Electrocardiographic findings associated with early clinical deterioration in acute pulmonary embolism

Anthony J. Weekes MD, MSc1 | Jaron D. Raper MD1 | Alyssa M. Thomas MD, MPH1 | Kathryn Lupez MD1 | Carly A. Cox MD1 | Dasia Esener MD, MS2 | Jeremy S. Boyd MD3 | Jason T. Nomura MD4 | Jillian Davison MD5 | Patrick M. Ockerse MD6 | Stephen Leech MD, MPH5 | Eric Abrams MD2 | Christopher Kelly MD6 | Nathaniel S. O’Connell PhD7

1Department of Emergency Medicine, Atrium Health’s Carolinas Medical Center (Carolinas Medical Center is the Central Site of the Pulmonary Embolism Short-term Outcomes Registry (PESCOR) consortium), Charlotte, North Carolina, USA
2Department of Emergency Medicine Kaiser Permanente, San Diego, California, USA
3Department of Emergency Medicine, Vanderbilt University Medical Center, Nashville, Tennessee, USA
4Department of Emergency Medicine, Christiana Care, Newark, Delaware, USA
5Department of Emergency Medicine, Orlando Health, Orlando, Florida, USA
6Division of Emergency Medicine, University of Utah Health, Salt Lake City, Utah, USA
7Department of Biostatistics and Data Science, Wake Forest School of Medicine, Winston-Salem, North Carolina, USA

Correspondence
Anthony J. Weekes MD, MSc, 1000 Blythe Blvd, Medical Education Building, 3rd Floor, Charlotte, North Carolina 28203.
Email: anthony.weekes@atriumhealth.org

Present address
Jaron D. Raper, Department of Emergency Medicine, University of Alabama at Birmingham, Birmingham, Alabama, USA
Alyssa M. Thomas, Emergency Department, Houston Methodist Baytown Hospital, Houston, Texas, USA
Kathryn Lupez, Department of Emergency Medicine, Tufts Medical Center, Boston, Massachusetts, USA
Carly A. Cox, Emergency Medicine of Idaho, Meridian, Idaho, USA

Abstract

Objectives: We sought to determine associations of early electrocardiogram (ECG) patterns with clinical deterioration (CD) within 5 days and with RV abnormality (abnormal RV) by echocardiography in pulmonary embolism (PE).

Methods: In this prospective, multicenter study of newly confirmed PE patients, early echocardiography and initial ECG were examined. Initial ECG patterns included lead-specific ST-segment elevation (STE) or depression (STD), T-wave inversion (TWI), supraventricular tachycardia (SVT), sinus tachycardia, and right bundle branch block as complete (cRBBB) or incomplete (iRBBB). We defined CD as respiratory failure, hypotension, dysrhythmia, cardiac arrest, escalated PE intervention, or death within...
5 days. We calculated odds ratios (ORs) for CD and abnlRV with univariate and full multivariate models in the presence of other variables.

**Results:** Of 1676 patients, 1629 (97.2%) had both ECG and GDE; 415/1676 (24.7%) had CD, and 529/1629 (32.4%) had abnlRV. AbnlRV had an OR for CD of 4.25 (3.35, 5.38). By univariable analysis, the absence of abnormal ECG patterns had OR for CD and abnlRV of 0.34 (0.26, 0.44; \(p < 0.001\)) and 0.24 (0.18, 0.31; \(p < 0.001\)), respectively. By multivariable analyses, one ECG pattern had a significant OR for CD: SVT 2.87 (1.66, 5.00). Significant ORs for abnlRV were: TWI V_{2,4} 4.0 (2.64, 6.12), iRBBB 2.63 (1.59, 4.38), STE aVR 2.42 (1.58, 3.74), S1-Q3-T3 2.42 (1.70, 3.47), and sinus tachycardia 1.68 (1.14, 2.49).

**Conclusions:** SVT was an independent predictor of CD. TWI V_{2,4}, iRBBB, STE aVR, sinus tachycardia, and S1-Q3-T3 were independent predictors of abnlRV. Finding one or more of these ECG patterns may increase considerations for performance of echocardiography to look for RV abnormalities and, if present, inform concerns for early clinical deterioration.

**INTRODUCTION**

The electrocardiogram (ECG) is one of the first tests performed in the emergency department (ED) on patients with symptoms of chest discomfort or dyspnea. Abnormal ECG findings can be found in cardiopulmonary conditions outside of acute coronary syndrome. Although no specific ECG pattern is diagnostic of acute PE, abrupt pulmonary arterial occlusion may provoke abnormalities of the right ventricle (RV) due to increased pressure, dilatation, or myocardial injury. Abnormal ECG patterns may also occur due to left ventricular (LV) ischemia from combinations of hypoxia or hypotension (decreased oxygen supply) and due to tachycardia (increased oxygen demand). Abnormal ECG patterns associated with RV abnormality (abnlRV) in PE include rightward axis shifts, repolarization abnormalities, ST-segment myocardial injury patterns, conduction delays, and rhythm disturbances.1,2

There is a lack of agreement in the literature on ECG changes in PE and their relation to abnlRV, and one study found no relationship whatsoever.3 Prior research in this area has been limited due to retrospective analyses, heterogeneous cohorts of PE patients, small sample sizes, and ECGs not immediately performed at time of PE presentation.4–12 Few reports involve immediate and contemporaneous evaluations of ECG and echocardiography at PE presentation. Still fewer focus on clinical outcomes that require hospital-based monitoring or support.

The primary objective of this study was to prospectively determine differences in proportions of predefined ECG patterns between ED patients with PE who experience early clinical deterioration and those who do not. The null hypothesis was there is no difference in proportions of ECG patterns between these two groups of patients. Our secondary objective was to determine the proportions of specific ECG patterns in PE patients with and without abnlRV by goal-directed echocardiography (GDE). The secondary null hypothesis was there is no difference in the proportions of ECG patterns between those with and without abnlRV. For exploratory objectives, we evaluated (1) the association of mean number of ECG patterns with classifications by PE triaging strategies, and (2) the association of specific ECG patterns with abnormal heart features (LV dysfunction, abnlRV by computed tomography [CT], and elevated biomarkers).

**METHODS**

**Study design and setting**

This study was a preplanned analysis from a previously reported registry database of ED patients with confirmed acute PE (clinicaltrials.gov NCT02883491 and NCT03915925).13 All components of this registry were collected at six urban, academic EDs with emergency medicine residency and advanced emergency ultrasound fellowship programs. A central institutional review board (IRB) approved this federally funded multi-site study.

**Study population**

Adult ED patients (18 years or older) with image confirmed acute PE diagnosed within 12 hours of ED presentation were eligible for enrollment. PE was confirmed by the presence of filling defects in the pulmonary arteries with contrast-enhanced chest CT or detection of high probability on nuclear ventilation perfusion scan. We excluded patients who refused consent for clinical follow-up and those who had a ventricular paced rhythm or poor quality ECG tracings that were considered uninterpretable.
Study protocol

As previously reported, the primary goal of the PE registry was to include early imaging, ECGs, and cardiac biomarkers of ED patients with confirmed PE in the database and determine their association with clinical deterioration.\(^1\)\(^2\) Research associates at each study site maintain awareness of all ED patients with positive PE studies. In addition, physicians caring for these patients routinely notified the research team and performed GDE. Clinical characteristics and demographic information were collected on each patient. ECGs were performed during the index PE ED visit as part of routine care. Emergency physicians performed GDE during the index PE ED visit, which was subsequently interpreted by faculty of the emergency ultrasound program. ECG and echocardiography were performed independently of each other. The clinical course of patients was monitored by thorough review of the electronic medical record (EMR) or direct communication with inpatient health care providers.

