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Baseline plasma lipid levels in patients with acute coronary syndrome: Association with 20-year mortality. The ABC-5a* Study on Heart Disease

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Abstract

Background: The relationship between baseline plasma lipid levels during acute coronary syndrome and the outcome has clinical relevance.

Methods: To evaluate their long-term prognostic value, we examined 589 patients admitted with acute coronary syndrome at three hospitals. Baseline plasma lipids were assessed on days 1 and 7. Patients were followed for 20 years or until death. Results: Virtually, all patients completed follow-up; 437 (74%) had died: 24% due to coronary artery disease/heart failure (CAD/HF), 21% sudden cardiac death (SCD), 16% due to other cardiovascular causes and 39% due to non-cardiac death. The incidence rate (IR) of all-cause mortality was not different among patients with baseline plasma lipids less or greater than the median value. The IR of CAD/HF mortality was not significantly higher among patients with greater than median low-density lipoprotein (LDL) cholesterol and triglyceride (TG) levels. The IR of non-cardiac death tended to be lower among patients with greater than median total cholesterol (TC) and LDL levels. Using three levels of adjusted Cox survival models, baseline plasma lipids had no consistent independent or inverse association with all-cause mortality, even after excluding patients who received statins. Competitive risk survival models for each cause of death revealed that the only hazard of non-cardiac death was consistently higher among patients with less than or equal to median TC and LDL levels. Conclusion: In the present prospective long-term study, after acute coronary syndrome, baseline plasma lipid levels seem not to be associated with long-term global mortality. Only an independent inverse association between TC and LDL and noncardiac death has been observed.

1 | INTRODUCTION

Ischaemic heart disease (IHD) is the world's biggest killer, accounting for 9.4 million deaths (16.6% of total deaths) in 2016 and has been the leading cause of death globally over the last 15 years.¹

High plasma lipid levels, particularly total serum cholesterol (TC) and low-density lipoprotein (LDL), have long been recognised as a major risk factor for atherosclerosis and IHD.²⁻⁴ However, this causal relationship has been debated in recent reports.⁵⁻⁷ The prognostic value of admission plasma lipid levels in patients presenting with acute coronary syndrome has rarely been discussed in the literature.⁸⁻¹⁰ In the present study, we aimed to report the association

*ABC is an acronym for Adria, Bassano, Conegliano, and Padova Hospitals.

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between the in-hospital plasma lipid profile and the subsequent long-term mortality risk over 20 years of follow-up in an unselected sample of patients discharged alive from three different centres after an index hospitalisation with acute coronary syndrome.

2 | METHODS

2.1 | Patients

The ABC Study on Heart Disease is an ongoing prospective investigation designed to represent, as closely as possible, an unbiased population of patients with acute coronary syndrome (www.abche artdiseasestudy.org/en/). The cohort includes Caucasian patients with definite acute coronary syndrome, including ST-elevation myocardial infarction (STEMI), non-ST elevation myocardial infarction (NSTEMI) or unstable angina, who were admitted to the intensive care units of Adria, Bassano, and Conegliano General Hospitals in Italy between June 1995 and January 1998. The original aim of the ABC study was to monitor these patients with regard to natural longterm history, both non-fatal and fatal events, and causes of death. Another study aim was to investigate the prognostic value of multiple baseline clinical variables. Criteria for acute coronary syndrome diagnosis included clinical presentation, electrocardiogram findings and the presence of serum biochemical markers of necrosis.^{11,12}

A total of 741 patients were considered eligible upon admission, 84 of whom were excluded because they had diseases other than acute coronary syndrome, and 23 were excluded due to a lack of baseline data. Among the 634 enrolled patients with acute coronary syndrome, 45 died during the index hospitalisation. Therefore, the postdischarge follow-up study included 589 patients (Figure 1). Each patient received an anonymous code, and no personal data or identifiers were included in the baseline or follow-up database. The study has been performed in accordance with the Declaration of Helsinki and it was approved by Adria, Bassano del Grappa and Conegliano General Hospitals ethics committee. All enrolled patients gave their written informed consent.

