Artificial Intelligence-Enabled Electrocardiogram Improves the Diagnosis and Prediction of Mortality in Patients With Pulmonary Hypertension

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ABSTRACT

BACKGROUND Pulmonary hypertension is a disabling and life-threatening cardiovascular disease. Early detection of elevated pulmonary artery pressure (ePAP) is needed for prompt diagnosis and treatment to avoid detrimental consequences of pulmonary hypertension.

OBJECTIVES This study sought to develop an artificial intelligence (AI)-enabled electrocardiogram (ECG) model to identify patients with ePAP and related prognostic implications.

METHODS From a hospital-based ECG database, the authors extracted the first pairs of ECG and transthoracic echocardiography taken within 2 weeks of each other from 41,097 patients to develop an AI model for detecting ePAP (PAP > 50 mm Hg by transthoracic echocardiography). The model was evaluated on independent data sets, including an external cohort of patients from Japan.

RESULTS Tests of 10-fold cross-validation neural-network deep learning showed that the area under the receiver-operating characteristic curve of the AI model was 0.88 (sensitivity 81.0%; specificity 79.6%) for detecting ePAP. The diagnostic performance was consistent across age, sex, and various comorbidities (diagnostic odds ratio > 8 for most factors examined). At 6-year follow-up, the patients predicted by the AI model to have ePAP were independently associated with higher cardiovascular mortality (HR: 3.69). Similar diagnostic performance and prediction for cardiovascular mortality could be replicated in the external cohort.

CONCLUSIONS The ECG-based AI model identified patients with ePAP and predicted their future risk for cardiovascular mortality. This model could serve as a useful clinical test to identify patients with pulmonary hypertension so that treatment can be initiated early to improve their survival prognosis. (JACC: Asia 2022;2:258–270) © 2022 The Authors. Published by Elsevier on behalf of the American College of Cardiology Foundation. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).
Pulmonary hypertension (PH), a condition of elevated pulmonary artery pressure (ePAP), affects more than a million individuals worldwide and causes premature disability, heart failure, and death. Left heart and lung diseases are the most common etiologies of PH. The presence of PH can increase the 5-year mortality rate to more than 30% if left untreated. The transthoracic echocardiography (TTE) is recommended for measuring pulmonary artery pressure (PAP) via the estimation of peak tricuspid regurgitation velocity. However, TTE is highly operator-dependent, and requires a good acoustic window and flow tracings for correct PAP measurements. As such, delays of up to 2 years between the onset of symptoms and diagnosis of ePAP are frequently observed. Alternative tests, such as the pulmonary function, serum N-terminal pro-B-type natriuretic peptide, and uric acid level tests, have been developed to increase the sensitivity for the early diagnosis of ePAP. Although their sensitivity remains unsatisfactory, ranging from 55.9% to 71.0%, these tests can identify patients with disease progression and future mortality, which is a key requirement for a qualified clinical test. Diagnosing PH at a stage when it may be more amenable to treatment is important. However, the current screening approaches for PH are resource-intensive, and population-based screening remains unavailable.

Recently, researchers have successfully trained artificial intelligence (AI) models to correlate electrocardiogram (ECG) signals to echocardiographic phenotypes and automatically screen for cardiac structural derangement (e.g., cardiac contractile dysfunction, hypertrophy). Using the same approach, a semiautomatic AI algorithm was recently created to detect ePAP, which requires a variety of clinical (e.g., age, sex, height, weight, body mass index) and ECG (e.g., P axis, PR interval, QRS duration, QT interval, presence of atrial fibrillation) characteristics, as well as predetermined arrhythmias from medical records or physician diagnoses. As numerous clinical and ECG predetermined parameters are still needed, the workload of manpower is expected not to be reduced.

The assessment of associations between a medical test or biomarker and relevant clinical outcomes is a prerequisite for a qualified diagnostic test. Although automatic detection of cardiac abnormality by AI is an encouraging new technology, AI algorithms have not been correlated to cardiovascular outcomes until very recently. In the present study, we intended to develop a completely automated AI-enabled ECG model that can detect ePAP and identify patients at risk of cardiovascular disease mortality. This qualified test could enable early diagnosis, risk assessment, and therapeutic intervention in patients with ePAP or suspicious PH.