The 12-lead ECG was completed during the ED management phase. ECGs were performed at the discretion of the treating health care team in response to the patient’s symptoms, clinical course, or PE diagnosis. Site investigators interpreted the 12-lead ECGs using predefined criteria discussed in pre-enrollment training modules. The 12-lead ECG was evaluated for each of the following previously reported patterns in PE: sinus tachycardia ($\geq 100$ beats per minute); incomplete right bundle branch block (IRBBB) if QRS complex $\geq 0.10$ and $< 0.12$ seconds and complete right bundle branch block (CRBBB) if QRS complex width $\geq 0.12$ seconds; complications of S wave in lead I, Q wave in lead III, and T wave inversion (TWI) in lead III, referred to as S1-Q3-T3 (the McGinnes-White sign); TWIs in precordial leads V2-4; TWI in leads II, III, and aVF; supraventricular tachycardia (SVT) including atrioventricular [AV] nodal reentry tachycardia, AV reentrant tachycardia, atrial tachycardia [unifocal or multifocal], junctional tachycardia, sinus nodal reentry tachycardia, atrial fibrillation, and atrial flutter); ST-segment depressions (STDs) in V4-6; ST-segment elevation (STE) in lead aVR, left bundle branch block (LBBB); and LV hypertrophy (LVH) with associated TWI.\(^2\)\(^4\)\(^5\)\(^8\) The ECG was considered positive for abnormal patterns if one or more of the above patterns were present. A TWI wave was defined as a negative deflection of the T wave greater than 0.5 mV below the T-P segment, which represents the isoelectric line (except in aVR or coexisting RBBB). TWI or STE associated with LBBB or RBBB patterns were considered normal. We used predefined criteria and guidelines. STE was determined 1 mm above the J point relative to the TP segment in the absence of early repolarization signs, LVH, or LBBB. ST depression was determined $> 0.5$ mm below the J point relative to the TP segment. TWI or STE in precordial leads associated with LBBB or RBBB patterns were not considered independent ECG abnormalities for this study. In addition, we reported the total number of mutually exclusive abnormal ECG patterns in the patient’s initial ECG as a discrete predictor variable.

Key outcome measures

The primary outcome was the incidence of one or more discrete clinical deterioration events that were considered PE-related complications, including death. Clinical deterioration included any episode that required hospital-based monitoring or hospital-based support or intervention. As previously reported, clinical deterioration endpoints included respiratory failure, cardiac arrest, new dysrhythmia, sustained hypotension requiring intravenous volume expansion or adrenergic medication, and escalated PE interventions.

In this prospective study, each site had institutional guidelines on PE management. We defined respiratory failure as respiratory distress associated with emergent or unplanned mechanical ventilation (intubation, non-invasive positive pressure ventilation, or cricothyrotomy). We defined cardiac arrest as new episodes of unstable cardiac rhythm or absent electrical activity with advanced cardiac life support for asystole, pulseless electrical activity, ventricular fibrillation, or unstable ventricular tachycardia. We defined new dysrhythmia as the onset of atrial fibrillation with rapid ventricular response, atrial flutter, SVT, stable ventricular tachycardia, or bradycardia that was not present at the ED presentation. We defined sustained hypotension as systolic blood pressure $< 90$ mm Hg (or $> 40$ mm Hg decrease from baseline) or shock index greater than 1.0 for $> 15$ minutes with administration of $> 499$ ml of intravenous fluids bolus for volume expansion or an infusion of nor-epinephrine, dopamine, or epinephrine. Escalated PE interventions included reperfusion interventions (systemic or catheter-directed thrombolysis, mechanical catheter directed aspiration, and surgical thrombectomy) and extracorporeal membrane oxygenation interventions.\(^1\)\(^3\) Clinical deterioration episodes were considered as single discrete episodes if separated by 12 hours or more. Individual patients could have more than one episode of clinical deterioration within the 5-day window. We followed patients via electronic medical record review during the hospital stay and up to 30 days later, as previously published.\(^1\)\(^3\)

The secondary outcome was the presence of abnlRV as determined by GDE. GDE was performed during the ED visit by emergency medicine physicians trained in GDE image acquisition and subsequently interpreted by site investigators (advanced emergency ultrasound fellowship directors). As previously reported, abnlRV included the presence of one or more of the following echocardiography findings: severe RV dilatation detected as qualitative assessment of RV:LV ratio $\geq 1:0$ and RV apex blunting in two or more views; severe RV systolic dysfunction as qualitatively estimated by diminished longitudinal contraction of the RV (estimated tricuspid annular plane systolic excursion [TAPSE] $\leq 1.0$ cm in subcostal or apical four-chamber views); and flattening or leftward bowing of the interventricular septum.\(^1\)\(^3\)\(^4\) (Rationale for using TAPSE $< 1.0$ cm as cut-off: The American Society of Echocardiography provides a normal reference range for RV measurements but no stratified classification for mild, moderate, or severe RV dilatation or systolic dysfunction.\(^1\)\(^5\)\(^-\)\(^7\) In previous reports, using TAPSE $< 1.0$ cm to
estimate severe RV systolic dysfunction had strong agreement with comprehensive echocardiography and strong inter- and intra-rater agreement.)

Site investigators (advanced emergency ultrasound fellowship directors) reviewed and interpreted the images according to interpretation guidelines discussed in training modules before the start of the study. GDE for abnRV in PE was scored on a 0–3 scale, where 0 = no severe RV dilatation, 1 = severe RV dilatation only, 2 = severe RV dilatation with either septal deviation or severe RV systolic dysfunction, and 3 = severe RV dilatation with both septal deviation and severe RV systolic dysfunction. The presence of abnRV was defined by a score of 1, 2, or 3. The frequency of ECG patterns was compared to abnRV responses as an ordinal variable (determined by the GDE severity score). As previously reported in the main manuscript from the PE registry database, the ordinal scale showed increased odds of clinical deterioration as scores increased.13 The GDE scale showed high intra- and inter-rater agreement and accuracy compared to comprehensive echocardiography.14,18

We compared the mean number of ECG patterns between assigned risk groups within a PE triaging strategy. We stratified each PE triaging strategy into low-risk and not low-risk groups. We used three PE triaging strategies: simplified pulmonary embolism severity index (sPESI), modified European Society of Cardiology (ESC), and the pulmonary embolism short-term clinical outcomes estimator (PE-SCORE).13,19,20 The sPESI was dichotomized as low-risk (0 points) versus not low-risk (>0 points). ESC was dichotomized as low-risk versus not low-risk. Low-risk ESC was defined as sPESI = 0 points and being without abnormal cardiac biomarkers or abnormal RV by CT or echocardiography. Not low-risk PE involves abnormal vital signs, RV abnormality by cardiac biomarkers, or abnormal RV by imaging (CT or echocardiography). PE-SCORE, which has a scale of 0 to 10 total points, was stratified as low-risk (0 points), intermediate-risk (1 to 4 points), and high-risk (>4 points) groups.

