Sympathetic activity–associated periodic repolarization dynamics predict mortality following myocardial infarction

Konstantinos D. Rizas, …, Georg Schmidt, Axel Bauer

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**Background.** Enhanced sympathetic activity at the ventricular myocardium can destabilize repolarization, increasing the risk of death. Sympathetic activity is known to cluster in low-frequency bursts; therefore, we hypothesized that sympathetic activity induces periodic low-frequency changes of repolarization. We developed a technique to assess the sympathetic effect on repolarization and identified periodic components in the low-frequency spectral range (≤0.1 Hz), which we termed periodic repolarization dynamics (PRD).

**Methods.** We investigated the physiological properties of PRD in multiple experimental studies, including a swine model of steady-state ventilation ($n = 7$) and human studies involving fixed atrial pacing ($n = 10$), passive head-up tilt testing ($n = 11$), low-intensity exercise testing ($n = 11$), and beta blockade ($n = 10$). We tested the prognostic power of PRD in 908 survivors of acute myocardial infarction (MI). Finally, we tested the predictive values of PRD and T-wave alternans (TWA) in 2,965 patients undergoing clinically indicated exercise testing.

**Results.** PRD was not related to […]

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Sympathetic activity–associated periodic repolarization dynamics predict mortality following myocardial infarction

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Background. Enhanced sympathetic activity at the ventricular myocardium can destabilize repolarization, increasing the risk of death. Sympathetic activity is known to cluster in low-frequency bursts; therefore, we hypothesized that sympathetic activity induces periodic low-frequency changes of repolarization. We developed a technique to assess the sympathetic effect on repolarization and identified periodic components in the low-frequency spectral range (≤0.1 Hz), which we termed periodic repolarization dynamics (PRD).

Methods. We investigated the physiological properties of PRD in multiple experimental studies, including a swine model of steady-state ventilation (n = 7) and human studies involving fixed atrial pacing (n = 10), passive head-up tilt testing (n = 11), low-intensity exercise testing (n = 11), and beta blockade (n = 10). We tested the prognostic power of PRD in 908 survivors of acute myocardial infarction (MI). Finally, we tested the predictive values of PRD and T-wave alternans (TWA) in 2,965 patients undergoing clinically indicated exercise testing.

Results. PRD was not related to underlying respiratory activity (P < 0.001) or heart-rate variability (P = 0.002). Furthermore, PRD was enhanced by activation of the sympathetic nervous system, and pharmacological blockade of sympathetic nervous system activity suppressed PRD (P ≤ 0.005 for both). Increased PRD was the strongest single risk predictor of 5-year total mortality (hazard ratio 4.75, 95% CI 2.94–7.66; P < 0.001) after acute MI. In patients undergoing exercise testing, the predictive value of PRD was strong and complementary to that of TWA.

Conclusion. We have described and identified low-frequency rhythmic modulations of repolarization that are associated with sympathetic activity. Increased PRD can be used as a predictor of mortality in survivors of acute MI and patients undergoing exercise testing.

Trial registration. ClinicalTrials.gov NCT00196274.

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Introduction

Sudden cardiac death (SCD) is the single most common cause of death in the industrialized world (1). A substantial proportion of SCD cases occur in patients after myocardial infarction (MI). Randomized trials have demonstrated that in high-risk patients after MI, mortality can be effectively reduced by prophylactic implantation of a cardioverter-defibrillator (ICD) (2). Consequently, identification of high-risk individuals is a major objective in cardiology. Current guidelines recommend the assessment of left ventricular ejection fraction (LVEF) as the gold standard risk predictor (3, 4); however, this approach lacks both sensitivity and specificity (1, 5). Therefore, development of novel risk markers is of great clinical interest.

