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Holkeri, Arttu

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1 Prognostic significance of flat T-waves in the lateral leads in
2 general population

3

4 **First author:**

5 Holkeri

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7 The significance of flat T-waves

8 **Authors:**

9 Arttu Holkeri, MD^a; Antti Eranti, MD^b; M. Anette E. Haukilahti, MD^c; Tuomas Kerola, MD^a;
10 Tuomas V. Kenttä, PhD^c; Kai Nojonen, MSc^d; Tapio Seppänen, PhD^d; Harri Rissanen, MSc^e;
11 Markku Heliövaara, MD^e; Paul Knekt, PHD^e; M. Juhani Junttila, MD^c; Heikki V. Huikuri,
12 MD^c; Aapo L. Aro, MD^f

13 **Affiliations:**

14 ^aDepartment of Internal Medicine, Päijät-Häme Central Hospital, Lahti, Finland. Address:
15 Päijät-Hämeen keskussairaala, Keskussairaalankatu 7, 15850, Lahti, Finland.

16 ^bHeart Center, Central Hospital of North Karelia, Joensuu, Finland. Address: Tikkamäentie
17 16, 80210, Joensuu, Finland.

18 ^cResearch Unit of Internal Medicine, Medical Research Center, Oulu University Hospital and
19 University of Oulu, Oulu, Finland. Address: Faculty of Medicine, PO Box 5000, FI-90014,
20 University of Oulu, Finland.

21 ^dCenter for Machine Vision and Signal Analysis, University of Oulu, Oulu, Finland.
22 Address: Center for Machine Vision and Signal Analysis, PO Box 4500, FI-90014, University
23 of Oulu, Finland.

24 ^eFinnish Institute for Health and Welfare, Helsinki, Finland. Address: PO Box 30, FI-00271,
25 Helsinki, Finland.

26 ^fDivision of Cardiology, Heart and Lung Center, University of Helsinki and Helsinki
27 University Hospital, Helsinki, Finland. Address: Meilahti Tower Hospital, PL 340, 00029
28 HUS, Finland.

29 **Conflict of Interest**

30 None

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34 **Corresponding author:**

35 Arttu Holkeri, arttu.holkeri@helsinki.fi, Department of Internal Medicine, Päijät-Häme
36 Central Hospital, Lahti, Finland. Telephone number: +358381911 Address: Päijät-Hämeen
37 keskussairaala, Keskussairaalankatu 7, 15850, Lahti, Finland.

38 **Keywords:**

39 Electrocardiography, sudden cardiac death, T-wave, repolarization

40 Abstract

41 **Background**

42 Negative T-waves are associated with sudden cardiac death (SCD) risk in the general
43 population. Whether flat T-waves also predict SCD is not known. The aim of the study was to
44 examine the clinical characteristics and risk of SCD in general population subjects with flat T-
45 waves.

46 **Methods**

47 We examined the electrocardiograms of 6750 Finnish general population adults aged ≥ 30
48 years and classified the subjects into 3 groups: 1) negative T-waves with an amplitude ≥ 0.1
49 mV in ≥ 2 of the leads I, II, aVL, V4–V6, 2) negative or positive low amplitude T-waves with
50 an amplitude < 0.1 mV and the ratio of T-wave and R-wave $< 10\%$ in ≥ 2 of the leads I, II,
51 aVL, V4–V6, and 3) normal positive T-waves (not meeting the aforesaid criteria). The
52 association between T-wave classification and SCD was assessed during a 10-year follow-up.

53 **Results**

54 A total of 215 (3.2%) subjects had negative T-waves, 856 (12.7%) flat T-waves, and 5679
55 (84.1%) normal T-waves. Flat T-wave subjects were older and had more often cardiovascular
56 morbidities compared to normal T-wave subjects, while negative T-wave subjects were the
57 oldest and had most often cardiovascular morbidities. After adjusting for multiple factors,
58 both flat T-waves (hazard ratio [HR] 1.81; 95% confidence interval [CI] 1.13–2.91) and
59 negative T-waves (HR 3.27; 95% CI 1.85–5.78) associated with SCD.

60 **Conclusions**

61 Cardiovascular risk factors and disease are common among subjects with flat T-waves, but
62 these minor T-wave abnormalities are also independently associated with increased SCD risk.