We compared proportions of the specific ECG patterns to the following abnormal heart features: abnormal LV systolic function (<30%), RV abnormality as measured by CT RV:LV ratio ≥1.0, and elevated cardiac biomarkers (definitions follow). As a surrogate of myocardial stretch, brain natriuretic peptide (BNP) elevation was defined by BNP ≥90 pg/ml or N-terminal BNP ≥500 pg/ml. We used elevated troponin levels as a surrogate of myocardial injury. Toward the last quarter of the enrollment phase, the central site had an institution-wide change from troponin I to high sensitivity troponin. As previously reported in the first manuscript of the PE registry, at each participating institution, an elevated troponin I was defined as ≥0.07 ng/ml (99% upper reference limit ≥0.028 ng/ml). An elevated high sensitivity troponin was defined as ≥20 pg/ml for males and 12 pg/ml for females (99% upper reference limit range between 24 and 30 pg/ml). We used i-STAT BNP test cartridge measured in pg/ml and i-STAT cardiac troponin test cartridge measured in pg/ml (both from Abbott Point of Care, Abbott Park, IL) for troponin I or high sensitivity troponin assay.

Data analysis

We conducted univariate analyses for the primary and secondary outcomes. For the secondary outcome, GDE score (ordinal from 0–3) was dichotomized to 0 vs. >0 for ease of interpretation and presentation of results. In addition, we stratified ECG patterns by abnormal LV systolic function, CT RV:LV ratio elevation, troponin elevation, and BNP elevation as binary dependent variables. We stratified the number of abnormal ECG patterns by the three PE triaging strategy classifications as dependent variables. Descriptive statistics included means, standard deviations, median, and interquartile ranges (IQRs) for continuous variables, as well as frequencies and percentages for binary variables. For univariable statistical comparisons, we used two-sample t-tests or analysis of variance (ANOVA) for continuous variables and chi-square tests for categorical variables.

For each primary and secondary outcome, we fit a multivariable logistic regression model with all ECG variables included as primary main effects of interest, and with the following as confounders: race, gender, ethnicity, age, initial heart rate, shock index, initial respiratory rate, initial oxygen saturation, preceding episode of syncope, prior diagnosis of PE, and whether patients had one or more abnormal ECG patterns or not. In addition, in the model for primary composite outcome, we included RV:LV ratio ≥1 as determined by CT, troponin elevation, echocardiography showing abnRV, and natriuretic peptide elevation as additional predictors. We displayed results for each full regression model in tables with odds ratios (ORs) with 95% confidence intervals (CIs). All p values <0.05 were considered statistically significant.

RESULTS

As shown in Figure 1, 1736 PE patients met inclusion criteria for the registry databases between August 2016 and November 2020. Of these, 1680 (96.8%) patients had an initial ECG performed in the ED. Four of these patients had external cardiac devices leading to ventricular paced rhythms and were excluded from this analysis. We therefore had 1676 patients with initial native ECG tracings available for statistical analyses for the primary outcome. ECGs were performed within a mean of 1.90 ± 3.38 hours of measuring initial vital signs. Enrollment numbers for the six clinical sites were: Site 1 (695, 41.5%), Site 2 (248, 14.8%), Site 3 (206, 12.3%), Site 4 (156, 9.3%), Site 5 (185, 11.0%), and Site 6 (186, 11.1%).

All ECGs and GDE were performed during the immediate ED course and before anticoagulation or escalated PE intervention. Anticoagulation or escalated PE intervention was initiated either during the ED course or immediately on arrival to the hospital floor in 97.2% (1630/1676) of the PE patients. We followed patients via EMR review during the hospital stay and up to 30 days, as previously published.13 All patients included in this report had 5-day follow up information completed.

Demographics (Table 1) included the mean age of 59.6 ± 16.6 years, 48.2% female, 65.5% Caucasian, 29.1% African
American, and 1% Asian. The mean shock index was 0.77 ± 0.25 beats per minute/mmHg, respiratory rate 19.9 ± 4.6 breaths per minute, and oxygen saturation 95.4 ± 4.6%. Of the 1676 patients, 160 (9.5%) had a preceding episode of syncope and 420 (25.1%) had prior diagnosis of PE or deep venous thrombosis (DVT). For each PE triaging strategy, clinical deterioration was experienced by a lower proportion of patients classified as low-risk compared to those classified as non-low-risk.

Of the 1676 patients, 415 (24.7%) had one or more clinical deterioration events or hospital-based intervention within 5 days of PE diagnosis. Forty-seven of the 1676 patients did not have GDE. Of the 1629 patients with ECG and GDE, 529 (32.4%) had abnlRV by GDE. Table 1 shows numbers and proportions with abnlRV by GDE stratified by clinical deterioration, leading to an OR of 4.25 (3.35, 5.38, p < 0.0001).

Table 2 shows the prevalence of each specific ECG pattern. By univariate analysis, the absence of abnormal ECG patterns had an OR for clinical deterioration of 0.34 (0.25, 0.44; p < 0.001). The most common ECG patterns were sinus tachycardia (38.9%), S1–Q3–T3 pattern (16.3%), incomplete or complete RBBB (15.1%), and TWI in V2–4 (14%). In comparison to patients without the specific ECG pattern, those with the following patterns were significantly more likely to experience clinical deterioration: sinus tachycardia, TWI in V2–4, STE in aVR, STE in V1, SVT, TWI in II and III and aVF, and STD in V4–6. LVH with TWI (3%) and LBBB were uncommon and not significantly different between outcome groups.

The results of full model multivariable logistic regression analysis are shown in Table 3. SVT was the only ECG pattern that was an independent predictor of CD. Absence of abnormal ECG patterns was not an independent predictor. Abnormal RV by GDE, elevated troponin measurements, elevated natriuretic peptide measurements, CT derived RV-LV ratio ≥1.0, shock index >1.0 and preceding episode of syncope were independent predictors of clinical deterioration. Table 4 shows univariable analysis results for all ECG patterns, except LBBB and LVH, had significant ORs for abnlRV by GDE (our secondary outcome). Sinus tachycardia, TWI in II III, aVF and V2–4, S1–Q3–T3, iRBBB, and STE in aVR had ORs with the lower boundary of 95% CI above 1.0. For presence and absence of abnlRV by
### Table 1 Patient characteristics

