



AperTO - Archivio Istituzionale Open Access dell'Università di Torino

Impact of aspirin on takotsubo syndrome: a propensity score-based analysis of the InterTAK Registry

This is a pre print version of the following article:				
Original Citation:				
Availability:				
This version is available	http://hdl.handle.net/2318/1725723	since	2020-01-29T09:42:07Z	
Published version:				
DOI:10.1002/ejhf.1698				
Terms of use:				
Open Access				
Creative Commons licen	s the full text of works made available as "Ope se can be used according to the terms and co ight holder (author or publisher) if not exempte	nditions of sa	aid license. Use of all other works	
•				

(Article begins on next page)

Impact of Aspirin on Takotsubo Syndrome: A Propensity Score Based

2 Analysis of the InterTAK Registry

3

Running title: Aspirin in Takotsubo syndrome

5

6Fabrizio D'Ascenzo¹, MD, PhD; Sebastiano Gili², MD; Maurizio Bertaina¹, MD; Mario Iannaccone¹, MD; 7Victoria L. Cammann³, MD; Davide Di Vece³, MD; Ken Kato³, MD, PhD; Andrea Saglietto¹, MD; Konrad A. 8Szawan³, MD; Antonio H. Frangieh^{4,5}, MD; Beatrice Boffini²; Margherita Annaratone⁶, MS; Annahita Sarcon⁷, 9MD; Rena A Levinson^{3,8}, BS; Jennifer Franke⁹, MD; L. Christian Napp¹⁰, MD; Milosz Jaguszewski¹¹, MD, 10PhD; Michel Noutsias¹², MD; Thomas Münzel¹³, MD; Maike Knorr¹³, MD; Susanne Heiner¹³, MD; Hugo A. 11Katus⁹, MD; Christof Burgdorf¹⁴, MD; Heribert Schunkert^{4,5}, MD; Holger Thiele¹⁵, MD; Johann Bauersachs¹⁰, 12MD; Carsten Tschöpe¹⁶, MD; Burkert M. Pieske¹⁶, MD; Lawrence Rajan¹⁷, MD; Guido Michels¹⁸, MD; Roman 13Pfister¹⁸, MD; Alessandro Cuneo¹⁹, MD; Claudius Jacobshagen²⁰, MD; Gerd Hasenfuß²⁰, MD; Mahir 14Karakas^{21,22}, MD; Wolfgang Koenig^{4,5}, MD; Wolfgang Rottbauer²³, MD; Samir M. Said²⁴, MD; Ruediger C. 15Braun-Dullaeus²⁴, MD; Adrian Banning²⁵, MD; Florim Cuculi²⁶, MD; Richard Kobza²⁶, MD; Thomas A. 16Fischer²⁷, MD; Tuija Vasankari²⁸, MD; K.E. Juhani Airaksinen²⁸, MD, PhD; Grzegorz Opolski²⁹, MD, PhD; 17Rafal Dworakowski³⁰, MD; Philip MacCarthy³⁰, MD, PhD; Christoph Kaiser³¹, MD; Stefan Osswald³¹, MD; 18Leonarda Galiuto³², MD, PhD; Filippo Crea³², MD; Wolfgang Dichtl³³, MD, PhD; Wolfgang M. Franz³³, MD; 19Klaus Empen^{34,35}, MD; Stephan B. Felix^{34,35}, MD; Clément Delmas³⁶, MD; Olivier Lairez³⁶, MD, PhD; Ibrahim 20El-Battrawy^{37,38}, MD; Ibrahim Akin^{37,38}, MD; Martin Borggrefe^{37,38}, MD; John D. Horowitz³⁹, MBBS, PhD; 21Martin Kozel⁴⁰, MD; Petr Tousek⁴⁰, MD; Petr Widimský⁴⁰, MD, PhD; Ekaterina Gilyarova⁴¹, MD; Alexandra 22Shilova⁴¹, MD, PhD; Mikhail Gilyarov⁴¹, MD, PhD; Giuseppe Biondi-Zoccai⁴², MD, MStat; David E. 23Winchester⁴³, MD; Christian Ukena⁴⁴, MD; Michael Neuhaus⁴⁵, MD; Jeroen J. Bax⁴⁶, MD, PhD; Abhiram 24Prasad⁴⁷, MD;, MD; Carlo Di Mario⁴⁹, MD, PhD; Michael Böhm⁴⁴, MD; Mauro Gasparini⁶, PhD; Frank 25Ruschitzka³, MD; Eduardo Bossone⁵⁰, MD, PhD; Rodolfo Citro⁵¹, MD, PhD; Mauro Rinaldi¹, MD; Gaetano 26Maria De Ferrari⁴⁸; Thomas Lüscher^{52,53}, MD; Jelena R. Ghadri³, MD; Christian Templin³, MD, PhD