METHODS

In our study, ePAP was defined as PAP > 50 mm Hg by TTE, which indicates a high probability of PH according to established guidelines. A reproducibility analysis of the TTE measurements in the database we used has been reported, which showed that interobserver variability (4.4% to 6.4%) was just a bit higher than intra-observer variability (4.0% to 4.8%). From the TTE data, PAP was estimated as the equation of estimated right atrium pressure. Deep learning AI models were trained to identify patients with ePAP. The design of the AI model is shown in Supplemental Figure S1 and the Supplemental Methods. The overall process is illustrated in the Central Illustration.

STUDY POPULATION. A total of 63,767 patients with both ECGs (Philips Healthcare) and TTEs between 2010 and 2017 were obtained from the ECG database at Taipei Veterans General Hospital, Taiwan. Among those, 229,787 ECG-TTE pairs were performed within a 2-week interval. We selected the 2-week interval following a previous study design. The first such ECG-TTE pair for each patient (n = 41,097) was selected to form the main analysis data to create the AI model (Group 1). To examine whether the AI model could predict the long-term incidence of developing cardiac abnormality in patients whose hearts were initially diagnosed as normal, we retrieved a TTE follow-up cohort of 10,818 patients from Group 1. These patients received a follow-up TTE after the first ECG-TTE pair at a mean interval of 2.6 ± 1.7 years (IQR from first ECG-TTE pair: 1.2 to 3.7 years). This subset of Group 1 patients was designated as Group 1'. Another 18,373 ECG-TTE pairs had intervals of more than 2 weeks between the ECG and TTE. These patients (designated as Group 2) were not in the main analysis group (Group 1) and thus also not in the follow-up analysis group (Group 1'). Group 2 was used as an independent group for ancillary validation of the AI model for cardiovascular mortality. An external cohort of Japanese patients (Group 3), described in the following text, was used to further test the model. Figure 1 shows the study’s flow diagram.

MAIN ANALYSIS TO CREATE THE AI MODEL (GROUP 1). Ten AI models were derived from 10-fold cross-validation neural network deep learning. In this AI learning, all ECG-TTE pairs from Group 1 were...
randomly assigned to 10 independent segments (8 to the training set, 1 to the validation set, and 1 to the test set). Because any given patient's data were exclusively present in only 1 of the 10 segments, the data could not be used simultaneously for model training, validation, or testing. This 8-1-1 training, validation, and test data segmentation scheme yielded 32,875 to 32,879 ECG-TTE pairs in the training set, 4,109 to 4,111 pairs in the validation set, and 4,109 to 4,111 pairs in the test set for each of the 10 cross-validation models. We used a rolling data-segment selection scheme (Figure 1) to ensure all pairs in Group 1 were used once for validation and tested once in 1 of the 10 models, thereby avoiding data selection bias. Note the cross-validation procedure used here is not a conventional one because the validation set is for the selection of the best model instead of the best hyperparameters (Supplemental Methods).

**FOLLOW-UP ANALYSIS FOR PATIENTS DEVELOPING ePAP (GROUP 1).** Group 1 patients included 2 subgroups: AI-predicted non-ePAP patients and AI-predicted ePAP patients. AI-predicted non-ePAP patients were those predicted in the test sets as not having an ePAP who also had a confirmatory PAP <50 mm Hg by TTE (n = 6,532). AI-predicted ePAP patients were those identified by AI as having ePAP but had a contradictory PAP <50 mm Hg by TTE examined within 2 weeks of the ECG (n = 2,984). We followed up the incidences of developing ePAP in patients of both subgroups.

**ANCILLARY ANALYSIS OF PATIENTS WHOSE ECG AND TTE OCCURRED MORE THAN 2 WEEKS APART (GROUP 2).** Group 2 patients had their TTEs more than 2 weeks after their initial ECGs. The mean interval from ECG to TTE was 2.6 ± 2.1 years (IQR: 0.7-4.1 years). For the ancillary test on Group 2
Patients, we intentionally used the worst-performing AI model from the 10-fold cross-validation of the main (Group 1) analysis. Because of the lack of the ECG-TTE pairs within an interval of 2 weeks, we applied the AI model to merely evaluate its prognostic value for Group 2 patients.

