Benefits and safety of transdermal glyceryl trinitrate in acute stroke: a systematic review and meta-analysis of randomised trials (protocol)

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ABSTRACT

Introduction
High blood pressure (BP) in acute stroke has adverse outcomes. Transdermal glyceryl trinitrate (GTN) has beneficial properties in controlling BP. The 2016 meta-analysis and 2017 Cochrane review showed that transdermal GTN was beneficial in a small patient subgroup with stroke onset ≤6 hours. Larger studies focusing on this patient subgroup have since been conducted. We report the protocol for an updated systematic review and meta-analysis on the safety and benefits of transdermal GTN in acute stroke.

Methods and analysis
We will search Medline, Pubmed, Embase, CINAHL and Cochrane Library from inception until June 2020 for randomised trials that report the efficacy and safety of transdermal GTN versus placebo/control therapy among adult patients with acute stroke. Primary outcomes include in-hospital mortality, BP lowering and late functional status. Secondary outcomes include early, late, resource utilisation and surrogate outcomes. Safety outcomes include reported adverse events. Reviewers will first screen titles and abstracts, and then full texts, to identify eligible studies. Independently and in duplicate, they will extract data, assess risk of bias (RoB) using a modified Cochrane RoB tool and quality of evidence using Grading of Recommendations, Assessment, Development and Evaluation. Disagreement will be resolved by discussion and consultation with an external reviewer if necessary. Using a random-effects model, we will report effect sizes using relative risks and 95% CIs. We will perform predefined subgroup analyses: intracerebral haemorrhage versus ischaemic stroke; minor (NIHSS ≤5) versus major (NIHSS >5) ischaemic stroke; ischaemic stroke with versus without thrombolysis; prehospital versus non-prehospital settings; time from stroke to randomisation ≤6 versus >6 hours and high versus low overall RoB studies. We will also perform trial sequential analysis for the primary outcomes.

Ethics and dissemination
Ethics board approval is unnecessary. PROSPERO registration has been obtained. The results will be disseminated through publication in a peer-reviewed journal.

PROSPERO registration number CRD42020173093.

INTRODUCTION
High blood pressure (BP) is present in greater than 70% of patients with acute ischaemic stroke.1 It is associated with poor outcomes including acute stroke recurrence, death within a few weeks or combined death and dependency after several months.1–4 High BP is similarly common in acute intracerebral haemorrhage (ICH)5 and may be associated with haematoma expansion and increased mortality.6–8

It is recommended to lower BP in ICH9 10 although controversy exists regarding optimal BP target in patients with ICH and there is no current literature on the role of prehospital BP reduction. The management of high BP in acute ischaemic stroke and the decision to treat or not to treat has been a constant debate since 1985. Current available guidelines recommend withholding antihypertensive therapy in the early poststroke period unless there is markedly elevated BP (>220/120 mm Hg) or with BP >185/110 mm Hg for patients eligible for

Strengths and limitations of study

This is an updated meta-analysis which includes more recent larger trials.

This study will examine an important gap on the benefits of transdermal glyceryl trinitrate (GTN) in ultra-acute stroke (<≤6 hours) identified by previous reviews.

Other strengths include a comprehensive search strategy, an extensive predefined subgroup analysis plan and inclusion of Grading of Recommendations, Assessment, Development and Evaluation methodology to assess certainty of evidence.

This study will be the first to use trial sequential analysis on important primary outcomes.

Limitations include high clinical heterogeneity given the different subtypes of acute stroke, variation in timing of randomisation from onset of stroke to transdermal GTN or placebo/control therapy and reporting of outcome measures across trials.
thrombolysis or BP >180/105 mm Hg during the 24-hour period following reperfusion.\textsuperscript{11–13}

Nitric oxide (NO) donors are candidate agents to lower BP in acute stroke because NO is a cerebral and systemic vasodilator, modulates vascular and neuronal function, is neuroprotective and inhibits apoptosis.\textsuperscript{14} In addition, vascular NO concentrations are low in acute stroke which are associated with increased severity of stroke, mortality and institutionalisation.\textsuperscript{15} These observations support that NO supplementation might be beneficial.

