Cohort profile

Scheitz, Jan F.

2018-09


http://hdl.handle.net/10138/272990
https://doi.org/10.1136/bmjopen-2018-023265

Downloaded from Helda, University of Helsinki institutional repository.

This is an electronic reprint of the original article.

This reprint may differ from the original in pagination and typographic detail.

Please cite the original version.
Cohort profile: Thrombolysis in Ischemic Stroke Patients (TRISP): a multicentre research collaboration

Jan F Scheitz,¹ Henrik Gensicke,² Sanne M Zinkstok,³ Sami Curtze,⁴,⁵ Marcel Arnold,⁶ Christian Hametner,⁷ Alessandro Pezzini,⁸ Guillaume Turc,⁹ Andrea Zini,¹⁰ Visnja Padjen,¹¹ Susanne Wegener,¹² Annika Nordanstig,¹³ Lars Kellert,¹⁴ Georg Kägi,¹⁵ Yannick Bejot,¹⁶ Patrik Michel,¹⁷ Didier Leys,¹⁸ Christian H Nolte,¹ Paul J Nederkoorn,³ Stefan T Engelter,²,¹⁹ on behalf of the TRISP collaboration

ABSTRACT

Purpose The Thrombolysis in Ischemic Stroke Patients (TRISP) collaboration aims to address clinically relevant questions about safety and outcomes of intravenous thrombolysis (IVT) and endovascular thrombectomy. The findings can provide observational information on treatment of patients derived from everyday clinical practice.

Participants TRISP is an open, investigator-driven collaborative research initiative of European stroke centres with expertise in treatment with revascularisation therapies and maintenance of hospital-based registries. All participating centres made a commitment to prospectively collect data on consecutive patients with stroke treated with IVT using standardised definitions of variables and outcomes, to assure accuracy and completeness of the data and to adapt their local databases to answer novel research questions.

Findings to date Currently, TRISP comprises 18 centres and registers >10 000 IVT-treated patients. Prior TRISP projects provided evidence on the safety and functional outcome in relevant subgroups of patients who were excluded, under-represented or not specifically addressed in randomised controlled trials (ie, pre-existing disability, cervical artery dissections, stroke mimics, prior statin use), demonstrated deficits in organisation of acute stroke care (ie, IVT during non-working hours, effects of onset-to-door time on onset-to-needle time), evaluated the association between laboratory findings on outcome after IVT and served to develop risk estimation tools for prediction of haemorrhagic complications and functional outcome after IVT.

Future plans Further TRISP projects to increase knowledge of the effect and safety of revascularisation therapies in acute stroke are ongoing. TRISP welcomes participation and project proposals of further centres fulfilling the outlined requirements. In the future, TRISP will be extended to include patients undergoing endovascular thrombectomy.

INTRODUCTION

Intravenous thrombolysis (IVT) with recombinant tissue plasminogen activator (rtPA) for ischaemic stroke markedly improves functional outcome with a strong impact of time to treatment.¹ The pivotal National Institute of Neurological Disorders and Stroke (NINDS) trial in 1995 and more recently the individual patient data meta-analyses of subsequent randomised controlled trials (RCTs) proved safety and efficacy of IVT in acute ischaemic stroke.² ³

Implementation of IVT in acute medical treatment for ischaemic stroke has profoundly changed clinical routine. However, selection of patients for IVT in RCTs was based on strict criteria in order to exclude patients with presumably (1) low probability of favourable outcome per se or (2) high bleeding risk.⁴ ⁵ The current exclusion criteria listed in the American Heart Association and American Stroke Association 2013 acute stroke management guidelines are still based largely on...
the criteria listed in the NINDS trial, with only minor modifications over time. Many criteria addressed in the guidelines were adapted from the cardiologic literature or from basic science publications. This overall conservative approach in selecting patients suitable for IVT is controversial as it is often unseen whether patients meeting the established eligibility criteria do not benefit from IVT. As a consequence, the practice pattern of IVT usage has been shown to vary largely across centres. Furthermore, a restrictive approach in using IVT is likely to contribute to low IVT rates, especially in centres with reduced service levels. Systematically ascertained, comprehensive and high-quality observational data are useful to both (1) challenge or (2) confirm the clinical usefulness of commonly used but often arbitrary eligibility criteria. Of note, approximately half of patients have at least one contraindication or warning in large dedicated stroke centres. This underlines that clinical reality often differs from clinical trial settings. Register-based observational data can add clinically focused information whether certain warnings from licence or variables that have not been well-studied in RCTs—for example, comedications—matter or not with regard to outcome or complications of IVT. Ideally, the results from such observational studies will be verified or falsified by RCTs. However, with few exceptions (eg, age limit) this is unlikely to happen. Thus, register-based data will reflect the highest level of evidence, available currently and in the foreseeable future. In the last few years, endovascular thrombectomy (EVT) in patients with proximal vessel occlusion has emerged as another evidence-based treatment option, and acute multimodal imaging methods challenge the classical time-window concept. This makes acute stroke care even more complex. As with IVT, the benefits of EVT in subgroups of patients underrepresented in clinical trials deserve further research.