Assessment of repolarization instability may more directly estimate the risk of fatal cardiac arrhythmias (6). It is well known from experimental and clinical studies that enhanced sympathetic activity is a key factor leading to the destabilization of myocardial repolarization (7–14). However, without directly recording neural activity, which is impractical in the clinical setting, assessment of the sympathetic effect on myocardial repolarization has not been possible to date. As sympathetic activity is organized in a series of low-frequency bursts (15–19), we postulated that repolarization changes induced by the sympathetic nervous system would exhibit low-frequency periodic features. In the present study, we propose what we believe is a novel way to assess the sympathetic effect on cardiac repolarization. We developed a technology and uncovered periodic components of repolarization in the low-frequency spectral range (≤0.1 Hz), which we termed periodic repolarization dynamics (PRD). The first part of
this article focuses on the physiological properties of PRD, including activation and blockade of the sympathetic nervous system. In the second part of this investigation, we assess the prognostic meaning of enhanced PRD in patients surviving acute MI (post-MI cohort; Figure 1A) and patients undergoing clinically indicated exercise testing (stress-test cohort; Figure 1B). In the stress-test cohort we also tested the prognostic meaning of exercise-induced T-wave alternans (TWA), which is presently considered to be the strongest existing marker of repolarization instability.

Results

Repolarization is subject to low-frequency periodic modulations. We developed a technique to dynamically track repolarization dynamics and to quantify their periodic components. Details of the methodology are reported in Methods. Briefly, we used standard, high-resolution, surface ECG recorded in or converted to the orthogonal Frank lead configuration. As electrocardiographic repolarization is a phenomenon occurring in both space and time, we integrated the spatiotemporal information of each T-wave into a single vector, \( T° \). We used the angle \( dT° \) between successive repolarization vectors as an estimate of the instantaneous repolarization instability (Figure 2, A–C). We observed characteristic low-frequency oscillations in \( dT° \) in health and disease (Figure 2D). In order to quantify these low-frequency \((≤0.1 \text{ Hz})\) periodic patterns, we employed wavelet analysis (Figure 2E).

PRD is not an epiphenomenon of underlying heart rate variability. We tested whether PRD was present in the absence of heart rate variability (HRV). We studied 10 individuals (median age 52 [interquartile range (IQR) 32] years, 5 females), who underwent a clinically indicated electrophysiological (EP) study at our institution. Patient characteristics are provided in Methods. We compared 5-minute episodes of spontaneous sinus rhythm to 5-minute episodes during fixed atrial stimulation, which was set above the spontaneous heart rate. Fixed atrial pacing almost abolished HRV \((P < 0.001;\text{ Supplemental Table 1; supplemental material available online with this article; doi:10.1172/JCI70085DS1)}\), but exerted only minimal, non-significant effects on PRD (ratio of PRD after provocation to PRD before provocation \([\text{PRD ratio}] = 0.75, 95\% \text{ CI } 0.50–1.17, P = 0.193;\text{ Figure 3A, Supplemental Figure 1A, and Supplemental Table 1)}\).

PRD is not an epiphenomenon of underlying respiratory activity. To test whether PRD was present in the absence of spontaneous breathing, we performed an experimental study in a swine model. Seven female domestic pigs were mechanically ventilated and sedated with \(\alpha\)-chloralose, which has been shown to induce only minimal effects on the cardiac autonomic nervous system (20). Respiratory frequency and tidal volume were maintained constant by means of volume-controlled ventilation. Details of the experimental design are provided in Methods. PRD occurred independently of respiratory activity, as illustrated in Figure 4A. There was no interference between respiratory activity and PRD in any animal, as confirmed by spectral and crossspectral analysis (Figure 4, B and C; median coherence 0.044 [IQR 0.026]; \(P < 0.001\) for the difference from the threshold of 0.5).

PRD is enhanced by sympathetic activation and suppressed by sympathetic blockade. We tested the effects of sympathetic activation on PRD in 11 healthy male volunteers (median age 24 [IQR 3] years). Sympathetic activation was achieved by means of head-up tilt testing and low-intensity exercise. Both tilt-table testing (PRD ratio 1.80, 95% CI 1.35–2.58, \(P = 0.005\); Figure 3B, Supplemental Figure 1B, and Supplemental Table 1) and low-intensity exercise (PRD ratio 3.85, 95% CI 2.49–5.61, \(P = 0.001\); Figure 3C, Supplemental Figure 1B, and Supplemental Table 1) led to substantial enhancement of PRD.
Conversely, we tested the effects of antiadrenergic intervention in 10 patients (median age 57 [IQR 21] years, 7 females) undergoing an EP study at our institution. Antiadrenergic intervention was achieved by pharmacological beta blockade. The diagnostic protocol is described in Methods. Beta blockade caused a striking suppression of PRD in all patients (PRD ratio 0.41, 95% CI 0.28–0.61, \( P = 0.002 \); Figure 3D, Supplemental Figure 1C, and Supplemental Table 1).