Glyceryl trinitrate (GTN) is an example of a NO donor. Transdermal GTN administration offers a constant release of the drug across the skin into the systemic circulation for 24 hours which achieves sustained steady-state plasma concentrations.\textsuperscript{16} Transdermal GTN offers a formulation which is easily administered in many clinical settings (prehospital, Emergency Department (ED) and inpatient) managing acute stroke which may help to minimise fluctuations in drug concentrations and hence BP.

The latest meta-analysis published in 2016\textsuperscript{17} and Cochrane review in 2017,\textsuperscript{18} using data from five completed transdermal GTN trials (n=4197), reported no improvement in outcomes across a range of domains, such as death, disability, cognition, mood and quality of life, with transdermal GTN versus placebo or control therapy. However, in a prespecified subgroup analysis of patients with time from stroke to randomisation ≤6 versus >6 hours (n=312), these two reviews reported a favourable functional outcome as measured by modified Rankin Scale (mRS) at 90 days with transdermal GTN.

There were important limitations in these reviews. Four of all patients and 86.9% of those randomised within 6 hours of onset (as reported by the authors). In addition, all the included trials were conducted by a single research group and it is important that other research groups study the role of transdermal GTN in acute stroke. Finally, a relatively small number of patients (n=312) were treated within 6 hours of stroke onset and these patients came from just two of the five trials.

The recently published multicentre RIGHT-2 trial randomised 1149 participants with acute stroke within 4 hours of onset to receive transdermal GTN versus sham therapy.\textsuperscript{20} The data from this study more than triple that used to examine the role of transdermal GTN in ultra-early stroke (onset ≤6 hours). There is an urgent need to update the evidence behind the efficacy and safety of transdermal GTN in acute stroke especially among those patients with ultra-early (≤6 hours) presentation.

The aim of this systematic review and meta-analysis is to examine, using recent data, whether transdermal GTN improves important patient-centred outcomes and is safe among patients with acute stroke in the prehospital and inpatient settings compared with placebo or control therapy by reviewing randomised controlled trials (RCTs).

METHODS
Study registration
This systematic review and meta-analysis protocol has been registered in the International Prospective Register of Systematic Reviews. We will adhere to the Preferred Reporting Items for Systematic Reviews and Meta-Analysis statement for reporting systematic review and meta-analysis.\textsuperscript{21}

Eligibility criteria
We will include randomised trials investigating the efficacy and safety of transdermal GTN versus placebo or control therapy among adult patients presenting with acute stroke.

Patients aged ≥16 years presenting with either acute ischaemic or haemorrhagic stroke in the prehospital, ED and inpatient clinical settings will be considered. Acute stroke patients are defined as those with presentation within 5 days of onset of symptoms. We select 5 days since onset of symptoms as the cut-off criterion for this review because there can be significant delay in presentation after an acute stroke; especially for less severe ischaemic strokes.\textsuperscript{22} Patients with ischaemic stroke are included regardless of whether they receive thrombolysis. The comparator arms will include transdermal GTN patch, sham patch and control with existing standard therapy.

Primary outcomes are important patient-centred outcomes including in-hospital mortality, lowering of BP measurements and late functional status. BP parameters will include systolic BP, diastolic BP and mean arterial pressure measured at intervals stated by the authors. Late functional status will involve assessment using the mRS within 3 months of stroke onset or later (as reported by the authors); the preferred outcome measurement for acute stroke trials.\textsuperscript{23} The hierarchical mRS scores range from 0 to 6, with a score of 0 indicating no symptoms, 1 indicating some symptoms but no significant disability, 2–5 indicating increasing levels of disability and dependency and 6 indicating death.\textsuperscript{24}

Secondary outcomes are classified as early, late, resource utilisation and surrogate outcomes. Early secondary outcomes include development of ICH, recurrent stroke and change in calculated National Institutes of Health Stroke Scale (NIHSS). Late secondary outcomes include reported changes in activities of daily living, cognition, quality of life and mood. Resource utilisation secondary outcomes include length of hospital stay and discharge destination. Surrogate secondary outcomes include changes in cerebral haemodynamics and laboratory parameters like platelet aggregation.

Safety outcomes include any adverse events reported by the authors.