28 1 Division of Cardiology, Department of Medical Sciences, AOU Città della Salute e della Scienza, University of Turin, 29 Turin, Italy 30 ² Centro cardiologico Monzino, IRCCS, Milan, Italy 31 ³ University Heart Center, Department of Cardiology, University Hospital Zurich, Zurich, Switzerland 32 ⁴ Deutsches Herzzentrum München, Technische Universität München, Munich, Germany 33 ⁵ DZHK (German Centre for Cardiovascular Research), partner site Munich Heart Alliance, Munich, Germany ⁶ Department of Mathematical Sciences, Politecnico di Torino, Turin, Italy 35 ⁷ Section of Cardiac Electrophysiology, Department of Medicine, University of California-San Francisco, San Francisco, 36 CA, USA 37 ⁸ Division of Biological Sciences, University of California San Diego, San Diego, CA, USA 38 ⁹ Department of Cardiology, Heidelberg University Hospital, Heidelberg, Germany ¹⁰ Department of Cardiology and Angiology, Hannover Medical School, Hannover, Germany 39 ¹¹ First Department of Cardiology, Medical University of Gdansk, Gdansk, Poland 41 ¹² Department of Internal Medicine III, Division of Cardiology, Angiology and Intensive Medical Care, University Hospital 42 Halle, Martin-Luther-University Halle, Halle (Saale), Germany 43 ¹³ Center for Cardiology, Cardiology 1, University Medical Center Mainz, Mainz, Germany ¹⁴ Heart and Vascular Centre Bad Bevensen, Bad Bevensen, Germany 44 45 ¹⁵ Heart Center Leipzig - University Hospital, Department of Internal Medicine/Cardiology, Leipzig, Germany ¹⁶ Department of Cardiology, Charité, Campus Rudolf Virchow, Berlin, Germany 46 47 ¹⁷ TJ Health Partners Heart and Vascular, Glasgow, KY, USA ¹⁸ Department of Internal Medicine III, Heart Center University of Cologne, Cologne, Germany 48 49 19 Krankenhaus "Maria Hilf" Medizinische Klinik, Stadtlohn, Germany ²⁰ Clinic for Cardiology and Pneumology, Georg August University Goettingen, Goettingen, Germany 50 51 ²¹ Department of General and Interventional Cardiology, University Heart Center Hamburg, Hamburg, Germany 52 53 54 ²² DZHK (German Centre for Cardiovascular Research), partner site Hamburg/Kiel/Luebeck, Hamburg, Germany ²³ Department of Internal Medicine II - Cardiology, University of Ulm, Medical Center, Ulm, Germany ²⁴ Internal Medicine/Cardiology, Angiology, and Pneumology, Magdeburg University, Magdeburg, Germany 55 ²⁵ Department of Cardiology, John Radcliffe Hospital, Oxford University Hospitals, Oxford, United Kingdom 56 ²⁶ Department of Cardiology, Kantonsspital Lucerne, Lucerne, Switzerland ²⁷ Department of Cardiology, Kantonsspital Winterthur, Winterthur, Switzerland 57 58 ²⁸ Heart Center, Turku University Hospital and University of Turku, Turku, Finland ²⁹ Department of Cardiology, Medical University of Warsaw, Warsaw, Poland

1	30 Department of Cardiology, Kings College Hospital, Kings Health Partners, London, United Kingdom
2	³¹ Department of Cardiology, University Hospital Basel, Basel, Switzerland
3	32 Fondazione Policlinico Universitario A. Gemelli IRCCS, Università Cattolica del Sacro Cuore, Rome, Italy
4	³³ University Hospital for Internal Medicine III (Cardiology and Angiology), Medical University Innsbruck, Innsbruck,
5	Austria
6	³⁴ University Medicine Greifswald, Department of Internal Medicine B, Greifswald, Germany
7	35 DZHK (German Centre for Cardiovascular Research), partner site Greifswald, Greifswald, Germany
8	36 Department of Cardiology and Cardiac Imaging Center, University Hospital of Rangueil, Toulouse, France
9	³⁷ First Department of Medicine, Faculty of Medicine, University Medical Centre Mannheim (UMM) University of
10	Heidelberg, Mannheim, Germany
11	³⁸ DZHK (German Center for Cardiovascular Research), partner site, Heidelberg-Mannheim, Mannheim, Germany
	³⁹ Department of Cardiology, Basil Hetzel Institute, Queen Elizabeth Hospital, University of Adelaide, Adelaide, Australia
13	
	⁴⁰ Cardiocenter, Third Faculty of Medicine, Charles University in Prague and University Hospital Královské Vinohrady,
14	Prague, Czech Republic
15	⁴¹ Intensive coronary care Unit, Moscow City Hospital # 1 named after N. Pirogov, Moscow, Russia
16	⁴² Department of Medico-Surgical Sciences and Biotechnologies, Sapienza University of Rome, Latina, Italy
17	⁴³ Department of Medicine, College of Medicine, University of Florida, Gainesville, FL, USA
18	44 Klinik für Innere Medizin III, Universitätsklinikum des Saarlandes, Homburg/Saar, Germany
19	45 Department of Cardiology, Kantonsspital Frauenfeld, Frauenfeld, Switzerland
20	⁴⁶ Department of Cardiology, Leiden University Medical Centre, Leiden, The Netherlands
21	⁴⁷ Division of Cardiovascular Diseases Mayo Clinic, Rochester, MN, USA
22	⁴⁸ Department of Molecular Medicine University of Pavia, and Cardiac Intensive Care Unit and Laboratories for
23	Experimental Cardiology, IRCCS Fondazione Policlinico San Matteo, Pavia, Italy
24	⁴⁹ Structural Interventional Cardiology, University Hospital Careggi, Florence, Italy
25	50 Division of Cardiology, "Antonio Cardarelli" Hospital, Naples, Italy
26	⁵¹ Heart Department, University Hospital "San Giovanni di Dio e Ruggi d'Aragona", Salerno, Italy
27	⁵² Center for Molecular Cardiology, Schlieren Campus, University of Zurich, Zurich, Switzerland
28	⁵³ Royal Brompton and Harefield Hospitals Trust and Imperial College, London, United Kingdom
29	
رد	
30	
31	
32	
33	
2.4	Correspondence to
34	Correspondence to:
35	Fabrizio D'Ascenzo, MD, PhD
36	Interventional Cardiologist
37	Division of Cardiology, Department of Medical Sciences
38	AOU Citta della Salute e della Scienza, University of Turin
	\cdot
39	Corso Bramante 88-90,
40	10126, Turin, Italy
41	Phone: +39 0116334446
42	E-mail: fabrizio.dascenzo@gmail.com
43	
44	Word count: 1894 (without title, abstract, references, tables, and figure legends)

1ABSTRACT

Aims: The aim of the present study was to investigate the impact of aspirin on prognosis in 3takotsubo syndrome (TTS).

