



AperTO - Archivio Istituzionale Open Access dell'Università di Torino

Antemortem characterization of sudden deaths as first-manifestation in Italy

This is the author's manuscript				
Original Citation:				
Availability:				
This version is available http://hdl.handle.net/2318/1814202 since 2022-05-28T17:07:18Z				
Published version:				
DOI:10.1007/s10840-021-00949-5				
Terms of use:				
Open Access				
Anyone can freely access the full text of works made available as "Open Access". Works made available under a Creative Commons license can be used according to the terms and conditions of said license. Use of all other works requires consent of the right holder (author or publisher) if not exempted from copyright protection by the applicable law.				

(Article begins on next page)

1	Ante-mortem characterization of sudden deaths as
2	first-manifestation in Italy.
3	
4	Frontera A, MD; ^{1,4} Anselmino M, MD ^e ; Matta M, MD ^e ; Baccelli A, MD ¹ ; Vlachos K, MD ⁴ ; Bonsignore A, MD,
5	PhD [°] ; Camaioni C, MD [°] , Notarstefano P, MD [°] ; Mahida S, MD, PhD [°] ; Nesti M, MD [°] ; Sacher F, MD, PhD [*] ; Tunzi
6	R. M, MD [*] ; Landoni G, MD [*] ; Aschieri D, MD ¹⁰ ; Castelli V, MD; ¹¹ M. Hocini, MD ⁺ ; Jaïs P, MD ⁺ ; Gaita F, MD [*] ;
7	Derval N, MD ⁺ ; Haïssaguerre M, MD. ⁺
8	
9	¹ Arrhythmology Department, IRCCS San Raffaele Scientific Institute and Vita-Salute San Raffaele University,
10	Milan, Italy
11	² Cardiology Division, "Città Della Salute e della Scienza di Torino" Hospital, Department of Medical Sciences,
12	University of Turin, Turin, Italy
13	³ Department of Legal and Forensic Medicine, University of Genova, Genova, Italy
14	*Electrophysiology Department, LIRYC Institute, Bordeaux University Hospital, Bordeaux, France
15	⁵ Cardiology Department, Istituto clinico Città Studi, Milan, Italy
16	° Cardiology Department, San Donato Hospital, Arezzo, Italy
17	⁷ Department of Cardiac Electrophysiology and Inherited Cardiac Diseases, Liverpool Heart and Chest Hospital,
18	United Kingdom
19	[*] Cardiology Department, University of Bari, Bari, Italy
20	⁹ Anesthesia and Intensive Care Department, IRCCS San Raffaele Scientific Institute and Vita-Salute San Raffaele
21	University, Milan, Italy
22	¹⁰ Cardiology Department, Castel San Giovanni Hospital, Piacenza, Italy
23 24	'' Fondazione Giorgio Castelli ONLUS, Roma, Italy
25	Corresponding author:
26	Dr. Andrea Baccelli, MD. Arrhythmology Department, San Raffaele Hospital, Via Olgettina 60, 20132 Milan,
27	Italy. E-mail: andrea.baccelli@unimi.it
28	
29	Words: 2508
30	
31	Keywords: Sudden death; young; cardiopulmonary resuscitation; idiopathic VF, young adults.
32	
33	

34 Abstract

- 35 Purpose. There is a relative paucity of data on ante-mortem clinical characteristics of young (age 1 36 to 35 years) sudden death (SD) victims. Aim of the study was to characterize ante-mortem 37 characteristics of SD victims, in a selected national cohort identified by web search.
- 38

39 Methods. A dataset of all SD, between January 2010 and December 2015 was built from the national 40 forensic and medical records, integrated with GoogleTM search module. Families were contacted to 41 obtain consent for interviews. Data were obtained on ante-mortem symptoms. ECG and autopsy data 42 were collected where available.

43

44 Results. Out of 301 SD cases collected, medical and family history was available in 132 (43.9%) 45 patients. Twenty-eight (21.1%) had a positive family history for SD. SD occurred during sport/effort 46 in 76 (57.6%). One hundred and twelve (85%) SD cases had no prior symptoms. Autopsy data was 47 available in 100 out of 132 (75,8%) cases: extra cardiac cause was identified in 20 (20%). Amongst the 48 61 cases with a cardiac diagnosis, 21 (34%) had hypertrophic cardiomyopathy. Among the 19 (19%) 49 victims without no structural abnormalities, molecular autopsy identified pathogenic variants for 50 channelopathies in 9 cases. Ten (10%) victims had no identifiable cause. 51 52 **Conclusions**. Most SD were due to cardiac causes and occurred in previously asymptomatic patients,

52 **Conclusions**. Most SD were due to cardiac causes and occurred in previously asymptomatic patients, 53 mainly during strenuous activity. In a minority of cases, no cause was identified. The web-based 54 selection criteria, and incomplete data retrieval, need to be carefully taken into account for data 55 interpretation and reproducibility.