|                          | No clinical deterioration (N = 1261) | Clinical deterioration (N = 415) | Overall (N = 1676) |
|--------------------------|--------------------------------------|----------------------------------|---------------------|
| **Gender**               |                                       |                                  |                     |
| Female                   | 601 (47.7%)                          | 206 (49.6%)                      | 807 (48.2%)         |
| Male                     | 660 (52.3%)                          | 209 (50.4%)                      | 869 (51.8%)         |
| **Race**                 |                                       |                                  |                     |
| Caucasian                | 819 (64.9%)                          | 279 (67.2%)                      | 1098 (65.5%)        |
| African/American         | 376 (29.8%)                          | 112 (27.0%)                      | 488 (29.1%)         |
| American Indian/Alaskan Native | 10 (0.8%)                             | 2 (0.5%)                         | 12 (0.7%)           |
| Asian                    | 12 (1.0%)                            | 4 (1.0%)                         | 16 (1.0%)           |
| Hispanic                 | 91 (7.2%)                            | 25 (6.0%)                        | 116 (6.9%)          |
| Asian                    | 12 (1.0%)                            | 4 (1.0%)                         | 16 (1.0%)           |
| **Mean Age, years**      | 58.76 ± 16.8                          | 62.2 ± 15.7                      | 59.6 ± 16.6         |
| **Systolic blood pressure, mm Hg** | 133.5 ± 23.1                          | 122.2 ± 25.5                     | 132.2 ± 24.4        |
| **Heart rate, beats per minute** | 95.4 ± 20.05                           | 106.2 ± 23.6                     | 98.1 ± 21.5         |
| **Shock index**          | 0.73 ± 0.22                           | 0.91 ± 0.29                      | 0.77 ± 0.25         |
| **Respiratory rate, breaths per minute** | 19.6 ± 4.2                             | 21.0 ± 5.54                      | 19.9 ± 4.60         |
| **Oxygen saturation, %** | 95.8 ± 4.2                            | 94.2 ± 5.6                       | 95.4 ± 4.63         |
| **Cancer**               | 302 (23.9%)                          | 109 (26.3%)                      | 411 (24.5%)         |
| **COPD**                 | 176 (14.0%)                          | 75 (18.1%)                       | 251 (15.0%)         |
| **Previous PE/DVT**      | 313 (24.8%)                          | 107 (25.8%)                      | 420 (25.1%)         |
| **Abnormal RV by GDE (N = 1629)** | 296 (24.2%)                           | 233 (57.5%)                      | 529 (43.2%)         |
| Present                  | 928 (75.8%)                          | 172 (42.5%)                      | 1100 (89.9%)        |
| **PE-Score**             |                                       |                                  |                     |
| Low-risk (0 points)      | 294 (23.3%)                          | 15 (3.6%)                        | 309 (18.4%)         |
| Intermediate-risk (1–4 points) | 864 (68.5%)                           | 295 (71.1%)                      | 1159 (69.2%)        |
| High-risk (>4 points)    | 26 (2.1%)                            | 75 (18.1%)                       | 101 (6.0%)          |
| Missing                  | 77 (6.1%)                            | 30 (7.2%)                        | 107 (6.4%)          |
| **ESC**                  |                                       |                                  |                     |
| Low-risk                 | 166 (13.2%)                          | 5 (1.2%)                         | 171 (10.2%)         |
| Not low-risk             | 1095 (86.8%)                         | 410 (98.8%)                      | 1505 (89.8%)        |
| Missing                  | 0(0%)                                | 0 (0%)                           | 0 (0%)              |
| **sPESI**                |                                       |                                  |                     |
| Low-risk                 | 518 (41.1%)                          | 71 (17.1%)                       | 589 (35.1%)         |

Abbreviations: COPD, chronic obstructive pulmonary disease; DVT, deep venous thrombosis; ESC, European Society of Cardiology PE dichotomized into low-risk versus not low-risk classifications; GDE, goal-directed echocardiography; PE, pulmonary embolism; PE-Score, pulmonary embolism short-term clinical outcomes risk estimator; RV, right ventricle; sPESI, simplified Pulmonary Embolism Severity Index.

---

GDE, the mean number of abnormal ECG patterns was 2.20 ± 1.69 and 0.93 ± 1.13, respectively (p < 0.0001). The absence of abnormal ECG patterns had an OR for abnlRV by GDE of 0.24 (0.18, 0.31; p < 0.001). The results of full multivariable logistic regression analysis are shown in Table 5. Independent ECG predictors of abnlRV in order of ORs (high to low) were: STE in aVR, TWI V2–4, incomplete RBBB, S1-Q3-T3, and sinus tachycardia. LVH with associated TWI lowered the odds of abnlRV. With multivariable analysis, the absence of any of the ECG patterns was not an independent predictor of clinical deterioration. Clinical predictors were age, shock index, initial heart rate, initial respiratory rate, and preceding episode of syncope.

Of the 1676 patients, 1090 (65.0%) had at least one of the abnormal ECG patterns. The mean number of abnormal ECG patterns was 1.33 ± 1.4. For patients with and without the primary outcome, the mean numbers of abnormal ECG patterns were 1.97 ± 1.65 and 1.20 ± 1.30, respectively (Table 2). Supporting Information Table S1, which is available online, shows patients with low-risk classifications by all three PE triaging strategies had a lower number of abnormal ECG patterns compared to those in the higher risk classifications.

Supporting Information Tables S2–S9 show results of analyses of specific ECG patterns stratified by LV systolic dysfunction, RV dilatation by CT, and elevated cardiac biomarkers. Of 1629 with GDE, 1620 had a report on LV systolic function, with 127 (7.8%) having severe LV dysfunction. SVT, LBBB, and LVH with TWI were significantly associated with severe LV systolic dysfunction by univariable analysis (Supporting Information Table S2), whereas only SVT and LBBB were significant by multivariable analysis (Supporting Information Table S3). Supporting Information Table S4 shows 450 of 1659 patients (27.1%) had troponin elevations. All ECG patterns, except iRBBB, were associated with troponin elevation by univariable analysis. However, Supporting Information Table S5 shows sinus tachycardia, S1-Q3-T3, TWI V2–4, and STE aVR were independent predictors of troponin elevations by multivariable analysis. Supporting Information Table S6 shows 642 of 1608 patients (39.9%) with BNP measurements had BNP elevations. All ECG patterns, except iRBBB, were associated with natriuretic peptide elevation by univariable analysis. Supporting Information Table S7 shows S1-Q3-T3, STE V2–4, and SVT were significant by multivariable analysis. Supporting Information Table S8 shows...
**TABLE 2** ECG patterns stratified by primary clinical outcomes

| ECG Patterns | No clinical deterioration \((N = 1261)\) | Clinical deterioration \((N = 415)\) | Overall \((N = 1676)\) | OR | p-Value |
|--------------|-----------------------------------------|---------------------------------|----------------------|----|---------|
| **Mean number of abnormal ECG patterns** | \(1.20 \pm 1.31\) | \(1.99 \pm 1.67\) | \(1.34 \pm 1.44\) | \(0.34\) \((0.26, 0.44)\) | \(<0.0001\) |
| **Number of abnormal ECG patterns** | | | | \(0.34\) \((0.26, 0.44)\) | \(<0.0001\) |
| Zero abnormal patterns | 509 (40.4%) | 77 (18.6%) | 586 (35.0%) | 0.34 \((0.26, 0.44)\) | \(<0.0001\) |
| >0 abnormal patterns | 751 (59.6%) | 338 (81.4%) | 1089 (65.0%) | 0.34 \((0.26, 0.44)\) | \(<0.0001\) |
| Missing | 1 (0.1%) | 0 (0%) | 1 (0.1%) | 0.34 \((0.26, 0.44)\) | \(<0.0001\) |
| Complete RBBB | | | | | |
| Absent | 1180 (93.6%) | 369 (88.9%) | 1549 (92.4%) | 1.82 \((1.24, 2.66)\) | 0.003 |
| Present | 81 (6.4%) | 46 (11.1%) | 127 (7.6%) | | |
| Incomplete RBBB | | | | | |
| Absent | 1175 (93.2%) | 376 (90.6%) | 1551 (92.5%) | 1.42 \((0.95, 2.11)\) | 0.104 |
| Present | 86 (6.8%) | 39 (9.4%) | 125 (7.5%) | | |
| Sinus tachycardia | | | | | |
| Absent | 835 (66.2%) | 189 (45.5%) | 1024 (61.1%) | 2.34 \((1.87, 2.94)\) | \(<0.001\) |
| Present | 426 (33.8%) | 226 (54.5%) | 652 (38.9%) | | |
| S1-Q3-T3 pattern | | | | | |
| Absent | 1080 (85.6%) | 322 (77.6%) | 1402 (83.7%) | 1.72 \((1.30, 2.28)\) | \(<0.001\) |
| Present | 181 (14.4%) | 93 (22.4%) | 274 (16.3%) | | |
| STE V1 | | | | | |
| Absent | 1162 (92.1%) | 362 (87.2%) | 1524 (90.9%) | 1.72 \((1.21, 2.45)\) | 0.003 |
| Present | 99 (7.9%) | 53 (12.8%) | 152 (9.1%) | | |
| T wave inversions V2-4 | | | | | |
| Absent | 1112 (88.2%) | 330 (75.9%) | 1442 (86.0%) | 1.92 \((1.43, 2.58)\) | \(<0.001\) |
| Present | 149 (11.8%) | 53 (20.5%) | 202 (14.0%) | | |
| T wave inversions II, III, aVF | | | | | |
| Absent | 1153 (91.4%) | 354 (85.3%) | 1507 (89.9%) | 1.84 \((1.32, 2.57)\) | \(<0.001\) |
| Present | 108 (8.6%) | 61 (14.7%) | 169 (10.1%) | | |
| ST-segment depression V4-6 | | | | | |
| Absent | 1183 (93.8%) | 354 (85.3%) | 1537 (91.7%) | 2.61 \((1.83, 3.73)\) | \(<0.001\) |
| Present | 78 (6.2%) | 61 (14.7%) | 139 (8.3%) | | |
| STE aVR | | | | | |
| Absent | 1146 (90.9%) | 324 (78.1%) | 1470 (87.7%) | 2.82 \((2.09, 3.82)\) | \(<0.001\) |
| Present | 114 (9.0%) | 91 (21.9%) | 205 (12.2%) | | |
| Missing | 1 (0.1%) | 0 (0%) | 1 (0.1%) | | |
| SVT (including atrial fibrillation with rapid ventricular response) | | | | | |
| Absent | 1218 (96.6%) | 361 (87.0%) | 1579 (94.2%) | 4.24 \((2.80, 6.43)\) | \(<0.001\) |
| Present | 43 (3.4%) | 54 (13.0%) | 97 (5.8%) | | |
| LBBB associated with TWI | | | | | |
| Absent | 1238 (98.2%) | 412 (99.3%) | 1650 (98.4%) | 0.39 \((0.11, 1.31)\) | 0.179 |
| Present | 23 (1.8%) | 3 (0.7%) | 26 (1.6%) | | |
| LVH with TWI | | | | | |
| Absent | 1233 (97.8%) | 402 (96.9%) | 1635 (97.6%) | 1.42 \((0.73, 2.78)\) | 0.39 |
| Present | 28 (2.2%) | 13 (3.1%) | 41 (2.4%) | | |