63

64 Introduction

65 The T-wave in the electrocardiogram (ECG) coincides with the repolarization of the
66 ventricles. Numerous physiological and pathological factors, e.g. age, sex, autonomic tone,
67 drugs, and the presence of a cardiac disease, may affect the T-wave morphology, and T-wave
68 abnormalities can be used as diagnostic tools in certain acute and chronic cardiac conditions
69 [1]. There has been also growing interest on the long-term prognostic significance of T-wave
70 changes, as there is an ongoing search for novel inexpensive and large-scale applicable
71 prediction tools for sudden cardiac death (SCD). Several T-wave abnormalities, with varying
72 complexity and applicability to clinical practice, have been associated with increased risk of
73 SCD. Negative T-waves are easy to notice in clinical practice and are shown to predict SCD
74 [2]. Other relatively simple T-wave abnormalities associated with SCD risk include wide
75 frontal QRS-T angle and prolonged T_{peak-to-Tend} interval [3–5]. On the other hand, more
76 complex T-wave risk markers, including T-wave alternans, repolarization heterogeneity
77 markers, and three-dimensional T-wave parameters usually require special computer analysis
78 [5–8]. T-waves with low amplitude are easily assessed from the ECG, and they have been
79 associated with cardiovascular mortality in the general population [9–11]. However, whether
80 these flat T-waves are also linked to SCD risk in the general population is not known.
81 Therefore, we investigated the characteristics of flat T-waves in a large Finnish general
82 population cohort, and assessed the risk for SCD associated with these T-wave abnormalities.

83 Methods

84 *Study population*

85 The study population consisted of 7217 participants of the Social Insurance Institution's Mini-
86 Finland Health Survey conducted in 1978–1980. Participants were aged ≥ 30 years and the
87 survey population was a representative sample of the Finnish population. Participants were
88 interviewed regarding their health status, medications, diseases, symptoms, and lifestyle.
89 Furthermore, participants underwent health examinations including measurements of blood
90 pressure, body mass index (BMI), and total serum cholesterol, in addition to the recording of
91 an ECG. Moreover, plasma potassium levels were measured from a prespecified subset of
92 participants. Participants' baseline diagnoses were assessed using structured criteria based on
93 the interviews and the health examination findings (Supplemental Methods). The Mini-
94 Finland Health Survey methods have been reported more extensively previously [12].

95 All participants of the Mini-Finland Health Survey were fully informed about
96 the study, they participated in the study voluntarily, and the use of information for medical
97 research was explained to them. Agreeing to participate in the baseline health examination
98 was taken to indicate informed consent. The participants were free to unconditionally
99 withdraw their consent at any time, in which case their data were deleted. The study was
100 carried out following ethical guidelines and principals of the Declaration of Helsinki. The
101 study protocol and the practice of the subjects' voluntary participation indicating informed
102 consent were approved by the Institutional Review Board (IRB) of National Institute for
103 Health and Welfare (IRB 00007085, Federalwide Assurance (FWA) 00014588).

104

105 *Electrocardiographic measurement and analyses*

106 A standard 12-lead ECG with a paper speed of 50 mm/s and calibration of 1 mV/10 mm was
107 recorded from all study participants during the health examination in 1978–1980. After a few
108 months, a second ECG was recorded from a subgroup of participants who had signs of
109 cardiovascular disease in the health examinations, which were separately evaluated to assess
110 the permanence of flat T-waves as an ECG finding over time. The recorded paper ECGs were
111 digitized and digitally assessed in 2015–2016. Digital analysis was conducted by three
112 examiners with a custom-made ECG analysis software. Using median beats, T-wave
113 amplitude was assessed digitally with respect to the true baseline with 0.01 mV accuracy from
114 each lead. The paper ECG digitizing and digital assessment method has been described in
115 more detail previously [13]. After excluding subjects with missing health examination data,
116 subjects with missing or unreadable ECGs, and subjects with atrial fibrillation, atrial flutter,
117 left or right bundle branch block, second or third-degree atrioventricular block, pre-excitation
118 pattern, pacemaker rhythm, or ECG findings not representing the general population, a total
119 of 6750 subjects remained for the analyses.