Search strategy
We will search Medline, Pubmed, Embase, CINAHL and Cochrane Library from inception until June 2020 without language restrictions. We will review reference lists for
eligible new trials and also search ClinicalTrials.gov for ongoing or unpublished trials and for additional data from published trials. The search strategy will include the following keywords: stroke, ischaemic stroke, haemorrhagic stroke, intracerebral haemorrhage, transdermal glyceryl trinitrate, transdermal nitroglycerin, glyceryl trinitrate patch, nitroglycerin patch, trinitroglycerin, prehospital, mortality, blood pressure, functional outcome, humans and randomised clinical trials. Medical Subject Heading terms will include acute stroke, brain infarction, brain haemorrhage, prehospital emergency care, nitroglycerin, nitric oxide donors, blood pressure, haemodynamics and cerebral haemodynamics. A proposed search strategy on Medline using the Pubmed interface is attached as online supplementary appendix 1.

**Study selection**

Reviewers (LBL, CWL, LWF and NWM) will independently and in duplicate screen the titles and abstracts of all identified studies to generate a list of eligible trials from which full texts will be obtained. Subsequently, the same reviewers will independently assess eligibility of these full texts of published trials to decide on the final included studies. Discrepancies between reviewers will be resolved through discussion and consensus or, if needed, by adjudication from an external reviewer and/or contact with authors of the original trials for clarification.

**Data extraction**

Two pairs of reviewers (LBL and CWL; LWF and NWM) will extract data from included studies both independently and in duplicate. Data will be extracted using a predesigned data extraction form adapted from the Cochrane Collaboration.24 The data collection form is attached as online supplementary appendix 2. Data extracted will include the following: general study information (authors, publication year and study location(s)); study population details (clinical setting—prehospital versus ED versus inpatient, sample size, types of strokes—ischaemic versus haemorrhagic; subgroup of ischaemic strokes with thrombolysis); details on the comparator arms (different doses and duration of GTN patch; sham patch and control) as well as the primary, secondary and safety outcomes as listed above.

In randomised trials that included more than one arm of GTN dosing and duration, we will extract data from the arm closest to a single dose regimen that is comparable to other primary studies to be used for analysis. Discrepancies in data extraction will be resolved through discussion and consensus or, if needed, via consultation with an external reviewer.

**Risk of bias assessment**

We will assess the risk of bias (RoB) for each outcome of the individual studies using a modified Cochrane RoB instrument.25 The instrument assesses biases in the following five domains: selection bias (random sequence generation and allocation concealment); performance bias (blinding of participants and researchers); detection bias (blinding of outcome assessment); attrition bias (incomplete outcome data) and reporting bias (selective reporting). Within each domain, we will classify the RoB as high, unclear or low. Reviewers will also judge to determine whether any particular domain is impossible to achieve in any of the primary studies (like blinding in trials comparing GTN patch versus existing standard therapy) and likely or unlikely to affect the reported effect size of the outcome.

Primary studies will be classified as having an overall high RoB when they have been rated at least one domain as having high risk after exclusion of certain domain that is judged to be logistically impossible to achieve for that particular trial and unlikely to affect reported effect size of outcome. The overall RoB for each individual trial will be considered low if RoB is judged to be low in all domains and unclear if RoB is judged to be unclear in any of the domains.

**Quality of evidence**

We will also assess the quality of evidence for each outcome using the GRADE (Grading of Recommendations, Assessment, Development and Evaluation) approach that classifies evidence as high, moderate, low or very low quality based on considerations of RoB, consistency, directness, precision and publication bias.26 We attach a summary of findings table (online supplementary appendix 3) which is adapted using the GRADEpro software to demonstrate how we will present our GRADE assessment for the main outcomes.

Assessment of the individual and overall RoB categories as well as the quality of evidence will be performed independently by the two pairs of reviewers (LBL and CWL; LWF and NWM) with any discrepancies resolved by discussion and consensus or if necessary, via consultation with an external reviewer.

**Data analysis**

All analyses will be performed using RevMan V.5.3 (Cochrane Collaboration, Oxford) software. We will use DerSimonian and Laird random-effects model a priori to conduct the data analysis and meta-analysis. We chose the random-effects model as it produces more conservative CIs and it considers both within-study and between-study variability.21

For continuous outcomes, we will calculate the mean difference and its corresponding 95% CIs whenever possible. For dichotomous outcomes, we will calculate the relative risk and its corresponding 95% CIs. We will generate forest plots to demonstrate the individual and pooled effect sizes for the outcome of interest if there are at least two studies. We will assess for heterogeneity between studies by first visual inspection of the forest plots and then using the I² statistic. I² measures the percentage of the total variation in estimated effects of the outcome across studies that is due to heterogeneity rather than to...
chance. A P value of 0% indicates no observed heterogeneity, and larger values show increasing heterogeneity.