For comparison, the effects of sympathetic activation and blockade on the low-frequency component of heart-rate variability are shown in Supplemental Table 1.

Increased PRD predicts total and cardiovascular mortality after MI. We tested the prognostic significance of PRD in a cohort of 908 patients from the Autonomic Regulation Trial (median age 61 [IQR 17] years, 174 females) who survived an acute MI (Figure 1A and Table 1) (21, 22). Sixty-nine patients died within the first 5 years of follow-up. Representative resting \( dT° \) signals in a patient who survived the follow-up period and in a patient who suddenly died 8 months after index MI are depicted in Figure 5, A and B, respectively. Although low-frequency oscillations in \( dT° \) were evident in both patients, the amplitudes of PRD were much higher in the nonsurviving patient. The level of PRD was significantly associated with 5-year mortality (6.67 [IQR 8.58] deg\(^2\) vs. 2.66 [IQR 3.93] deg\(^2\); \( P < 0.001 \)). For subsequent survival analyses, we dichotomized PRD at the upper quartile of the study population. The 227 patients with PRD greater than or equal to 5.75 deg\(^2\) (Figure 5C) had a 5-year risk of death of 18.2% compared with 4.1% in the 681 patients with PRD of less than 5.75 deg\(^2\) (\( P < 0.001 \)). Both uni- and multivariable analyses for the prediction of 5-year total mortality indicated that PRD greater than or equal to 5.75 deg\(^2\) was the strongest single risk predictor in the study cohort (Table 2 and Supplemental Figure 2). The predictive value of PRD greater than or equal to 5.75 deg\(^2\) was independent of that of established risk markers, including reduced LVEF of 35% or less (3, 4), the Global Registry of Acute Coronary Events (GRACE) score (23), the presence of diabetes mellitus, elevated mean heart rate, reduced HRV, and increased QT variability index (QTVI) (24). Subsequently, we assessed the incremental prognostic value of PRD to established risk-prediction models (Supplemental Table 2). PRD significantly improved all tested risk-prediction models based on the combination of LVEF, GRACE score, respiratory rate, and HRV parameters. Subgroup analyses revealed that increased PRD was a particularly strong predictor of mortality in the 179 post-MI patients who also suffered from diabetes mellitus, identifying a group of 179 patients with a cumulative 5-year mortality rate of 38.8% (Figure 5D).

Finally, we also investigated whether PRD predicted cardiovascular mortality. Of the 69 deaths, 36 were cardiovascular deaths. As shown in Table 2, PRD greater than or equal to 5.75 deg\(^2\) was also a strong and independent predictor of 5-year cardiovascular mortality.

The predictive value of PRD is complementary to that of exercise-induced TWA. To test the predictive values of PRD and TWA, we studied 2,965 patients (median age 57 [IQR 16] years, 1,187 females) from the Finnish Cardiovascular Study (median age 57 [IQR 16], 1,187 females; Figure 1B and Table 1) who underwent a clinically indicated exercise test (25). During a median follow-up of 6 years, 309 patients died. In all patients, TWA was measured during exercise by the modified moving average (MMA) method (25–27). PRD was assessed in the preexercise period with patients sitting on a bicy-
cle ergometer. As expected, PRD levels in this cohort were higher than those in the post-MI cohort, where PRD was estimated in the supine resting position (9.2 [IQR 13.37] deg² vs. 2.82 [IQR 4.34] deg², respectively; \( P < 0.001 \)). Both PRD and TWA were significantly associated with mortality (8.96 [IQR 13.23] deg² vs. 12.04 [IQR 16.32] deg², \( P < 0.001 \), and 23.00 [IQR 15] μV vs. 26.00 [IQR 17] μV, \( P < 0.001 \), respectively). Univariable Cox regression analysis showed that both markers were strong predictors of total mortality (standardized coefficients 0.203, 95% CI 0.113–0.293, \( P < 0.001 \) for PRD; and 0.256, 95% CI 0.164–0.348, \( P < 0.001 \) for TWA). This remained true on multivariable analysis, which also included age, sex, previous MI, presence of diabetes mellitus, and treatment with beta-blockers (Table 3). The significant crossterm between TWA and PRD (TWA \( \times \) PRD) indicated that the relationship between the outcome and one predictor was dependent on the levels of the other predictor. We therefore tested the additive prognostic value of PRD for different levels of TWA. As illustrated in Supplemental Figure 3, PRD provided incremental prognostic information at all levels of TWA.