56

57

59 Introduction

Sudden death (SD) is a significant public health concern, particularly devastating in young patients. In cohorts of subjects aged 1 to 40 years, the incidence of SD has been reported in the range of 0.9 to 8.5 per 100,000 person-years,¹⁻³ with cardiac disease accounting for the vast majority of non-traumatic cases.⁴⁻⁷

64 Sudden arrhythmic death occurs more commonly at rest or during sleep and up to 90% of the 65 victims do not present prior symptoms or identifiable risk factors for Sudden Arrhythmic 66 Death Syndrome (SADS).⁸ An autopsy-based series of a US military population (18 years of 67 age and over) undergoing active surveillance, however, reported that 68.4% of SD were 68 witnessed; within the latter, 40% were temporally associated with exertion, and a prodrome 69 was documented within one week from the terminal event in half of the victims.⁹ Therefore, 70 despite being, by definition, unexpected and rapid, SD may be seen, from a temporal 71 perspective, as a sequence of events that may manifest weeks before the terminal event.

72 Due to the lack of data on ante-mortem characteristics of young SD victims, our study aimed 73 to analyze the ante-mortem characteristics of SD as the first manifestation in young adults 74 (<35 years old) in a nationwide population-based analysis study.</p>

75

76 Methods

Patient cohort. A dataset of young SD victims (age 1 to 35 years) who suffered SD as a first clinical manifestation, between January 1st 2010 and December 31st 2015, was created. In the absence of a national registry on sudden death, the dataset was built based on the national forensic and medical records, integrated by GoogleTM search module (thirty-one Italian search items were used, see full details in the *Supplemental table X*). 82 SD was defined as follows: non-traumatic, unexpected fatal event occurring within one hour 83 from the onset of symptoms in a healthy subject (if death was not witnessed, the definition 84 applies when the victim was in good health 24 hours before the event).

Inclusion criteria were the absence of known disease (cardiac or extracardiac), and positive toxicology screening (Figure 1). Of the total 412 SD victims identified, after careful review by two independent researchers (A.F. and M.A.), 111 did not satisfy inclusion criteria (see details in *Supplemental figure* 1).

89 SD was defined as unexplained when no cause was identified after a complete and90 comprehensive autopsy examination, including histologic and toxicological studies.

91

92 Data collection. The investigators contacted the local hospitals where the SD was 93 registered. Family members and/or 1st-degree relatives of the SD victim were offered 94 participation to the study and contacted by the investigators to obtain formal consent for 95 interviews. A standardized questionnaire which focused on ante-mortem, as well as post-96 mortem (autopsy and genetic tests) data, was used (details int the Supplemental data). The 97 questionnaire specifically focused on: 1) SD prodromes, 2) past medical history, including 98 congenital abnormalities and abnormal anthropometric measurements, 3) detailed breakdown 99 of the sports participation, 4) first assistance given to the victim (e.g. cardio-pulmonary 100 resuscitation), presence of on-site medical personnel, Basic Life Support and early 101 defibrillation with an automatic external defibrillator (BLSD), proficiency among rescuers, 102 AED availability, first recorded rhythm at AED arrival, number of shocks. Where available, 103 ECG and autopsy data were collected from the relatives, as well as from forensic and medical 104 records and sport medicine physicians distributed along the Italian territory. Each case was carefully investigated, and available witnesses were interviewed personally by the 105 106 investigators. Additionally, primary data was retrieved via the EMS run sheets. Any missing 107 data prevented the enrollment of the patient in the study. Autopsies were performed at the

108 respective centres according to Italian pathologists guidelines, recommending collection of 109 histological and toxicological data (including a panel of most common abuse substances 110 and alcohol). The study protocol was approved by the local ethics committee at the "Città 111 della Salute e della Scienza di Torino" Hospital and Bordeaux.

112

Statistics. Continuous variables were expressed as mean +/- standard deviation (SD) and categorical variables as number and percentage. A t-test or a Chi-square test was performed for comparison of continuous and categorical variables, respectively. P values < 0.05 were considered statistically significant. Data were analyzed using SPSS software 20.