120 Subjects were classified into 3 groups according to the T-wave polarities and
121 amplitudes in leads I, II, aVL, V4–V6: 1) negative T-waves (negative T-wave with an
122 amplitude of ≥ 0.1 mV in ≥ 2 of the leads), 2) flat T-waves (positive or negative T-wave with
123 and amplitude < 0.1 mV and the ratio of T-wave and R-wave $< 10\%$ in ≥ 2 of the leads and not
124 meeting the criterion for negative T-waves group), and 3) normal T-waves (not meeting the
125 criteria for negative T-waves or flat T-waves groups).

126 QRS duration, QT interval and Tpeak-to-Tend interval were measured from lead
127 V5. Bazett's formula was used for QT interval correction for heart rate. Frontal QRS axis and
128 T axis were calculated automatically by the digital measurement software. QTc > 450 ms in
129 men and > 460 ms in women, QRS-T angle $> 90^\circ$, and Tpeak-to-Tend > 90 ms were used as

130 cut-offs for abnormal values when the parameters were used as dichotomous categorical
131 variables. ST-segment depressions were defined as negative ST-segments of ≥ 0.1 mV at 60
132 ms from the J point in ≥ 2 of the leads I, II, III, aVL, aVF, and V1–V6.

133

134 *Follow-up*

135 Survey participants were followed from the baseline examination in 1978–1980 until the end
136 of 2011 using the nation wide Causes of Death Register maintained by Statistics Finland.
137 During the complete follow-up time of the Mini-Finland Health Survey, 1077 subjects (27%
138 of all deceased) of all the Mini-Finland participants were autopsied, of which 194 were SCD
139 cases (48% of SCD cases). SCD cases were determined by 2 cardiologists based on the death
140 certificates, hospital records, and autopsy records using the modified Cardiac Arrhythmia
141 Suppression Trial (CAST) criteria [14]. In cases of disagreement, a third cardiologist
142 reviewed the case and made the final classification. The primary endpoint was SCD, and the
143 secondary endpoints were cardiac death and death from any cause. Follow-up time was
144 limited to 10 years in the primary analyses, as the cardiovascular risk profile could change
145 during a longer follow-up. The complete follow-up time of the Mini-Finland Health Survey
146 was used in the secondary analyses.

147

148 *Statistical analysis*

149 Age and sex adjusted mean values \pm standard deviation for continuous variables and the
150 prevalence of categorical variables were compared using the general linear model. Hazard
151 ratios (HRs) and 95% confidence intervals (CIs) were calculated using the Cox proportional
152 hazards model. The multivariate models were adjusted with age, sex, systolic blood pressure,
153 heart rate, total serum cholesterol, BMI, diabetes, active smoking, beta blocker medication,

154 left ventricular hypertrophy (LVH) on ECG based on the Sokolow-Lyon criterion, and
155 presence of cardiac disease. The survival of subjects in T-wave groups was compared using
156 Kaplan–Meier plots. The statistical significances of effect modification by baseline
157 characteristics were tested using the Wald test by entering an interaction term of T-wave class
158 and the respective baseline characteristic. All reported p-values are two-sided and $p < 0.05$ was
159 considered statistically significant. All statistical analyses were performed using IBM SPSS
160 Statistics (version 25).

161 Results

162 A total of 215 (3.2%) subjects presented with negative T-waves, 856 (12.7%) with flat T-
163 waves, and 5679 (84.1%) with normal upright T-waves. Examples of different T-wave
164 morphologies are demonstrated in Figure 1. The distribution of T-wave morphologies in
165 individual leads is presented in the Supplemental Figure.

166 The baseline characteristics of the subjects with different T-wave morphologies
167 are displayed in Table 1. The age distributions of subjects with normal, flat, and negative T-
168 waves are displayed in the Supplemental Table 1. Subjects with flat or negative T-waves were
169 older, had higher blood pressure and heart rate, and had more often diabetes and cardiac
170 morbidities ($p < 0.05$ for all) than subjects with normal T-waves. However, subjects with
171 negative T-waves had higher systolic blood pressure and had more often cardiac disease,
172 coronary artery disease (CAD), diabetes, and beta blocker medication compared to subjects
173 with flat T-waves ($p < 0.05$ for all). Serum potassium was measured from a prespecified group
174 of 2798 subjects (mean 4.5 ± 0.4 mmol/l), with no significant differences between the T-wave
175 groups.