In the absence of RCT-based evidence, comprehensive observational data will also be useful for individual treatment decisions about IVT and EVT, and in evaluating processes of stroke triage and care for IVT or EVT. As a prerequisite, such data have to be based on well-maintained registries containing a large number of detailed, clearly defined and well-characterised variables. TRISP (ThRombolysis in Ischemic Stroke Patients) meets these prerequisites.

COHORT DESCRIPTION
Aims and objectives of TRISP
TRISP started as a joint initiative of 11 European stroke centres in 2010. Initiated by the stroke research team Basel, an explorative, research collaborative study was designed to study the impact of prior statin treatment on outcomes of patients with stroke receiving IVT. Currently, TRISP comprises 18 European stroke centres (please see online supplement for a list of contributors to the TRISP collaboration). The TRISP centres cover a distance from Scandinavia to the Mediterranean area and from Western to Eastern Europe. TRISP centres constitute high-volume IVT centres with a record of comprehensive stroke treatment for several years. All TRISP centres offer treatment fulfilling the criteria of Stroke Centres or Stroke Units as proposed by the European Stroke Organisation (ESO). The major aim of TRISP is to address clinically important questions about safety and outcomes of patients with ischemic stroke treated with IVT who are neither covered by RCTs nor other large-scale IVT registers such as the Safe Implementation of Treatment in Stroke-International Stroke Thrombolysis Register (SITS-ISTR) and Get-with-the-guidelines (GWTG) Stroke.

Design, structure and policy
TRISP operates as an investigator-driven, open platform with a clear emphasis on clinical research and data quality. TRISP is independent from industry and a non-profit collaboration. TRISP particularly aims at supporting young researchers. Thus, with the exception of the very first paper, first authors of the publications invariably were young stroke physicians or PhD students.

There is no official chair and no official administration, board or secretary within the TRISP collaboration. Currently, TRISP is coordinated by STE (Basel) and PJN (Amsterdam) who are primary contact persons.

Prior to the start of a novel project, standardised research proposals stating hypotheses and a statistical analysis plan are circulated to all participating centres and discussed for scientific content and feasibility. If a centre agrees to participate, then the centre contributes data for the time period for which they have data on all consecutive patients for the key variables of interest (eg, usage of a certain comedication). The information of the distribution of patients stratified to centre and time period is provided as online supplemental material in the respective manuscript. If a novel variable of interest needs to be added to answer a research question, the research proposal needs to describe a definition that allows to operationalise retrospective inclusion within the existing databases.

Lead of distinct project and authorship
The researcher or researches—usually 1 or 2 (rarely 3)—who had the idea and rendered the analysis proposal take(s) the lead and will get first and senior authorships. Coauthorships are attributed taking into account all aspects important for the success of a research project. This includes not only mere quantitative means (ie, number of patients contributed) but also quality of data (eg, completeness, though usually high across TRISP centres), handling and pooling of the multicentre data; maintenance of the pooled data set (including data cleaning); statistics, contribution to TRISP in general, and intellectual input in details of the design or the analyses of the research project. These criteria are suggestions and the researches taking the lead in each project take the final responsibility for the distribution of authorships.
Data collection and definitions