117

Patient and public involvement. This research was performed with the participation of the victims' relatives, who represented the largest source of ante-mortem data of the victims. Relatives were not invited to comment on the study design or interpret the results.

122 **Results**

Study population. During the study period (from 2010 through 2015), Italy had a mean population of 60.2 million residents, of whom 21.1 million were in the age group 1 to 35 years.¹⁰ The number of deaths in this age group and period was 7304.

In the present study we identified 301 cases of SD, yielding an estimated annual incidence of
sudden unexpected death in Italy of 0.24 cases per 100,000 persons (95% confidence interval,
0.13 to 0.45). Males had a higher incidence of SD (0.4 vs. 0.1 cases per 100,000 persons, P <
0.001).

130

131 Majority of the victims were males (81%) and SD occurred at a mean age of 22 ± 7.9 years 132 (Figure 2). 28 / 132 (21.1%) cases had a family history of SD (first-degree relatives in 9% of

133	cases; second/third degree relatives in 12.1%). 101 / 132 (76.5%) practiced sport; 67 of which
134	(50.8%) at high-level (Table 1)
135	
136	
137	
138	
139	

Age, mean (± SD), years	<i>22 ± 7.9</i>
Male, n (%)	107 (81%)
Ethnicity, n (%)	
Caucasian	126 (95.4%)
African	4 (3.0%)
Hispanic	1 (0.8%)
Asiatic	1 (0.8%)
BMI, mean (± SD)	24 ± 6.5
Distinctive features, n (%)	
Multiple nevi	5 (3.8%)
Marfanoid habitus	3 (2.3%)
Pectus excavatum	1 (0.8%)
Pectus carinatum	1 (0.8%)
Region, n (%)	
North	35~(26.6%)
Centre	46 (34.8%)
South	51 (38.6%)
Occupational Status, n (%)	
Student	64~(48.5%)
Employee	17 (12.9%)
Athlete	8 (6.0%)
Other	43 (32.6%)

Sport activity, n (%)	
Active (>5 hr/wk)	43 (32.6%)
Amateur	34(25.8%)
None	31 (23.5%)
Competitive	24 (18.2%)

142 Table 1. Demographic characteristics of the sudden death cohort

143

144 Ante-mortem characteristics. 112/132 (84.8%) victims had no prior symptoms, 13 (9.8%) 145 reported intense fatigue in the 2 weeks preceding the SD event, 3 (2.3%) suffered from 146 palpitations in the last 3 months, 2 (1.5%) had a history of syncope, and 2 (1.5%) of pre-147 syncope. Ten SD victims (7.6%) were on antibiotic therapy at the time of death (penicillin n= 148 4; fluoroquinolones n=3; macrolides n=2; sulfonamides n=1). Fourteen (10.6%) victims had 149 associated pyrexia on the day or days before the event (Table 2). One victim, a male 150 professional bodybuilder, reported regular use of anabolic steroids; post-mortem examination 151 revealed a previously undiagnosed hypertrophic cardiomyopathy (with a heart weight of 780 152 g). 4 victims, all male, reported use of protein powder; of these, 2 did not undergo post-153 mortem examination, 1 had a final diagnosis of HCM and 1 of a ruptured cerebral aneurysm 154 (Table 2).

Symptoms before SD, n (%)	
No prior symptoms	112 (84.8%)
Fatigue	13 (9.8%)
Palpitations	3 (2.3%)
Syncope	2 (1.5%)
Pre-syncope	2 (1.5%)
Allergies, n (%)	9 (6.8%)
Pulmonary disease, n (%)	
History of pneumonia	8 (6%)
Asthma	3(2%)
Tuberculosis	1 (0.8%)
Previous surgery, n (%)	19 (14.4%)

History of seizures, n (%)	3 (2.3%)		
Associated pyrexia, n (%)	14 (10.6%)		
Recent trauma, n (%)	5 (3.8%)		
Medical therapy at the time of SD, n (%)			
NSAID	4 (3%)		
Antibiotics	10 (7.6%)		
Oral Contraceptives	3 (12%)ተ		
Corticosteroids	2 (1.5%)		
Insulin	1 (1%)		
Anti-epileptic drugs	1 (1%)		
No therapy	111 (84.1%)		
Cardiovascular Risk factors, n (%)			
Smoke	10 (7.6%)		
Hypertension	0		
Type 1 Diabetes Mellitus	2(1.5%)		
Drugs addiction, n (%)	5 (3.8%)		
Energy drinks routine use, n (%)	12 (9.1%)		
Anabolic steroids, n (%)	1 (1%)		
Protein Powder, n (%)	4 (3%)		
Sudden increase in size in the last 3 months	5 (3.7%)+		
(when applicable), n (%)			
≁among female victims			
‡ among victims < 20 years of age			

