#### 2a-i) Problem and the type of system/solution

Describe the problem and the type of system/solution that is object of the study: intended as stand-alone intervention vs. incorporated in broader health care program? Intended for a particular patient population? Goals of the intervention, e.g., being more cost-effective to other interventions, replace or complement other solutions? (Note: Details about the intervention are provided in "Methods" under 5)

1 2 3 4 5

subitem not at all important O O O essential

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#### Does your paper address subitem 2a-i? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Therefore, many research efforts have been expended to detect AKI early on and to manage high-risk patients. Nevertheless, the reported incidence of AKI is 17%–25% in the hospital setting, and it has continued to rise globally during the recent decades.

#### 2a-ii) Scientific background, rationale: What is known about the (type of) system Scientific background, rationale: What is known about the (type of) system that is the object of the study (be sure to discuss the use of similar systems for other conditions/diagnoses, if appropiate), motivation for the study, i.e. what are the reasons for and what is the context for this specific study, from which $stakeholder\ viewpoint\ is\ the\ study\ performed,\ potential\ impact\ of\ findings\ [2].\ Briefly\ justify\ the\ choice\ of\ potential\ impact\ of\ findings\ [2]$ the comparator. 3 5 0 0 0 • subitem not at all important essential 선택해제 Does your paper address subitem 2a-ii? \* Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study Acute kidney injury (AKI) is a common clinical condition that can be attributed to multiple causes in various clinical settings. AKI increases mortality, morbidity, length of hospital stay, and healthcare costs . According to the National Confidential Enquiry on patient outcomes and death reports, approximately 17% of AKI is estimated to be avoidable and preventable . Machine learning methods can incorporate tremendously large number of features in the model compared with conventional regression models and thus enable the use of nonlinear algorithms. 2b) In INTRODUCTION: Specific objectives or hypotheses Does your paper address CONSORT subitem 2b? \* Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study we propose an externally validated RNN-based prediction model for in-hospital AKI and aimed to provide a framework to link the developed model with clinical decision support. **METHODS** 3a) Description of trial design (such as parallel, factorial) including allocation ratio Does your paper address CONSORT subitem 3a? \* Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study external validation 3b) Important changes to methods after trial commencement (such as eligibility criteria), with reasons Does your paper address CONSORT subitem 3b? \*

Not applicable for this study (This is not a RCT)

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

3b-i) Bug fixes, Downtimes, O Bug fixes, Downtimes, Content Chang changes to methods therefore also in during the trial (e.g., major bug fixes "unexpected events" that may have in failures/downtimes, etc. [2].	jes: eheal icludes im or change	th systems nportant ches in the fu	are often anges ma nctionality	ide on the or conter	interventiont) (5-iii) ar	on or comparator ad other
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All demographics, laboratory value he EHR of each hospital. The feat elated literature or are correlated levelopment. We developed two algorithms. Model 1 had a many- prediction output. The outcome vieven days.	atures th d with Al differen to-one a	at are co KI develo t models rchitectu	nsidered pment w (models re with s	as risk f ere selec 1 and 2) even seq	actors of ted for m with stac uential in	AKI from the odel ked RNN put and one
evisions and updating evisions and updating. Clearly menti and comparator, if applicable) evalua uring the evaluation process, or whe escribe dynamic components such a ne replicability of the intervention (fo	ted, or de ther the d as news f	escribe wh levelopme eeds or ch	ether the nt and/or anging co	interventic content w intent which	n underwe as "frozen"	nt major changes during the trial.
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and/or providing flowcharts of the all principle be able to replicate the stud	gorithms (	used. Repl	icability (i	e., other r		n-capture video, s should in
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We described the detailed mode distributed to other researchers				and techi	niques co	uld be
5-vi) Digital preservation Digital preservation: Provide the URL disappear over the course of the year webcitation.org, and/or publishing th pages behind login screens cannot b without login.	rs; also ma e source o	ake sure th code or sc	ne interver reenshots	ntion is arc /videos alc	hived (Inte	ernet Archive, e article). As
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5-vii) Access						
Access: Describe how participants ar or were paid) or not, whether they ha participants obtained "access to the editors/reviewers/readers, consider t eviewers/readers to explore the app	ad to be a platform a to provide	member o and Interne a "backdo	f specific et" [1]. To o or" login a	group. If k ensure acc account or	nown, des ess for demo mod	cribe how de for
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5-viii) Mode of delivery, features/functionalities/components of the intervention and comparator, and the theoretical framework