Data on the characteristics of patients treated with IVT are collected prospectively by all participating centres using standardised definitions and a standardised form. Not all centres have to provide data on all variables but have given a commitment to add missing variables retrospectively, if considered relevant to answer a specific research question. The dataset includes information on patients’ age, sex and treatment modality (IVT alone, IVT plus EVT, EVT alone). Date and time of stroke onset as well as onset-to-needle time and door-to-needle time are systematically collected. If the exact time of symptom onset is unknown or un witnessed, time from last seen normal until application of IVT will be provided. Stroke aetiology will be documented in distinct categories in accordance with the classification of the Trial of Org 10172 in Acute Stroke Treatment (TOAST) and classified as due to cardioembolic sources, large-artery atherosclerosis, small-vessel disease, other determined causes, concurrent aetiology or undetermined causes. The identified ‘other determined cause’ will include dissections. Other causes will be specified as precisely as possible as free text. Patients with final diagnosis other than stroke are classified as ‘stroke mimics’. Dose of rtPA as well as weight and body mass index (BMI) of the patients are documented in the majority of centres. Stroke severity at baseline is measured using the National Institutes of Health Stroke Scale (NIHSS) by experienced stroke physicians. If available, NIHSS at 24 hours after treatment and at discharge from the acute care hospital will be recorded. The functional status of the patient before the index stroke will be estimated by the stroke physicians who also indicated and applied IVT using the modified Rankin Scale (mRS). This prestroke mRS scores are based on information provided by patients and next-of-kin. Pre-treatment systolic and diastolic blood pressure (mm Hg) are also entered in the dataset. In case of more than one measurement, the values as close to application of IVT as possible will be noted.

Cardiovascular and stroke risk factors include atrial fibrillation (known from medical history, detected on baseline ECG or during in-hospital cardiac monitoring), diabetes mellitus (known from medical history, fasting plasma glucose ≥7.0 mmol/L or 126 mg/dL, HbA1c (haemoglobin A1c)≥6.5% or pre-existing treatment with antidiabetic drugs), hypercholesterolaemia (known from medical history, low-density lipoprotein >100mg/dL, (2.6 mmol/L) or treatment with cholesterol-lowering drugs), hypertension (known from medical history, repeated measurement of systolic blood pressure >140 mm Hg and/or diastolic blood pressure >90 mm Hg), previous stroke (known from medical history), smoking (patients who did not currently smoke were divided in non-smokers and past smokers≥22 years) and coronary artery disease (known from medical history). Medical history is provided by the patients, next-of-kin or family doctors.

Laboratory measures obtained on hospital admission include glucose (mmol/L), blood cell counts (platelets, haemoglobin, leucocytes), international normalised ratio and partial thromboplastin time. In case of use of novel oral anticoagulants, specific measurements of factor Xa activity, thrombin time or plasma levels of the respective drug are recorded, if available. Renal function is quantified by the estimated glomerular filtration rate (according to Chronic Kidney Disease Epidemiology Collaboration formula). Comedication used at the time of hospital admission prior to IVT is recorded precisely (eg, use of antithrombotic drugs and statins). Moreover, neuroimaging findings before and after treatment are systematically ascertained and comprise imaging modality (CT versus MRI) as well as specific imaging findings like hyperdense artery sign, presence and extent of early ischaemic signs according to the Alberta Stroke Program Early CT Score, site of vessel occlusion, presence of tandem occlusion of ipsilateral carotid artery, recanalisation status immediately after EVT and on follow-up imaging (quantified according Thrombolysis-In-Cerebral-infarction (TICI) score, TICI 2b/3 indicate successful recanalisation), white matter disease severity, presence and burden of cerebral microbleeds.

All patients are monitored for occurrence of haemorrhagic transformation. Follow-up imaging is usually scheduled at 24–36 hours after treatment or earlier in case of clinical worsening. Some centres perform follow-up imaging only in case of clinical worsening. The definition of symptomatic intracerebral haemorrhage (sICH) is in accordance with the definition used in the European Cooperative Acute Stroke Study (ECASS) II (blood at any site in the brain on the CT scan together with documentation of clinical deterioration or adverse events indicating clinical worsening or causing a decrease in the NIHSS score of 4 or more points). The majority of centres additionally evaluate type of haemorrhagic transformation (haemorrhagic infarction, parenchymal haemorrhage), indicate whether the bleeding occurred remotely from the infarcted area and document sICH according to NINDS, ECASS-II and SITS-MOST (Safe Implementation of Thrombolysis in Stroke-Monitoring Study) definitions. Functional outcome at 3 months is assessed using the mRS. The mRS is obtained by telephone calls, postal questionnaire or outpatient visits. If patients cannot be interviewed, close relatives, nurses or family doctors are asked for disability status. More recently, TRISP has been extended to include data on EVT, too.

Ethical considerations and data protection

Each study project within TRISP has to be approved by the ethics committee in the coordination and leading centres according to local regulations. It is in the responsibility of each participating centre to fulfill the local, regional and national legal and ethical requirements. Participating centres are responsible to obtain local ethics committee approval to collect and pool anonymised data with other
centres and to keep a local patient identifier. Once a novel project has been approved, data are forwarded to the centre leading the respective project. Only strictly anonymised data prepared to match the universal data collection sheet are accepted.