176 The baseline ECG features are shown in Table 2. QRS and Tpeak-to-Tend
177 intervals were longer in subjects with negative T-waves than subjects with either normal or
178 flat T-waves ($p < 0.05$ for both). LVH and ST-segment depressions were most common among
179 negative T-wave subjects while being more common among flat T-wave subjects than normal
180 T-wave subjects ($p < 0.05$ for all). The frontal QRS-T angle was narrowest among normal T-
181 wave subjects and widest in negative T-wave subjects ($p < 0.05$). QTc was longest in subjects
182 with flat T-waves and shortest in subjects with negative T-waves ($p < 0.05$).

183 A second ECG was recorded few months after the baseline examinations from
184 247 subjects with flat T-waves (28.9% of the subjects with flat T-waves). Flat T-waves were

185 again observed in 149 (60.3%) of the subjects, while 24 subjects (9.7%) had negative T-
186 waves, and 3 (1.2%) subjects had LBBB in the repeat ECG.

187 During the 10-year follow up, 131 subjects (60.9%) with negative T-waves, 267
188 subjects (31.2%) with flat T-waves, and 547 subjects (9.6%) with normal T-waves died. Of
189 these, 25 (19.1%) in negative T-wave subjects, 32 (12.0 %) in flat T-wave subjects, and 60
190 (11.0%) in normal T-wave subjects were SCDs. The survival curves for SCD and all-cause
191 mortality according to the T-wave group are presented in Figure 2. Both flat T-waves and
192 negative T-waves associated with SCD, cardiac death, and all-cause mortality when compared
193 to subjects with normal T-waves. Subjects with negative T-waves had worse prognosis
194 compared to subjects with flat T-waves. After multiple adjustments, flat T-waves subjects had
195 HR 1.81 (95% CI 1.13–2.91) and negative T-wave subjects had HR 3.27 (95% CI 1.85–5.78)
196 for SCD, compared to subjects with normal T-waves. The prognoses associated with
197 individual T-wave abnormalities are displayed in Table 3.

198 No significant effect modification was found between T-wave class and sex, age
199 ≤ 50 or >50 years, BMI ≤ 25 or >25 kg/m², heart rate ≤ 70 or >70 bpm, presence of LVH,
200 presence of hypertension diagnosis, or presence of cardiac disease diagnosis for SCD risk.

201 When ECG repolarization parameters wide QRS-T angle, prolonged QTc, and
202 prolonged Tpeak-to-Tend interval were used as dichotomous categorical variables and entered
203 simultaneously with ST-segment depressions and T-wave class into an age and sex adjusted
204 model, only ST-segment depressions and T-wave class associated with increased SCD risk,
205 with both flat T-waves and negative T-waves associating with risk of SCD (Supplemental
206 Table 2). When further adjusted with multiple factors, only T-wave class associated with SCD
207 risk, with both flat T-waves and negative T-waves remaining associated with risk of SCD
208 ($p < 0.05$).

209 In the secondary analyses using the complete follow-up of the Mini-Finland
210 Health Survey (mean 24.5±10.3 years), flat T-waves and negative T-waves similarly
211 associated with all endpoints, albeit not as strongly as in the primary analyses. During the
212 complete follow-up, flat T-waves had HR 1.40 (95% CI 1.05–1.85) for SCD when compared
213 to normal T-wave subjects after multivariate adjustments. The risks of endpoints associated
214 flat T-waves and negative T-waves during the complete follow-up are presented in the
215 Supplemental Table 3.

216 Discussion

217 In this large, prospective study, subjects with flat T-waves were more likely to have
218 cardiovascular risk factors and cardiac disease compared to subjects with normal T-waves.
219 Furthermore, flat T-waves independently associated with increased risk of SCD, in addition to
220 cardiac death, and death from any cause. Still, cardiovascular morbidities were most common
221 and the prognosis poorest among subjects with negative T-waves.

222 Normally, T-waves are upright in most of the ECG leads, and negative T-waves
223 in leads I, II, V3–V6 are considered abnormal [1]. Multiple factors can have an effect on T-
224 waves polarity and amplitude, for example age, sex, heart rate, autonomic nervous system,
225 electrolyte disturbances, and drugs [1]. Furthermore, T-wave changes have been associated
226 with cardiovascular conditions, e.g. hypertension, acute and chronic manifestations of CAD,
227 left ventricular hypertrophy, and cardiomyopathies [1].