4Methods and Results: Patients from the International Takotsubo Registry (InterTAK Registry) 5were categorized into two groups based on aspirin prescription at discharge. A comparison of 6clinical outcomes between the two groups was performed using an adjusted analysis with 7propensity score stratification; results from the unadjusted analysis were also reported to note the 8effect of the PS adjustment. Major adverse cardiac and cerebrovascular events (MACCE: a 9composite of death, myocardial infarction, TTS recurrence, stroke or transient ischemic attack 10[TIA]) were assessed at 30-day and 5-year follow-up. A total of 1533 TTS patients with known 11status regarding aspirin prescription at discharge were included. According to the adjusted analysis 12based on PS stratification, aspirin was not associated with a lower hazard of MACCE at 30-day 13(Hazard ratio [HR] 1.24, 95% confidence interval [CI] 0.50-3.04, P=0.64) or 5-year follow-up (HR 141.11, 95% CI 0.78-1.58, P=0.58). These results were confirmed by sensitivity analyses performed 15with alternative PS based methods, i.e. covariate adjustment and inverse probability of treatment 16weighting.

Conclusion: In the present study, no association was found between aspirin use in TTS patients 18and a reduced risk of MACCE at 30-day and 5-year follow-up. These findings should be confirmed 19in adequately powered randomized controlled trials. (ClinicalTrials.gov number: NCT01947621)

Keywords: Takotsubo syndrome; acute heart failure; outcome; medical therapy; aspirin

INTRODUCTION

- Takotsubo syndrome (TTS) mostly affects postmenopausal women and is usually preceded 3by an emotional or physical trigger.¹⁻³ Clinical symptoms and signs at presentation, along with 4electrocardiographic (ECG) and laboratory changes, may mimic acute coronary syndrome (ACS) 5or acute heart failure.^{1, 4-6} Although TTS has long been considered a benign condition, recent 6studies reported that it can be associated with significant adverse events both during 7hospitalization and after discharge.^{1, 7-12} Therefore, there is a compelling need for an optimal 8preventive therapy to reduce the incidence of adverse events following TTS. According to recent 9data, angiotensin-converting-enzyme inhibitors or angiotensin-receptor blockers have been 10reported to reduce mortality¹ and recurrence of TTS,¹³ while beta-blockers have not shown 11beneficial effects.^{1, 14} However, data on therapeutic management of TTS are mainly based on small 12case series,¹⁵ meta-analyses,^{13, 16} or retrospective data. There is still a lack of knowledge available 13on optimal treatment strategies.
- During the acute phase of TTS, a thrombogenic state may arise as a consequence of 15catecholamine-dependent ventricular dysfunction, platelet activation, and/or vasoconstriction.¹⁷ 16While anticoagulation therapy in the presence of left ventricular thrombus seems to be an 17appropriate choice, a recent retrospective study has also suggested a protective effect of anti-18platelet therapy during index TTS hospitalization.¹⁸ However, uncertainty persists regarding an 19association between aspirin use and adverse events in TTS patients post-discharge. Therefore, 20the present study aimed to investigate the impact of aspirin use in a large TTS patient cohort 21[International Takotsubo Registry (InterTAK Registry, www.takotsubo-registry.com)].

1METHODS

2Data collection

- The InterTAK Registry is an observational, prospective, and retrospective registry 4established at the University Hospital Zurich in 2011 in collaboration with 25 cardiovascular centers 5across 9 countries.^{1, 3} Patients were included in the registry between 2011 and 2014 based on 6modified Mayo Clinic Diagnostic Criteria as previously reported: 1, 19 i) transient abnormality of left 7ventricular wall-motion extending beyond a single coronary artery perfusion territory, ii) absence of 8 sobstructive coronary artery disease (CAD) or evidence of acute plaque rupture, iii) presence of new 9electrocardiographic abnormalities or elevation in troponin iv) absence of and 10pheochromocytoma/myocarditis. Exceptions included coexisting CAD in whom the wall motion 11abnormality was congruent with a single coronary artery territory or death during the acute phase 12before documentation of wall motion recovery. Data on demographics, triggering factors, 13cardiovascular risk factors, haemodynamic and angiographic findings, ECG and echocardiography 14parameters, laboratory values, use of medications, in-hospital complications, and management 15were collected through standardized forms on admission or during revision of clinical charts.
- For the purpose of the present analysis, patients were divided into two groups according to 17the prescription of aspirin at hospital discharge. Patients with unknown status regarding aspirin at 18discharge were excluded from the present study.
- The local ethics committee or institutional review board at each participating site reviewed 20the study protocol. Most ethics committees waived the need for informed consent due to the partly 21retrospective nature of the study. Formal written consent was obtained from patients or their 22surrogates at participating centers whose ethics committees or institutional review boards required 23informed consent or if patients were included prospectively.