2.2 | Measurements and follow-up

At enrolment, a thorough patient history was collected from medical 42 43 records and patient interviews. All presently analysed baseline clinical and laboratory data were obtained during the first 7 days of hos-44 45 pitalisation in the intensive coronary care unit. The acute coronary syndrome diagnosis criteria were the fulfilment of at least two of the 46 47 following: central chest pain lasting >30 minutes, typical changes in 48 serum enzymes, including total creatine kinase (CK) and creatine 49 kinase MB (CK-MB), and typical electrocardiogram changes with pathological Q waves and/or localised ST-T changes in at least two 50 contiguous leads.¹³ Two fasting venous blood samples (12 hours after 51 52 admission and before discharge) were drawn for TC, LDL, HDL, and 53 triglyceride (TG) measurements. LDL concentrations were estimated

What's known

 In many studies, increased serum lipids are reported to be associated with a higher risk of mortality.

What's new

- After 20 years of follow-up of acute coronary syndrome survivors, baseline plasma lipid levels were not associated with all-cause mortality.
- Baseline plasma lipid levels were significantly inversely associated with death due to non-cardiac causes.



FIGURE 1 Flow diagram of the study population and progress during follow-up. ACS, acute coronary syndrome; CAD/HF, coronary artery disease/heart failure; Other CV, other cardiovascular causes; SCD, sudden cardiac death

using the modified Friedewald formula (MFF): LDL (mg/dL) = non-HDL \times 90% – TG \times 10%. In all three hospitals, plasma lipid measurements were performed using an enzymatic colorimetric method.¹⁴ Details of the measured variables were published previously.^{11,12}

Each patient underwent a clinical check-up 1, 3, 5, 7, 10, 12, 15, 17 and 20 years after recruitment. At each recruitment hospital, two

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cardiologists were responsible for monitoring the cohort of patients throughout the follow-up. Data were obtained from scheduled examinations, public administrations, hospital records, family doctors, postmortem examinations and death certificates. The medications received during index hospitalisation and follow-up treatments were also recorded.

All data after enrolment were recorded prospectively following the protocol of the ABC Study on Heart Disease.¹¹ According to the protocol, baseline data and follow-up data were recorded in two different datasheets. For the present analysis, the datasheets were merged after completion of 20 years of follow-up.

2.3 | Causes of death

16 The primary clinical outcome was 20-year all-cause mortality. The sec-17 ondary outcomes were modes of death, which were classified into the 18 following categories: coronary artery disease and/or heart failure pro-19 gression (CAD/HF); sudden cardiac death (SCD), defined as witnessed, 20 out-of-hospital death within 1 hour after the onset of acute symptoms 21 or unwitnessed, unexpected death (eg, during sleep) in patients with 22 good health 24 hours before the event¹⁵; other cardiovascular (other 23 CV) causes; and non-cardiac death causes. The patients were exam-24 ined and the cause of death was determined by two clinical research-25 ers with no knowledge of the patients' baseline data.

2.4 | Statistical analysis

30 The accrued variables were analysed as continuous variables or pro-31 portions. Log transformations were applied to correct for positively 32 skewed distributions as appropriate. Measured variables were analysed using the unpaired Student's t test, and categorical variables using Pearson's χ^2 test. If a patient dropped out prior to 20 years of 34 35 follow-up, her/his data were censored at that time. Survival curves were constructed using cumulative hazards.¹⁶ We compared inci-36 37 dence rates (IRs) using Mantel-Haenszel estimates of the rate ratio. Interaction analysis for lipid versus age and gender was also per-38 formed.¹⁷ The risk ratios (RRs) estimated in the analysis and graphic 39 representations of the mortality were obtained using lipid values as 40 41 dichotomous variables, with the 50th percentile as a cut-off.