Patient and public involvement
Patients and public were not involved in the study.

Findings to date
Currently, data on IVT treatments of more than 10000 patients are available. Previous publications of TRISP (1) provided insight into safety and efficacy of IVT in subgroups of patients who were excluded (eg, patients dependent on the help of others prior to stroke), under-represented or not specifically addressed (eg, dissection as cause, impaired renal function, low platelet count, BMI, prior use of statins, serotonin reuptake inhibitors, prior use of novel oral anticoagulants) in RCTs, (2) were helpful to evaluate processes of acute stroke care such as the meaning of the ‘off-hour-thrombolysis’, IVT during ‘working hours’ or the variable ‘time’ in clinical practice, and (3) served to derive, validate and compare risk scores for sICH or functional 3-month outcome.

The idea of TRISP is that experienced stroke centres with a record and expertise in both (1) usage of thrombolysis and (2) maintenance of hospital-based thrombolysis databases pool their data. Usually this means, 50 IVT treatments per year and an IVT rate of >20% of all stroke admissions. Completeness of data regarding age, sex, initial NIHSS, sICH and mRS at 3 months in >95% of consecutive IVT-treated patients is warranted by all participating centres. In addition to the characteristics of the TRISP centres stated above, an advantage of TRISP is the availability of more additional variables than in other large-scale registers and the commitment by the collaborators to (1) accuracy and completeness of the data and to (2) the willingness to adapt the local databases and add quickly new variables retrospectively and prospectively. This enables gaining explorative insights in the putative prognostic importance of additional variables with unknown impact on outcome or risk of complications such as sICH. Strengths and limitations of TRISP in general and compared with other existing IVT registers are discussed below.

Strengths and limitations
Strengths of TRISP include (1) the high completeness of data with few missing data on 3-month outcome (all centres aim to keep this below 5%), (2) large sample sizes (several thousands) which reduces the risk of bias and allows adjustments for confounders, (3) the systematic and standardised data ascertainment which increases the homogeneity of the study population, (4) the intrinsic motivation of the study personal leads to a high rate of completeness of ascertained data sets, contributing to a high-quality registry and (5) the dynamic nature of the TRISP database due to the commitment of the centres to adapt the local database and add variables retrospectively and prospectively. In addition, (6) a large number of variables is gathered including those with unknown prognostic importance. This allows addressing novel yet unmet research questions. Moreover, (7) pooling of individual patient data increases generalisability compared with single centre studies and (8) the fact that variables and outcomes have been collected irrespective of the present research question reduces the risk of a bias.

There are other large-scale registers such as the international SITS-ISTR and the GWTG stroke registry. These registers have an unquestioned value and have served to address several research questions with high clinical impact. The TRISP collaboration does not intend to replace these registers but rather to refine the knowledge gained by them, to provide additive information and to fill gaps not covered yet. An example is the report about the usefulness of IVT in patients dependent on the help of others prior to stroke, which adds to the SITS-ISTR report about pre-existing disability. Compared with the GWTG registry that provides short-term outcomes only, TRISP provides systematic assessment of functional outcome at 3 months and systematic ascertainment of haemorrhagic complications. GWTG is mainly intended to be a database for quality control of acute stroke care. The possibility of participating centres in TRISP to develop individual research projects yields an intrinsic motivation to maintain a high data quality. Another difference of TRISP compared with SITS-ISTR (and also GWTG) are the characteristics of the sites that contribute data. In TRISP, solely experienced centres participate, while in SITS-ISTR and GWTG also low-volume centres with fewer IVT treatments per year enter their data.