24

25**Study outcomes**

Follow-up data were collected from clinical visits, medical charts, or telephone interviews as 27previously described.¹ The incidence of major adverse cardiovascular and cerebrovascular events 28(MACCE: a composite of all cause death, TTS recurrence, stroke or transient ischemic attack

1[TIA], or myocardial infarction [MI]) at 30-day and 5-year follow-up were the co-primary outcomes 2in the present analysis. Additionally, single components of MACCE at 5-year follow-up were 3analysed.

4

5Statistical analysis

6In the unadjusted analysis, continuous variables were summarized as mean ± standard deviation 7(SD) or median (1st-3rd quartile), and frequencies of categorical variables are presented as 8numbers with percentages. Categorical variables were compared with the Pearson chi-square test, 9continuous variables with the Student t-test.

10An adjusted analysis based on propensity score (PS) was performed. PS is the probability that 11each individual patient is included in the treatment group and is usually estimated via logistic 12regression based on the available baseline covariates. PS methods are used to compensate for 13the lack of proper statistical design and randomization in observational studies, like the present 14one. All variables expected to be associated with the outcomes of interest, or with both 15aspirin prescription and outcomes, are listed in SupplementaryTable 1 and were used to 16construct the PS model.

17The first step of the adjusted analysis was the treatment of missing data, which were present for a 18high number of variables (69 covariates out of 136). Assuming that data were missing at random 19and considering only the variables with less than 50% of missing data (the other covariates were 20excluded from the analyses)²⁰, we used polytomus logistic regression, logistic regression and 21predictive mean matching as multiple imputation techniques to fill in missing values, using the R 22mice package (version 3.6.0). We imputed five different datasets and the same statistical analyses 23were performed on each of them. After that, Rubin's rule²¹ was used to get pooled propensity score 24adjusted HR estimates and confidence intervals for each endpoint (primary and secondary), 25according to each of the three methods described below: stratification, and covariate adjustment 26and inverse probability of treatment weighting (IPTW) as sensitivity analysis.

27With the method based on stratification, the total dataset is divided into mutually exclusive groups 28(strata), based on quantiles (in our case, tertiles) of the estimated PS; in this way, subjects from

1both arms are stratified in subsets that are defined by specific thresholds in PS.^{22, 23} Then, all strata 2are included in a stratified proportional hazard Cox model to get an estimate of the HR for 3treatment, as previously described by Austin.²⁴

4In the case of the covariate adjustment method, a Cox model is built with two predictors, given by 5the treatment indicator and PS itself.²⁵ An estimate of the treatment effect is then obtained based 6on the Cox model.

7Finally, the IPTW technique involves assigning to each patient a stabilised weight equal to 8(1-p)/(1-PS) if a control, or equal to p/PS if a treated patient,²⁰ where p is the probability of 9treatment without any covariate and PS is the value of the PS for that patient. The choice of 10stabilised weights allowed us to work with a pseudo-sample (as large as the sum of the weights) 11that has approximately the same size as the actual one.²⁶ Then, the weights were included in the 12survival analysis to estimate two adjusted Kaplan-Meier curves²⁷ (one for each treatment). The 13weights were also used to estimate the parameters of the Cox model, and in particular the HR.²⁸ 14No association of any continuous predictor and aspirin prescription departed from linearity, 15as assessed through the statistical significance of quadratic and cubic terms.

16The adjusted statistical analysis was performed using R 3.5.1 and some of its packages^{29,30}, 17notably *mice* package (version 3.6.0), *rms* (version 5.1-3.1) and *survival* (version 2.44-1.1).

1RESULTS

2Study population

Out of 1750 patients in the InterTAK Registry, 1533 with documented status regarding 4aspirin at discharge were included in the present analysis (Figure 1). The mean age was 66.4±13.1 5years and 1382 (90.2%) were females. 989 (65.8%) patients had hypertension, 221 (14.7%) 6diabetes mellitus, and 480 (32.0%) hypercholesterolemia. ST-segment elevation was observed in 7606 (43.5%) patients on admission. An emotional trigger was identified in 447 (29.2%) patients and 8a physical trigger in 533 (34.8%). Patients' characteristics of the total study cohort and of TTS 9patients with and without aspirin at discharge are summarized in Table 1. Unadjusted outcomes 10are reported in Table 2, showing a higher risk of 5 years death for patients in aspirin and no 11difference for the other endpoints.

12

13Adjusted comparison using PS with the stratification method

According to PS stratification method, aspirin was not associated with a reduced hazard of 15MACCE at 30-day (HR 1.24, 95% CI 0.50-3.04, P=0.64) or 5-year follow-up (HR 1.11, 95% CI 160.78-1.58, P=0.58). Furthermore, no significant differences were observed for the single 17components of MACCE, including death (HR 1.36, 95% CI 0.79-2.34, P=0.27), TTS recurrence 18(HR 0.53, 95% CI 0.27-1.03 P=0.06), stroke/TIA (HR 1.52, 95% CI 0.65-3.54, P=0.33), or MI (HR 193.28, 95% CI 0.38-28.28, P=0.28) (Table 2).