Of the 309 deaths, 138 were cardiovascular deaths. Increased preexercise PRD was also a significant predictor of cardiovascular mortality in univariable (standardized coefficient 0.256, 95% CI 0.126–0.385; \( P < 0.001 \)) and multivariable analysis (Table 3).

**Discussion**

In the present study, we identified periodic oscillations of repolarization that were localized in the low-frequency spectral range and were detectable by conventional surface ECG. PRD was evident in health and disease without provocations and occurred autonomously from underlying HRV and respiratory activity. PRD was augmented by physiological provocations leading to activation of the sympathetic nervous system and was suppressed by pharmacological adrenergic blockade. Increased PRD obtained under resting conditions was a very strong predictor of total and cardiovascular mortality in survivors of acute MI and patients undergoing a clinically indicated exercise test. The prognostic value of PRD was incremental to that of established risk markers, including LVEF and TWA.
Noninvasive assessment of the sympathetic effect on myocardial repolarization is of great clinical interest. A wealth of evidence supports the widely held belief that increased sympathetic nervous system activity is associated with increased cardiac vulnerability (11–14). In human subjects, noninvasively measured parameters, including HRV and baroreflex sensitivity, have been employed to study sympathetic activity under routine clinical conditions (28). This approach is based on the principle that activation of the sympathetic nervous system evokes several physiological effects, including increasing systolic contractility rate and vasomotor tone as well as accelerating heart rate and atrioventricular conduction (10). However, these measurements provide only an indirect probe of the sympathetic effect on repolarization; they reflect influences on the sinoatrial node and blood vessels, not on the ventricular myocardium.

At the level of cardiomyocytes, stimulation of β-adrenergic receptors alters intracellular calcium dynamics (28) and shortens action potential duration (10). Importantly, the effect of sympathetic stimulation on the 3 cell types of the ventricular myocardium (epicardial cells, M cells, and endocardial cells) is nonuniform (29, 30). Adrenergic activation abbreviates the action potential duration of epicardial and endocardial cells to a greater degree than the action potential duration of M cells (31), leading to an increased transmural dispersion of repolarization (28).

It is well known that sympathetic activity is organized in series of low-frequency bursts (15–19). We therefore assumed that phasic sympathetic activation induces phasic changes in repolarization localized in the low-frequency spectral range, and we developed a method to track the sympathetic effect on transmural dispersion of repolarization. Our findings confirmed this hypothesis. For what we believe is the first time, we detected periodic changes in repolarization in the same range as those of the sympathetic nervous system. PRD was significantly enhanced by sympathetic activation and was substantially suppressed by sympathetic block-