Regardless of the observed statistical heterogeneity (I² values), we plan to conduct the following a priori subgroup analyses for each outcome when each subgroup is represented by at least two studies. These subgroup analyses will be: ICH versus ischaemic stroke; minor (NIHSS ≤ five) versus major (NIHSS > five) ischaemic stroke; ischaemic stroke with versus without thrombolysis; prehospital versus non-prehospital (ED and inpatient) settings; time from stroke to randomisation ≤ 6 versus > 6 hours; time from stroke to randomisation ≤ 2 versus > 2 hours and high versus low overall RoB studies.

Missing data in the primary studies will be addressed in several ways. We will evaluate for rates of missing data in these primary studies, reasons for missing data and to contact primary authors for clarification if necessary. We will determine whether authors of these primary studies attempted to address the impact of missing data by using intention-to-treat analysis and performing sensitivity analyses through methods like imputation, best-case and worst-case scenario analyses to investigate how their reported effect size estimates had changed. We will then make judgement independently, through consensus and/or consultation with an external reviewer whether the reported effect size estimates (including any sensitivity analyses) by the primary authors will likely or unlikely be affected by their missing data. We will perform separate sensitivity analyses of our pooled results by including and excluding those studies that are judged likely to be affected by missing data to investigate how the pooled effect size estimates will be affected. Finally, we will also assess the risk of missing data (attrition bias) of the primary studies through our RoB and GRADE assessments.

Meta-analyses may result in type I errors due to an increased risk of random error when sparse data are collected and repeated significance testing when a cumulative meta-analysis is updated with new trials. We will perform trial sequential analysis (TSA) using a random-effects model for the primary outcomes (in-hospital mortality, BP lowering and late functional status). In the TSA, we will use a statistical significance level of 5%, a power of 80% and an estimated effect size difference (mean difference for continuous outcomes) between transdermal GTN versus placebo or control therapy as reported by the included trials. TSA generates the required information size calculated as diversity-adjusted information size (DIS) suggested by the estimated effect size difference; thereby providing important information on how many more patients need to be included in further trials. TSA also creates adjusted thresholds for statistical significance (trial sequential monitoring boundaries) with addition of each new trial. The cumulative Z curve which includes the selected trials; if it crosses the trial sequential monitoring boundary, will signify that a sufficient level of evidence has been reached and no further trials are needed. If the Z curve fails to cross the trial sequential monitoring boundary, the required information size is not reached and there is insufficient evidence to reach a conclusion.

TSA will be performed using TSA V.0.9.5.10 beta (Copenhagen Trial Unit, Centre for Clinical Intervention Research, Rigshospitalet, Copenhagen, Denmark, www.ctu.dk/tsa).

**DISCUSSION**

Acute ischaemic and haemorrhagic strokes are frequently managed in various clinical settings; from prehospital, ED to inpatient. High BP is common in both types of strokes and is associated with short-term poor outcomes (acute stroke recurrence, death within a few weeks and haematomata expansion) and adverse effects in the longer term (delayed death and dependency after several months). BP control is an essential part of the management of acute ischaemic and haemorrhagic strokes.

NO donors are candidate agents to lower BP in acute stroke because of its various beneficial properties ranging from vasodilatation to neuroprotection and inhibition of apoptosis. GTN is an example of a NO donor. Transdermal GTN offers an easily administered formulation which is valuable especially in the prehospital and ED settings to provide constant drug release.

There was a meta-analysis in 2016 and Cochrane review in 2017 that investigated the effects of transdermal GTN in acute stroke which reported no overall benefits. However, they reported a favourable functional outcome (improvement in mRS at 90 days) with transdermal GTN versus placebo or control therapy in a prespecified subgroup of patients with ultra-acute stroke (time from stroke to randomisation ≤ 6 hours). The meta-analysis and Cochrane review had important limitations. Apart from the ENOS trial, the remaining four included trials had small sample sizes (n=590) and all these trials were conducted by a single research group. In addition, that subgroup analysis involving ultra-acute stroke patients also suffered from a small sample size (n=312).