- **Table 2**. Clinical and behavioural variables of the sudden death victims
- 158

Circumstances of death. A significantly higher number of SD events occurred in the winter months (p=0.1, **Table 3**). In 116/132 (87.9%), the SD event occurred in an urban setting. 78 (57.6%) of SD events occurred in the context of high adrenergic tone (sport n=37, physical effort n=31; immediate post-effort recovery n=10). In 5 cases (3.8%), SD took place during intense emotional stress. In 30 (22.2%) cases SD occurred at rest and in 12 (8.9%) while sleeping. 101 (76.5%) of SD events were witnessed (in sporting facilities n= 45; at home n= 20; at school n = 12; other settings n= 24).

Time, n (%)	
Morning (6am – 12pm)	32 (24.2%)
Afternoon/evening (12pm – 11pm)	71 (53.8%)
Night (11pm – 6am)	33 (25%)
Season, n (%)	
Winter	42(31.8%)
Spring	33 (25%)
Summer	27 (20.5%)
Autumn	30 (22.7%)
Location, n (%)	
Urban	116 (87.9%)
Rural	16 (12.1%)
Witnessed, n (%)	101 (76.5%)
SD occurred during, n (%)	
Sport/ Effort/ Post-effort recovery	76 (57.6%)
Rest/ Sleep	42 (31.8%)
Emotional stress	5 (3.8%)
Miscellanea	9~(6.8%)
Unknown	
SD took place at	
Sport facility	49 (37.1%)
Home	43 (32.6%)
School	12 (9.1%)
Other (i.e. public streets, bars, beaches etc.)	28 (21.2%)

Table 3. Circumstances of sudden death

Resuscitation data. Amongst the 101 witnessed SD events, CPR began promptly (within 1
minute) in 28 cases (27.7%). Overall, 120/132 (91%) SD cases received CPR. An Automatic
External Defibrillator (AED) was available onsite in 19 cases (14.4%). 45 of 120 rescuers
(37.5%) were BLS-certified. 41 (31%) victims presented with gasping, and 31 (23.5%) had
their eyes wide open with a fixed gaze. The first recorded rhythm obtained by AED analysis

175 at arrival was available in 101 victims (56 [55.4%] VF; 39 [38.6%] asystole; 6 [6%] pulseless

- electrical activity). 55/132 (41.7%) victims received at least one shock. Complete data on the
 immediate management by bystanders and emergency medical personnel are included in
- 178 **Table 4**.

Behaviour of the victim, n (%)	
Gasping	41 (31%)
Eyes wide open	31 (23.5%)
Seizure-like movements	28 (21.2%)
Urine/stool emission	24(18.2%)
Vomit	13 (9.8%)
AED available onsite, n (%)	19 (14.4%)
AED onsite (only sports centers, $n=49$)	7 (14.3%)
CPR began promptly (within 1 minute), n (% out of witnessed SD)	28~(27.7%)
CPR performed by BLS certified bystander, n (% out of total nr of CPRs)	45 (37.5%)
Ventilation performed, n (%)	30 (22,7%)
Rhythm at AED arrival, n (% among available tracings, n= 101)	
VF	56(55.4%)
ASY	39~(38.6%)
PEA	6 (6%)
Shocked at least once, n (%)	55 (41,7%)

180 Table 4. Resuscitation data

181

182 Electrocardiographic data. ECGs were available in 45/ out of 132 (34%) subjects, and in 183 8% of the cases it was performed as routine preparticipation sports screening. 4 presented 184 incomplete right bundle block, 9 had early repolarization, 1 short QT interval, 1 long QT 185 interval. Mean PR interval was 171 ± 14 ms. Mean QRS duration was 93 ± 11 ms. The patient 186 with a short QT interval had 289 ms (*Figure 3, supplementary material*). The patient with 187 long QT interval had 540 ms and was undertaking an antibiotic therapy (penicillin). The site 188 of early repolarization was inferior with horizontal slope in all cases. 189

190 Post-mortem characteristics. Autopsy data were available in 100 cases. Cardiac structural
191 disease accounted for 61 (61%) of all autopsied SD events. Extracardiac causes were identified