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**Table 1**

<table>
<thead>
<tr>
<th>Study characteristics</th>
<th>Post-MI cohort</th>
<th>Stress-test cohort</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of patients, n</td>
<td>908</td>
<td>2,965</td>
</tr>
<tr>
<td>Median follow-up (IQR), months</td>
<td>60 (0)</td>
<td>75.1 (47.7)</td>
</tr>
<tr>
<td>Total deaths, n (%)</td>
<td>69 (7.6)</td>
<td>309 (10.4)</td>
</tr>
<tr>
<td>Cardiovascular deaths, n (%)</td>
<td>36 (4.0)</td>
<td>138 (4.7)</td>
</tr>
<tr>
<td>Patient characteristics</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median age (IQR), years</td>
<td>61 (17)</td>
<td>57 (16)</td>
</tr>
<tr>
<td>Females, n (%)</td>
<td>174 (19.2)</td>
<td>1187 (40.0)</td>
</tr>
<tr>
<td>Diabetes mellitus, n (%)</td>
<td>179 (19.7)</td>
<td>344 (11.6)</td>
</tr>
<tr>
<td>History of previous MI, n (%)</td>
<td>86 (9.5)</td>
<td>552 (18.6)</td>
</tr>
<tr>
<td>Median LVEF (IQR), %</td>
<td>53 (15)</td>
<td>66 (16)</td>
</tr>
<tr>
<td>Known CAD (%)</td>
<td>908 (100)</td>
<td>883 (29.8)</td>
</tr>
<tr>
<td>Treatment</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PCI, n (%)</td>
<td>848 (93.4)</td>
<td>NA</td>
</tr>
<tr>
<td>Thrombolysis, n (%)</td>
<td>13 (1.4)</td>
<td>NA</td>
</tr>
<tr>
<td>CABG, n (%)</td>
<td>6 (0.7)</td>
<td>NA</td>
</tr>
<tr>
<td>Beta blockers, n (%)</td>
<td>864 (95.1)</td>
<td>1709 (57.6)</td>
</tr>
</tbody>
</table>

CABG, coronary artery bypass graft; CAD, coronary artery disease; NA, not available; PCI, percutaneous coronary intervention.

In conclusion, PRD constitutes an electrocardiographic phenomenon that most likely reflects the myocardial response to sympathetic activation. Increased PRD is a strong predictor of total and cardiovascular mortality and provided incremental prognostic information to that of exercise-induced TWA. This indicates that PRD can be used to detect high-risk patients who are not identified by TWA. The complementary prognostic information provided by PRD and TWA implies that these 2 markers capture different aspects of repolarization instability. While PRD most probably reflects low-frequency oscillations related to sympathetic activity, TWA is mainly caused by high-frequency action potential oscillations provoked by abnormal calcium handling (34). TWA is an important predictor of cardiovascular mortality, including sudden death (6, 27, 35, 36). However, it needs to be provoked by exercise (37) or invasive procedures (38). Assessment of PRD is inexpensive, easily obtainable under resting conditions, and noninvasive and significantly improves available risk-stratification strategies.

Our study has several limitations. First, high-resolution ECG is required in order to measure PRD. It remains to be demonstrated whether our results are reproducible with lower-resolution tracings. Second, in both cohorts, risk markers were only assessed at enrollment. Therefore, we cannot comment on the immediate and long-term reproducibility of PRD as well as the effect of treatment on PRD. Third, the prognostic value of PRD needs to be validated in independent cohorts. Fourth, as PRD is dependent on the patient’s body position and activity level, the proposed cutoff value is only valid for recordings obtained in the supine resting position. Fifth, we confirmed the prognostic value of PRD for prediction of total mortality and cardiovascular mortality. Although it is plausible to assume that increased levels of PRD are also associated with arrhythmic mortality, this needs to be tested in future studies. Finally, although we have shown that increased PRD is a powerful risk predictor, we have no data to show that specific treatments based on the use of this predictor will improve patient outcome.
of total and cardiovascular mortality, and its use significantly improves established risk-stratification concepts. Future studies are needed to test whether high-risk patients identified by PRD benefit from prophylactic therapies.

Methods
Participants. The physiological properties of PRD were studied in 3 cohorts at the University Hospital of Tübingen. We tested the effects of fixed atrial pacing (atrial-pacing cohort) in 10 individuals (median age 52 [IQR 42] years, 5 females) undergoing a clinically indicated diagnostic EP study. Indications for EP studies were paroxysmal supraventricular tachycardia in 7 patients and evaluation of unexplained syncope in 3 patients.