**FURTHER VALIDATION OF THE AI MODEL ON A JAPANESE COHORT (GROUP 3).** A total of deidentified 279 ECGs (114 PH and 165 non-PH patients confirmed by TTE within a 2-week interval of ECG-TTE pairs) with the recordings of age, sex, and history of diseases, as well as follow-up of cardiovascular mortality and all-cause mortality, were randomly retrieved from a registry compiled between 2009 and 2020 at Makiminato Central Hospital and Nakagami Hospital, Japan. The 10-second ECGs were acquired from Cardiofax V ECG-2400 (Nihon Kohden Corporation) and analyzed by the same AI model tested on Group 2 without retraining.

The methods of survival and sensitivity analyses and using conventional ECG characteristics to diagnose ePAP were provided in the Supplemental Appendix.
AI MODEL FOR ePAP. Our model was adapted with permission from the deep learning neural network model of our previous work designed for the detection and classification of cardiac arrhythmias (Supplemental Figure S1, Supplemental Methods).

ETHICAL APPROVAL. This study was approved by the institutional review board at Taipei Veterans General Hospital, Taipei, Taiwan.

STATISTICAL ANALYSIS. All analyses were performed using SAS (SAS Institute) statistical proprietary software 9.4 (TS1M3 DBCS3170). The statistical methods used are described in the Supplemental Appendix.

RESULTS

MAIN ANALYSIS OF DIAGNOSTIC PERFORMANCE OF THE AI MODEL FOR ePAP. The patients in the main analysis (Group 1) were 60.0 ± 18.6 years of age, and 6.9% of them were diagnosed as having ePAP by TTE. Table 1 shows the baseline clinical characteristics. The patients with ePAP were older, more often male, and had more comorbidities.

In the test sets of the main analysis (Group 1), the area under the curves (AUCs) of the 10 cross-validation AI models for detecting ePAP were consistent (mean 0.88; 95% CI: 0.87–0.89) (Figure 2). Consistency was also observed for sensitivity (81.0%, 95% CI: 77.6%–84.4%), specificity (79.6%, 95% CI: 77.4%–81.7%), and accuracy (79.7%, 95% CI: 77.8%–81.5%) (Supplemental Table S1). The AUCs of the 10 cross-validation AI models for detecting ePAP were consistent across the different individual test sets, and that these test sets were randomly segmentalized, the possible bias caused by different models tested on different patients could be minimized. These results demonstrated the robustness of our AI models, which required no pre-determined clinical or ECG characteristics as input.

The AI performance was strong across age, sex, and various comorbidities (Figure 3). For comorbidities, the diagnostic odds ratios all exceeded 7.84, indicating that the AI’s diagnostic performance was consistently predictive for all these different diseases. Interestingly, the presence of comorbidity was associated with a decreased diagnostic odds ratio for some diseases with statistical significance (interaction P < 0.05), suggesting a potential interference of ECG changes from these diseases in the AI diagnosis.

The observed diagnostic odds ratio decreased with age but remained powerful, and even in patients older than 80 years, it was as high as 5.86. No sex difference was observed in the performance of the AI model.

The power of the AI model was explained by V1, V2, and V3 being identified as the most important individual leads for AI to predict ePAP (Supplemental Figure S3A). This is consistent with features of right ventricular hypertrophy that physicians observed in these 3 leads to diagnose ePAP (Supplemental Figures S3B to S3E).

For comparison, we analyzed the diagnostic performance of using conventional ECG characteristics to detect ePAP. The sensitivity of using conventional ECG characteristics to identify ePAP was too low for effective clinical use, in agreement with previous reports.

The diagnostic performance of the AI model was also better than a logistic regression model utilizing traditional ECG variables, as shown in the Supplemental Results and Supplemental Figure S4.

