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Impact of age and sex on the long-term prognosis associated with early repolarization in the general population

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25 **Declarations of interest:**

None 26

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31 Abstract

32 Background

33 Early repolarization (ER) has been linked to the risk of sudden cardiac death (SCD) in the general

34 population, although controversy remains regarding risks across various subgroups.

35 *Objective*

36 We investigated whether age and sex influence the prognostic significance of ER.

37 Methods

38 We evaluated the 12-lead electrocardiograms of 6631 Finnish general population subjects aged \geq 30

39 years (mean age 50.1 \pm 13.9 years, 44.5% men) for the presence of ER (J-point elevation \ge 0.1 mV

40 in \geq 2 inferior/lateral leads), following them for 24.4 ± 10.3 years. We analyzed the association

41 between ER and the risk of SCD, cardiac death, and all-cause mortality in subgroups according to

42 age (<50 or \geq 50 years) and sex.

43 Results

ER was present in 367 of the 3305 subjects under 50 and in 426 of 3326 subjects aged \geq 50 years. ER was not associated with any of the endpoints in the entire study population. After adjusting for clinical factors, ER was associated with SCD (hazard ratio [HR] 1.88; 95% confidence interval [CI] 1.16–3.07) in subjects under 50, but not in older subjects (interaction between ER and age group, *P* = .048). Among the younger subgroup, women with ER had a high risk of SCD (HR 4.11; 95% CI 1.41–12.03), whereas among men ER was not associated with SCD. Finally, ER was not associated with cardiac mortality or all-cause mortality in either age group.

51 Conclusion

52 ER associates with SCD in subjects younger than 50 years, particularly in women, but not in
53 subjects 50 years and older.

55 Keywords

56 Electrocardiography; Sudden cardiac death; Early repolarization; Epidemiology; Age groups

ound

57 Introduction

An early repolarization (ER) pattern was previously considered a benign electrocardiogram (ECG) 58 59 pattern, until it was shown to be associated with idiopathic ventricular fibrillation in three separate case-control studies in 2008.^{1–3} Subsequently, researchers found that ER was also associated with 60 all-cause mortality, cardiac death, and sudden cardiac death (SCD) in the general population.^{4–8} 61 However, some studies found no link between ER and adverse events.⁷ Consequently, researchers 62 attempted to distinguish benign ER patterns from patterns that associate with more unfavorable 63 prognoses.^{5,9–12} Furthermore, other studies examined whether the prognosis associated with ER 64 varies across different patient subgroups.^{6,9,11} In some studies ER was associated with cardiac 65 66 mortality, particularly among younger middle-aged subjects, whereas in studies among older subjects ER was not associated with an excess risk.^{6,13} In young adult populations, however, ER is a 67 prevalent finding and considered a benign phenomenon.^{14–16} Whether age affects the risk of SCD 68 69 associated with ER in adult subjects remains unclear. 70 Here, we present our investigation of the association between ER and SCD, cardiac mortality, and all-cause mortality in a Finnish general population cohort and examine whether this 71 association differs between subjects younger than 50 years old and those \geq 50 years. Furthermore, 72

73 we assess whether sex impacted the risk associated with ER in these age groups.

74 Methods

75 Study population

The study population consisted of participants of the Mini-Finland Health Survey, a representative 76 77 sample of the Finnish population, conducted in 1978–1980. The survey consisted of health 78 interviews regarding the subjects' health status, diseases, medications, symptoms, and lifestyle, 79 together with health examinations that measured blood pressure, body mass index, and serum 80 cholesterol, and included an electrocardiogram (ECG). In total, 8000 subjects aged \geq 30 years were 81 invited to take part, among whom 7217 participated in the health examination. The extensive survey methods are reported elsewhere.¹⁷ In total, 17 survey participants in this study also participated in a 82 previous cohort study by Tikkanen et al.⁴ 83

84

85 Electrocardiographic measurement and analyses

A standard 12-lead ECG was recorded with a paper speed of 50 mm/s for all study subjects during the health examinations conducted in 1978–1980, and subsequently stored for later assessment. The presence of an ER pattern was assessed manually from the original paper ECGs by three physicians in 2016–2018, with assistance from a cardiologist when needed.

90 An ER pattern was defined and assessed based on a slightly modified version of the recommendations from a published consensus paper.¹⁸ Briefly, we defined an ER pattern as an end-91 92 QRS notch or slur on the downward slope of the prominent R-wave at the J-point, with an 93 amplitude of ≥ 0.1 mV measured with respect to the true baseline determined as the T–P segment. The presence of a pathological Q-wave in the lead with an end-QRS notch or slur was considered a 94 possible peri-infarction block and not classified as an ER pattern.¹⁹ A subject's ECG was 95 considered positive for ER if an ER pattern was present in either ≥ 2 of the inferior (II, III, or aVF) 96 or ≥ 2 of the lateral (I, aVL, V4, V5, or V6) leads. An ER amplitude was classified as ≥ 0.1 mV, but 97

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98	<0.2 mV or ≥ 0.2 mV. Each ECG positive for ER was classified according to the configuration of
99	the ER patterns as a slur, notch, or undetermined (no predominant form). The ST-segment was
100	regarded horizontal or descending if the amplitude of the ST-segment 100 ms after the J-point was
101	less than or equal to the amplitude at the J-point end, and ascending if the amplitude was greater
102	than the amplitude at the J-point end. ¹⁸ An ECG was classified as a low amplitude T-wave if any T-
103	wave in leads I, II, or V4–V6 was inverted, biphasic, or had an amplitude $\leq 0.1 \text{ mV}$ and $\leq 10\%$ of the
104	R-wave amplitude in the same lead. ¹²
105	We excluded subjects ($n = 248$) with missing or unreadable ECGs and subjects ($n = 248$)

bundle branch block, left anterior or posterior fascicular block, QRS duration >110 ms, a pacemaker rhythm, or rare ECG findings not representing the general population. We also excluded subjects (n = 7) with missing data.