42 Log-transformed continuous variables were used for global 43 mortality risk estimation using Cox regression models and scaled Schoenfeld residuals were used to test the proportionality assump-44 45 tion with 95% confidence intervals (CIs). Competitive risk regression 46 models with the Fine-Gray method using the same log-transformed 47 continuous variables were set-up to analyse each of the four modes 48 of death. The strength of an association was expressed as β coeffi-49 cient ± SE. The predictive power of lipid values as continuous vari-50 ables with regard to mortality was also estimated. Based on survival 51 regression, a postestimation prediction of mortality was carried 52 out and graphically presented along with the appropriate P-values 53 calculated by the survival regression. Unless otherwise indicated,

two-tailed *P*-values < .05 were considered significant. Statistical analyses were performed using STATA 14.

3 | RESULTS

All enrolled patients completed the follow-up unless pre-empted by death, except three patients for whom survival time was censored before 20 years; two withdrew consent and one moved overseas (Figure 1). Table 1 compares the demographic and baseline clinical characteristics of the patients according to plasma TC level at admission in relation to the median value of 208 mg/dL.

At enrolment, the two groups did not differ in most of the clinical characteristics. However, females were more frequent among patients with higher TC levels. In addition, these patients had higher BMI, used alcohol less frequently and had higher blood haemoglobin values. Patients who had TC >208 mg/dL at admission also had higher LDL and TG levels and lower HDL levels.

By the end of 20 years of follow-up, 437 (74%) of the patients discharged alive had died: 105 (24%) due to CAD/HF, 90 (21%) SCD, 71 (16%) other CV causes and 171 (39%) non-cardiac death (Figure 1). The IR of all-cause mortality throughout follow-up after acute coronary syndrome was approximately 66.7 (95% CI 60.7-73.3) cases/1000 person-years. Unexpectedly, the IR was not associated with the baseline plasma lipids (Table S1). Cumulative hazard estimate curves for 20-year all-cause mortality risk according to baseline plasma lipid levels are illustrated in Figure 2 and Figure S1.

The final survival analysis accounted for competitive risks for each of the main four causes of death, revealing that the positive association of baseline TC, LDL and TG levels and the negative association of HDL level with long-term CAD/HF mortality were not statistically significant in the unadjusted model, a model adjusted for age and gender or the fully adjusted model (Table 2). The analysis also showed a lack of consistently positive or negative association between TC and LDL and other causes of cardiac mortality. Only the hazard of non-cardiac death was consistently higher among patients with TC or LDL values less than or equal to median values at admission in all models (Table 2B).

4 | DISCUSSION

This prospective three-centre study of an unselected real-world patient discharged alive after acute coronary syndrome showed that baseline plasma TC, LDL, HDL and TG levels are not associated with long-term global mortality. The association with modes of death showed an independent inverse association between TC and LDL and non-cardiac death. The same findings were reported in our research letter.¹⁸

The results are consistent for lipid measurements at admission or after 1 week of hospitalisation. Furthermore, the associations were independent of anti-lipid treatment, suggesting that the core prognostic issue lies on the lipid levels per se.