228 Since the T-wave corresponds to the vulnerable repolarization period of the
229 cardiac cycle, multiple T-wave parameters have been studied and linked with increased risk
230 for arrhythmic death. T-wave inversions, abnormal T-wave axis, and wide QRS-T angle have
231 been associated with SCD in the general population [2,3,5]. Furthermore, prolonged Tpeak-
232 to-Tend interval has been associated with SCD risk in the general population, although not all
233 studies have had similar findings [4,5]. In addition, several complex T-wave parameters,
234 including T-wave alternans, repolarization heterogeneity markers, and spatial T-wave
235 parameters, have been shown to associate with increased SCD risk, but efficient analysis of
236 these parameters requires special computer software [5–8].

237 Low amplitude or flat T-waves have been associated with sudden cardiac arrest
238 in subjects with hypertrophic cardiomyopathy [15], and they are also more prevalent among
239 subjects with early repolarization who will suffer ventricular fibrillation, compared to subjects
240 with early repolarization with benign prognosis [16]. Low amplitude T-waves have been

241 linked also to ventricular tachycardia and ventricular fibrillation risk in some patients with
242 implantable cardioverter defibrillator [17]. Moreover, when flat T-waves were classified
243 together with negative T-waves in middle-aged men, this group of subjects was at increased
244 risk for SCD, compared to subjects with normal positive T-waves [18]. The present study is
245 the first to demonstrate that flat T-waves independently associate with increased SCD risk in
246 the general population.

247 After relevant exclusions, the prevalence of flat T-waves was 13% in the Mini-
248 Finland Health Survey cohort of the present study. The definition used in the present study
249 included both negative and positive T-waves with an amplitude <0.1 mV and the ratio of T-
250 wave and R-wave $<10\%$ as flat T-waves. The amplitude cut-off of ≥ 0.1 mV has been
251 commonly used for negative T-waves, as more minor negative T-waves can be difficult to be
252 definitely determined as negative in comparison to isoelectric or low amplitude biphasic or
253 positive T-wave [1,2]. Previous studies have commonly used the Minnesota Code definitions
254 to assess minor T-wave abnormalities, with prevalences ranging from 2% to 13% [9–11].
255 Although there are small differences between the definitions used in the present and the
256 previous studies, flat T-waves seem to be a relatively common finding in the general
257 population.

258 In the present study, abnormal T-waves were more prevalent among older
259 subjects. However, no significant effect modification was noted between the T-wave class and
260 age group, suggesting that age does not affect the prognosis associated with abnormal T-
261 waves. Subjects with flat T-waves also had more often cardiovascular morbidities than
262 subjects with normal T-waves. Nonetheless, flat T-waves predicted SCD, cardiac mortality
263 and all-cause mortality also independently from cardiovascular risk factors and diseases.
264 Furthermore, no significant effect modification between the T-wave class and presence of

265 diagnosed cardiac disease was observed. Finally, negative T-wave subjects had the highest
266 prevalence of cardiovascular morbidities and even worse prognosis.

267 T-wave morphology changes are caused by local or diffuse voltage gradient and
268 spatial changes in the ventricular repolarization [1]. T-wave changes after myocardial
269 infarction are assumed to be caused by nonradial potential gradients or decreased transmural
270 conduction velocity [19]. Other plausible mechanisms causing T-wave changes in heart
271 diseases include alterations in the autonomic nervous system and ventricular myocyte
272 hypertrophy [1]. Accordingly, it could be speculated that flat T-waves may in some cases act
273 as markers of underlying electrical or structural cardiac pathology, that may be less severe
274 than in subjects with negative T-waves. Moreover, myocardial fibrosis is also associated with
275 T-wave inversion and T-wave changes among SCD victims, especially in subjects with
276 ischemic heart disease [20]. Non-specific ST-T abnormalities have been also linked to
277 impaired left ventricular relaxation causing abnormal repolarization, with cardiac fibrosis
278 being a possible underlying mechanism [21]. Consequently, flat T-waves could also be a
279 direct sign of increased vulnerability for fatal ventricular arrhythmias.