331) with II/III-degree atrioventricular block, ventricular pre-excitation, complete or incomplete

110

106

111 Follow-up

Subjects were followed from the baseline examinations in 1978–1980 until the end of 2011 using the Causes of Death Register maintained by Statistics Finland. SCDs likely caused by terminal arrhythmias were determined by two cardiologists. These cardiologists reviewed the data for all deaths from cardiovascular causes from death, hospital, and autopsy records using the SCD definitions based on the modified Cardiac Arrhythmia Suppression Trial (CAST) criteria.²⁰ In cases of disagreement, a third cardiologist reviewed and classified the case. The primary endpoint was SCD, and the secondary endpoints were cardiac death and death from any cause.

119 The Mini-Finland Health Survey preceded the current legislation on ethics in medical 120 research. All participants were fully informed about the survey and its implications, participated in 121 the study voluntarily, and were advised that their information would be used for medical research. 122 Agreeing to participate in the baseline health examination was taken to indicate their informed

- 123 consent. Record linkage with national health registers to the survey data was approved by the
- 124 register authorities.
- 125

126 Statistical analysis

127 Continuous data are presented as the mean \pm standard deviation, while categorical data appear as 128 the number of cases and prevalence in the study population in parentheses. We used the general linear model to compare the age- and sex-adjusted mean values for continuous variables, and the 129 130 prevalence of categorical variables in cross-sectional baseline data. Hazard ratios (HRs), 95% 131 confidence intervals (95% CIs), and P values were calculated using the Cox proportional hazards 132 model. We tested the assumption for proportional hazards for each covariate in the final Cox 133 regression model. Age, sex, systolic blood pressure, total serum cholesterol, smoking, diabetes, and 134 coronary artery disease (CAD) were used as covariates in the multivariate models. The statistical 135 significance of the effect modification by age group (subjects aged <50 years and ≥ 50 years, 136 respectively) and sex were tested using the Wald test by entering an interaction term for ER and age 137 group, and ER and sex, respectively. We considered P < .05 as statistically significant. All 138 statistical analyses were performed using IBM SPSS Statistics (version 24) and R (version 3.6.1, 139 https://www.r-project.org/).

140 Results

141 Baseline characteristics of subjects

142 Table 1 summarizes the baseline characteristics. ER was slightly more prevalent among subjects 143 aged \geq 50 years compared to subjects under 50 years (12.8% vs 11.1%; P = .033). Subjects with ER 144 were more likely male than subjects without ER among subjects aged <50 (68.1% vs 45.7%; P <145 .001). Yet, we found no significant sex difference in subjects \geq 50 years. Subjects under 50 with ER 146 had a lower systolic blood pressure, a lower heart rate, and a shorter QRS duration and QTc interval 147 compared to subjects without ER after adjusting for age and sex. Subjects \geq 50 years with ER had a lower heart rate and were less likely to have diabetes, but were more likely to take beta blocker 148 149 medication compared to subjects without ER after adjusting for age and sex.

150

151 Impact of age and sex on ER prognosis

152 Among 3305 subjects under 50, 748 (22.6%) died during a mean follow-up of 30.2 ± 6.4 years,

among whom 237 died from cardiac causes (31.7% of all deaths), and 95 from SCD (12.7% of all deaths). Among those \geq 50 years old, 2819 of 3326 subjects (84.8%) died during a mean follow-up of 18.7 ± 10.2 years. Among those who died, 1283 deaths resulted from cardiac causes (45.5% of all deaths) and 251 from SCD (8.9% of all deaths).

157 Across the entire study population, ER was not associated with any of the endpoints

158 (see Supplemental Material). Furthermore, from the different ER patterns, only ER with a low

amplitude T-wave (n=158 [19.9% of ER subjects], multivariate-adjusted HR 1.75; 95% CI 1.06–

160 2.87; P = .027) was associated with SCD in the entire study population when compared to subjects

161 without ER (see Supplemental Material). Table 2 shows the risk for SCD and the secondary

- 162 endpoints associated with ER in the age subgroups, and the interaction between ER and age group.
- 163 ER was not associated with cardiac death or all-cause mortality in either age group. During the

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164	follow-up period, 5.7% of subjects with ER and 2.5% of subjects without ER under 50 suffered an
165	SCD, compared to 7.7% with and 7.5% without ER, respectively, \geq 50 years old. We detected a
166	significant interaction between ER and the age group in SCD (multivariate-adjusted $P = .048$). In
167	addition, ER was associated with an increased risk of SCD in subjects under 50 in the multivariate
168	analysis (HR 1.88; 95% CI 1.16–3.07; $P = .011$), whereas among subjects ≥ 50 years ER was not
169	associated with an increased SCD risk. Figure 1 provides the survival plots according to age group
170	for SCD adjusted for confounders.