FOLLOW-UP ANALYSIS ON FUTURE INCIDENTS OF ePAP. Of the Group 1 patients identified by the AI model as having a normal PAP who also had a confirmatory normal PAP by TTE within a 2-week window, 6,532 had a follow-up TTE. Of these AI-predicted non-ePAP patients, 372 (5.7%) went on to
develop ePAP. By contrast, for the 2,984 patients labeled by the AI model as having ePAP but with a normal PAP by TTE (AI-predicted ePAP), 697 (23.4%) went on to develop ePAP (Supplemental Figure S5A). Compared with the AI-predicted non-ePAP patients, this represents a 5.04-fold risk of developing ePAP for the AI-predicted ePAP patients when the AI model defined their ECG as abnormal (multivariate-adjusted HR: 3.74 [95% CI: 3.28–4.26]). The AI model thus identified ECG abnormalities before overt ePAP manifested.

ANCILLARY ANALYSIS OF FUTURE INCIDENTS OF ePAP FOR AN ADDITIONAL COHORT OF PATIENTS. We conducted further analysis on future incidents of ePAP for an independent group of patients (Group 2) identified by the AI model as having ePAP or normal PAP. The baseline characteristics of Group 2 patients were shown in Supplemental Table S2. Of the 4,269 AI-predicted ePAP patients, the incidence of ePAP as examined by TTE at more than 2 weeks after the ECG was 27.3%. In comparison, of the 14,104 patients identified by the AI model as having normal PAP, the incidence of an ensuing TTE-determined ePAP was 4.9% (Supplemental Figure S5B). This represents a 6.61-fold risk of developing ePAP for patients with AI-defined abnormal ECG (multivariate-adjusted HR: 4.32 [95% CI: 3.91–4.78]) (Supplemental Figure S5B). This result for an independent group of patients further demonstrates the robustness and ability of the AI algorithm to identify ePAP even in the absence of a TTE diagnosis.

CARDIOVASCULAR OUTCOMES PREDICTED BY THE AI MODEL. Cardiovascular mortality and all-cause mortality were analyzed for Group 1 patients. During the 6-year follow-up, Kaplan-Meier survival analysis showed that in comparison to patients stratified as non-ePAP by the AI model, those stratified as ePAP were associated with higher cardiovascular mortality (AI-predicted ePAP vs AI-predicted non-ePAP 1,027 [16.2%] vs 389 [2.0%] patients; \( P < 0.001 \)) and higher all-cause mortality (3,581 [45.5%] vs 2,422 [10.9%] patients; \( P < 0.001 \)), as shown in Figure 4.

Using multivariate Cox regression analysis to adjust for potential confounding factors (age, sex, and various comorbidities), the AI stratification remained independent and the strongest predictor of cardiovascular mortality (HR: 3.69; 95% CI: 3.27–4.17; \( P < 0.001 \)) and all-cause mortality (HR: 2.63; 95% CI: 1.79; 95% CI: 1.67–1.91; \( P < 0.001 \)), as shown in Table 3. Further analysis of patients stratified into groups according to presence/absence of a variety of comorbidities showed that the AI model’s predictive power for cardiovascular death was consistent across diabetes mellitus, hypertension, heart failure, myocardial infarction, stroke, lung diseases, kidney diseases, thromboembolism, and rheumatic and other diseases (Figure 5). Similar results were obtained for all-cause mortality (Supplemental Figure S6). The mortality analysis was also conducted for Group 2 patients and similar results were obtained. AI-predicted ePAP was associated with higher cardiovascular mortality (Supplemental Figure S7) (AI-predicted ePAP vs AI-predicted non-ePAP, 444 [14.9%] vs 277 [7.7%] patients; \( P < 0.001 \)) and higher all-cause mortality (1,645 [44.5%] vs 1,900 [16.3%] patients; \( P < 0.001 \)). Multivariate analysis also showed that AI-predicted ePAP was an independent predictor of cardiovascular mortality (HR: 2.63; 95% CI: 2.27–3.04; \( P < 0.001 \)) and all-cause mortality (HR: 1.79; 95% CI: 1.67–1.91; \( P < 0.001 \)), as shown in Supplemental Table S3. Further analysis showed that the AI model’s predictive power for cardiovascular death and all-cause death in Group 2 patients also was consistent across a variety of diseases (Supplemental Figures S8 and S9). The results from the 2 independent data sets (Groups 1 and 2) thus suggest the AI model reliably predicted cardiovascular outcomes.
EXTERNAL VALIDATION OF THE AI MODEL. The baseline characteristics of the Japanese cohort were provided in Supplemental Table S4. A comparison of baseline characteristics among Group 1, 2, and 3 patients were presented in Supplemental Table S5. The comparison showed that the Japanese cohort (Group 3) exhibited a higher age, more females, and more comorbidities in chronic kidney disease, chronic lung diseases, and pulmonary embolism. In comparison, a higher incidence of hypertension, diabetes mellitus, and prior myocardial infarction was seen in Group 2, and the incidence of congestive heart failure and rheumatic diseases was higher in Group 1.