Abbreviations: ECG, electrocardiography; LBBB, left bundle branch block; LVH, left ventricular hypertrophy; OR, odds ratio; STE, ST-segment elevation; SVT, supraventricular tachycardia; RBBB, right bundle branch block; TWI, T-wave inversion (0.5 mV negative deflection).

*The total number of ECG patterns present was greater than the total number of unique patients because any single patient could have more than one of the ECG patterns.

*For those with either iRBBB/cRBBB without TWI in V2-4 \((n = 168)\) versus those with TWI V2-4 \((n = 82)\), 56 (33.3%) and 29 (35.2%) had clinical deterioration, respectively. The OR for iRBBB without TWI was 0.91 \((0.52, 1.59)\); \(p = 0.75\).
### Table 3 Multivariable analysis stratified by clinical deterioration within 5 days

| Predictors                                    | OR     | CI       | p-Value |
|-----------------------------------------------|--------|----------|---------|
| (Intercept)                                   | 0.56   | 0.03–13.59 | 0.722   |
| Complete RBBB                                 | 1.06   | 0.64–1.73  | 0.825   |
| Incomplete RBBB                               | 1.03   | 0.61–1.69  | 0.918   |
| Sinus tachycardia                             | 1.43   | 0.96–2.15  | 0.081   |
| S1-Q3-T3 pattern                              | 0.85   | 0.58–1.22  | 0.375   |
| ST elevation V1                               | 0.90   | 0.55–1.45  | 0.670   |
| T wave inversions V2-4                        | 1.01   | 0.66–1.52  | 0.981   |
| T wave inversions II, III, aVF                | 1.08   | 0.67–1.71  | 0.754   |
| ST depression in V4-6                         | 1.14   | 0.69–1.85  | 0.614   |
| STE aVR                                       | 1.16   | 0.75–1.78  | 0.494   |
| SVT                                           | 2.87   | 1.66–5.00  | <0.001  |
| LBBB with associated TWI                     | 0.21   | 0.03–0.86  | 0.060   |
| LVH with associated TWI                       | 1.49   | 0.60–3.47  | 0.369   |
| Echocardiography showing abnormal RV          | 1.68   | 1.17–2.41  | 0.005   |
| Elevated troponin level                       | 1.47   | 1.08–2.00  | 0.014   |
| RV:LV ratio 1.0 or greater by CT              | 1.49   | 1.07–2.08  | 0.019   |
| Elevated natriuretic peptide level            | 1.26   | 0.93–1.70  | 0.130   |
| Male                                          | 1.01   | 0.77–1.32  | 0.952   |
| African American/Black                        | 0.66   | 0.36–1.23  | 0.182   |
| White                                         | 0.81   | 0.46–1.48  | 0.486   |
| Ethnicity                                     | 1.00   | 1.00–1.00  | 0.322   |
| Age                                           | 1.01   | 1.00–1.02  | 0.025   |
| Initial heart rate                            | 0.99   | 0.97–1.00  | 0.008   |
| Initial shock index                           | 13.07  | 5.69–30.65 | <0.001  |
| Initial respiratory rate                      | 1.01   | 0.98–1.04  | 0.534   |
| Initial oxygen saturation on room air         | 0.97   | 0.95–1.00  | 0.060   |
| Preceding episode of syncope                  | 1.75   | 1.14–2.66  | 0.010   |
| Prior history of PE or DVT                    | 1.02   | 0.75–1.39  | 0.885   |
| No abnormal ECG pattern                      | 0.84   | 0.54–1.32  | 0.460   |
| Observations                                  | 1472   |           |         |
| $R^2$ $\text{Tjur}$                           | 0.197  |           |         |

Abbreviations: CI, confidence interval; DVT, deep venous thrombosis; ECG, electrocardiogram; LBBB, left bundle branch block; LVH, left ventricular hypertrophy; OR, odds ratio; PE, pulmonary embolism; STE, ST-segment elevation; SVT, supraventricular tachycardia; TWI = T-wave inversion (0.5 mV negative deflection).

Table S8 shows elevated CT RV:LV ratios in 552 of 1640 patients (33.7%) who had CT performed. All ECG patterns, except SVT, LBBB, and LVH with TWI, were associated with elevated CT RV:LV ratio by univariable analysis. Supporting Information Table S9 shows complete and incomplete RBBB, S1-Q3-T3, TWI V1–4, STE aVR, and SVT were significant by multivariable analysis. Finally, Supporting Information Table S10 shows the prognostic metrics of the following categorical variables: abnormal ECG patterns, elevated troponin, elevated natriuretic peptide, initial shock index (>1.0), CT RV:LV ratio, hypotension (<100 mmHg), hypoxia (<92%), and preceding syncope. Predictor positive likelihood ratios ordered high to low were: hypotension, initial shock index, preceding syncope, hypoxia, elevated troponin, increased RV:LV ratio, elevated natriuretic peptide, and abnormal ECG pattern. The number of abnormal ECG patterns and the CT RV:LV ratio demonstrated the highest negative predictive values, followed by elevated natriuretic peptide, initial shock index, elevated troponin, hypotension, hypoxia, and preceding syncope.

### Limitations

This study had several limitations. First, preexisting ECG findings were not reviewed to determine if index PE-associated ECG abnormalities were acute or preexisting. Second, there are multiple causes of tachycardia that are unrelated to and not directly provoked by the pathophysiology of PE. Third, TWIs may be associated with normal variants and can be associated with RBBB. Furthermore, the RBBB pattern can exist in the asymptomatic population without PE.