We investigated the effect of beta blockade in 10 patients (median age 57 [IQR 21] years, 7 females) undergoing an EP study for paroxysmal supraventricular tachycardia (adrenergic-blockade cohort). In both EP studies, all patients were in sinus rhythm, had normal LVEF, were not suspected of suffering from coronary artery disease, and had no significant valve stenosis or insufficiency on echocardiography. We also studied the effects of passive head-up tilt and low-intensity exercise in 11 healthy male volunteers (median age 24 [IQR 3] years, adrenergic-activation cohort).

The prognostic power of PRD was tested in 908 survivors (median age 61 [IQR 17] years, 174 females) of acute MI (post-MI cohort; Figure 1A and Table 1) and 2,965 patients (median age 57 [IQR 16] years, 1,187 females) undergoing clinically indicated exercise testing (stress-test cohort; Figure
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In 908 survivors of acute MI (post-MI cohort)...

**Table 2**

Univariable and multivariable association of risk markers with 5-year all-cause and cardiovascular mortality in 908 survivors of acute MI (post-MI cohort)

<table>
<thead>
<tr>
<th>Risk variable</th>
<th>Univariable Cox regression</th>
<th>Multivariable Cox regression</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Hazard ratio (95% CI)</td>
<td>P value</td>
</tr>
<tr>
<td>LVEF ≤ 35%</td>
<td>3.81 (2.23 to 6.51)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>GRACE* score ≥120</td>
<td>5.54 (3.24 to 9.46)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>2.61 (1.61 to 4.23)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Mean HR &gt; 75 bpm</td>
<td>1.98 (1.11 to 3.55)</td>
<td>0.020</td>
</tr>
<tr>
<td>SDNN ≤ 70 ms</td>
<td>2.01 (1.22 to 3.33)</td>
<td>0.007</td>
</tr>
<tr>
<td>QTc &gt; –0.47</td>
<td>2.54 (1.55 to 4.19)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>PRD ≥ 5.75 deg²</td>
<td>4.75 (2.94 to 7.66)</td>
<td>&lt; 0.001</td>
</tr>
</tbody>
</table>

For the post-MI cohort, we used 30-minute high-resolution (1,600 Hz) digital ECG (TMS; Porti System) recorded in Frank leads configuration. For the pre-exercise period of at least 2.5 minutes was recorded using 12-channel digital ECG (500 Hz), which was converted into the Frank leads configuration by means of the inverse Dower matrix (39).

Assessment of PRD. The technique used to calculate PRD is illustrated in Figure 2 and in Supplemental Figures 4–6. The prerequisite for computing \( dT^° \) is an ECG tracing recorded in or converted to the 3 orthogonal axes X, Y, and Z. The time positions of the T-waves were identified using previously published algorithms (40, 41). The end of each T-wave was set as the reference point (amplitude = 0 mV). The first step in calculating the new parameter was to transform the Cartesian coordinates X, Y, and Z (Supplemental Figure 4A) into a time series of polar coordinates defined by 2 angles (elevation and azimuth) and the resultant-force amplitude XYZ (Supplemental Figure 4B). For example, we selected a time point t1 (Supplemental Figure 4B) and decomposed the XYZ vector into 2 orthogonal vectors on the y axis and the transverse (XZ) plane. The angle between the vector and the y axis was termed the elevation (42) (Supplemental Figure 4D), with an angle of 0° defined as the vector pointing to the causal direction. The angle between the vector on the transverse plane and the x axis was termed the azimuth (42).

On the basis of the 3 new time signals of the polar coordinates, we defined the weight-averaged direction of repolarization, which can be...
described by a set of 2 polar coordinates that we called the weight-averaged azimuth (WAA) and the weight-averaged elevation (WAE). WAA and WAE can be calculated using Equations 1 and 2, respectively. For each time point $t$, the resultant-force amplitude $XYZ$, represented as $\text{Amp}(t)$ in Equations 1 and 2, was multiplied by the corresponding values for azimuth and elevation at the same time point. The $(\text{Amp}(t) \times \text{Angle}(t))$ products were initially summed for the entire duration of the T-wave and thereafter divided by the sum of all resultant-force amplitudes. The result of each equation represents the "with-the-amplitude weighted" average angle, which is measured by means of the same units ($\text{deg rad}$) as the angle in Equations 1 and 2. Using the repolarization wave of Supplemental Figure 4 as a backbone, exemplary WAA and WAE values were calculated as illustrated in Supplemental Figure 5.