192 in 20 cases (20%). There were no identified structural abnormalities in 19 (19%) of the 193 available autopsies (Figure 4-A). Among the sudden cardiac deaths, 21/ out of 61 (34.4%) 194 documented Hypertrophic Cardiomyopathy (HCM), 16 (26.2%) Arrhythmogenic Right 195 Ventricular Cardiomyopathy (ARVC), 14 (23%) Ischemic Heart Disease (IHD), 7 (11.5%) 196 Myocarditis, 2 Aortic Dissection (3.3%), and 1 (1.6%) an anomalous origin of the coronary 197 arteries (Figure 4-B). Extracardiac causes of SD included cerebrovascular events (n=15, 75% 198 among extracardiac SD), possible drowning (n=1), sepsis (n=1), and aortic dissection (n=3). 199 Among the 61 cases presenting structural cardiac disease, the final diagnosis was achieved by 200 gross and histopathologic studies, without genetic testing. Among the 19 victims without 201 signs of macroscopic or microscopic cardiac structural disease, genetic testing for variants 202 implicated in heritable cardiomyopathies and/or channelopathies (long OT syndrome, short 203 OT syndrome, Brugada syndrome, and catecholaminergic polymorphic ventricular 204 tachycardia) was performed as per consensus guidelines. In the aforementioned 19 cases, a 205 final molecular diagnosis of Brugada Syndrome (BrS) was reached in 3, Long QT Syndrome 206 (LQTS) in 5 patients, and Catecholaminergic Polymorphic Ventricular Tachycardia (CPVT) 207 in 1 patient. The cause of death was not identified in 10 cases (Sudden Unexplained Death).

208

The cause of death varied according to age group; considering the four most common causes of SCD, HCM and myocarditis were the most frequent causes of SD in the 16 to 20 age group, ARVC was the most frequent cause of SD in the 26 to 30 age group. The incidence of IHD as a cause of SD increased progressively from the ages of 21 to 35, with no reported incidences in the 16 to 20-year age group (**Figure 5**).

214

215 Clinical screening among relatives of the victims. Screening data among relatives (parents 216 and siblings) was available in 70 SD cases. In each of these, at least an ECG or echocardiogram 217 was performed. Genetic testing was performed in only 5 cases. A definite clinical diagnosis was established in 6 of the 70 (8.5%) screened families (CPVT, n=1; BrS, n=1; ARVC, n=2; HCM, n=1). There was one diagnosis of aortic stenosis in the mother of a victim. No deaths occurred amongst relatives of the victims over 41 ± 11 months following the index event in the proband.

222

223 **Discussion**

224 In the present study, based on a national cohort identified by web search, we report ante-225 mortem and post-mortem data from a selection of SD cases aged 1 - 35 years. The findings 226 are as follows: 1) Only 15% of subjects had symptoms before the SD event, with fatigue, 227 palpitations, syncope and pre-syncope being the most commonly reported symptoms from 228 patient relatives. 2) Close to two-thirds of SD events occurred in the context of high 229 adrenergic tone. 3) The age-related distribution of SD showed a bimodal trend, with the first 230 peak at 15-20 years and a second peak at 30-35, with the underlying causes differing 231 significantly between age groups.

232 The estimated annual incidence of sudden unexpected death in Italy in the study period was 0.24. However, a recent nationwide study¹¹ has reported estimates up to $\sim 1.3/100,000$ -233 234 person-years. Web-based population selection, together with difficulties in complete data 235 retrieval, therefore need to be carefully considered for data interpretation and reproducibility. 236 However, given the paucity of data on ante-mortem clinical characteristics of young SD 237 victims, in our opinion, the present study provides interesting insights on the terminal event 238 leading to SD, in which, despite an attempted resuscitation in almost every victim, there was 239 not, in any cases and at any moment, a return of spontaneous circulation (ROSC).

240

The majority of SD events in the present study were associated with high adrenergic tonerelated activities. These findings contrast with the results of Bagnall et al. ¹¹, where most of

the deaths occurred at rest or during sleep. There are several potential explanations for this 243 244 discrepancy. In our study, the leading causes of SD were ARVC, HCM and IHD, conditions 245 that have a strong correlation with physical exercise-induced SD. The most frequent causes 246 of death in the study by Bagnall et al., instead, were Sudden Unexplained Cardiac Death (with 247 an overall diagnostic yield of genetic testing of 27%, mostly for channelopathies), CAD and 248 DCM. The presumably higher rate of primary arrhythmic syndromes may then account for 249 the higher rate of parasympathetic and non-stress-related adrenergic circumstances of death. 250 Another potential explanation is population selection bias, with SD events occurring in the 251 context of sporting activity being more likely to be covered by media and therefore identified 252 by the Google search[™] module. In this respect, the abovementioned study by Bagnall and 253 colleagues, in which all cases of sudden death in subjects 1 to 35 years of age were 254 prospectively collected by forensic pathology centers, probably represents a more accurate 255 and representative data.