Describe mode of delivery, features/functionalities/components of the intervention and comparator, and the theoretical framework [6] used to design them (instructional strategy [1], behaviour change techniques, persuasive features, etc., see e.g., [7, 8] for terminology). This includes an in-depth description of the content (including where it is coming from and who developed it) [1]," whether [and how] it is tailored to individual circumstances and allows users to track their progress and receive feedback" [6]. This also includes a description of communication delivery channels and – if computer-mediated communication is a component – whether communication was synchronous or asynchronous [6]. It also includes information on presentation strategies [1], including page design principles, average amount of text on pages, presence of hyperlinks to other resources, etc. [1].

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#### Does your paper address subitem 5-viii? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

We developed two different models (models 1 and 2) with stacked RNN algorithms. Model 1 had a many-to-one architecture with seven sequential input and one prediction output. The outcome variable for model 1 was the occurrence of AKI in the next seven days. In model 2, we constructed a prediction model of the trajectory of Cr values after 24 h, 48 h, and 72 h with available Cr values during the observation window. Model 2 had a many-to-many structure with an output length of three on seven input sequences. To improve the predictive accuracy of model 2, we developed a two-stage hierarchical RNN prediction model. That is, based on the results of model 1, different types of model 2 were applied.

#### 

#### Does your paper address subitem 5-ix?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

we applied a class weight parameter to the loss function to handle class imbalances. Alpha drop-out, L2 regularization, and early stopping approaches were implemented to prevent overfitting. Batch normalization was applied to each RNN layer for efficient and effective learning. The learning rate was set to 10-4 for pretraining, then to 10-6 for early-stop learning.

5-x) Clarify the level of huma Clarify the level of human involvemen in the e-intervention or as co-interven as well as "type of assistance offered medium by which the assistance is di human involvement required for the t	t (care pro tion (deta I, the timir elivered". I	oviders or il number ng and fred It may be i ne level of	and exper quency of necessary human in	tise of pro the suppo to disting olvement	fessionals t, how it is uish betwe required f	involved, if any, s initiated, and the een the level of
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During data collection, error valu were examined and removed afte variables at baseline were exclud	er review	by doma	in exper	ts. Subje	cts with r	
5-xi) Report any prompts/rer Report any prompts/reminders used: use the application, what triggered th level of prompts/reminders required tapplication outside of a RCT setting (	Clarify if t em, freque or the tria	there were ency etc. I al, and the	t may be r level of pr	ecessary ompts/rer	to distinguninders fo	ish between the
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5-xii) Describe any co-intervont Describe any co-interventions (incl. traddition to the targeted eHealth interintervention. This includes training set the level of training required for the tract RCT setting (discuss under item 21 –	aining/su vention, as ssions an ial, and th	pport): Cle s ehealth i d support le level of	early state nterventio [1]. It may	any interv n may not be neces	be design sary to dis	ed as stand-alone tinguish between
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#### Does your paper address subitem 5-xii? \* Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study To identify the association between multiple features and response (the occurrence of AKI), we examined the following approaches using model-agnostic methods in model 1. (1) Global interpretation with SHapley Additive exPlanations (SHAP), partial dependence plots (PDP), and accumulated local effects (ALE) plots, and (2) instance-wise interpretation with individual conditional expectation (ICE) plots and SHAP 6a) Completely defined pre-specified primary and secondary outcome measures, including how and when they were assessed Does your paper address CONSORT subitem 6a? \* Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study AKI was defined according to the Kidney Disease: Improving Global Outcomes (KDIGO) Clinical Practice Guideline for AKI [30]. Because urine volume data were not available, AKI stage was defined based on serum Cr levels. Baseline Cr levels were determined by searching the minimum serum Cr level within a period of 2 weeks before admission. 6a-i) Online questionnaires: describe if they were validated for online use and apply CHERRIES items to describe how the questionnaires were designed/deployed If outcomes were obtained through online questionnaires, describe if they were validated for online use and apply CHERRIES items to describe how the questionnaires were designed/deployed [9]. 2 3 0 0 0 0 subitem not at all important essential 선택해제

apply CHERRIES items to describe how the questionnaires were designed/deployed

If outcomes were obtained through online questionnaires, describe if they were validated for online use and apply CHERRIES items to describe how the questionnaires were designed/deployed [9].