The AUC of the AI model for ePAP in the external cohort was 0.88 (sensitivity 93.9%, specificity

| Age groups   | SENs | SPEc | DOR  | L95CI | U95CI | Interaction P |
|--------------|------|------|------|-------|-------|---------------|
| <40          | 0.73 | 0.94 | 45.27| 28.27 | 72.50 |                |
| 40-60        | 0.79 | 0.89 | 29.64| 23.25 | 37.80 |                |
| 60-80        | 0.79 | 0.77 | 12.20| 10.51 | 14.16 |                |
| >80          | 0.84 | 0.52 | 5.86 | 4.98  | 6.90  |                |
| Gender       |      |      |      |       |       | 0.73          |
| Female       | 0.76 | 0.85 | 17.92| 15.59 | 20.59 |                |
| Male         | 0.85 | 0.74 | 15.37| 13.39 | 17.65 |                |
| Hypertension |      |      |      |       |       | <0.01         |
| Yes          | 0.81 | 0.73 | 11.01| 9.69  | 12.51 |                |
| No           | 0.81 | 0.85 | 23.81| 20.50 | 27.67 |                |
| Diabetes mellitus | |      |      |       |       | <0.01         |
| Yes          | 0.82 | 0.68 | 9.89 | 8.21  | 11.92 |                |
| No           | 0.80 | 0.82 | 18.46| 16.47 | 20.69 |                |
| Congestive heart failure | |      |      |       |       | 0.21          |
| Yes          | 0.91 | 0.51 | 9.72 | 5.51  | 17.15 |                |
| No           | 0.81 | 0.80 | 16.66| 15.09 | 18.39 |                |
| Prior stroke |      |      |      |       |       | <0.01         |
| Yes          | 0.89 | 0.53 | 8.85 | 7.36  | 10.65 |                |
| No           | 0.75 | 0.83 | 14.35| 12.76 | 16.14 |                |
| Prior myocardial infarction | |      |      |       |       | 0.12          |
| Yes          | 0.83 | 0.63 | 8.19 | 6.04  | 11.11 |                |
| No           | 0.81 | 0.81 | 17.55| 15.84 | 19.45 |                |
| Chronic kidney disease | |      |      |       |       | 0.15          |
| Yes          | 0.86 | 0.56 | 7.84 | 5.97  | 10.28 |                |
| No           | 0.80 | 0.81 | 16.80| 15.14 | 18.66 |                |
| Chronic lung diseases | |      |      |       |       | <0.01         |
| Yes          | 0.83 | 0.68 | 10.49| 8.64  | 12.75 |                |
| No           | 0.80 | 0.82 | 17.87| 15.97 | 20.00 |                |
| Pulmonary embolism | |      |      |       |       | 0.57          |
| Yes          | 0.77 | 0.71 | 8.03 | 3.66  | 17.65 |                |
| No           | 0.81 | 0.80 | 16.69| 15.13 | 18.41 |                |
| Rheumatic diseases | |      |      |       |       | 0.94          |
| Yes          | 0.80 | 0.75 | 12.15| 8.87  | 16.64 |                |
| No           | 0.81 | 0.80 | 17.00| 15.35 | 18.83 |                |
| Atrial fibrillation/flutter | |      |      |       |       | 0.19          |
| Yes          | 0.94 | 0.40 | 11.28| 7.91  | 16.07 |                |
| No           | 0.77 | 0.83 | 16.13| 14.55 | 17.88 |                |
| Idiopathic PAH | |      |      |       |       | 0.30          |
| Yes          | 0.83 | 0.62 | 7.96 | 3.41  | 18.61 |                |
| No           | 0.81 | 0.80 | 16.58| 15.01 | 18.27 |                |
| Overall      | 0.81 | 0.80 | 16.58| 15.01 | 18.27 |                |