280 We compared the T-wave classification to other easily assessable ECG
281 repolarization markers that have been associated with SCD. Some of the other analyzed
282 markers have independent clinical value, e.g. ST-segment depression may indicate an
283 underlying CAD. ST-segment depressions were most common in subjects with negative T-
284 waves, but were rarely observed in subjects with normal T-waves. This was in concordance
285 with the prevalence of CAD diagnosis in the health examinations, with the highest prevalence
286 of CAD observed in subjects with negative and lowest prevalence in subjects with normal T-
287 waves. Prolonged QTc, a known marker of increased risk of SCD, was notably longer among
288 subjects with flat T-waves. Flat T-waves are known to decrease the certainty of determining
289 the end of T-wave, which may explain the longer QT intervals [22]. Nonetheless, when the

290 repolarization markers were analyzed simultaneously in a multivariate model, the T-wave
291 classification was the only one that remained associated with SCD risk. This could indicate,
292 that the simple analysis of the polarities and amplitudes of the T-waves could relay major
293 information about the association between repolarization and SCD risk in clinical practice.
294

295 *Limitations*

296 Due to the inclusion criterion of age ≥ 30 years, these findings may not be directly applicable
297 to younger adult populations. As a further limitation, although the baseline examinations were
298 extensive, an echocardiographic study was not performed to survey participants, and,
299 consequently, no data was available on the cardiac structure or systolic function of the heart.
300 Moreover, although flat T-waves were a stable finding in the majority of the cases, in some
301 cases flat T-waves seems to be observable only in some of the ECG recordings. Finally, the
302 results may not be directly applicable to modern populations, as diagnostic tests and
303 treatments of cardiovascular disease have improved since the baseline examinations of the
304 Mini-Finland Health Survey in 1978–1980. However, this limitation is inevitable in this kind
305 of cohort studies with long follow-up periods.
306

307 *Conclusions*

308 Flat T-waves are a relatively common finding in the general population, but they are often a
309 sign of underlying cardiac disease. Furthermore, flat T-waves are independently associated
310 with increased SCD risk, although the prognosis is better than observed with negative T-
311 waves. More focus should be placed on these minor T-wave abnormalities in the future, as
312 individuals with these ECG patterns may benefit from a careful clinical evaluation and closer
313 monitoring.

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- 397
- 398

399 **Figure legends**

400 Figure 1 legend

401 Demonstration of different T-wave morphologies. A) A normal T-wave, B) a flat T-wave, and
402 C) a negative T-wave.

403

404 Figure 2

405 Figure 2 legend

406 Survival curves for A) SCD and B) all-cause mortality according to the T-wave morphology.

407 The Y-axis of A) graph is scale broken to better illustrate the SCD survival curves. Survival
408 curves were compared with Logrank test.

409 SCD = sudden cardiac death.

410

411 Tables

412 Table 1

413 Baseline characteristics

	Normal T-waves n=5679 (84.1%)	Flat T-waves n=856 (12.7%)	Negative T-waves n=215 (3.2%)	Flat vs normal T-waves p-value	Negative vs normal T- waves p-value	Flat vs negative T-waves p-value
Male sex (%) †	2656 (46.8%)	312 (37.5%)	84 (39.1%)	0.002	ns	ns
Age (yr) ‡	48.9±13.0	60.6±12.8	68.4±10.5	<0.001	<0.001	<0.001
Systolic blood pressure (mmHg) §	140±21	156±25	167±29	<0.001	<0.001	0.02
Diastolic blood pressure (mmHg) §	86±11	91±12	90±13	<0.001	0.03	ns
Hypertension (%) §	2995 (52.7%)	698 (81.5%)	204 (94.9%)	<0.001	<0.001	ns
Heart rate (bpm) §	67±13	73±15	72±14	<0.01	0.01	ns
Total serum cholesterol (mmol/l) §	6.9±1.4	7.3±1.4	7.4±1.6	ns	ns	ns
Body mass index (kg/m ²) §	25.6±3.9	27.7±4.7	26.6±4.3	<0.001	ns	<0.001
Cardiac disease (%) §	635 (11.2%)	341 (39.8%)	174 (80.9%)	<0.001	<0.001	<0.001
Coronary artery disease (%) §	380 (6.7%)	219 (25.6%)	103 (47.9%)	<0.001	<0.001	<0.001
Diabetes (%) §	164 (2.9%)	128 (15.0%)	60 (27.9%)	<0.001	<0.001	<0.001
Smoking (%) §	1348 (23.7%)	145 (16.9%)	36 (16.7%)	ns	ns	ns
β-blocker medication (%) §	305 (5.4%)	93 (10.9%)	50 (23.3%)	<0.001	<0.001	<0.001