With the inclusion of the recently published multicentre RIGHT-2 trial which recruited 1149 patients with acute stroke within 4 hours of onset, our systematic review and meta-analysis will significantly increase the sample size available for pooling of studies; especially so when it will more than triple that used to examine the role of transdermal GTN in ultra-acute stroke (onset ≤ 6 hours). Our planned subgroup analysis of patients with ultra-acute stroke will address a significant gap in the literature that arose from these previous reviews.

In addition, our TSA for the important primary outcomes will reduce type I error. Our TSA will determine whether the DIS and trial sequential monitoring boundaries for these outcomes have indeed been reached in our
meta-analysis; signifying that a sufficient level of evidence has been attained to reach a conclusion.

Other strengths of our protocol include a comprehensive search strategy of published and unpublished literature, extensive subgroup analyses involving clinically important patient subgroups and using GRADE methodology to assess certainty of evidence.

Limitations to our protocol include the anticipated high clinical heterogeneity given the haemorrhagic and ischaemic subtypes of acute stroke as well as variation in timing of randomisation from stroke onset to transdermal GTN or placebo/control therapy and reporting of outcome measures across trials even within a subtype of acute stroke. We will address clinical heterogeneity by evaluating for statistical heterogeneity, explore predefined clinically important subgroup analyses and to account for inconsistencies in our GRADE evaluation. In order to address for differences in reporting of outcome measures across included trials, we will include a spectrum of primary and secondary outcomes. We will assess reporting of these outcomes independently and in duplicate and if there are discrepancies, we will resolve through discussion, consensus, potentially involving an external reviewer and contacting the primary authors for clarification.

In conclusion, this protocol describes the details and methodology of a planned systematic review and meta-analysis addressing the safety and benefits of transdermal GTN in acute stroke. The results of this meta-analysis are expected to fill the gap in the literature on the subgroup of patients with ultra-acute stroke (onset ≤6 hours), inform daily practice, clinical practice guidelines and guide areas of investigation for future RCTs.

Correction notice This article has been corrected since it first published. The provenance and peer review statement has been included.

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Appendix 1. Medline Search Strategy Through Pubmed Interface

Filters:
Time period: 1st Jan 1966 to 30th June 2020
Article type: Clinical Study, Clinical Trial Phases 1 to IV, Controlled Clinical Trial, Pragmatic Clinical Trial
Species: Humans
Language restrictions: No
Age limits: 13 to 80+ years

| #  | Searches                                      |
|----|----------------------------------------------|
| 1  | Acute stroke/                               |
| 2  | Ischaemic stroke (tw)                       |
| 3  | Haemorrhagic stroke (tw)                    |
| 4  | Intracerebral haemorrhage (tw)              |
| 5  | Brain infarction/                           |
| 6  | Brain haemorrhage/                          |
| 7  | Prehospital emergency care/                 |
| 8  | 1-7 or                                      |
| 9  | Transdermal glyceryl trinitrate (tw)        |
| 10 | Nitroglycerin/                              |
| 11 | Trinitroglycerin (tw)                       |
| 12 | GTN (tw)                                    |
| 13 | Nitric oxide donor/*                        |
| 14 | 9-13 or                                    |
| 15 | Mortality (tw)                              |
| 16 | Death (tw)                                  |
| 17 | Blood pressure/                             |
| 18 | Functional outcome*/ (tw)                   |
| 19 | Haemodynamic*/                              |
| 20 | Cerebral haemodynamic*/                     |
| 21 | 15-20 or                                   |
| 22 | 8 and 14 and 21                            |
Appendix 2. Data Collection Form (Adapted from Cochrane Collaboration)

General Information

| Date form completed |   |
|---------------------|--|
| Name of reviewer extracting data |   |
| Contact details of reviewer extracting data |   |
| Title of publication |   |
| Publication ID (first author and year of publication) |   |
| Country in which study was conducted |   |
| Study funding source |   |
| Possible conflicts of interests for study authors |   |

Primary Study Details

1. Methods

| Study Characteristics | Review Inclusion Criteria |
|-----------------------|---------------------------|
| Design (Type of randomized trial) | Blinded vs non-blinded |
|                        | Cross-over present |
| Method(s) of recruitment of participants |   |
| Unit of allocation (individual vs cluster/group) |   |
| Clinical setting | Pre-hospital vs Emergency Department vs Hospital |
| Types of intervention | Different dosing regimens of transdermal GTN |
### Types of comparator

| Types of comparator | Standard therapy vs placebo |
|---------------------|-----------------------------|