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Does your paper address subitem 6a-i?
Copy and paste relevant sections from manuscript text

Not applicable for this study.

6a-ii) Describe whether and how "use" (including intensity of use/dosage) was defined/measured/monitored
Describe whether and how "use" (including intensity of use/dosage) was defined/measured/monitored (logins, logfile analysis, etc.). Use/adoption metrics are important process outcomes that should be reported in any ehealth trial.

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<b>7a) How sample size was de</b> NPT: When applicable, details of whe addressed			ıstering by	/ care prov	rides or ce	nters was
7a-i) Describe whether and had calculating the sample size						
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#### Does your paper address CONSORT subitem 7b? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Not applicable for this study (This is not a RCT)

#### 8a) Method used to generate the random allocation sequence

NPT: When applicable, how care providers were allocated to each trial group

#### Does your paper address CONSORT subitem 8a? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

The SNUBH dataset was divided into a training set (90%) and an internal validation set (10%) using the stratified random split.

#### 8b) Type of randomisation; details of any restriction (such as blocking and block size)

#### Does your paper address CONSORT subitem 8b? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Not applicable for this study.

9) Mechanism used to implement the random allocation sequence (such as sequentially numbered containers), describing any steps taken to conceal the sequence until interventions were assigned

#### Does your paper address CONSORT subitem 9? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Not applicable for this study.

10) Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions

#### Does your paper address CONSORT subitem 10? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Not applicable for this study (This is not a RCT)

11a) If done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how NPT: Whether or not administering co-interventions were blinded to group assignment

11a-i) Specify who was blinde	ed, and	who wa	sn't			
Specify who was blinded, and who wa participants [1, 3] (this should be clea assessors, those doing data analysis	arly ackno	wledged),	but it may	be possib	le to blind	
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Does your paper address CONSORT subitem 12a? \*

#### Does your paper address subitem X26-i? Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study This study was performed in accordance with the recommendations laid out in the World Medical Association Declaration of Helsinki. The study protocol was approved by the institutional review boards (IRB) of the Seoul National University Bundang Hospital (SNUBH) (IRB No. B-1912/583-406) and Seoul National University Hospital (SNUH) (IRB No. H-1911-043-1076). x26-ii) Outline informed consent procedures Outline informed consent procedures e.g., if consent was obtained offline or online (how? Checkbox, etc.?), and what information was provided (see 4a-ii). See [6] for some items to be included in informed consent documents. 3 5 $\circ$ 0 0 subitem not at all important essential 선택해제 Does your paper address subitem X26-ii? Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Written consent was waived by the IRB because of the retrospective nature of the study, and all data were completely anonymized.

# X26-iii) Safety and security procedures Safety and security procedures, incl. privacy considerations, and any steps taken to reduce the likelihood or detection of harm (e.g., education and training, availability of a hotline) 1 2 3 4 5 subitem not at all important O ● ● ● essential

#### Does your paper address subitem X26-iii?

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all data were completely anonymized.

#### **RESULTS**

13a) For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcome

NPT: The number of care providers or centers performing the intervention in each group and the number of patients treated by each care provider in each center

#### Does your paper address CONSORT subitem 13a? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

A total of 482,467 patients (182,976 in SNUBH and 299,491 in SNUH) were initially screened from EHR data obtained from each participating hospital.

13b) For each group, losses and exclusions after randomisation, together with reasons

#### Does your paper address CONSORT subitem 13b? (NOTE: Preferably, this is shown in a CONSORT flow diagram) \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

After considering the exclusion criteria, 69,081 patients were finally included in the SNUBH training dataset, 7,675 in the SNUBH internal validation dataset, and 72,352 in the SNUH external validation dataset.

#### 13b-i) Attrition diagram

Strongly recommended: An attrition diagram (e.g., proportion of participants still logging in or using the intervention/comparator in each group plotted over time, similar to a survival curve) or other figures or tables demonstrating usage/dose/engagement.

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#### Does your paper address subitem 13b-i?

Copy and paste relevant sections from the manuscript or cite the figure number if applicable (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Multimedia Appendix 2. Flow diagram for study participants.

#### 14a) Dates defining the periods of recruitment and follow-up

#### Does your paper address CONSORT subitem 14a? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Study populations were all patients aged  $\geq$  18 years who were hospitalized for more than 48 h between 2013 and 2017 in two tertiary hospitals in Korea (Seoul National University Bundang Hospital and Seoul National University Hospital).