$$
\sum_{t=1}^{T} (\text{Amp}_{t} \times \text{Azimuth}_{t}) \\
\sum_{t=1}^{T} \text{Amp}_{t}
$$

(Equation 1)

$$
\sum_{t=1}^{T} (\text{Amp}_{t} \times \text{Elevation}_{t}) \\
\sum_{t=1}^{T} \text{Amp}_{t}
$$

(Equation 2)

We used the angle $dT^{\circ}$ between successive repolarization vectors as an estimate of the instantaneous degree of repolarization instability (Figure 2, A–C). The angle $dT^{\circ}$ is calculated using the dot product (scalar product) equation (43), which by 2 vectors of the same length $r$ can be simplified to Equation 3 as illustrated in Supplemental Figure 5.2. In brief, the MMA algorithm separates odd from even beats. The average morphologies of both the odd and even beats were calculated separately and continuously updated by a weighting factor of 1/8 or 1/32 of the difference between the ongoing average and the new incoming beats. The update was calculated for every incoming beat, resulting in continual moving averages of the odd and even beats. This approach makes the MMA suitable for TWA analysis during the period of activity or during periods of fluctuating heart rates (non–steady state periods) (45). In addition, algorithms were incorporated to reduce the influence of noise and artifacts, such as those caused by pedaling and respiration (46). The TWA values were calculated continuously during the entire exercise test, from rest to recovery, using all precordial leads. Finally, the maximum TWA value at heart rates of less than 125 bpm was derived.

**Assessment of TWA.** We assessed TWA by the time-domain MMA method according to previously established technologies (26) (GE Healthcare version 5.2). In brief, the MMA algorithm separates odd from even beats. The average morphologies of both the odd and even beats were calculated separately and continuously updated by a weighting factor of 1/8 or 1/32 of the difference between the ongoing average and the new incoming beats. The update was calculated for every incoming beat, resulting in continual moving averages of the odd and even beats. This approach makes the MMA suitable for TWA analysis during the period of activity or during periods of fluctuating heart rates (non–steady state periods) (45). In addition, algorithms were incorporated to reduce the influence of noise and artifacts, such as those caused by pedaling and respiration (46). The TWA values were calculated continuously during the entire exercise test, from rest to recovery, using all precordial leads. Finally, the maximum TWA value at heart rates of less than 125 bpm was derived.

**Assessment of other risk predictors.** We assessed LVEF by echocardiography or angiography. We obtained short-term and 24-hour HRV in time and frequency domains as previously proposed (47). Since the standard deviation of all normal-to-normal intervals (SDNN) provided the strongest prognostic power of all HRV measures, we used SDNN as a marker of HRV. We assessed QTc from the resting ECGs according to previously published technological methods (24). We calculated the GRACE score, which combines several clinical risk factors, specifically patient age, history of previous MI and congestive heart failure, ST-segment deviation, elevated cardiac enzymes, renal impairment, systolic blood pressure and heart rate upon admission, and percutaneous coronary interventions during the hospital stay (23).

**Animal study.** Seven female domestic pigs (60–78 kg) were preanesthetized with propofol (2 mg/kg i.v.) and anesthetized with α-chloralose (150 mg/kg i.v. with supplemental doses of 600 mg in 60 ml saline as required), which has been shown to induce only minimal effects on the cardiac autonomic nervous system (20). Immediately after induction of anesthesia, the trachea was cannulated and the lungs were mechanically ventilated with room air. Constant respiratory frequency and tidal volume were maintained by means of volume-controlled ventilation with a fixed tidal volume (6 ml/kg) and a fixed respiratory rate. In each ani-