256

257 Concerning acute management of SD events, even though the majority of SD events were 258 witnessed, less than a third received immediate CPR (defined as CPR delivery within 1 259 minute). Furthermore, only a minority of resuscitation providers were BLS trained, and 260 AEDs were available in an even smaller proportion of cases.

These findings further underscore the importance of BLS-D training promotions and campaigns for more widespread availability of AED to improve the outcome of resuscitation.

SD was associated with a bimodal trend in the present study, with the first peak at 15-20 and a second peak at 30-35 years of age. The underlying causes are significantly different between age groups, with myocarditis and HCM representing the most common findings in the first age group, and IHD and HCM accounting for more than half of SCD in the latter. The most common finding at autopsy was a cardiac structural abnormality, followed by an extracardiac 269 structural pathology. However, a fifth of the cases had no evidence of structural anomalies. 270 Genetic analysis carried out on the latter group reached a diagnostic yield of 47%. Therefore, 271 autopsy combined with molecular analysis was associated with a substantially higher 272 likelihood of determining the cause of SD. Acknowledging the potential for misclassification 273 of abuse substances overdose sudden deaths as cardiac in the absence of a toxicology 274 screening at autopsy, as reported by two recent large reports^{12,13}, in Italy, in cases of SDs, the 275 Public Prosecutor commonly regests complete forensic investigation, including the 276 collection of toxicological data. Accordingly, in our Country no particular risks about this misclassification exist. 277

278

279 Clinical perspectives

280 It is still controversial whether mass screening programs hold a favorable cost-effectiveness 281 to prevent SD among children and young adults. However, our study shows that a significant 282 proportion of victims had a positive family history for SD, thus suggesting the importance of 283 a targeted and thorough cardiological and genetic examination among first and second-284 degree relatives by a multidisciplinary team. Since approximately 40% of SCD in the young 285 remain unexplained after the autopsy and given the genetic basis of most underlying 286 cardiovascular causes, a specific screening should always be preceded by a complete and 287 accurate autopsy alongside DNA storage of the victim. In this setting, autopsy plays an 288 essential role, and should be performed in any case of SD in the young, always associated to 289 genetic testing, not only in unexplained cases but also in the structurally abnormal hearts, in 290 search of genotype-phenotype associations. Standard molecular autopsy panels will typically 291 include the four main genes accounting for a significant number of previously unexplained SCDs, including KCNQ1 (LQT1), KCNH2 (LQT2), SCN5A (LQT3/BrS1), and RYR2 292 293 (CPVT1).

Finally, the results of our investigation call for further AED diffusion and BLS-D trainingcampaigns among the general population.

296

297 Limitations

298 The present study has several limitations. The authors acknowledge the fact that the dataset 299 is not based on a prospective national registry, and that poorer and minority populations not 300 covered by the media might have been excluded. We also acknowledge that only 132 of 301 301 SD cases were included for analysis. Thus a large majority of cases were not thoroughly 302 examined and the cohort analyzed may substantially be biased relative to the (unknown) 303 young SD population in Italy during the study period. Secondly, autopsies and genetic 304 testing, when performed, were carried out at different Institutions throughout the Country, 305 with significant methodological differences and accuracy. Lastly, this was not a 306 geographically or ethnically diverse population, and the results might not apply to other 307 geographic regions and other populations.

308

309 **Conclusions**

The majority of sudden unexpected deaths in patients under 35 years are due to cardiac causes. Cardiomyopathies prevail in younger age, while ischemic heart diseases in the older. The vast majority of the SD occur in previously asymptomatic patients, mainly during strenuous activity, further highlighting the importance of screening programs. In 10% of the cases no clear cause of SD could be identified. Web-based population selection, together with difficulties in complete data retrieval, need to be carefully considered for data interpretation and reproducibility.