The 2 dashed vertical lines indicate the diagnostic odds ratio reference (odds ratio: 1.00) and the overall diagnostic odds ratio (here, odds ratio: 16.58), respectively. DOR = diagnostic odds ratio; L95CI = lower limit of 95% CI; PAH = pulmonary artery hypertension; SENs = sensitivity; SPEc = specificity; U95CI = upper limit of 95% CI.
58.7%, accuracy 72.4%) (Figure 6A). Similarly, the patients stratified as ePAP by AI model were consistently associated with higher cardiovascular mortality (Figure 6B) (AI-predicted ePAP vs AI-predicted non-ePAP 32 [29.8%] vs 1 [1.4%] patients; \( P < 0.001 \)) and higher all-cause mortality (Supplemental Figure S10) (52 [46.7%] vs 3 [5.1%] patients; \( P < 0.001 \)). The AI-predicted ePAP was an independent predictor for cardiovascular mortality (HR: 21.08; 95% CI: 2.51–176.88; \( P < 0.01 \)) and all-cause mortality (HR: 9.15; 95% CI: 2.52–33.27; \( P < 0.01 \)) after multivariate analysis (Supplemental Table S6). These results suggest the AI model could accommodate ECG data for patients from a different hospital in a different country.

### DISCUSSION

In this study, we developed an automated AI model capable of identifying ePAP patients and predicting their risk for cardiovascular and all-cause mortality. The performance of this AI model was shown to be robust for both early diagnosis of ePAP and prognosis of mortality risk in independent patient groups, including an external cohort of Japanese patients. These results support the potential of this AI model as a qualified and valid clinical test to screen for patients at risk of developing ePAP so that treatment can be initiated early to improve their odds of survival.

The ability to detect ePAP is crucial for clinical suspicion of PH, early diagnosis, and prompt treatment because most patients with PH have minimal or no symptoms at an early stage.7,17,23,24 Early detection of PH and prompt therapy would translate into better 3-year survival rates for more than 50% of patients.25 The limitations of TTE have led to efforts devoted to developing alternative clinical tests. However, the traditional ECG criteria used to identity ePAP are not considered a reliable screening tool caused by their low sensitivity of 34% to 55%, as observed in this study and others.12,22 The sensitivities of the pulmonary function, serum N-terminal pro-B-type natriuretic peptide, and uric acid level tests are slightly higher but still only 71%, 56% to 69%, and 68%, respectively.9-12,26,27 Therefore, screening algorithms (eg, DETECT [Early, Simple and Reliable Detection of Pulmonary Arterial Hypertension in Systemic Sclerosis] or ASCS [Australian Scleroderma Cohort Study])

### TABLE 2

| Diagnostic Performance of Conventional ECG Characteristics for Detecting ePAP Patients in Group 1 | Sensitivity, % | Specificity, % |
|---|---|---|
| Conventional ECG characteristics for elevated PAP\(^a\) | 34.1 | 78.7 |
| P pulmonale\(^b\) | 2.6 | 98.1 |
| Right axis deviation\(^c\) | 11.3 | 93.0 |
| Right ventricular hypertrophy\(^d\) | 5.8 | 96.6 |
| Right ventricular strain\(^e\) | 2.4 | 97.5 |
| Right bundle branch block mimics\(^f\) | 19.2 | 89.9 |

\(^a\)Integrated electrocardiogram characteristics to diagnose elevated PAP using any of the following abnormalities: \(^b\)P pulmonale indicates right atrial enlargement, right atrial abnormality, or biatrial abnormalities in ECG annotation; \(^c\)right axis deviation in ECG annotation; \(^d\)right ventricular hypertrophy in ECG annotation; \(^e\)right ventricular strain indicates T-wave abnormalities or ST-segment depression over anterior precordial leads in ECG annotation; \(^f\)right bundle branch block mimics indicates right bundle branch block or rSR pattern in V1 or V2 without fulfilling the criteria of right bundle branch block in ECG annotation.