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Variable	Overall sample (n = 589)	TC ≤208 mg/ dL (n = 295)	TC >208 mg/dL (n = 294)	P values	
Median age, years	67 (58-74)	67 (59-75)	66 (58-73)	.47	
Gender, female	29.2	22	36	<.0001	
Education above primary school	26	29	23	.37	
1edian body mass index, kg/m ²	26 (24-28)	25 (23-27)	26 (24-29)	.0001	
Smoking habit ^a	67	71	64	.09	
lcohol use	74	80	68	.001	
lypertension	47	51	44	.08	
viabetes mellitus	22	22	23	.68	
/ledian systolic blood pressure, mm Hg	120 (110-130)	120 (110-130)	120 (110-135)	.19	
1edian diastolic blood pressure, mm Hg	80 (70-80)	75 (70-80)	80 (70-80)	.50	
∕ledian heart rate, beats/min	71 (60-82)	72 (60-81)	70 (61-82)	.54	
Ion-ST elevation ACS	38	38	39	.79	
Killip class >1	33	34	33	.75	
VEF (%) (n = 500)	52 (45-61)	53 (45-60)	52 (45-61)	.73	
lb, g/dL	14 (13-15)	13 (12-15)	14 (13-15)	.0001	
ilood glucose level, mg/dL	119 (100-158)	117 (98-153)	122 (103-172)	.06	
Gerum creatinine level, mg/dL	0.9 (0.8-1.1)	0.9 (0.8-1.1)	0.9 (0.8-1.1)	.95	
CK-MB peak ^b , U/L	102 (43-203)	100 (42-186)	104 (34-214)	.15	
Serum lipids ^b , mg/dL					
Total cholesterol					
At admission	208 (179-243)	179 (159-193)	243 (225-264)	<.0001	
At discharge	198 (174-224)	177 (155-192)	220 (204-245)	<.0001	
LDL ^c					
At admission	133 (108-157)	108 (91-158)	157 (144-178)	<.0001	
At discharge	125 (106-147)	109 (93-122)	143 (128-163)	<.0001	
HDL					
At admission	44 (37-51)	42 (36-51)	45 (39-52)	.005	
At discharge	39 (33-47)	37 (32-46)	40 (33-49)	.01	
TGs					
At admission	127 (93-178)	109 (81-147)	155 (112-218)	<.0001	
At discharge	146 (111-188)	129 (102-162)	162 (124-212)	<.0001	

TABLE 1Baseline characteristics ofpatients with acute coronary syndromeaccording to the baseline total cholesterollevel

Note: The values are presented as medians and interquartile ranges or percentages.

Abbreviations: ACS, acute coronary syndrome; CK-MB, creatine kinase-MB isoenzyme; Hb,

⁺ haemoglobin; HDL, high-density lipoprotein cholesterol; LDH, lactate dehydrogenase-1 isoenzyme;

⁵ LDL, low-density lipoprotein cholesterol; LVEF, left ventricular ejection fraction; TC, total

5 cholesterol; TG, triglyceride.

⁷ ^aPrevious smokers and currently smoking patients.

48 ^bP-values were calculated using log-transformed data.

49 ^cCalculated using modified Friedewald formula.

51 The general view is that an elevated level of plasma of TC and 52 LDL is a major risk factor for developing atherosclerotic cardio-53 vascular disease (CVD).²⁻⁴ Interestingly, our results demonstrated that neither total nor LDL-cholesterol was elevated at the time of admission with acute coronary syndrome as it should be expected according to that view. Low or normal TC and LDL-C in patients



FIGURE 2 Cumulative hazard estimate curves for 20-year all-cause mortality risk. A, Cumulative hazard estimate according to day 1 plasma lipid levels. B, Estimated prediction. HDL, high-density lipoprotein cholesterol; LDL, low-density lipoprotein cholesterol; TC, total cholesterol; TG, triglyceride. *P*-values were calculated by unadjusted Cox regression models using log-transformed continuous variables

with CVD has been reported by other researchers.^{19,20} Recently,
the general view has been questioned.⁵⁻⁷ As reported by Ravnskov
et al, ⁷ there are many observations and experiments contradicting

that high TC or LDL-C causes atherosclerosis and CVD, and several recent observations have supported their view. For instance, six recent follow-up studies including almost 700 000 individuals **TABLE 2** Cox regression analysis of all-cause mortality risk (A) and competing risk regression analysis for the main causes of death (B) according to baseline plasma lipid levels