171 Among subjects under 50, we detected a significant interaction between ER and sex in 172 SCD after adjusting for age (P = .024), which did not remain significant in the multivariate analysis 173 (P = .092). When women under 50 were analyzed separately, ER was associated with a high risk of 174 SCD in both the age-adjusted (HR 5.34; 95% CI 1.88–15.19; P = .002) and multivariate-adjusted (HR 4.11; 95% CI 1.41–12.03; P = .010) analyses when compared to subjects without ER. Figure 2 175 provides an example ER pattern from a woman under 50 years old. In comparison, ER was not 176 177 associated with SCD among men under 50. Neither men nor women under 50 with ER exhibited an 178 increased risk for cardiac death or all-cause mortality.

179

180 Risk of SCD based on the ER pattern in subjects under 50

181 Table 3 summarizes the risks of SCD based on the ER pattern among subjects under 50 in the

182 multivariate analyses. When assessed by ER localization, both inferior (HR 1.92; 95% CI 1.04–

183 3.56; P = .038) and lateral (HR 2.08; 95% CI 1.10–3.95; P = .024) ER localizations were associated

- 184 with SCD risk among subjects under 50. Furthermore, a slurred ER (HR 2.09; 95% CI 1.19–3.67; P
- 185 = .010), ER with a horizontal or descending ST-segment (HR 3.12; 95% CI 1.56–6.26; P = .001),
- and ER with a low amplitude T-wave (HR 4.47; 95% CI 1.75–11.42; P = .002) were associated
- 187 with SCD risk among subjects under 50 years old.

188 Discussion

189	We evaluated the prognosis associated with ER based on sex and age groups in a large
190	representative population cohort with a long follow-up period. We found that ER was associated
191	with an increased risk of SCD among adults aged 30-50 years, whereas no increased SCD risk was
192	observed among subjects with ER aged \geq 50 years. Furthermore, among subjects under 50, women
193	with ER exhibited a high SCD risk, whereas ER was not associated with SCD among men.
194	In this study, ER prevalence reached 12.0% in the entire study population. In previous
195	studies, the prevalence of ER ranged from 0.9% to 23.9%. ^{7,8,21} We defined ER following minor
196	adjustments as recommended in a recent consensus paper. We measured the ER amplitude with
197	respect to the true baseline determined as the T-P segment, compared to with respect to the QRS
198	onset suggested by the consensus paper. This difference could have had an effect on the ER
199	amplitude measurements, especially on tachycardic subjects. The ER definition used in the present
200	study is similar albeit somewhat modified to that used in a previous Finnish middle-aged general
201	population cohort study in which ER prevalence was 5.8%. ^{4,10,18} A possible explanation for the
202	difference in the ER prevalence between these studies may lie in the improved ECG quality in the
203	present study given the more modern recording device, as borderline cases would be determined ER
204	positive in the present study and negative in the previous study. Concordant with previous studies,
205	ER was more prevalent among men younger than 50, whereas no sex difference was identified
206	among older subjects. ¹⁶ One possible explanation for this may lie in the association between ER and
207	testosterone levels in men, which begin declining before the age of 50. ²²
208	Previously, few studies examined ER prognosis in different age groups. In a German
209	cohort study, ER was associated with all-cause and cardiac mortality among subjects aged 35-54,
210	while ER was not associated with an adverse prognosis in older age groups. ⁶ Similarly, in a
211	Japanese cohort study, subjects aged <60 years with ER exhibited an increased risk for cardiac

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death, while subjects with ER \geq 60 years carried no such increased risk.¹³ However, in young adults 212 213 aged 18-30 in the United States, ER with an ascending ST-segment was not associated with adverse outcomes, and the prevalence of ER markedly decreased during the follow-up period.¹⁴ 214 215 Interestingly, the prevalence of ER with an ascending ST-segment appears to change in male and 216 female subjects throughout puberty, while the prevalence of ER with a horizontal or descending STsegment remains fairly constant among children, adolescents, and middle-aged subjects.^{10,15,23} 217 218 To our knowledge, no previous studies examined the impact of age on SCD risk related 219 to ER. In our study, subjects aged 30–50 years exhibited an increased risk of SCD, while ≥50-year-220 old subjects with ER showed no increase in SCD risk. Previous studies demonstrated that ER may predispose an individual to a fatal arrhythmia during ischemic or nonischemic events.^{11,24} This 221 222 vulnerability could manifest after a longer time period, perhaps explaining why ER was associated 223 with SCD only among the younger subjects in our study. Furthermore, older subjects may have died due to other comorbidities before a critical event occurred. A plausible explanation could then be 224 225 that ER in young adults, particularly with an ascending ST-segment, represents a benign ECG 226 finding that normally disappears before middle age. This stands in contrast to a more constant and unchanging ER with a descending or horizontal ST-segment, which associates with a long-term 227 vulnerability to more nefarious arrhythmias.^{14–16} It may also be that the most malign ER phenotypes 228 229 manifest at a younger age and, thus, the more benign ER phenotypes may be overrepresented 230 among the very old. Moreover, as the risk of SCD increases with age, other factors may associate 231 with SCD risk more strongly than ER in older individuals.