ECG — electrocardiogram; ePAP — elevated pulmonary artery pressure; PAP — pulmonary artery pressure.

### FIGURE 4

**Kaplan-Meier Survival Curves for AI-Classified ePAP or Non-ePAP Patients**

**A** Cardiovascular mortality. **B** All-cause mortality. Abbreviations as in Figure 1.

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AI-predicted ePAP: Group 1 patients classified by the AI model as having ePAP. AI-predicted non-ePAP: Group 1 patients classified by the AI model as having non-ePAP. **(A)** Cardiovascular mortality. **(B)** All-cause mortality. Abbreviations as in Figure 1.
that incorporate clinical characteristics (eg, the presence of telangiectasia) and a variety of tests (eg, ECG, pulmonary function, serum N-terminal pro-B-type natriuretic peptide, and uric acid level) were developed to increase the chance of detection by relying heavily on expert opinions.8,9

Compared with these tests, our AI model automatically detected patients with ePAP and a high probability of PH with compatible, if not better, sensitivity and specificity. The results suggest the AI model could be used as a standalone test or incorporated into screening algorithms for early diagnosis of patients with PH. Notably, our AI model is noninvasive and needs only the information from ECG signals without any input of clinical characteristics or predetermined ECG parameters and diagnosis, which is a significant improvement to the recently reported AI algorithm.15 The low cost, low operator dependence, and worldwide usage of ECG examination confer an extraordinary opportunity for our AI-enabled ECG model to be a cost-effective and widely applicable tool to change clinical practice in screening patients at risk of PH.

We applied the same threshold used in the worst-performing validation model (Supplemental Figure S2) for the Group 1 patients to compute the AI model’s sensitivity and specificity for the external cohort of Japanese patients (Group 3). Although the diagnostic performance in the AUC of the AI model was the same (0.88) between the 2 cohorts, the specificity was lower and sensitivity higher for the external cohort caused by a tradeoff between specificity and sensitivity. In future application, a different threshold value could be used to meet specific clinical considerations for Japanese patients. In addition, because all the 10 AI models from the 10-fold cross validation produced similar test results, it is likely that, for future clinical practice, no particular model of the 10 could significantly outperform others, although the best validation model could be a logical choice for this purpose at present.

We also performed an analysis of the Akaike Information Criterion (AIC)28 for Group 1 patients. The AIC is used to evaluate the quality of each model relative to each of the other models in a multiple models analysis. The AIC analysis showed that the model combining baseline characteristics and AI-predicted ePAP had the best quality to assess the risk of cardiovascular mortality (Supplemental Table S7). The analysis also showed that, although the AI-predicted ePAP was the factor of the highest HR, it alone would not be a better model than that of all the baseline characteristics combined for predicting mortality risk. To be able to use these results in clinical practice, we devised an index score based on the HR for each parameter from the multivariate analysis of the combined model to calculate the risk of cardiovascular mortality and all-cause mortality for Group 1 patients, as shown in Supplemental Tables S8 and S9, respectively. By summing over the index scores, the 1-year cardiovascular mortality and 1-year all-cause mortality can be estimated from the high correlation between the index score sum, now called risk score, and mortality, as shown in Supplemental Figures S11 and S12, respectively. Note that the precipitating disruption of the correlation at the risk score of 16 for all-cause mortality was caused by there being only 2 patients with that highest risk score, and both survived at the 1-year point (Supplemental Figure S12).