<table>
<thead>
<tr>
<th>Risk variable</th>
<th>All-cause mortality</th>
<th>Cardiovascular mortality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, continuous</td>
<td>0.055 (0.042 to 0.068)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Sex (yes, no)</td>
<td>-0.535 (-0.790 to -0.280)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Diabetes mellitus (yes, no)</td>
<td>-0.393 (-0.675 to -0.111)</td>
<td>0.006</td>
</tr>
<tr>
<td>Previous MI (yes, no)</td>
<td>0.205 (-0.049 to 0.459)</td>
<td>0.115</td>
</tr>
<tr>
<td>Beta blocker (yes, no)</td>
<td>0.350 (0.085 to 0.614)</td>
<td>0.010</td>
</tr>
<tr>
<td>TWA, continuous</td>
<td>0.175 (0.007 to 0.284)</td>
<td>0.002</td>
</tr>
<tr>
<td>PRD, continuous</td>
<td>0.198 (0.103 to 0.292)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>TWA × PRD</td>
<td>-0.091 (-0.160 to -0.022)</td>
<td>0.010</td>
</tr>
</tbody>
</table>

Beta, standardized coefficients.

\[dT^{\circ} = \cos(\text{WAE}_1) \times \cos(\text{WAA}_1) \times \sin(\text{WAE}_1) \times \sin(\text{WAA}_1)\]

\[+ \cos(\text{WAE}_2) \times \cos(\text{WAE}_2) \times \sin(\text{WAE}_1) \times \sin(\text{WAE}_2) \times \sin(\text{WAA}_2)\]

(Equation 3)
nal, the respiratory rate was set individually (respiratory rate 12–18 per minute corresponding to a frequency of 0.20–0.30 Hz) to maintain normal end-tidal CO₂. Respiratory activity was recorded by a piezoelectric thoracic sensor (48). Two hours after the administration of Propofol, a high-resolution 30-minute ECG (2,048 Hz) was recorded in the Frank leads configuration.

Statistics. We present continuous variables as medians with IQRs. Categorical data are presented as proportions. Results are presented as mean values with 95% CI. Results of physiological and EP studies are presented as the ratio of the value after provocation to the corresponding value before provocation with 95% CI. Differences in the logarithmic ratios were assessed by means of a paired Wilcoxon signed-rank test. To evaluate the effects of respiratory activity on PRD, we estimated the square coherence function between respiratory and dT° signals using the cross-spectral method (49). The coherence function (range 0–1) expresses the linear coupling between 2 signals in the frequency domain (50). A square coherence function greater than 0.5 was considered significant (19, 51, 52). The end points of both prognostic studies were all-cause and cardiovascular mortality. We used the standardized definition of cardiovascular death (53, 54). The median follow-up time in the post-MI cohort was 5 years. PRD was dichotomized at the upper quartile of the study population. For dichotomization of other risk markers, we used established cutoff values of 35% or less for LVEF (55), –0.47 or more for QTVI (24), greater than 75 for mean heart rate (56), 70 ms or less for SDNN (56), and 120 or more for the GRACE score (23). We estimated survival curves using the Kaplan-Meier method. Multivariable analyses were implemented by the adaptation of Cox and multinominal logistic regression models. The latter method was used to calculate integrated discrimination improvement (IDI) scores (57). The median follow-up time in the stress-test cohort was 6 years. The prognostic powers of PRD and TWA were tested with univariable and multivariable Cox regression models. The latter method was used to calculate integrated discrimination improvement (IDI) scores (57). The median follow-up time in the stress-test cohort was 6 years. The prognostic powers of PRD and TWA were tested with univariable and multivariable Cox regression analysis, including age, sex, previous MI, the presence of diabetes mellitus, treatment with beta blockers, and the cross-term between TWA and PRD (TWA x PRD). PRD, TWA, and age were normalized by subtraction of their mean value and division by their SD and were included as scaled factors in the multivariable model. Differences were considered statistically significant when the 2-sided P value was less than 0.05. All statistical analyses were performed using CRAN R, version 2.15.1.

Study approval. The ethics committees of Tübingen, Tampere, and Munich approved the studies performed in the physiological, stress-test, and post-MI cohorts, respectively. Written informed consent was obtained from each participant. The animal protocol was in accordance with the German guidelines for use of living animals and was approved by the local governmental commission for animal research (K 5/10, Regierungssprasium Tübingen, Baden-Wuerttemberg, Germany).

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