Previous studies have provided contradictory results on the impact of sex on ER prognosis. For example, a German study found that ER was associated with cardiac mortality in a subgroup of men, but not among women.⁶ In contrast, a cohort study from the United States demonstrated an association between ER and SCD only among women.⁹ However, in the same study population, automatically detected ER was associated with cardiovascular mortality only

among men.²⁵ We, however, observed an association between ER and the risk of SCD in women
under 50, but not among men.

239 Various studies provide a large degree of heterogeneous results in their examinations 240 of the risk associated with ER among general populations, with several studies finding no link between ER and an increased risk of adverse outcomes.^{7,9,21} In the present study, we found that ER 241 was not associated with SCD, cardiac mortality, or all-cause mortality across the entire study 242 population. Possible explanations for these contradicting results across studies include the different 243 study population characteristics, follow-up periods, and ER definitions applied. In addition, only a 244 245 small minority of subjects with ER will eventually experience SCD, while the majority will enjoy 246 benign prognoses. Therefore, future research should continue to refine or better define the specific patient characteristics and ER pattern features to more accurately identify that minority of 247 individuals who will suffer an SCD. Better identifying such individuals will ultimately serve to 248 improve their prognosis. 249

250

251 Limitations

Although the subjects underwent extensive health interviews and examinations at the beginning of the survey, the subjects' health status or the presence of ER in ECG were not reassessed during the follow-up period. Therefore, we had no information on whether participants' health status, comorbidities, or ER status changed during the follow-up period. Yet, ER has been shown to be relatively stable ECG finding among middle-aged subjects.⁴ A further limitation to this study lies in the study population, which consisted of only Caucasian subjects. Thus, these results are not directly generalizable to other ethnicities.

260 *Conclusions*

In conclusion, among adults aged 30–50 years, ER associates with SCD. In particular, women under 50 years old with ER exhibited a higher risk of SCD, while ER was not associated with SCD among men <50 years old. In addition, we found that among subjects \geq 50 years old, ER was not associated with an adverse prognosis at all. Future research should focus on identifying factors that account for the differences between age groups, and improving the risk stratification in younger patient populations with ER.

267

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277 Conflicts of Interest

None declared.

References 279

280	1.	Haïssaguerre M, Derval N, Sacher F, et al.: Sudden cardiac arrest associated with early
281		repolarization. N Engl J Med 2008; 358:2016–2023.
282	2.	Nam G-B, Kim Y-H, Antzelevitch C: Augmentation of J waves and electrical storms in patients
283		with early repolarization. N Engl J Med 2008; 358:2078–2079.
284	3.	Rosso R, Kogan E, Belhassen B, et al.: J-point elevation in survivors of primary ventricular
285		fibrillation and matched control subjects: incidence and clinical significance. J Am Coll
286		Cardiol 2008; 52:1231–1238.
287	4.	Tikkanen JT, Anttonen O, Junttila MJ, et al.: Long-term outcome associated with early
288		repolarization on electrocardiography. N Engl J Med 2009; 361:2529–2537.
289	5.	Rollin A, Maury P, Bongard V, et al.: Prevalence, prognosis, and identification of the malignant
290		form of early repolarization pattern in a population-based study. Am J Cardiol 2012;
291		110:1302–1308.
292	6.	Sinner MF, Reinhard W, Müller M, et al.: Association of early repolarization pattern on ECG
293		with risk of cardiac and all-cause mortality: a population-based prospective cohort study
294		(MONICA/KORA). PLoS Med 2010; 7:e1000314. 7. Cheng Y-J, Lin X-X, Ji C-C, et
295		al.: Role of Early Repolarization Pattern in Increasing Risk of Death. J Am Heart Assoc 2016;
296		5:e003375.
297	8.	Haruta D, Matsuo K, Tsuneto A, et al.: Incidence and prognostic value of early repolarization

pattern in the 12-lead electrocardiogram. Circulation 2011; 123:2931–2937. 298

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299	9. Olson KA, Viera AJ, Soliman EZ, Crow RS, Rosamond WD: Long-term prognosis associated
300	with J-point elevation in a large middle-aged biracial cohort: the ARIC study. Eur Heart J
301	2011; 32:3098–3106.
302	10. Tikkanen JT, Junttila MJ, Anttonen O, et al.: Early repolarization: electrocardiographic
303	phenotypes associated with favorable long-term outcome. Circulation 2011; 123:2666–2673.
304	11. Adler A, Rosso R, Viskin D, Halkin A, Viskin S: What do we know about the "malignant form"
305	of early repolarization? J Am Coll Cardiol 2013; 62:863–868.
306	12. Roten L, Derval N, Maury P, et al.: Benign vs malignant inferolateral early repolarization:
307	Focus on the T wave. Heart Rhythm 2016; 13:894–902.
308	13. Hisamatsu T, Ohkubo T, Miura K, et al.: Association between J-point elevation and death from
309	coronary artery disease15-year follow up of the NIPPON DATA90. Circ J 2013; 77:1260-
310	1266.
311	14. Ilkhanoff L, Soliman EZ, Prineas RJ, et al.: Clinical characteristics and outcomes associated
312	with the natural history of early repolarization in a young, biracial cohort followed to middle
313	age: the Coronary Artery Risk Development in Young Adults (CARDIA) study. Circ
314	Arrhythm Electrophysiol 2014; 7:392–399.
315	15. Sager SJ, Hoosien M, Junttila MJ, et al.: Comparison of inferolateral early repolarization and its
316	electrocardiographic phenotypes in pre- and postadolescent populations. Am J Cardiol 2013;
317	112:444–448.
318	16. Lanza GA, Mollo R, Cosenza A, et al.: Prevalence and clinical correlates of early repolarization
319	and J wave in a large cohort of subjects without overt heart disease. J Electrocardiol 2012;
320	45:404–410.