20

21 Sensitivity analysis: PS covariate adjustment and PS IPTW method

- PS IPTW and PS covariate adjustment methods did not show any association between 23aspirin and a risk reduction for MACCE or its single components (Table 2), except for TTS 24recurrence, which shows some weak association. The survival analysis for MACCE and death 25based on IPTW results confirmed these findings as reported in Figure 2, which depicts the Kaplan-26Meier curves of the two groups crossing each other.
- In order to verify that the application of the IPTW method allowed to achieve a gain in 28similarity between the active and control groups, we plotted two "mirrored" histograms showing the

1 distribution of PS (averaged on 5 imputed datasets) within each treatment group on the true and 2 the pseudo populations (see Supplementary Figure 2). After the application of the IPTW method, 3 the distribution of PS between the two groups looks more symmetrical: treated PSs are "shifted" 4 towards 0, while untreated PSs towards 1. The difference of frequency within each PS interval 5 between the active and control groups is due to the different sizes of the two groups, 1031 treated 6 subjects and 502 untreated ones.

1 DISCUSSION

- The increased awareness of TTS has resulted in a higher recognition of TTS among aphysicians. The increased awareness of TTS has resulted in a higher recognition of TTS among aphysicians. The increased awareness of TTS has resulted in a higher recognition of TTS among approximately approximately
- The present study found that aspirin at hospital discharge did not relate to short- nor longsterm prognosis in a large population of TTS patients. Incidence of MACCE in patients discharged 6with aspirin, who were not randomized but were adjusted for a higher burden of comorbidities with 7PS methods, was not significantly different compared to patients without aspirin, both at short and 8long-term follow-up. Furthermore, single components of MACCE were similar at 5 years. Presence 9of CAD at baseline did not affect these results.
- TTS pathophysiology is hypothesized to be mediated by an abrupt surge of catecholamines 11leading to ventricular dysfunction.³² An increased cardiac sympathetic activity is known to be 12associated with unfavourable outcomes in cardiovascular diseases.³³⁻³⁵ Of note, the 13catecholaminergic surge may activate platelets and proinflammatory pathways, setting the stage 14for the use of antiplatelet agents such as aspirin. The protective effect of aspirin in acute 15cardiovascular diseases, however, is mainly related to the reduction of thrombotic events induced 16by platelet activation following plaque erosion or rupture. These mechanisms do not appear to play 17a significant role in TTS, as it appears that TTS mainly involves the microcirculatory system, thus 18this explains the lack of potential benefit associated with aspirin in this syndrome.³⁶
- Aspirin acts both as an antithrombotic as well as an anti-inflammatory agent, suppressing 20the production of prostaglandins, thromboxane, and decreasing plasma levels of several 21inflammatory biomarkers, posing a potential prognostic benefit in TTS. Nevertheless, a negative 22interaction has been shown between aspirin (related to dose) and survival benefit of ACE-inhibitors 23therapy in patients admitted for heart failure and could have implications in TTS patients as well.³⁷ 24In a recent study of Dias et al. a beneficial effect of aspirin on an in-hospital combined endpoint 25has been reported when given on TTS index event.¹⁸ However, this effect may result from the 26combined therapy of aspirin and clopidogrel together. Moreover, the authors evaluated only 27hospital events in a relatively low sample size, which may have produced incidental findings.

- In line with our results, Fazio et al. demonstrated a lack of benefit of in-hospital aspirin 2administration on both hospitalisation length and ejection fraction improvement in a relatively small 3number of TTS patients.³⁸ Of note, we focused on aspirin use after hospital discharge, also 4adjusting for major confounding factors with PS-stratified analysis, and similarly we could not 5demonstrate an association between aspirin and improved outcome at follow-up. We found some 6evidence of weak association between aspirin and only TTS recurrence; such weak association is 7detected by the covariate adjustment and the IPTW methods and not by the stratification method. 8Therefore, this potential association should be interpreted carefully, considering the lack of a 9supporting pathophysiological mechanism. Since any potential benefit should be pondered with the 10inevitable higher bleeding risk in patients taking aspirin on a long-term basis, the routine use of 11aspirin should not be encouraged especially in patients at high risk for bleeding.³⁹
- Our results suggest that TTS *per se* does not represent an indication for treatment with 13aspirin. Aspirin treatment might be withdrawn even during hospitalisation once the clinical picture 14of TTS has been unmasked, unless there are coexisting comorbidities that confer a high 15atherosclerotic risk and require antiplatelet therapy according to current guidelines.

16

17**Study limitations**

- The present study is not a randomized controlled trial, but we tried to address this 19shortcoming using PS, which may nonetheless adjust only for recorded variables and not for the 20missing ones. Given the low prevalence of TTS it is challenging to obtain robust data on treatment 21or to conduct comparative randomized controlled trials. Therefore, the application of PS methods is 22currently state of the art in this setting.
- A methodological limitation of the study is that we mostly observed the absence of aspirin 24effects. As it is well known, absence of evidence is not evidence of absence, and a statistical proof 25of the lack of aspirin effect should properly be conducted within an equivalence approach using 26appropriately designed clinical trials, whereas it is not possible to do so using only observational 27studies.

Performance of PS was tested by assessing the standardized differences before and after 2 propensity score using IPTW on the covariates used, with satisfactory results (Supplementary 3 Table 2): in fact, the computation of standardized differences (SD) demonstrate that even though 4 some of the SDs increased from the unadjusted to the adjusted population, this led to an overall 5 decrease in all SDs adjusted with IPTW, so that almost all variables have a SD lower than 0.1 6 between treatment groups. Regarding non linearity, residuals are symmetrically distributed around 70 and lowess interpolation within each plot do not show any particular non-linear relationships (see 8 supplementary Figure 1). Moreover, in the stratification analysis, we used 3 strata, with a potential 9 higher risk of bias: however, the results are consistent with the other two analyses, confirming the 10 overall strength of our model.