A qualified biomarker needs to meet adequate diagnostic performance and be associated with the clinical endpoints of diseases. Although various AI models have been developed to detect cardiovascular anomalies, their relevance to cardiovascular
outcomes remains unknown. The present work showed that an AI-enabled ECG can be an independent predictor of long-term cardiovascular and all-cause mortality. As shown in Figure 4, our model identified patients at high risk of cardiovascular mortality, with the risk estimated at 4.2%, 9.4%, and 14.0%, and of all-cause mortality, estimated at 14.4%, 29.2%, and 40.6%, during their 1-, 3-, and 5-year follow-ups, respectively.

Stratification based on mortality risk is pivotal in optimizing therapeutic strategies to treat patients with PH. Currently, PH is clinically categorized as pulmonary arterial hypertension (PAH), or as PH caused by left heart disease, lung disease or hypoxia, chronic thromboembolism, or multifactorial mechanisms. The current guideline recommends using a comprehensive assessment of patient prognosis to administer therapies according to disease categories. For example, in PH patients diagnosed with PAH, the mortality risk can be classified as low, intermediate, or high, with an estimated 1-year all-cause mortality of below 5%, 5–10%, or above 10%, respectively.17,29-31

Doctors use this risk stratification along with patients’ clinical characteristics, exercise test results, serum N-terminal pro-B-type natriuretic peptide levels, cardiac images, and hemodynamic determinants to determine an adequate treatment strategy (eg, single or combination therapy, intravenous or oral therapy). 17 A similar strategy of decision-making is applied to patients with PH caused by left heart disease or chronic thromboembolism.32,33

The prognostic power of our AI model suggests its usefulness in providing accurate risk stratification to direct differential therapeutic interventions, although a randomized clinical trial is needed to justify this application. Moreover, according to our subgroup sensitivity analysis, the AI model could be consistently applied to estimate mortality risk for...
different PH categories, including PAH, PH caused by left heart disease, lung disease, or hypoxia, and chronic thromboembolism. The expected wide applicability to patients with different diseases is a significant asset of our AI model.

**STUDY LIMITATIONS.** The ePAP affirmed by TTE was selected as a surrogate marker for highly suspicious PH based on an established guideline, but no other marker was tested. This is a limitation due mainly to a limited number of qualified patients available in our database. For example, right heart catheterization for definitive diagnosis of PH was not used in this study because only 17 such patients (0.04% of Group 1) had an ECG within the 14-day interval of our study design. Likewise, the number of patients tested (279 patients) for the external cohort was relatively small. Finally, the ability of the AI model to detect dynamic changes, such as disappearing PH or PH developed at follow-up, is likely limited because the model was not designed to predict those, and the small number of patients under those conditions prevented a thorough analysis. Further study on larger cohorts is necessary to fully validate the AI model’s performance.

**CONCLUSIONS**

In this work, we showed that the deep learning neural network AI model we previously developed for detecting cardiac arrhythmias could be extended to accurately identify patients with ePAP from their ECG data. We further showed that the AI model could predict future incidents of ePAP, as well as risk for cardiovascular and all-cause mortality, even for patients with various different diseases. These results suggest our AI model could be a useful clinical tool to identify patients with PH so that treatment can be initiated early to improve their survival prognosis.

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COMPETENCY IN MEDICAL KNOWLEDGE: Pulmonary hypertension affects more than a million individuals around the world and causes premature disability and heart failure, and increases the 5-year mortality rate to more than 30% if left untreated. Early detection of ePAP is needed for prompt diagnosis and treatment to avoid detrimental consequences of pulmonary hypertension. We developed the ECG-based AI model to identify patients with ePAP and predicted their future risk for cardiovascular mortality, which was validated in independent patient groups, including an external cohort of Japanese patients. The diagnostic performance (AUC: 0.88) and risk prediction for cardiovascular mortality (HR: 3.6-21.1) satisfied clinical standards, and outperformed conventional ECG diagnosis by cardiologists.

TRANSLATIONAL OUTLOOK: The AI-enabled ECG model can serve as a first automated, qualified, and valid clinical test for early diagnosis of ePAP and prognosis of mortality risk.

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KEY WORDS all-cause mortality, artificial intelligence, cardiovascular mortality, deep learning, electrocardiogram, pulmonary hypertension

APPENDIX For expanded Methods and Results sections, and supplemental figures and tables, please see the online version of this paper.