Adjusted for age and gender		Fully adjusted ^a			Fully adjusted ^b				
) (aviable		7			7		β ± Robust	7	-
	p ± Robust SE	2	Р	p ± Kobust SE	Z	٢	SE	Z	P
(A) Cox regression ana									
All-cause mortality (n	= 437)								
	0.44 + 0.00	0.07	0.4	0.50 . 0.00	0 (0	04	0.44 + 0.04	4.07	0(
At admission	0.46 ± 0.22	-2.07	.04	0.58 ± 0.22	-2.62	.01	0.44 ± 0.24	-1.86	.06
At discharge	0.55 ± 0.27	-2.02	.04	0.50 ± 0.27	-1.86	.06	0.32 ± 0.28	-1.14	.25
	0.22 + 0.17	2.02	0.4	0.42 + 0.47	2.50	01	0.00 + 0.10	1.00	07
At admission	0.33 ± 0.16	-2.02	.04	0.43 ± 0.17	-2.58	.01	0.32 ± 0.18	-1.82	.07
At discharge	0.32 ± 0.22	-1.46	.14	0.34 ± 0.22	-1.58	.12	0.20 ± 0.23	-0.86	.39
HDL									
At admission	0.41 ± 0.18	-2.20	.03	0.36 ± 0.19	-1.89	.06	0.35 ± 0.19	-1.87	.06
At discharge	0.64 ± 0.20	-3.22	<.0001	0.46 ± 0.21	-2.21	.03	0.41 ± 0.21	-1.98	.05
TGs									
At admission	0.11 ± 0.10	1.04	.30	0.06 ± 0.11	0.58	.56	0.10 ± 0.11	0.90	.37
At discharge	0.11 ± 0.14	0.81	.42	0.02 ± 0.14	0.16	.87	0.07 ± 0.14	0.49	.62
(B) Competing risk reg	ression								
CAD/HF mortality (n =	= 105)								
Total cholesterol									
At admission	0.78 ± 0.48	1.63	.10	0.78 ± 0.48	1.63	.10			
At discharge	0.33 ± 0.57	0.57	.57	0.57 ± 0.56	1.01	.31			
LDL									
At admission	0.66 ± 0.37	1.76	.08	0.69 ± 0.39	1.77	.08			
At discharge	0.55 ± 0.46	1.20	.23	0.74 ± 0.46	1.60	.11			
HDL									
At admission	-0.66 ± 0.38	-1.74	.08	-0.51 ± 0.41	-1.24	.22			
At discharge	-0.72 ± 0.39	-1.85	.06	-0.54 ± 0.42	-1.29	.20			
TGs									
At admission	0.41 ± 0.20	2.01	.05	0.33 ± 0.21	1.59	.11			
At discharge	0.19 ± 0.27	0.69	.49	0.05 ± 0.27	0.19	.85			
SCD (n = 90)									
Total cholesterol									
At admission	0.58 ± 0.52	1.12	.26	0.83 ± 0.46	1.83	.07			
At discharge	0.27 ± 0.64	0.42	.67	0.74 ± 0.59	1.25	.21			
LDL									
At admission	0.44 ± 0.41	1.09	.28	0.62 ± 0.38	1.62	.10			
At discharge	0.46 ± 0.52	0.89	.38	0.76 ± 0.50	1.52	.13			
HDL									
At admission	-0.41 ± 0.44	-0.94	.35	-0.35 ± 0.46	-0.76	.45			
At discharge	-1.11 ± 0.44	-2.53	.01	-0.76 ± 0.46	-1.63	.10			
TGs									
At admission	0.10 ± 0.23	0.45	.65	0.25 ± 0.22	1.17	.24			
At discharge	0.06 ± 0.31	0.20	.85	0.12 ± 0.31	0.37	.71			

 (Continues)

TABLE 2 (Continued)