Proper sample size calculation showed that this study is formally underpowered for main 12outcomes, although it should be remembered that the present is the largest available registry on 13this topic. This is particularly true for MI, which occurred only for 9 patients leading to large CI after 14PS adjustement.

The dose-dependent detrimental interaction of aspirin on ACE-Inhibitors therapy survival 16benefit makes the net effect of aspirin alone not completely predictable in TTS patients where both 17therapies are usually co-administered.

18

19

20CONCLUSIONS

In the present analysis, we found no evidence, after adjusting for potential confounding 22factors, that aspirin at discharge is associated with a reduced risk of MACCE at short- or long-term 23follow-up in TTS patients. These findings should be confirmed in adequately powered randomized 24controlled trials.

25

26

27

1ACKNOWLEDGEMENTS: none

2

3CONFLICTS OF INTEREST: none declared

9the manuscript; and decision to submit the manuscript for publication.

4

5FUNDINGS: CT has been supported by the H.H. Sheikh Khalifa bin Hamad Al-Thani Research 6Programme and the Swiss Heart Foundation. The InterTAK Registry is supported by The Biss 7Davies Charitable Trust. The funding sources had no role in the design and conduct of the study; 8collection, management, analysis, and interpretation of the data; preparation, review, or approval of

1REFERENCES

- 1. Templin C, Ghadri JR, Diekmann J, Napp LC, Bataiosu DR, Jaguszewski M, Cammann VL, Sarcon A, Geyer V, Neumann CA, Seifert B, Hellermann J, Schwyzer M, Eisenhardt K, Jenewein J, Franke J, Katus HA, Burgdorf C, Schunkert H, Moeller C, Thiele H, Bauersachs J, Tschope C, Schultheiss HP, Laney CA, Rajan L, Michels G, Pfister R, Ukena C, Bohm M, Erbel R, Cuneo A, Kuck KH, Jacobshagen C, Hasenfuss G, Karakas M, Koenig W, Rottbauer W, Said SM, Braun-Dullaeus RC, Cuculi F, Banning A, Fischer TA, Vasankari T, Airaksinen KE, Fijalkowski M, Rynkiewicz A, Pawlak M, Opolski G, Dworakowski R, MacCarthy P, Kaiser C, Osswald S, Galiuto L, Crea F, Dichtl W, Franz WM, Empen K, Felix SB, Delmas C, Lairez O, Erne P, Bax JJ, Ford I, Ruschitzka F, Prasad A, Luscher TF. Clinical Features and Outcomes of Takotsubo (Stress) Cardiomyopathy. N Engl J Med 2015;373(10):929-38.
- 2. Kato K, Lyon AR, Ghadri JR, Templin C. Takotsubo syndrome: aetiology, presentation and treatment. Heart 2017;103(18):1461-1469.
- 3. Ghadri JR, Cammann VL, Templin C. The International Takotsubo Registry: Rationale, Design, Objectives, and First Results. Heart Fail Clin 2016;12(4):597-603.
- 4. Gianni M, Dentali F, Grandi AM, Sumner G, Hiralal R, Lonn E. Apical ballooning syndrome or takotsubo cardiomyopathy: a systematic review. Eur Heart J 2006;27(13):1523-9.
- 5. Ghadri JR, Ruschitzka F, Luscher TF, Templin C. Takotsubo cardiomyopathy: still much more to learn. Heart 2014;100(22):1804-12.
- 6. Ghadri JR, Wittstein IS, Prasad A, Sharkey S, Dote K, Akashi YJ, Cammann VL, Crea F, Galiuto L, Desmet W, Yoshida T, Manfredini R, Eitel I, Kosuge M, Nef HM, Deshmukh A, Lerman A, Bossone E, Citro R, Ueyama T, Corrado D, Kurisu S, Ruschitzka F, Winchester D, Lyon AR, Omerovic E, Bax JJ, Meimoun P, Tarantini G, Rihal C, S YH, Migliore F, Horowitz JD, Shimokawa H, Luscher TF, Templin C. International Expert Consensus Document on Takotsubo Syndrome (Part I): Clinical Characteristics, Diagnostic Criteria, and Pathophysiology. Eur Heart J 2018;39(22):2032-2046.
- 7. Ghadri JR, Kato K, Cammann VL, Gili S, Jurisic S, Di Vece D, Candreva A, Ding KJ, Micek J, Szawan KA, Bacchi B, Bianchi R, Levinson RA, Wischnewsky M, Seifert B, Schlossbauer SA,

Citro R, Bossone E, Munzel T, Knorr M, Heiner S, D'Ascenzo F, Franke J, Sarcon A, Napp LC, Jaguszewski M, Noutsias M, Katus HA, Burgdorf C, Schunkert H, Thiele H, Bauersachs J, Tschope C, Pieske BM, Rajan L, Michels G, Pfister R, Cuneo A, Jacobshagen C, Hasenfuss G, Karakas M, Koenig W, Rottbauer W, Said SM, Braun-Dullaeus RC, Banning A, Cuculi F, Kobza R, Fischer TA, Vasankari T, Airaksinen KEJ, Opolski G, Dworakowski R, MacCarthy P, Kaiser C, Osswald S, Galiuto L, Crea F, Dichtl W, Empen K, Felix SB, Delmas C, Lairez O, El-Battrawy I, Akin I, Borggrefe M, Horowitz J, Kozel M, Tousek P, Widimsky P, Gilyarova E, Shilova A, Gilyarov M, Winchester DE, Ukena C, Bax JJ, Prasad A, Bohm M, Luscher TF, Ruschitzka F, Templin C. Long-Term Prognosis of Patients With Takotsubo Syndrome. J Am Coll Cardiol 2018;72(8):874-882.