Journal Pre-proof

321	17. Knekt P, Rissanen H, Järvinen R, Heliövaara M: Cohort Profile: The Finnish Mobile Clinic
322	Health Surveys FMC, FMCF and MFS. Int J Epidemiol 2017; 46:1760–1761i.
323	18. Macfarlane PW, Antzelevitch C, Haissaguerre M, et al.: The Early Repolarization Pattern: A
324	Consensus Paper. J Am Coll Cardiol 2015; 66:470-477.
325	19. Castle CH, Keane WM: Electrocardiographic "Peri-Infarction Block" A Clinical and Pathologic
326	Correlation. Circulation 1965; 31:403–408.
327	20. Pratt CM, Greenway PS, Schoenfeld MH, Hibben ML, Reiffel JA: Exploration of the precision
328	of classifying sudden cardiac death. Implications for the interpretation of clinical trials.
329	Circulation 1996; 93:519–524.
330	21. Klatsky AL, Oehm R, Cooper RA, Udaltsova N, Armstrong MA: The early repolarization
331	normal variant electrocardiogram: correlates and consequences. Am J Med 2003; 115:171-
332	177.
333	22. Junttila MJ, Tikkanen JT, Porthan K, et al.: Relationship between testosterone level and early
334	repolarization on 12-lead electrocardiograms in men. J Am Coll Cardiol 2013; 62:1633–1634.
335	23. Rosso R, Glikson E, Belhassen B, et al.: Distinguishing "benign" from "malignant early
336	repolarization": the value of the ST-segment morphology. Heart Rhythm 2012; 9:225–229.
337	24. Holmström LTA, Haukilahti MA, Tikkanen JT, et al.: Inferolateral early repolarization among
338	non-ischaemic sudden cardiac death victims. Europace 2018; 20:f93-f98.
339	25. O'Neal WT, Wang YG, Wu H-T, et al.: Electrocardiographic J Wave and Cardiovascular
340	Outcomes in the General Population (from the Atherosclerosis Risk In Communities Study).
341	Am J Cardiol 2016; 118:811–815.

342 Tables

343 *Table 1*

344 Baseline characteristics

	All			Age <50 years			Age \geq 50 years		
	n = 6631		n = 3305			n = 3326			
	No ER	ER	<u>.</u>	No ER	ER	·	No ER	ER	
	n = 5838	n = 793		n = 2938	n = 367		n = 2900	n = 426	
	(88.0%)	(12.0%)	Р	(88.9%)	(11.1%)	Р	(87.2%)	(12.8%)	Р
Mala (0/)+	2522	432	< 001	1342	250	< 001	1180	182	405
Male (%)	(43.2%)	(54.5%)	< .001	(45.7%)	(68.1%)	< .001	(40.7%)	(42.7%)	.405
$\Delta qe (vears)$	51.0	52.1	006	39.3	39.6	131	62.7	62.9	680
Age (years) _‡	±13.9	±13.8	.000	±5.8	±5.9		± 8.8	±8.6	.060
Systolic blood	143.2	143.0	074	132.3	131.9	022	154.3	152.5	114
pressure (mmHg)§	±23.2	±22.3	.074	±16.6	±15.2	.033	±23.7	±23.0	.114
Diastolic blood	86.8	87.2	633	84.9	85.6	397	88.8	88.6	661
pressure (mmHg)§	±11.5	±11.5	1000	±11.2	±10.9		±11.5	±11.9	1001
Body mass index	25.9	26.2	082	25.0	25.3	583	26.8	26.9	425
(kg/m^2) §	±4.1	±4.1	.002	±3.8	±3.6	.505	±4.2	±4.4	.423
	6.9	7.1		6.6	6.7		7.3	7.4	
Cholesterol (mmol/l,	±1.4	±1.5	008	±1.3	±1.3	203	±1.4	±1.6	164
mg/dl)§	268	274	.008	254	260	.295	283	286	.164
	±53	±57		±48	±49		±53	±61	
Heart rate (hnm)8	69	65	< 001	67	63	< 001	71	66	< 001
ricari rate (opin)8	±14	±12	< .001	±12	±11	< .001	±15	±14	< .001
QRS duration (ms)§	85	85	.079	86	86	.001	85	85	.793

			Journa	l Pre-proc	of			
	±9	± 8		±9	± 8		±9	± 8
OT a internal (ma) 8	404	400	004	400	393	000	407	405
Q1c interval (ms)§	±24	±25	.004	±23	±25	.009	±24	±24
$\Omega_{\rm res}$ = $1 \frac{1}{2} m = \langle 0 \rangle $	1266	199	.291	779	118	544	487	81
Smoking (%)§	(21.7%)	(25.1%)		(26.5%)	(32.2%)	.544	(16.8%)	(19.1%)
Dishets: $(0/)$ S	320	27	.002	34	4	(())	286	23
Diabetes (%)§	(5.5%)	(3.4%)		(1.2%)	(1.1%)	.002	(9.9%)	(5.4%)
Coronary artery	603	76	0.67	55	8	007	548	68
disease (%)§	(10.3%)	(9.6%)	.067	(1.9%)	(2.2%)	.987	(18.9%)	(16.0%)
Beta blocker	370	73	007	88	17	126	282	56
medication (%)§	(6.3%)	(9.2%)	.007	(3.0%)	(4.6%)	.136	(9.7%)	(13.1%)