Fourth, we reported predetermined ECG patterns that were fully recorded and part of the medical record. It is plausible that ECG abnormalities were in evolution. STE was defined as the ST segment being ≥1 mm above the isoelectric line (TP segment). However, use of the J point relative to PQ junction, instead of TP segment, is currently advised for determining ST depression.

Several investigators witnessed important but transient changes to ECG patterns in higher acuity acute PE patients that were not fully recorded or available in the EMR for interpretation or analysis. Our report focused on the initial ECG in the ED; however, there were several instances in patients with abnormal RV by GDE where conduction, TWI, or ST-segment abnormalities were transient. We observed several instances of normalization of initial abnormal ECG patterns and improvements to RV dilatation or systolic function (determined by serial echocardiography) after administration of thrombolytic agents. Further supporting evidence for this observation is provided by Choi et al., who showed not only an association between TWI and abnlRV, but also found that reversals of T wave abnormalities were temporally associated with resolution of an abnormality. Our report focused on the initial ECG in the ED; however, there were several instances in patients with abnormal RV by GDE where conduction, TWI, or ST-segment abnormalities were transient. We observed several instances of normalization of initial abnormal ECG patterns and improvements to RV dilatation or systolic function (determined by serial echocardiography) after administration of thrombolytic agents. Further supporting evidence for this observation is provided by Choi et al., who showed not only an association between TWI and abnlRV, but also found that reversals of T wave abnormalities were temporally associated with resolution of an abnormality.
### Table 4: Univariable analysis of ECG patterns stratified by echocardiographic RV abnormalities

| Echocardiography showing RV abnormalities (47 missing) | No (N = 1100) | Yes (N = 529) | OR       | p-Value |
|------------------------------------------------------|---------------|---------------|----------|---------|
| **Mean number of ECG patterns**                      |               |               |          |         |
| (standard deviation)                                 | 0.93 (1.13)   | 2.20 (1.69)   | <0.0001  |         |
| **Number of abnormal ECG patterns**                  |               |               |          |         |
| None                                                 | 485 (44.1%)   | 84 (15.9%)    | 0.24 (0.18, 0.31) | <0.001 |
| One or more                                          | 614 (55.8%)   | 445 (84.1%)   |          |         |
| Missing                                              | 1 (0.1%)      | 0 (0%)        |          |         |
| **Complete RBBB**                                    |               |               |          |         |
| Absent                                               | 1043 (94.8%)  | 461 (87.1%)   | 2.70 (1.87, 3.9) | <0.001 |
| Present                                              | 57 (5.2%)     | 68 (12.9%)    |          |         |
| **Incomplete RBBB**                                  |               |               |          |         |
| Absent                                               | 1046 (95.1%)  | 458 (86.6%)   | 2.95 (2.04, 4.26) | <0.001 |
| Present                                              | 54 (4.9%)     | 71 (13.4%)    |          |         |
| **Sinus tachycardia**                                |               |               |          |         |
| Absent                                               | 759 (69.0%)   | 241 (45.6%)   | 2.67 (2.15, 3.30) | <0.001 |
| Present                                              | 341 (31.0%)   | 288 (54.4%)   |          |         |
| **S1-Q3-T3 pattern**                                 |               |               |          |         |
| Absent                                               | 991 (90.1%)   | 371 (70.1%)   | 3.87 (2.95, 5.08) | <0.001 |
| Present                                              | 109 (9.9%)    | 158 (29.9%)   |          |         |
| **STE V1**                                           |               |               |          |         |
| Absent                                               | 1026 (93.3%)  | 454 (85.8%)   | 2.29 (1.63, 3.22) | <0.001 |
| Present                                              | 74 (6.7%)     | 75 (14.2%)    |          |         |
| **T wave inversions V2-4**                           |               |               |          |         |
| Absent                                               | 1019 (92.6%)  | 381 (72.0%)   | 4.89 (3.64, 6.57) | <0.001 |
| Present                                              | 81 (7.4%)     | 148 (28.0%)   |          |         |
| **T wave inversions II, III, aVF**                   |               |               |          |         |
| Absent                                               | 1029 (93.5%)  | 434 (82.0%)   | 3.17 (2.29, 4.40) | <0.001 |
| Present                                              | 71 (6.5%)     | 95 (18.0%)    |          |         |
| **ST-segment depression V4-6**                        |               |               |          |         |
| Absent                                               | 1038 (94.4%)  | 453 (85.6%)   | 2.81 (1.98, 4.00) | <0.001 |
| Present                                              | 62 (5.6%)     | 76 (14.4%)    |          |         |
| **STE aVR**                                          |               |               |          |         |
| Absent                                               | 1023 (93.0%)  | 404 (76.4%)   | 4.16 (3.06, 5.55) | <0.001 |
| Present                                              | 76 (6.9%)     | 125 (23.6%)   |          |         |
| **SVT (including atrial fibrillation with rapid ventricular response)** | | | | |
| Absent                                               | 1049 (95.4%)  | 485 (91.7%)   | 1.87 (1.23, 2.83) | 0.004 |
| Present                                              | 51 (4.6%)     | 44 (8.3%)     |          |         |
| **LBBB associated with TWI**                          |               |               |          |         |
| Absent                                               | 1079 (98.1%)  | 524 (99.1%)   | 0.49 (0.18, 1.31) | 0.214 |
| Present                                              | 21 (1.9%)     | 5 (0.9%)      |          |         |
| **LVH with TWI**                                     |               |               |          |         |
| Absent                                               | 1070 (97.3%)  | 518 (97.9%)   | 0.76 (0.38, 1.52) | 0.54  |
| Present                                              | 30 (2.7%)     | 11 (2.1%)     |          |         |

Abbreviations: ECG, electrocardiogram; LBBB, left bundle branch block; LVH, left ventricular hypertrophy; OR, odds ratio; RBBB, right bundle branch block; RV, right ventricle; STE, ST-segment elevation; SVT, supraventricular tachycardia; TWI, T-wave inversion (0.5 mV negative deflection).
We prospectively evaluated PE patients in the ED to report on the associations of multiple specific ECG patterns for clinical deterioration and abnlRV by echocardiography using univariable and multivariable analyses. ECGs were performed early and contemporaneously with GDE assessments for abnlRV during the ED work-up. Abnormal RV (as determined by GDE) had an OR of 4.25 (3.35, 5.38) for clinical deterioration. The absence of abnormal ECG patterns on initial early ECG had ORs of 0.34 (0.26, 0.44) and 0.24 (0.18, 0.31) for clinical deterioration and abnlRV by GDE, respectively. However, on multivariable analysis, absence of any ECG patterns did not qualify as an independent predictor of clinical deterioration or abnlRV.

We found all abnormal ECG patterns had statistically significant increased ORS above unity for clinical deterioration within 5 days of PE diagnosis, except for LBBB and LVH with TWI. Of the ECG patterns, only SVT was significant as an independent predictor for clinical deterioration. For abnlRV by echocardiography, associations were significant for most ECG patterns. Sinus tachycardia, STE in aVR, incomplete RBBB, S1-Q3-T3, and TWI V2-4 were significant as independent predictors for abnlRV by GDE. Patients with acute pulmonary embolism may have one or more of the specific ECG abnormalities. In patients with confirmed acute pulmonary embolism, one or more of these ECG patterns should increase considerations for performance of echocardiography to look for RV abnormalities and, if present, increase concerns for early clinical deterioration. By univariable analysis, we found the absence of any of the abnormal ECG patterns was associated with lower probability of clinical deterioration. For abnlRV by echocardiography, associations of abnormal ECG patterns was not an independent predictor of clinical deterioration or abnlRV by echocardiography.