- 8. Elesber AA, Prasad A, Lennon RJ, Wright RS, Lerman A, Rihal CS. Four-year recurrence rate and prognosis of the apical ballooning syndrome. J Am Coll Cardiol 2007;50(5):448-52.
- 9. Sharkey SW, Windenburg DC, Lesser JR, Maron MS, Hauser RG, Lesser JN, Haas TS, Hodges JS, Maron BJ. Natural history and expansive clinical profile of stress (tako-tsubo) cardiomyopathy. J Am Coll Cardiol 2010;55(4):333-41.
- 10. Ghadri JR, Wittstein IS, Prasad A, Sharkey S, Dote K, Akashi YJ, Cammann VL, Crea F, Galiuto L, Desmet W, Yoshida T, Manfredini R, Eitel I, Kosuge M, Nef HM, Deshmukh A, Lerman A, Bossone E, Citro R, Ueyama T, Corrado D, Kurisu S, Ruschitzka F, Winchester D, Lyon AR, Omerovic E, Bax JJ, Meimoun P, Tarantini G, Rihal C, S YH, Migliore F, Horowitz JD, Shimokawa H, Luscher TF, Templin C. International Expert Consensus Document on Takotsubo Syndrome (Part II): Diagnostic Workup, Outcome, and Management. Eur Heart J 2018;39(22):2047-2062.
- 11. Di Vece D, Citro R, Cammann VL, Kato K, Gili S, Szawan KA, Micek J, Jurisic S, Ding KJ, Bacchi B, Schwyzer M, Candreva A, Bossone E, D'Ascenzo F, Sarcon A, Franke J, Napp LC, Jaguszewski M, Noutsias M, Munzel T, Knorr M, Wagner S, Katus HA, Burgdorf C, Schunkert H, Thiele H, Bauersachs J, Tschope C, Pieske B, Rajan L, Michels G, Pfister R, Cuneo A, Jacobshagen C, Hasenfubeta G, Karakas M, Koenig W, Rottbauer W, Said SM, Braun-Dullaeus RC, Banning A, Cuculi F, Kobza R, Fischer TA, Vasankari T, Airaksinen KEJ, Opolski G, Dworakowski R, MacCarthy P, Kaiser C, Osswald S, Galiuto L, Crea F, Dichtl W, Empen K, Felix

- SB, Delmas C, Lairez O, El-Battrawy I, Akin I, Borggrefe M, Gilyarova E, Shilova A, Gilyarov M, Horowitz J, Kozel M, Tousek P, Widimsky P, Winchester DE, Ukena C, Di Mario C, Prasad A, Bohm M, Bax JJ, Luscher T, Ruschitzka F, Ghadri JR, Templin C. Outcomes Associated With Cardiogenic Shock in Takotsubo Syndrome: Results From the International Takotsubo Registry. Circulation 2018.
- 12. Kato K, Sakai Y, Ishibashi I, Himi T, Fujimoto Y, Kobayashi Y. Predictors of in-hospital cardiac complications in patients with Takotsubo syndrome. Heart Vessels 2018;33(10):1214-1219.
- 13. Singh K, Carson K, Usmani Z, Sawhney G, Shah R, Horowitz J. Systematic review and meta-analysis of incidence and correlates of recurrence of takotsubo cardiomyopathy. Int J Cardiol 2014;174(3):696-701.
- 14. Isogai T, Matsui H, Tanaka H, Fushimi K, Yasunaga H. Early beta-blocker use and inhospital mortality in patients with Takotsubo cardiomyopathy. Heart 2016;102(13):1029-35.
- 15. Santoro F, Ieva R, Ferraretti A, Ienco V, Carpagnano G, Lodispoto M, Di Biase L, Di Biase M, Brunetti ND. Safety and feasibility of levosimendan administration in takotsubo cardiomyopathy: a case series. Cardiovasc Ther 2013;31(6):e133-7.
- 16. Santoro F, Ieva R, Musaico F, Ferraretti A, Triggiani G, Tarantino N, Di Biase M, Brunetti ND. Lack of efficacy of drug therapy in preventing takotsubo cardiomyopathy recurrence: a meta-analysis. Clin Cardiol 2014;37(7):434-9.
- 17. Cecchi E, Parodi G, Giglioli C, Passantino S, Bandinelli B, Liotta AA, Bellandi B, Cioni G, Costanzo M, Abbate R, Gensini GF, Antoniucci D, Mannini L. Stress-induced hyperviscosity in the pathophysiology of takotsubo cardiomyopathy. Am J Cardiol 2013;111(10):1523-9.
- 18. Dias A, Franco E, Koshkelashvili N, Bhalla V, Pressman GS, Hebert K, Figueredo VM. Antiplatelet therapy in Takotsubo cardiomyopathy: does it improve cardiovascular outcomes during index event? Heart Vessels 2016;31(8):1285-90.
- 19. Prasad A, Lerman A, Rihal CS. Apical ballooning syndrome (Tako-Tsubo or stress cardiomyopathy): a mimic of acute myocardial infarction. Am Heart J 2008;155(3):408-17.