345

346 ER = early repolarization; QTc = QT corrected for heart rate using Bazett's formula. Continuous

347 data are presented as means \pm standard deviation, while categorical data are presented as the

number of cases (% of study population). Statistical test for the difference between subjects with

and without ER in all subjects, subjects aged <50 years, and subjects aged ≥50 years.

350 †Adjusted for age.

351 ‡Adjusted for sex.

352 §Adjusted for age and sex.

354 *Table 2*

Risk of sudden cardiac death, cardiac death, and death from any cause associated with ER in

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356 subjects aged <50 years and subjects aged \geq50 years
```

	Age <50 years		Age	ER*age group	
	n =	n = 3305		n = 3326	
	No ER	ER	No ER	ER	
	n = 2938	n = 367	n = 2900	n = 426	Р
SCD					
# of SCDs	74	21	218	33	
(# of SCDs in men)	(62)	(16)	(123)	(19)	
Age- and sex-adjusted	1	1.72	1	1.01	045
HR (95% CI)	1	(1.05–2.80)	1	(0.70–1.46)	.045
Multivariate-adjusted	1	1.88	1	1.01	048
HR (95% CI)	1	(1.16–3.07)	1	(0.70–1.46)	.040
Cardiac death					
# of cardiac deaths	199	38	1112	171	
(# of cardiac deaths in men)	(150)	(31)	(507)	(82)	
Age- and sex-adjusted	1	1.20	1	1.03	170
HR (95% CI)	1	(0.85–1.70)	1	(0.88–1.21)	.170
Multivariate-adjusted	1	1.13	1	1.08	175
HR (95% CI)	1	(0.79–1.60)	1	(0.92–1.27)	.175
Death					
# of deaths	649	99	2442	377	
(# of deaths in men)	(404)	(75)	(1052)	(166)	
	1				

	Journal Pre-proof		
Age- and sex-adjusted	1.05	1.04	505
HR (95% CI)	1 (0.85–1.30)	1 (0.93–1.16)	.385
Multivariate-adjusted	1.03	1.07	620
HR (95% CI)	(0.83–1.28)	1 (0.96–1.19)	.620

- 358
- 359 ER = early repolarization; SCD = sudden cardiac death. The hazard ratios (HRs) and 95%
- 360 confidence intervals (CIs) were calculated using the Cox proportional hazards model. Variables
- 361 included in the multivariate analyses were age as a continuous variable, sex, systolic blood pressure,
- total serum cholesterol, coronary artery, diabetes, smoking, and ER. The effect modification was
- 363 tested by entering an interaction term for ER and the age group in the multivariate analysis.

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364 *Table 3*

365 Risk of sudden cardiac death according to the ER pattern in subjects aged <50 years

Age < 50	years
----------	-------

n = 3305

	# of	# of	Age- and sex-adjusted	Multivariate-adjusted
	subjects	SCDs	HR (95% CI)	HR (95% CI)
No ER	2938	74	1	1
Inferior/lateral ER	367	21	1.72 (1.05–2.80)	1.88 (1.16–3.07)
Inferior ER	213	12	1.72 (0.93–3.19)	1.92 (1.04–3.56)
Lateral ER	174	11	1.80 (0.95–3.39)	2.08 (1.10-3.95)
Slurred inferior/lateral ER	251	15	1.82 (1.04–3.18)	2.09 (1.19–3.67)
Notched inferior/lateral ER	74	4	1.59 (0.58–4.37)	2.28 (0.82-6.31)
Inferior/lateral ER, ascending ST-segment	253	12	1.34 (0.72–2.47)	1.45 (0.78–2.67)
Inferior/lateral ER, horizontal or descending ST-segment	114	9	2.74 (1.37–5.47)	3.12 (1.56–6.26)
Inferior/lateral ER ≥ 0.1 mV but < 0.2 mV	300	21	2.00 (1.23-3.25)	2.16 (1.33-3.52)
Inferior/lateral ER ≥0.2 mV	46	0	_	_
Low amplitude T-wave	29	5	6.79 (2.73–16.89)	4.47 (1.75–11.42)

366

367 The hazard ratios (HRs) and 95% confidence intervals (CIs) for sudden cardiac death were

368 calculated using the Cox proportional hazards model. Variables included in the multivariate

369 analyses were age, sex, systolic blood pressure, total serum cholesterol, diabetes, smoking, coronary

artery disease, and the ER pattern.

372 Figures

373 Figure 1



375 Figure 1 legend

- 376 Survival plots of A) subjects aged <50 years and B) subjects aged ≥50 years with and without ER
- 377 for sudden cardiac death (SCD), adjusted for age, sex, systolic blood pressure, total serum
- 378 cholesterol, smoking, diabetes, and coronary artery disease.

379

Journal Prevention