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17a-i) Presentation of process	s outco	mes sud	ch as me	etrics of	use and	intensity of
In addition to primary/secondary (clin metrics of use and intensity of use (do not only refer to metrics of attrition (1 metrics such as "average session leng metric like a "session" is defined (e.g.,	ose, expos 3-b) (ofte gth". Thes	sure) and n a binary e must be	their opera variable), accompa	ational def but also to nied by a t	initions is o more cor echnical d	critical. This does tinuous exposure escription how a
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Does your paper address sub Copy and paste relevant sections from indicate direct quotes from your manu- information not in the ms, or briefly ex Model performances were assess predictive value (PPV), negative properating characteristic (ROC) cur	n the man uscript), or xplain why sed base predictive	uscript (ir r elaborate r the item ed on acc e value (N	e on this it is not app curacy, se NPV), and	em by pro licable/rel nsitivity, I the area	viding add evant for y specificit under th	itional our study  y, positive e receiver
error (MSE) for model 2.						
17b) For binary outcomes, pr sizes is recommended	resenta	tion of	both ab	solute a	ind relat	ive effect
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18) Results of any other analadjusted analyses, distinguis				-		analyses and
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18-i) Subgroup analysis of cor A subgroup analysis of comparing onl stressed that this is a self-selected sa (see 16-iii).	y users is imple and	not unco	mmon in e r an unbias	sed sample	e from a ra	
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19) All important harms or un (for specific guidance see CONSORT			cts in ea	ach gro	up	
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19-i) Include privacy breache Include privacy breaches, technical pr but also incidents such as perceived o unexpected/unintended incidents. "Ur	oblems. Tor real priv	This does i vacy bread	not only in hes [1], te	chnical pr	oblems, ar	nd other
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19-ii) Include qualitative feed staff/researchers Include qualitative feedback from par		·				
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22) Interpretation consistent with results, balancing benefits and harms, and considering other relevant evidence

NPT: In addition, take into account the choice of the comparator, lack of or partial blinding, and unequal expertise of care providers or centers in each group

22-i) Restate study questions and summarize the answers suggested by the data, starting with primary outcomes and process outcomes (use)

Restate study questions and summarize the answers suggested by the data, starting with primary outcomes and process outcomes (use).

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subitem not at all important O O O essential

선택해제

#### Does your paper address subitem 22-i? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

In the present study, we developed a continuous prediction model for in-hospital AKI using the RNN algorithm with external validation and demonstrated its applicability for the support of clinical decision making. The developed model was constructed for all general inpatients based on the use of various dynamic and static clinical features, showing relatively good performance. External validation was performed with data from an independent center. Furthermore, we showed examples relevant to the presentation of feature information to help actual clinical decisions at the global and individual patient level using several model-agnostic interpretation methods.

#### 22-ii) Highlight unanswered new questions, suggest future research

Highlight unanswered new questions, suggest future research.

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#### Does your paper address subitem 22-ii?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

The interpretation methods themselves do not indicate causal inferences between features and the model output. Currently, various feature engineering techniques are actively being studied and constitute promising fields of artificial intelligence. Therefore, human-friendly interpretable models that reflect possible causal inferences will be developed in the near future.

20) Trial limitations, addressing sources of potential bias, imprecision, and, if relevant, multiplicity of analyses

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OTHER INFORMATION						
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25) Sources of funding and funders	other s	upport (	such as	supply	of drug	s), role of
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X27) Conflicts of Interest (n	ot a CC	ONSORT	item)			
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Does your paper address subitem X27-i? Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study No conflicts of interest. About the CONSORT EHEALTH checklist As a result of using this checklist, did you make changes in your manuscript?\* yes, major changes yes, minor changes O no What were the most important changes you made as a result of using this checklist? It contributes to the systemic structure of the article How much time did you spend on going through the checklist INCLUDING making changes in your manuscript \* 1 hour As a result of using this checklist, do you think your manuscript has improved? \* yes O no ○ 기타: Would you like to become involved in the CONSORT EHEALTH group? This would involve for example becoming involved in participating in a workshop and writing an "Explanation and Elaboration" document yes O no ○ 기타: 선택해제 Any other comments or questions on CONSORT EHEALTH This is a very structured item.

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