Limitations are inherent to the design of TRISP. (1) Data are derived from registers that are neither monitored nor randomised. Usually, there will be no control group without IVT which disallows the assessment of effectiveness of IVT in study populations. (2) As true for all observational studies, TRISP analyses have a higher risk of bias than RCTs. Thus, we urge to a cautious interpretation of findings and observations. (3) All TRISP centres are experienced in stroke treatment. This includes in particular the use of IVT and increasingly EVT, too. This expertise implies—as a downside—a limited generalisability of TRISP findings to all stroke providers with less expertise and less advanced setting. (4) The majority of our included patients are Caucasians. Thus, we cannot compare ethnical differences, but might have the potential to set up further collaborations in the future. (5) Centres from Eastern Europe are relatively under-represented in TRISP. We encourage participation of centres from Eastern Europe in future projects to overcome this limitation and further increase generalisability of findings from TRISP. (6) Currently, there is no ‘core lab’ to validate haemorrhagic complications and 3-month mRS ratings. As valid for other registries like SITS and GWTG, local interpretation of outcome data may differ between sites. Since TRISP centres are high-volume centres with long-standing
Box 1 Universal standards and requirements for contributing to TRISP*

- Prospective registry of consecutive patients with systematic check-up of missing cases.
- Comprehensive collection of baseline characteristics according to consensus definitions stated in this manuscript.
- Prospective assessment of haemorrhagic complications (symptomatic intracerebral haemorrhage according to ECASS II criteria) and functional outcome at 3 months (according to the modified Rankin Scale; either telephone interview, postal questionnaire or clinical follow-up visit).
- Approval of institutional review board to maintain the respective IVT database and to obtain 3-month follow-up data.
- TRISP centres are comprehensive stroke centres with high-volume IVT applications and usually University hospitals or affiliated to University hospitals.
- Treatment of patients with acute ischaemic stroke with IVT according to guidelines valid at the relevant time (http://www.eso-stroke.org/eso-stroke/education/guidelines.html) or documentation of deviation therefrom.

*TRISP welcomes participation and project proposals of further centres fulfilling the commitment and the outlined requirements.

Note: ECASS II, European Cooperative Acute Stroke Study II; IVT, intravenous thrombolysis; TRISP, Thrombolysis in Ischemic Stroke Patients

The experience in maintaining IVT databases, this bias is likely to be smaller than in these other registries.

Collaboration

The TRISP collaboration welcomes participation and project proposals of further centres fulfilling the requirements stated in box 1. The currently participating centres have agreed to fulfill the prerequisites that are summarised in box 1. Participation in other registries does not exclude from participation in the TRIPS collaboration. In order to participate or suggest a novel project, please contact the TRISP coordinating centres Basel or Amsterdam (see corresponding author or list of contributors in the online supplements). On request, the standardised data collection form and project proposal form will be forwarded. Only strictly anonymised data are used and can be included. Data sharing is restricted to non-commercial, purely academic purposes only. Furthermore, data ownership remains at the centre/physician by whom the data were originally obtained.

Outlook

Currently, several further TRISP projects are ongoing or planned. Given the potential of brain MRI to provide more prognostic information than plain CT (eg, presence of large infarct core, presence of cerebral microbleeds), centres with regular MRI-based application of IVT (ie, Berlin, Bern, Lille and Paris) can offer to pool data within the subsection ‘TRISP MRI’, while others (eg, Basel) can focus on advanced CT imaging (TRISP-CT).

In the light of the recent positive trials of EVT in anterior circulation stroke the subsection ‘TRISP endovascular’ is operating since 2018. A standardised form including additional procedure-related variables was set up and agreed on during the ESO conference in 2018.

An official meeting will be scheduled annually during the respective ESO Conference. The aim is to discuss progress of ongoing projects as well as feasibility and scientific value of new project proposals. Summarising minutes are circulated to all participating centres.

Summary

The TRISP collaboration is an open platform dedicated to conduct joint research projects in patients with ischaemic stroke treated with IVT and also with EVT in the future. TRISP aims to increase knowledge regarding safety and outcomes after IVT and EVT and to evaluate processes of care. As shown in previous publications, TRISP has the potential to provide observational information on treatment of patients derived from daily clinical practice. Prospective and standardised documentation of individual patient data according to consensus definitions is a major requirement to maintain the quality of registries that contribute to TRISP. TRISP welcomes participation and project proposals of further centres fulfilling the requirements stated above.