380 Figure 2



382 Figure 2 legend

Forty-five-year-old woman with typical inferior ER pattern with horizontal ST-segments. She died
of sudden cardiac death during the follow-up period. Paper speed is 50 mm/s. Arrows indicate the
ER patterns.

Table 1 Baseline characteristics

	All n = 6631			Age	Age <50 years n = 3305			Age ≥ 50 years n = 3326		
				t						
	No ER	ER		No ER	ER		No ER	ER		
	n = 5838	n = 793		n = 2938	n = 367		n = 2900	n = 426		
	(88.0%)	(12.0%)	Р	(88.9%)	(11.1%)	Р	(87.2%)	(12.8%)	Р	
Male (%)†	2522 (43.2%)	432 (54.5%)	<.001	1342 (45.7%)	250 (68.1%)	<.001	1180 (40.7%)	182 (42.7%)	.405	
Age (years)‡	51.0 ±13.9	52.1 ±13.8	.006	39.3 ±5.8	39.6 ±5.9	.434	$\begin{array}{c} 62.7 \\ \pm 8.8 \end{array}$	62.9 ±8.6	.68	
Systolic blood pressure (mmHg)§	143.2 +23.2	143.0 +22.3	.074	132.3 ±16.6	131.9 +15.2	.33	154.3 +23.7	152.5 + 23.0	.114	
Diastolic blood pressure (mmHg)§	86.8 ±11.5	87.2 ±11.5	.633	84.9 ±11.2	85.6 ±10.9	.397	88.8 ±11.5	88.6 ±11.9	.661	
Body mass index (kg/m ²)§	25.9 ±4.1	26.2 ±4.1	.082	25.0 ±3.8	25.3 ±3.6	.583	26.8 ±4.2	26.9 ±4.4	.425	
Cholesterol (mmol/l, <i>mg/dl</i>)§	$6.9 \pm 1.4, 265 \pm 53$	7.1 ±1.5, 274 ±57	.008	6.6 ±1.3, 254 ±48	$6.7 \pm 1.3, 260 \pm 49$.293	7.3 $\pm 1.4,$ 283 ± 53	7.4 ±1.6, 286 ±61	.164	
Heart rate (bpm)§	69 ±14	65 ±12	<.001	67 ±12	63 ±11	<.001	71 ±15	66 ±14	<.001	
QRS duration (ms)§	85 ±9	85 ±8	.079	86 ±9	86 ± 8	.001	85 ±9	85 ±8	.793	
QTc interval (ms)§	404 ±24	400 ±25	.004	400 ±23	393 ±25	.009	407 ±24	405 ±24	.254	
Smoking (%)§	1266 (21.7%)	199 (25.1%)	.291	779 (26.5%)	118 (32.2%)	.544	487 (16.8%)	81 (19.1%)	.301	
Diabetes (%)§	320 (5.5%)	27 (3.4%)	.002	34 (1.2%)	4 (1.1%)	.662	286 (9.9%)	23 (5.4%)	.002	
Coronary artery disease (%)§	603 (10.3%)	76 (9.6%)	.067	55 (1.9%)	8 (2.2%)	.987	548 (18.9%)	68 (16.0%)	.107	

Beta blocker	370 (6.3%)	73 (9.2%)	.007	88 (3.0%)	17 (4.6%)	.136	282 (9.7%)	56 (13.1%)	.004
metheation (%)§									

ER = early repolarization; QTc = QT corrected for heart rate using Bazett's formula. Continuous data are presented as means \pm standard deviation, while categorical data are presented as the number of cases (% of study population). Statistical test for the difference between subjects with and without ER in all subjects, subjects aged <50 years, and subjects aged \geq 50 years.

† Adjusted for age.

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Table 2

Risk of sudden cardiac death, cardiac death, and death from any cause associated with ER in subjects aged <50 years and subjects aged ≥50 years

	Age $<$ 50 years n = 3305		Ag	$e \ge 50$ years n = 3326	ER*age group interaction
	No ER n – 2938	ER n = 367	No ER n – 2900	ER n = 426	Р
SCD	n = 2950	n = 507	n = 2900	n = 420	1
# of SCDs	74	21	218	33	
(# of SCDs in men)	(62)	(16)	(123)	(19)	
Age- and sex-adjusted HR (95% CI)	1	1.72 (1.05–2.80)	1	1.01 (0.70–1.46)	0.045
Multivariate-adjusted HR (95% CI)	1	1.88 (1.16–3.07)	1	1.01 (0.70–1.46)	0.048
Cardiac death					
# of cardiac deaths	199	38	1112	171	
(# of cardiac deaths in men)	(150)	(31)	(507)	(82)	
Age- and sex-adjusted HR (95% CI)	1	1.20 (0.85–1.70)	1	1.03 (0.88–1.21)	0.170
Multivariate-adjusted HR (95% CI)	1	1.13 (0.79–1.60)	1	1.08 (0.92–1.27)	0.175
Death					
# of deaths	649	99	2442	377	
(# of deaths in men)	(404)	(75)	(1052)	(166)	
Age- and sex-adjusted HR (95% CI)	1	1.05 (0.85–1.30)	1	1.04 (0.93–1.16)	0.585
Multivariate-adjusted HR (95% CI)	1	1.03 (0.83–1.28)	1	1.07 (0.96–1.19)	0.620

intervals (CIs) were calculated using the Cox proportional hazards model. Variables included in the multivariate analyses were age as a continuous variable, sex, systolic blood pressure, total serum cholesterol, coronary artery, diabetes, smoking, and ER. The effect modification was tested by entering an interaction term for ER and the age group in the multivariate analysis.