317

319 Acknowledgements

320	This work is dedicated to all families who have lost their son and daughter. The authors
321	would like to thank Dr Andrea Ghidini Ottonelli for his precious collaboration.
322	
323	Conflicts of interest: none
324	
325	Funding: This research did not receive any specific grant from funding agencies in the public,
326	commercial, or not-for-profit sectors.
327	
328	
329	
330	
331	
332	
333	
334	
335	
336	
337	
338	
339	
340	
341	
342	
343	

344 **References**

- 1. Shen, W. K. *et al.* Sudden unexpected nontraumatic death in 54 young adults: a 30-year
- 347 population-based study. *Am. J. Cardiol.* **76**, 148–152 (1995).
- 348 2. Liberthson, R. R. Sudden Death from Cardiac Causes in Children and Young Adults.
- 349 New England Journal of Medicine **334**, 1039–1044 (1996).
- 350 3. Wisten, A., Andersson, S., Forsberg, H., Krantz, P. & Messner, T. Sudden cardiac death
- in the young in Sweden: electrocardiogram in relation to forensic diagnosis. *Journal of*
- 352 Internal Medicine **255**, 213–220 (2004).
- 353 4. Drory, Y. *et al.* Sudden unexpected death in persons <40 years of age. *American Journal*354 of Cardiology 68, 1388–1392 (1991).
- 355 5. Vaartjes, I. *et al.* Sudden death in persons younger than 40 years of age: incidence and
 356 causes. *European Journal of Cardiovascular Prevention & Rehabilitation* 16, 592–596
 357 (2009).
- 358 6. Neuspiel, D. R. & Kuller, L. H. Sudden and Unexpected Natural Death in Childhood and
 359 Adolescence. JAMA 254, 1321–1325 (1985).
- 360 7. Wren, C. Sudden death in children and adolescents. *Heart* **88**, 426–431 (2002).
- 361 8. Mellor, G. et al. Clinical Characteristics and Circumstances of Death in the Sudden
- 362 Arrhythmic Death Syndrome. *Circulation: Arrhythmia and Electrophysiology* 7, 1078–1083
 363 (2014).
- 364 9. Eckart, R. E. et al. Sudden Death in Young Adults: An Autopsy-Based Series of a
- 365 Population Undergoing Active Surveillance. Journal of the American College of Cardiology
- **366 58**, 1254–1261 (2011).
- 367 10. Annuario statistico italiano 2018. https://www.istat.it/it/archivio/225274.

368	11. Bagnall, R. D. et al. A Prospective Study of Sudden Cardiac Death among Children and
369	Young Adults. New England Journal of Medicine 374, 2441–2452 (2016).
370	12. Tseng, Z. H. et al. Prospective Countywide Surveillance and Autopsy Characterization
371	of Sudden Cardiac Death: POST SCD Study. Circulation 137, 2689–2700 (2018).
372	13. Bjune, T. et al. Post-mortem toxicology in young sudden cardiac death victims: a
373	nationwide cohort study. <i>Europace</i> 20 , 614–621 (2018).
374	
375	
376	
377	
378	
379	
380	
381	
382	
383	
384	
385	
386	
387	
388	
389	
390	
391	
392	

393 Contributorship statements

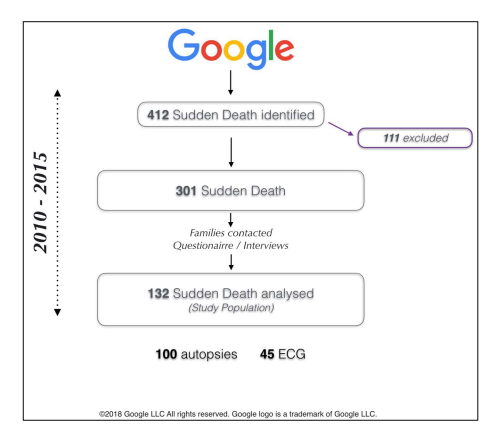
394	Dr Frontera	, Dr Anselm	ino, Dr Matta	, Dr Aschieri,	Dr Castelli,	Dr Gaita, Dr	· Notarstefano,

- 395 Dr Haissaguerre, Dr Baccelli, Dr Bonsignore and Dr Landoni contributed to the planning of
- 396 the work described in the article;
- 397 Dr Frontera, Dr Baccelli, Dr Nesti, Dr Tunzi, Dr Bonsignore, Dr Camaioni, Dr Notarstefano
- 398 contributed to the conduct of the work described in the article;
- 399 Dr Frontera, Dr Baccelli, Dr Sacher, Dr Mahida, Dr Vlachos, Dr Derval, Dr Jais, Dr Hocini

400 contributed to the reporting of the work described in the article;

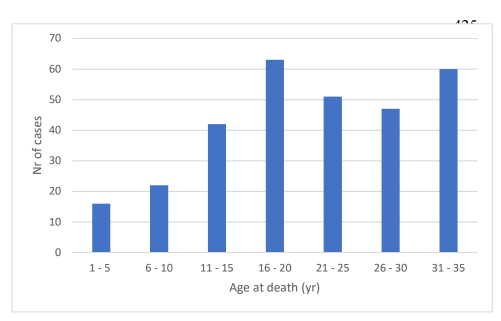
- 401 Dr Frontera is responsible for the overall content as guarantor.