	Adjusted for age	and gender		Fully adjusted ^a	Fully adjusted ^a		Fully adjusted ^b		
			_			_	β ± Robust		
Variable	$\beta \pm Robust SE$	Z	Р	$\beta \pm Robust SE$	Z	Р	SE	Z	Р
Other CV causes (n =	= 71)								
Total cholesterol									
At admission	-0.63 ± 0.46	-1.35	.18	-0.44 ± 0.50	-0.87	.38			
At discharge	-0.68 ± 0.60	-1.13	.26	-0.56 ± 0.65	-0.87	.39			
LDL									
At admission	-0.38 ± 0.31	-1.21	.23	-0.23 ± 0.36	-0.66	.51			
At discharge	-0.68 ± 0.46	-1.48	.14	-0.62 ± 0.51	-1.22	.22			
HDL									
At admission	0.15 ± 0.50	0.30	.77	0.34 ± 0.49	0.70	.49			
At discharge	0.22 ± 0.48	0.45	.65	0.49 ± 0.57	0.86	.39			
TGs									
At admission	-0.12 ± 0.24	-0.50	.61	-0.30 ± 0.26	-1.16	.25			
At discharge	0.27 ± 0.31	0.89	.37	0.08 ± 0.34	0.24	.81			
Non-cardiac death (n	n = 171)					~			
Total cholesterol									
At admission	-0.86 ± 0.36	-2.40	.02	-1.12 ± 0.39	-2.86	<.0001			
At discharge	-0.28 ± 0.42	-0.66	.51	-0.53 ± 0.46	-1.15	.25			
LDL									
At admission	-0.71 ± 0.27	-2.67	.01	-0.88 ± 0.28	-3.11	<.0001			
At discharge	-0.40 ± 0.33	-1.20	.23	-0.55 ± 0.36	-1.54	.12			
HDL									
At admission	0.38 ± 0.28	1.38	.17	0.25 ± 0.29	0.87	.39			
At discharge	0.64 ± 0.31	2.05	.04	0.35 ± 0.33	1.07	.29			
TGs									
At admission	-0.28 ± 0.18	-1.60	.11	-0.32 ± 0.19	-1.64	.10			
At discharge	-0.18 ± 0.22	-0.84	.40	-0.11 ± 0.23	-0.48	.64			

Note: P-values were calculated for log-transformed data.

Abbreviations: CAD, coronary artery disease; CV, cardiovascular; HDL, high-density lipoprotein cholesterol; HF, heart failure; LDL, low-density lipoprotein cholesterol; SCD, sudden cardiac death; SE, standard error; TG, triglyceride.

^aAdjusted for age, gender, smoking, diabetes mellitus, hypertension, heart failure at admission, STEMI and hospital site.

^bAdjusted for age, gender, smoking, diabetes mellitus, hypertension, heart failure at admission, STEMI, hospital site and the use of statins.

of all ages from various countries have shown that neither TC nor LDL-C is a risk factor of mortality, neither among statin-treated patients nor among untreated individuals.²¹⁻²⁵ In fact, as shown by our results as well, low values are associated with increased total mortality. This finding challenges the view that high cholesterol is the cause of early death from CVD as well because the commonest cause of death in most countries is CVD. Furthermore, in the study by Ravnskov et al,²⁶ CVD mortality was recorded as well in seven of the 19 studies they had included in their review, and in six of them, LDL-C was not associated with CVD mortality and in one of them, CVD mortality was highest among those with the lowest LDL-C.

In many studies, increased serum lipids and lipoproteins are
 associated with a higher risk of total and cardiac mortality.²⁷⁻³⁵

However, absence of an association, or even an inverse association, has been reported, ⁵⁻⁷ especially with increasing age.^{26,36} In accordance, prior observational studies have reported paradoxically better outcomes in hypercholesterolaemic patients who sustain acute coronary syndrome, the so-called 'cholesterol paradox'.^{20,37-39}

The occurrence of acute coronary syndrome is accompanied by transient changes in the plasma lipid profile, including decreases in TC, LDL and HDL levels and increases in plasma TG.^{40,41} The prognostic value of plasma lipids at admission was investigated in prior studies, and a hypercholesterolaemia paradox has also been observed.^{8-10,20,42-45}

Importantly, the majority of these reports were performed in the short- to intermediate term; the longest was approximately 3 years.