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Table 3

Risk of sudden cardiac death according to the ER pattern in subjects aged <50 years

	Age <50 years					
	n = 3305					
	# of	# of SCDs	Age- and sex-adjusted	Multivariate adjusted		
	subjects	" OI BCD3	HR (95% CI)	HR (95% CI)		
No ER	2938	74	1	1		
Inferior/lateral ER	367	21	1.72 (1.05-2.80)	1.88 (1.16-3.07)		
Inferior ER	213	12	1.72 (0.93-3.19)	1.92 (1.04-3.56)		
Lateral ER	174	11	1.80 (0.95-3.39)	2.08 (1.10-3.95)		
Slurred inferior/lateral ER	251	15	1.82 (1.04-3.18)	2.09 (1.19-3.67)		
Notched inferior/lateral ER	74	4	1.59 (0.58–4.37)	2.28 (0.82-6.31)		
Inferior/lateral ER, ascending ST segment	253	12	1.34 (0.72–2.47)	1.45 (0.78–2.67)		
Inferior/lateral ER, horizontal or descending ST segment	114	9	2.74 (1.37-5.47)	3.12 (1.56-6.26)		
Inferior/lateral ER ≥0.1 mV but <0.2 mV	300	21	2.00 (1.23-3.25)	2.16 (1.33-3.52)		
Inferior/lateral ER $\geq 0.2 \text{ mV}$	46	0	-	-		
Low amplitude T-wave	29	5	6.79 (2.73–16.89)	4.47 (1.75–11.42)		

The hazard ratios (HRs) and 95% confidence intervals (CIs) for sudden cardiac death were calculated using the Cox proportional hazards model. Variables included in the multivariate analyses were age, sex, systolic blood pressure, total serum cholesterol, diabetes, smoking, coronary artery disease, and the ER pattern.

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1 SUPPLEMENTAL MATERIAL

Journal Pre-proof

- 2 Supplemental Table 1
- 3 Risk of sudden cardiac death, cardiac death, and death from any cause associated with ER in the
- 4 entire study population

	All			
	n = 6631			
	No ER	ER		
	n = 5838	n = 793		
SCD				
# of SCDs	292	54		
Age- and sex-adjusted HR (95% CI)	1	1.22 (0.91-1.63)		
Multivariate-adjusted HR (95% CI)	1	1.23 (0.92-1.64)		
Cardiac death				
# of cardiac deaths	1311	209		
Age- and sex-adjusted HR (95% CI)	1	1.07 (0.93-1.24)		
Multivariate-adjusted HR (95% CI)	1	1.12 (0.96-1.29)		
Death				
# of deaths	3091	476		
Age- and sex-adjusted HR (95% CI)	1	1.05 (0.95-1.15)		
Multivariate-adjusted HR (95% CI)	1	1.07 (0.97-1.18)		
	1			

5

 $6 \quad ER = early repolarization; SCD = sudden cardiac death.$

7 The hazard ratios (HRs) and 95% confidence intervals (CIs) were calculated using the Cox

8 proportional hazards model. Variables included in the multivariate analyses were age as a

9 continuous variable, sex, systolic blood pressure, total serum cholesterol, coronary artery, diabetes,

10 smoking, and ER.

All

11 Supplemental Table 2

12 Risk of sudden cardiac death according to the ER pattern in the entire study population

			n = 6631	
	# of subjects	# of SCDs	Age- and sex- adjusted HR (95% CI)	Multivariate-adjusted HR (95% CI)
No ER	5838	292	1	1
Inferior/lateral ER	793	54	1.22 (0.91–1.63)	1.23 (0.92–1.64)
Inferior ER	392	27	1.29 (0.87–1.92)	1.26 (0.85–1.88)
Lateral ER	429	30	1.20 (0.82–1.75)	1.25 (0.86–1.82)
Slurred inferior/lateral ER	555	38	1.25 (0.89–1.75)	1.26 (0.90–1.76)
Notched inferior/lateral ER	138	10	1.26 (0.67–2.37)	1.29 (0.68–2.43)
Inferior/lateral ER, ascending ST segment	470	28	1.04 (0.71–1.54)	1.10 (0.75–1.63)
Inferior/lateral ER, horizontal or descending ST segment	323	26	1.49 (1.00–2.23)	1.39 (0.93–2.09)
Inferior/lateral ER $\geq 0.1 \text{ mV}$ but $< 0.2 \text{ mV}$	680	47	1.27 (0.93–1.73)	1.27 (0.93–1.73)
Inferior/lateral ER ≥0.2 mV	113	7	0.98 (0.46-2.07)	1.00 (0.47–2.12)
Low amplitude T-wave	158	17	1.85 (1.13–3.03)	1.75 (1.06–2.87)

13

14 The hazard ratios (HRs) and 95% confidence intervals (CIs) for sudden cardiac death (SCD) were

15 calculated using the Cox proportional hazards model. Variables included in the multivariate

16 analyses were age, sex, systolic blood pressure, total serum cholesterol, diabetes, smoking, coronary

17 artery disease and the ER pattern.