418 Figures

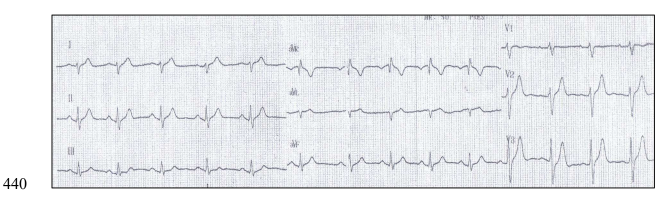


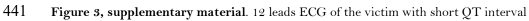
420 Figure 1. Cases of Sudden Death. Amongst the 412 SD identified through google search module, 111 were
421 excluded because were toxicology positive, or were cases of homicide/suicide. Our analysis focused on 301
422 sudden death: families were contacted and data collected.

423



435 Figure 2. Sudden death according to age group (N=301)





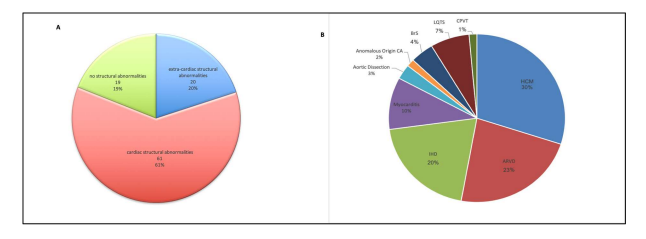
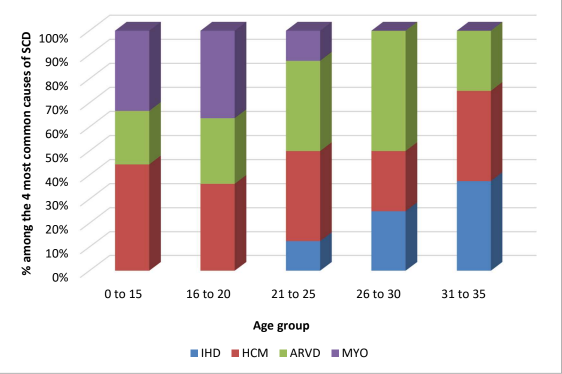


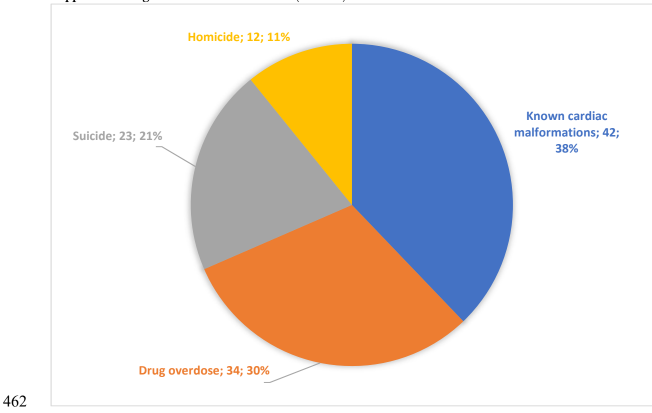
Figure 4. Left and right panel respectively show causes of sudden death and of sudden cardiac death. IHD: ischemic heart disease, HCM: hypertrophic Cardiomyopathy, ARVD: arrhythmogenic right ventricular dysplasia, MYO: myocarditis, BrS: Brugada Syndrome, LQTS: Long QT Syndrome, CPVT: Catecholaminergic Polymorphic Ventricular Tachycardia



454 Figure 5. Trends of the 4 most common causes of SCD by age group. IHD: ischemic heart disease, HCM: hypertrophic Cardiomyopathy, ARVD: arrhythmogenic right ventricular dysplasia, MYO: myocarditis

459 SUPPLEMENTAL FIGURE 1

460



461 Supplemental figure 1. Excluded SD cases (n = 111)