414

415 Continuous variables are presented as mean values \pm standard deviation and categorical variables as prevalences. Hypertension = systolic blood
416 pressure >140 mmHg, diastolic blood pressure >90 , or diuretic, beta-blocker or other hypertensive drug therapy.

417 Between group comparisons were: † adjusted for age, ‡ adjusted for sex, and § adjusted for age and sex.

418

419 *Table 2*

420 Electrocardiographic features

	Normal		Negative		Negative vs	
	T-waves		T-waves	Flat vs	normal T-	Flat vs
	n=5679	Flat T-waves	n=215	normal T-waves	waves	negative T-waves
	(84.1%)	n=856 (12.7%)	(3.2%)	p-value	p-value	p-value
QRS duration (ms)	84±11	84±12	88±15	ns	<0.001	<0.001
QTc (ms)	406±25	428±39	392±41	<0.001	<0.001	<0.001
LVH (%)	681 (12.0%)	147 (17.1%)	100 (46.8%)	0.006	<0.001	<0.001
Frontal QRS axis (degrees)	29±39	11±36	11±38	<0.001	ns	0.05
Frontal T-wave axis (degrees)	24±25	15±75	1±136	<0.001	<0.001	<0.001
Frontal QRS-T angle (degrees)	34±28	73±48	140±36	<0.001	<0.001	<0.001
Tpeak-to-Tend (ms)	80±13	71±22	86±34	<0.001	<0.001	<0.001
ST-segment depressions (%)	105	103 (12.0%)	104	<0.001	<0.001	<0.001

(1.8%)

(48.4%)

421

422 Continuous variables are presented as mean values \pm standard deviation and categorical variables as prevalences. Ns = not significant. LVH =
423 Left ventricular hypertrophy based on Sokolow-Lyon ECG criterion. QRS duration, QTc, and Tpeak-to-Tend were assessed from lead V5. ST-
424 segment depressions = negative ST-segment of ≥ 0.1 mV at 60 ms from the J point in ≥ 2 of the following leads: I, II, III, aVL, aVF, and V1–V6.
425 Between group comparisons were adjusted for age and sex.

426

427 *Table 3*

428 Prognostic significance of flat T-waves and negative T-waves during the 10-year follow-up

	Normal T-waves n=5679 (84.1%)	Flat T-waves n=856 (12.7%)	p-value	Negative T-waves n=215 (3.2%)	p-value
SCD					
No. of SCDs (% of subjects)	60 (1.0%)	32 (3.7%)		25 (11.6%)	
Age and sex adjusted HR (95% CI)	1	2.55 (1.62–4.00)	<0.001	6.50 (3.87–10.91)	<0.001
Multivariate adjusted HR (95% CI)	1	1.81 (1.13–2.91)	0.014	3.27 (1.85–5.78)	<0.001
Cardiac death					
No. of cardiac deaths (% of subjects)	194 (3.4%)	116 (13.6%)		79 (36.7%)	
Age and sex adjusted HR (95% CI)	1	2.25 (1.78–2.86)	<0.001	4.51 (3.40–5.98)	<0.001

Multivariate adjusted					
HR (95% CI)	1	1.57 (1.22–2.02)	<0.001	2.29 (1.68–3.11)	<0.001
Death from any cause					
No. of deaths					
(% of subjects)	547 (9.6%)	267 (31.2%)		131 (60.9%)	
Age and sex adjusted					
HR (95% CI)	1	1.87 (1.61–2.18)	<0.001	2.79 (2.28–3.41)	<0.001
Multivariate adjusted					
HR (95% CI)	1	1.54 (1.31–1.80)	<0.001	1.85 (1.48–2.30)	<0.001

429

430 M

431 u

432 l

433 t

434 i

435 v

436 27

437 r

438 i

439 a