Most previous studies have been retrospective analyses with ECGs completed 10 to 72 hours removed from the time of PE diagnosis. Strengths of our study include its prospective design, a large database of patients representing the full spectrum of PE severity, and early timing of ECG relative to ED diagnosis of PE.

**Clinical deterioration**

Previous reports have shown several ECG patterns (ST, STsegment depression, S1-Q3-T3, RBBB, TWI, and SVT) were associated with clinical deterioration.2,12 Similarly, by univariable analysis, our prospective study showed significant associations of these abnormal ECG patterns with clinical deterioration within 5 days. In addition, we noted the following ECG patterns were also associated with increased risk for clinical deterioration: STE in V1; TWI in leads II, III, and aVF; ST depression in V4-V6; STE in aVR; and SVT (Table 2). However, SVT was the only ECG pattern determined to be an independent predictor from multivariable analyses in the presence of other clinical variables.

**Abnormal RV by echocardiography**

In our study, early echocardiography findings of abnormal RV within the ED course were significantly associated with clinical

---

**TABLE 5** Full multivariable analysis model stratified by RV abnormalities (by echocardiography)

| Predictors                              | Echocardiography with abnlRV |
|-----------------------------------------|------------------------------|
|                                         | OR   | CI   | p-Value |
| (Intercept)                             | 30.78| 1.09-93.83 | 0.046 |
| Complete RBBB                          | 1.23 | 0.74-2.04 | 0.422 |
| Incomplete RBBB                        | 2.63 | 1.59-4.38 | <0.001|
| Sinus tachycardia                      | 1.68 | 1.14-2.49 | 0.009 |
| S1-Q3-T3 pattern                       | 2.42 | 1.70-3.47 | <0.001|
| STE V1                                 | 1.36 | 0.83-2.22 | 0.215 |
| T wave inversions V2-4                 | 4.00 | 2.64-6.12 | <0.001|
| T wave inversions II, III, aVF         | 1.26 | 0.78-2.02 | 0.349 |
| ST depression in V4-6                  | 0.79 | 0.47-1.31 | 0.364 |
| STE aVR                                | 2.42 | 1.58-3.74 | <0.001|
| SVT                                    | 1.25 | 0.71-2.18 | 0.441 |
| LBBB with associated TWI              | 0.30 | 0.07-0.97 | 0.069 |
| LVH with associated TWI                | 0.33 | 0.11-0.86 | 0.029 |
| Male                                   | 0.97 | 0.75-1.27 | 0.848 |
| African American/Black                 | 1.18 | 0.64-2.25 | 0.605 |
| White                                  | 1.05 | 0.58-1.95 | 0.883 |
| Ethnicity                              | 1.00 | 1.00-1.00 | 0.579 |
| Age                                    | 1.02 | 1.01-1.03 | <0.001|
| Initial heart rate                     | 1.00 | 0.99-1.01 | 0.664 |
| Initial shock index                    | 5.45 | 2.36-12.76| <0.001|
| Initial respiratory rate               | 1.04 | 1.02-1.08 | 0.003 |
| Initial oxygen saturation on room air  | 0.92 | 0.89-0.94 | <0.001|
| Preceding episode of syncope           | 3.09 | 2.00-4.82 | <0.001|
| Prior history of PE or DVT            | 1.07 | 0.79-1.44 | 0.651 |
| No abnormal ECG pattern                | 1.08 | 0.70-1.68 | 0.722 |
| Observations                           | 1472 |
| \( R^2 \) Tjur                         | 0.284 |
deterioration within 5 days. In turn, abnormal ECG patterns (sinus tachycardia, STE in aVR, incomplete RBBB, S1-Q3-T3, and TWI V$_{2-4}$) were independent predictors of abnlRV by echocardiography even in the presence of other clinical variables. By multivariable analyses, the number of abnormal ECG patterns was not an independent predictor of clinical deterioration or abnl RV by echocardiography.

Reports on the associations between ECG and abnlRV by echocardiography in PE patients are mixed, with important differences in timing of ECG performance studies, patient cohorts, and the ECG patterns selected. A report by Vanni et al. studied ECG and echocardiography performed within 1 hour of PE diagnosis and found that patients with RBBB, precordial repolarization abnormalities, or the S1-Q3-T3 pattern complex were twice as likely to have echocardiographic signs of abnlRV than patients without any of the evaluated ECG patterns. Punukollu et al. showed S1-Q3-T3 and TWI in V$_{2-3}$ were more likely in abnlRV, especially if echocardiography was performed at admission or within 24 hours of hospitalization. Daniel et al. showed that having multiple abnormal ECG patterns offered low sensitivity but high specificity for echocardiographic evidence of severe abnlRV. In contrast, another report showed having a few abnormal ECG patterns had moderate sensitivity and moderate specificity for abnormal RV. In one study, TWI V$_{2-4}$ was six times more frequent in PE patients with abnlRV than those without abnlRV. Another report showed TWI in V$_{1-2}$, tachycardia, and S wave in lead I were strong predictors of abnlRV by echocardiography, CT, or cardiac biomarkers. In contrast, a retrospective study of 289 patients by Stein et al. did not find ECG to be useful in predicting the presence or absence of RV dilatation. In our study, the independent predictors of abnlRV by echocardiography were sinus tachycardia, S1-Q3-T3, TWI V$_{2-4}$, and STE in aVR. Our study findings suggest the presence of one or more of these ECG patterns may be associated with the presence of concurrent abnlRV in suspected or confirmed acute PE patients, which in turn, warns providers the PE patient under their care has an increased risk of clinical deterioration within 5 days. The presence of an acute abnlRV may be determined by reviewing recently performed imaging (CT or echocardiography).

Reports on the prevalence of specific PE-associated ECG abnormalities vary. In a meta-analysis of 45 studies that involved 8209 PE patients, the most prevalent abnormal ECG patterns were sinus tachycardia (38%), TWI V$_{2-4}$ (38%), and STE in aVR (36%). Similarly, sinus tachycardia was the most prevalent ECG pattern in our study. Other similarities and differences between our results and previous studies are as follows. First, the prevalence of RBBB (complete or incomplete) in other studies has ranged from 4.8% to 32.0%. In our study, complete RBBB and incomplete RBBB were present in 7.6% and 7.5% of 1676 patients, respectively. Second, the prevalence of precordial TWI in other studies has ranged from 14% to 59%. In our study, TWI in V$_{2-4}$ was present in 11% of our patients. Third, the prevalence of TWI in inferior leads was as high as 48% of patients in other studies, whereas only 7% of our patients had inferior TWI. Fourth, previous reports had high prevalence of STE in aVR (34%-45%) in patients with more severe PE presentation, including hypotension, need for thrombolysis or adrenergic medication use. Although we found STE in aVR in less PE patients (12%) than previous studies, STE aVR had ORs of 2.82 for 5-day clinical deterioration and 4.16 for abnlRV in our study. We also found a prevalence of 18% for the S1-Q3-T3 pattern, whereas previous studies have reported a prevalence between 8.5% and 33.0%, and as high as 53% in PE patients with clinical shock. Like previous ECG scoring systems, sinus tachycardia and anterior precordial TWI (V$_{2-4}$) were strong predictors of clinical deterioration.

By univariate analysis, mean number of abnormal ECG patterns was significantly greater in those with clinical deterioration and abnl RV by echocardiography versus those without. The mean number of abnormal ECG patterns was lower in those with low-risk classification by any of the PE triaging strategies.