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Furthermore, a single LDL or TG level at admission was used, accordingly, the effect of the changes in their levels on clinical outcomes could not be assessed. Notably, these reports tended to focus on total mortality.^{8-10,20,42-45}

5 Although it was not significant, we also observed the negative 6 trend (hypercholesterolaemia paradox). Our different prognostic results support the assumption of the presence of other modifiable 7 8 risk factors, other than cholesterol levels, that deliberates this risk 9 in the majority of patients and may explain why patients develop myocardial infarction despite a low LDL level ^{19,39} or why a consid-10 erable proportion of major adverse events are still not prevented 11 6 by aggressive lipid-lowering treatment.⁴⁶ We also observed during 12 13 the long-term follow-up that non-cardiac death was more prevalent, 14 and an inverse association with plasma lipid levels was observed in concordance with previous reports.^{5,47,48} This observation is clini-15 cally relevant if we consider that the non-cardiac mortality rate after 16 17 acute coronary syndrome has progressively increased in recent 18 years.^{49,50} We could assume that the improvement in CV treatment tends to reduce CV mortality, allowing non-cardiac disorders and 19 20 their complications to prevail. In agreement with our findings, other authors have reported an unexplained absence of an independent 21 association between cholesterol and stroke mortality, especially at 22 older ages or in the presence of higher blood pressures.³² Age is also 23 24 a considerable factor, as the median age of our cohort at the time of 25 admission was approximately 67 years, and it is understandable that the prognostic role of plasma lipids changes with the ageing of the 26 population. In a systematic review, Ravnskov et al ²⁶reported a lack 27 28 of an association or inverse association between LDL and mortality 29 in elderly people. In short, our study provides new clinical insight 30 and perspective into the long-term prognostic significance of ad-31 mission plasma lipid levels in survivors of acute coronary syndrome.

The present prospective long-term study revealed that baseline plasma lipid levels during acute coronary syndrome seem not to be associated with long-term global mortality. Only an independent inverse association between TC and LDL and non-cardiac death has been observed.

5 | STUDY LIMITATIONS

A major limitation of the ABC study of acute coronary syndrome 41 was that, at the time of patient enrolment, percutaneous coronary 42 angioplasty was not yet used to reopen coronary arteries in pa-43 tients with STEMI. Thus, it remains uncertain whether the results 44 45 may have been altered by early mechanical reperfusion. Although Cho et al⁴³ reported that admission LDL-cholesterol level was not 46 47 an independent predictor of mortality at 12 months among 9571 patients with acute myocardial infarction who had a percutaneous 48 coronary intervention (PCI). Pres et al⁴⁵ also reported that LDL level 49 50 at admissions was not associated with increased in-hospital mor-51 tality in non-diabetic patients treated with PCI for STEMI, and Al-52 Mallah et al⁸ reported an inverse association, despite that more than 53 60% of their patients have been subjected to PCI or coronary artery

bypass graft surgery. Furthermore, statin treatment was much less commonly used at the beginning of the study period (1995-1998) and steadily increased from the 1st to the 20th year of follow-up in accordance with guideline revisions over that time period. Finally, as the patients in this study were all Caucasians, we cannot generalise the present findings to other populations and ethnic groups.

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DISCLOSURES

None.

AUTHOR CONTRIBUTIONS

Dr G. Berton and Dr F. Cavuto designed the study. Dr R. Cordiano and Dr F. Bagato contributed to the original data collection. Dr F. Cavuto and Dr R. Cordiano contributed to data handling and patient follow-up. Dr G. Berton and Dr HT Mahmoud contributed to the data analysis and interpretation, tables and figures preparation and manuscript preparation. All authors contributed to ensuring the accuracy of the data analysis.

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

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