Author affiliations

1Department of Neurology and Center for Stroke Research, Charité-Universitätsmedizin Berlin, Berlin, Germany
2Department of Neurology and Stroke Center, University of Basel and University Hospital Basel, Basel, Switzerland
3Department of Neurology, Academic Medical Center, Amsterdam, The Netherlands
4Department of Neurology, Helsinki University Hospital, Helsinki, Finland
5Department of Neurological Sciences, University of Helsinki, Helsinki, Finland
6Department of Neurology, Inselspital, Bern University Hospital, University of Bern, Bern, Switzerland
7Department of Neurology, University Hospital Heidelberg, Heidelberg, Germany
8Department of Clinical and Experimental Sciences, Neurology Clinic, University of Brescia, Brescia, Italy
9Université Paris Descartes Sorbonne Paris Cité, Centre Hospitalier Sainte-Anne, Paris, France
10Stroke Unit, Department of Neurosciences, Nove Ospedale Civile S. Agostino-Estense, University Hospital, Modena, Italy
11Neurology Clinic, Clinical Centre of Serbia, Belgrad, Serbia
12Department of Neurology, University of Zürich, Zürich, Switzerland
13Department of Neurology, Sahlgrenska University Hospital, Gothenburg, Sweden
14Department of Neurology, Klinikum der Universität München, Ludwig-Maximilians University, München, Germany
15Department of Neurology, Kantons spitäl St. Gallen, St. Gallen, Switzerland
16Dijon Stroke Registry, E44184, University Hospital and Medical School of Dijon, University of Burgundy, Dijon, France
17Department of Neurology, Centre Hospitalier Universitaire Vaudois and University of Lausanne, Lausanne, Switzerland
18Department of Neurology, University of Basel and University Hospital, Basel, Switzerland
19Neurorhabilitation Unit, University of Basel and University Center for Medicine of Aging and Rehabilitation, Felix Platter Hospital, Basel, Switzerland

Contributors All authors conceived the study and developed the protocol. JFS wrote the first draft. JFS, HG, SMZ, SC, MA, CH, AP, GT, AZ, VP, SW, AN, LN, KX, GK, YB, PM, DL, CHN, P, PN. STE critically revised the manuscript for important intellectual content, were involved in study concept and design, will be involved in acquisition of data and approved the final version of the manuscript. All contributors listed in the supplemental material approved submission of the manuscript.

Funding The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.
Competing interests JFS has received speaker honoraria from W L Gore & Associates GmbH and travel support from Bayer and Boehringer-Ingelheim; HG has received research support from the Swiss National Science Foundation; AZ has received funding for speaker honoraria and consulting fees from Boehringer-Ingelheim and Medtronic-Covidien and consulting fees from Nestec; GK received grants from Swiss Parkinson Association, Swiss Heart Association, Swiss National Science Foundation and served on advisory boards for Boehringer-Ingelheim, Bayer, Daiichi Sankyo (Switzerland) AG, Zambon, Nestle, CE Healthcare within the last two years; YB received honoraria or consulting fees from AstraZeneca France, Daiichi-Sankyo, BMS-Pfizer, Covidien, Bayer and MSD France; PM received research grants from the Swiss National Science Foundation, the Swiss Heart Foundation; speaker fees from Boehringer-Ingelheim, Bayer, Covidien and Stryker; honoraria from scientific advisory boards from Boehringer-Ingelheim, Bayer, Pfizer, Amgen; consulting fees from Pierre-Fabre and Astra-Zeneca; and travel support from Boehringer-Ingelheim and Bayer. All this support is received by the institution (CHUV) and is used for stroke education and research; DL participated during the last 5 years to 1 advisory boards, symposia or trials sponsored by Sanofi Aventis, BMS, AstraZeneca, Boehringer-Ingelheim, Servier, BMS, Pfizer et Allergan (honorarium paid to Adirond or research account of the hospital) and was an associated editor of the Journal of neurology, neurosurgery and psychiatry 2004-2010 (personal financial compensation); CHN has received funding for travel or speaker honoraria from Bayer, Boehringer-Ingelheim, Takeda, and BMS/Pfizer; PJN has received consulting fees from Boehringer-Ingelheim; STE has received funding for travel or speaker honoraria from Bayer and Boehringer-Ingelheim, he has served on scientific advisory boards for Bayer, Boehringer-Ingelheim, BMS/Pfizer and Covidien and on the editorial board of Stroke. He has received an educational grant from Pfizer and research support from the Science Funds (Wissenschaftsfonds) of the University Hospital Basel, the University Basel, the Swiss Heart Foundation and the Swiss National Science Foundation.

Patient consent Not required.

Provenance and peer review Not commissioned; externally peer reviewed.

Data sharing statement Data sharing will be restricted to non-commercial and academic purposes only. Data ownership remains at the center that originally obtained the data. The centers will have to give permission for re-use of the data. Data inquiries or further suggestions for analyses/participation in TRISP can be properly cited, appropriate credit is given, any changes made indicated, and the use and license their derivative works on different terms, provided the original work is academic purposes only. Data ownership remains at the center that originally

REFERENCES