- 20. Morris TP, White IR, Royston P. Tuning multiple imputation by predictive mean matching and local residual draws. BMC Medical Research Methodology 2014;14(1):75.
- 21. Leyrat C, Seaman SR, White IR, Douglas I, Smeeth L, Kim J, Resche-Rigon M, Carpenter JR, Williamson EJ. Propensity score analysis with partially observed covariates: How should multiple imputation be used? Statistical Methods in Medical Research 2017;28(1):3-19.
- 22. Austin PC. An Introduction to Propensity Score Methods for Reducing the Effects of Confounding in Observational Studies. Multivariate behavioral research 2011;46(3):399-424.
- 23. Haukoos JS, Lewis RJ. The Propensity Score. JAMA 2015;314(15):1637-1638.
- 24. Austin PC. A Tutorial and Case Study in Propensity Score Analysis: An Application to Estimating the Effect of In-Hospital Smoking Cessation Counseling on Mortality. Multivariate behavioral research 2011;46(1):119-151.
- 25. Elze MC, Gregson J, Baber U, Williamson E, Sartori S, Mehran R, Nichols M, Stone GW, Pocock SJ. Comparison of Propensity Score Methods and Covariate Adjustment: Evaluation in 4 Cardiovascular Studies. Journal of the American College of Cardiology 2017;69(3):345-357.
- 26. Xu S, Ross C, Raebel MA, Shetterly S, Blanchette C, Smith D. Use of stabilized inverse propensity scores as weights to directly estimate relative risk and its confidence intervals. Value in health: the journal of the International Society for Pharmacoeconomics and Outcomes Research 2010;13(2):273-277.
- 27. Xie J, Liu C. Adjusted Kaplan–Meier estimator and log-rank test with inverse probability of treatment weighting for survival data. Statistics in Medicine 2005;24(20):3089-3110.
- 28. Buchanan AL, Hudgens MG, Cole SR, Lau B, Adimora AA, Women's Interagency HIVS. Worth the weight: using inverse probability weighted Cox models in AIDS research. AIDS research and human retroviruses 2014;30(12):1170-1177.
- 29. R Core Team. R: A language and environment for statistical computing. R Foundation for Statistical Computing. https://www.R-project.org/.
- 30. Maechler. M., Rousseeuw. P., Struyf, A., Hubert, M., Hornik, K.(2013). cluster: Cluster Analysis **Basics** and Extensions. R package version 1.14.4.

- 31- Ghadri JR, Cammann VL, Napp LC, Jurisic S, Diekmann J, Bataiosu DR, Seifert B, Jaguszewski M, Sarcon A, Neumann CA, Geyer V, Prasad A, Bax JJ, Ruschitzka F, Luscher TF, Templin C, International Takotsubo R. Differences in the Clinical Profile and Outcomes of Typical and Atypical Takotsubo Syndrome: Data From the International Takotsubo Registry. JAMA Cardiol 2016;1(3):335-40.
- 32. Wittstein IS, Thiemann DR, Lima JA, Baughman KL, Schulman SP, Gerstenblith G, Wu KC, Rade JJ, Bivalacqua TJ, Champion HC. Neurohumoral features of myocardial stunning due to sudden emotional stress. N Engl J Med 2005;352(6):539-48.
- 33. Cohn JN, Levine TB, Olivari MT, Garberg V, Lura D, Francis GS, Simon AB, Rector T. Plasma norepinephrine as a guide to prognosis in patients with chronic congestive heart failure. N Engl J Med 1984;311(13):819-23.
- 34. Kaye DM, Lefkovits J, Jennings GL, Bergin P, Broughton A, Esler MD. Adverse consequences of high sympathetic nervous activity in the failing human heart. J Am Coll Cardiol 1995;26(5):1257-63.
- 35. Brunner-La Rocca HP, Esler MD, Jennings GL, Kaye DM. Effect of cardiac sympathetic nervous activity on mode of death in congestive heart failure. Eur Heart J 2001;22(13):1136-43.
- 36. Luscher TF, Templin C. Is takotsubo syndrome a microvascular acute coronary syndrome? Towards of a new definition. Eur Heart J 2016.
- 37. Guazzi M, Brambilla R, Reina G, Tumminello G, Guazzi MD. Aspirin-angiotensin-converting enzyme inhibitor coadministration and mortality in patients with heart failure: a dose-related adverse effect of aspirin. Arch Intern Med 2003;163(13):1574-9.
- 38. Fazio G, Pizzuto C, Barbaro G, Sutera L, Incalcaterra E, Evola G, Azzarelli S, Palecek T, Di Gesaro G, Cascio C, Novo G, Akashi YJ, Novo S. Chronic pharmacological treatment in takotsubo cardiomyopathy. Int J Cardiol 2008;127(1):121-3.
- 39. Fletcher RH. Review: Aspirin for CVD primary prevention increases gastrointestinal bleeding and hemorrhagic stroke. Ann Intern Med 2016;165(4):JC17.

1FIGURE LEGENDS

2Figure 1. Study design.

3MACCE denotes major adverse cardiac and cerebrovascular event; TIA transient ischemic attack, 4TTS takotsubo syndrome.

5

6Figure 2. Inverse probability of treatment weighting adjusted Kaplan-Meier Analysis.

7Colored bands represent the 95% pointwise confidence bands.

8MACCE denotes major adverse cardiac and cerebrovascular event; IPTW, inverse probability of 9treatment weighting.

10