CONCLUSIONS

Several ECG patterns are significantly more frequent in those with clinical deterioration versus those without. SVT was the independent predictor of subsequent clinical deterioration. Sinus tachycardia, STE in aVR, incomplete RBBB, S1-Q3-T3, and TWI V$_{2-4}$ were independent predictors of abnlRV by echocardiography. The absence of any of the abnormal ECG patterns was associated with reduced risk for both clinical deterioration and abnlRV by univariable analysis but not by multivariable analysis. By univariate analyses, higher mean number of abnormal ECG patterns was associated with presence of clinical deterioration, RV abnormality by echocardiography, and PE triaging strategy classification as “not low-risk” versus the groups without these characteristics.

FUNDING INFORMATION

This was supported by the Agency for Healthcare Research and Quality [grant number R01HS025979]. The content is solely the responsibility of the authors and does not necessarily represent the official views of the Agency for Healthcare Research and Quality.

AUTHOR CONTRIBUTIONS

A.J.W. designed the study. J.D.R., A.T., C.C., K.L., and A.J.W. supervised the conduct of the trial and data collection. All PESCOR investigators enrolled patients. N.O. and A.J.W. provided statistical analysis and interpretation of the data. A.J.W. managed the data, including quality control. A.J.W., J.D.R., K.L. drafted the manuscript. All authors contributed substantially to article revision for important intellectual content. A.J.W. takes responsibility for the paper as a whole.

CONFLICT OF INTEREST

All authors report no conflicts of interest.

ORCID

Anthony J. Weekes https://orcid.org/0000-0001-9309-7910
REFERENCES

1. Digby GC, Kukla P, Zhan ZQ, et al. The value of electrocardiographic abnormalities in the prognosis of pulmonary embolism: a consensus paper. Ann Noninvasive Electrocadiol. 2015;20(3):207-223.

2. Shopp JD, Stewart LK, Emmett TW, Kline JA. Findings from 12-lead electrocardiography that predict circulatory shock from pulmonary embolism: systematic review and meta-analysis. Acad Emerg Med. 2015;22(10):1127-1137.

3. Stein PD, Matta F, Sabra MJ, et al. Relation of electrocardiographic changes in pulmonary embolism to right ventricular enlargement. Am J Cardiol. 2013;112(12):1958-1961.

4. Daniel KR, Courteney DM, Kline JA. Assessment of cardiac stress from massive pulmonary embolism with 12-lead ECG. Chest. 2001;120(2):474-481.

5. Janata K, Hocht T, Wenzel C, et al. The role of ST-segment elevation in lead aVR in the risk assessment of patients with acute pulmonary embolism. Clin Res Cardiol. 2012;101(5):329-337.

6. Kostrubiec M, Hrynkiwicz A, Pedowska-Wloszek J, et al. Is it possible to use standard electrocardiography for risk assessment of patients with pulmonary embolism? Kardiol Pol. 2009;67(7):744-750.

7. Kukla P, Dlugopolinski R, Krupa E, et al. The prognostic value of ST-segment elevation in the lead aVR in patients with acute pulmonary embolism. Kardiol Pol. 2011;69(7):649-654.

8. Kukla P, McIntyre WF, Fijorek K, et al. Electrocardiographic abnormalities in patients with acute pulmonary embolism complicated by cardiogenic shock. Am J Emerg Med. 2014;32(6):507-510.

9. Punukollu G, Gowda RM, Vasavada BC, Khan IA. Role of electrocardiography in identifying right ventricular dysfunction in acute pulmonary embolism. Am J Cardiol. 2005;96(3):450-452.

10. Toosi MS, Merlino JD, Leeper KV. Electrocardiographic score and short-term outcomes of acute pulmonary embolism. Am J Cardiol. 2007;100(7):1172-1176.

11. Vanni S, Polidori G, Vergara R, et al. Prognostic value of ECG among patients with acute pulmonary embolism and normal blood pressure. Am J Med. 2009;122(3):257-264.

12. Qaddoura A, Digby GC, Kabali C, Kukla P, Zhan Z-Q, Baranchuk AM. The value of electrocardiography in prognosticating clinical deterioration and mortality in acute pulmonary embolism: a systematic review and meta-analysis [internet]. Clin Cardiol. 2017;40(10):814-824. doi:10.1002/ccd.22742

13. Weekes AJ, Raper JD, Lopez K, et al. Development and validation of a prognostic tool: pulmonary embolism short-term clinical outcomes risk estimation (PE-SCORE). PLoS One. 2021;16(11):e0260036.

14. Weekes AJ, Thacker G, Troha D, et al. Diagnostic accuracy of right ventricular dysfunction markers in normotensive emergency department patients with acute pulmonary embolism. Ann Emerg Med. 2016;68(3):277-291.

15. Rudski LG, Lai WW, Afifalo J, et al. Guidelines for the echocardiographic assessment of the right heart in adults: a report from the American Society of Echocardiography endorsed by the European Association of Echocardiography, a registered branch of the European Society of Cardiology, and the Canadian Society of Echocardiography. J Am Soc Echocardiogr. 2010;23(7):685-713. quiz 786–8.

16. Lang RM, Badano LP, Mor-Avi V, et al. Recommendations for cardiac chamber quantification by echocardiography in adults: an update from the American Society of Echocardiography and the European Association of Cardiovascular Imaging. Eur Heart J Cardiovasc Imaging. 2015;16(3):233-270.

17. Addetia K, Miyoshi T, Citro R, et al. Two-dimensional echocardiographic right ventricular size and systolic function measurements stratified by sex, age, and ethnicity: results of the world Alliance of societies of echocardiography study [internet]. J Am Soc Echocardiogr. 2021;34:1148-1157.e1. doi:10.1016/j.echo.2021.06.013

18. Weekes AJ, Oh L, Thacker G, et al. Interobserver and Intraobserver agreement on qualitative assessments of right ventricular dysfunction with echocardiography in patients with pulmonary embolism. J Ultrasound Med. 2016;35(10):2113-2120.

19. Jimenez D, Auyesky D, Moores L, et al. Simplification of the pulmonary embolism severity index for prognostication in patients with acute symptomatic pulmonary embolism. Arch Intern Med. 2010;170(15):1383-1389.

20. Konstantinides SV, Meyer G, Becattini C, et al. 2019 ESC guidelines for the diagnosis and management of acute pulmonary embolism developed in collaboration with the European Respiratory Society (ERS). Eur Heart J. 2020;41(4):543-603.

21. Bussink BE, Holst AG, Jespersen L, Jensen GB, Prescott E. Right bundle branch block: prevalence, risk factors, and outcome in the general population: results from the Copenhagen City heart study. Eur Heart J. 2013;34(2):138-146.

22. Kashou AH, Basit H, Malik A. ST Segment. StatPearls. StatPearls Publishing: 2021.

23. Choi B-Y, Park D-G. Normalization of negative T-wave on electrocardiography and right ventricular dysfunction in patients with an acute pulmonary embolism. Korean J Intern Med. 2012;27(1):53-59.

24. Harirahan P, Dudzinski DM, Okechukwu I, Takayesu JK, Chang Y, Kabrhel C. Association between electrocardiographic findings, right heart strain, and short-term adverse clinical events in patients with acute pulmonary embolism [internet]. Clin Cardiol. 2015;38(4):236-242. doi:10.1002/ccd.22383

SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

How to cite this article: Weekes AJ, Raper JD, Thomas AM, et al. Electrocardiographic findings associated with early clinical deterioration in acute pulmonary embolism. Acad Emerg Med. 2022;29:1185-1196. doi: 10.1111/acem.14554