### Types of outcome measures

| Types of outcome measures |
|---------------------------|
| Primary:                  |
| Secondary:                |
| Safety:                   |

## 2. Study Population and Setting

| Study population description | Stroke subtypes: haemorrhagic vs ischaemic  |
|-----------------------------|---------------------------------------------|
|                             | Stroke onset to randomization               |
|                             | Other stroke subgroups (like IV thrombolytics, etc) |

### Inclusion criteria

### Exclusion criteria

### Start date

### End date

### Duration of participation (recruitment to last follow-up)

## 3. Participants (in intervention vs control/placebo groups)

| Total number of individuals randomized | Intervention group: |
|----------------------------------------|---------------------|
|                                        | Control/ placebo group: |

| Total number of clusters randomized (if applicable) | Intervention group: |
|-----------------------------------------------------|---------------------|
|                                                     | Control/ placebo group: |

| Number of withdrawals/exclusions | Intervention group: |
|----------------------------------|---------------------|
|                                  | Control/ placebo group: |

| Number of cross-overs | Intervention group: |
|------------------------|---------------------|
| Control/ placebo group: |
|------------------------|
| Baseline imbalances     |
| Other treatments (apart from intervention vs control/placebo) |
| Intervention group:    |
| Control/ placebo group: |
| Subgroups measured     |
| Subgroups reported     |

### 4. Outcomes (create a separate section for each outcome)

| Outcome name |
|--------------|
| Outcome type (Primary vs secondary vs safety) |
| Time points when outcome was measured (from start or at end of intervention or control/placebo) |
| Time points reported |
| Outcome definition |
| Method(s) of outcome assessment (using any tool/scale, etc) |
| Is the outcome assessment tool validated? |
| Persons measuring and /or reporting outcome |
| Imputation of missing data |
| Analysis via intention-to-treat or per-protocol or both |

### 5. Results (create a separate section for each outcome)

| Outcome |
|---------|
| **Dichotomous or continuous** |  |
|-------------------------------|---|
| **Subgroup**                  |  |
| **Time point**                |  |
| **Results** (may have more than two arms)** | Intervention group:  |
|                               | Control/placebo group  |
| **Number of missing participants** | Intervention group:  |
|                               | Control/placebo group:  |
| **Number of cross-over**      | Intervention group:  |
|                               | Control/placebo group:  |
| **Statistical methods used and appropriateness of these methods** |  |

**Risk of Bias Assessment (create a separate section for each outcome)**

| **Domain** | **Risk of bias (High/Low/Unclear)** | **Support for Judgement** |
|------------|-------------------------------------|---------------------------|
| Random sequence generation (selection bias) |  |  |
| Allocation concealment (selection bias) |  |  |
| Blinding of participants and personnel (performance bias) |  |  |
| Blinding of outcome assessment (detection bias) |  |  |
| Incomplete outcome data (attrition bias) |  |  |
| Selective outcome reporting (reporting bias) |  |
| Other bias |  |
Appendix 3. Summary of findings table

| Certainty assessment | No of patients | Effect | Certainty | Importance |
|----------------------|----------------|--------|-----------|------------|
|                      | Transdermal GTN | Control/placebo therapy | Relative (95% CIs) | Absolute (95% CIs) |
| In-hospital mortality (follow up: range 1 day to 3 months) [Primary Outcome] | | | | |
| Mean arterial pressure (follow up: range 1 day to 10 days) [Primary Outcome] | | | | |
| Modified Rankin Scale (follow up: range 3 months to 12 months) [Primary Outcome] | | | | |
| Development of intracerebral haemorrhage (follow up: range 1 day to 10 days) [Secondary Outcome] | | | | |
| Deterioration of NIHSS scores by at least 4 points during hospitalization (follow up: range 1 day to 10 days) [Secondary Outcome] | | | | |
| Length of hospital stay (follow up: range 1 day to 3 months) [Secondary Outcome] | | | | |
| Number of hypotensive episodes requiring intervention* (follow up: range 1 day to 10 days) [Safety Outcome] | | | | |

GTN: Glyceryl trinitrate; CIs: Confidence intervals; NIHSS: National Institutes of Health Stroke Scale
*Interventions include discontinuing transdermal GTN, administration of intravenous fluids and/or inotropic drugs