Pressor therapy in acute ischaemic stroke: an updated systematic review

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Abstract

Background

Low blood pressure (BP) in acute ischaemic stroke (AIS) is associated with poor functional outcome, death, or severe disability. Increasing BP might benefit patients with post-stroke hypotension including those with potentially salvageable ischaemic penumbra. This updated systematic review considers the present evidence regarding the use of vasopressors in AIS.

Methods

We searched the Cochrane Database of Systematic Reviews, MEDLINE, EMBASE and trial databases using a structured search strategy. We examined reference lists of relevant publications for additional studies examining BP elevation in AIS.

Results

We included 27 studies involving 1886 patients. Nine studies assessed increasing BP during acute reperfusion therapy (intravenous thrombolysis, mechanical thrombectomy, intraarterial thrombolysis or combined). Eighteen studies tested BP elevation alone. Phenylephrine was the most commonly used agent to increase BP (n=16 studies), followed by norepinephrine (n=6), epinephrine (n=3) and dopamine (n=2). Because of small patient numbers and study heterogeneity, a meta-analysis was not possible. Overall, BP elevation was feasible in patients with fluctuating or worsening neurological symptoms, large vessel occlusion with labile BP, sustained post stroke hypotension and ineligible for intravenous thrombolysis or after acute reperfusion therapy. The effects on functional outcomes were largely unknown and close monitoring is advised if such intervention is undertaken.

Conclusion

Although theoretical arguments support increasing BP to improve cerebral blood flow and sustain the ischaemic penumbra in selected AIS patients, the data are limited and results largely inconclusive. Large, randomised controlled trials are needed to identify the optimal BP target, agent, duration of treatment and effects on clinical outcomes.

Key words: stroke, blood pressure, acute, elevation, induced hypertension, vasopressor, ischaemic

Introduction

Up to 25% of patients with acute ischaemic stroke (AIS) have a systolic blood pressure (BP) below 140 mm Hg within 48 hours of onset.^{1, 2} Low BP in acute stroke is associated with poor prognosis: observational data from a large randomised trial showed that early death (2-week) increased by ~18% for every 10 mm Hg below 150 mm Hg. Another prospective study of patients with first-ever stroke reported that the relative risk of death increased at 1-month and 1-year by 28% and 17% respectively, for every 10 mm Hg systolic BP below 130 mm Hg.³ Recently, post-hoc analysis from the Multicentre Randomised Clinical Trial of Endovascular Treatment for Acute Ischaemic Stroke in the Netherlands (MR CLEAN) found that hypotension during mechanical thrombectomy (MT) was strongly associated with death and poor functional outcome at 3 months.⁴ Similar results were reported in a prospective endovascular treatment registry.⁵ The reasons for these observations are unclear but might be explained by concomitant factors including poor collateral circulation, prior cardiac disease, arrhythmias or sepsis; hypotension per se; or damage to the autonomic centres in the brain from the stroke itself.^{6,7} Moreover, studies have reported that hypotension post-stroke is associated with reduced cerebral blood flow leading to infarct extension and neurological deterioration.^{8,9} Thus, increasing BP in AIS patients with low BP, confirmed large vessel occlusion (LVO) and salvageable penumbra could potentially increase brain perfusion and improve functional outcomes.

Interventions to increase BP in acute stroke have been examined for >40 years. Volume expansion/haemodilution is one approach that has been assessed in a number of clinical trials. However, a Cochrane Review (21 studies, >4,000 patients) found no significant benefit.¹⁰ Another approach is using pharmacological agents to raise BP, which is recognised for treating cerebral ischaemia in patients with vasospasm after subarachnoid haemorrhage.¹¹ However, little is known of the effects in AIS. A meta-analysis, led by the Blood Pressure in Acute Stroke Collaboration in 2014 found insufficient evidence to provide reliable guidance on deliberate pharmacological intervention in acute stroke.¹² An earlier review of pressor therapy in acute stroke concluded that there were few trials and the results were inconclusive.¹³ Clinical guidelines on management of post-stroke hypotension provide no objective clarification, reflecting the paucity of evidence in this area. However, guidelines advise that patients with low BP may need inotropic support with close cardiac and neurological monitoring.^{14, 15} This suggests that potential causes of low BP in stroke should be addressed and vasopressor therapy may be considered in selected patients. However, it is unclear when to increase or maintain BP, duration, the ideal agent and which group of patients might benefit.

In the last decade, there are increasing reports of patients treated with intravenous thrombolysis (IVT) and or during mechanical thrombectomy (MT) receiving vasopressors to increase or maintain BP. This updated systematic review considers the present evidence regarding the use of vasopressors in patients with AIS.

Methods

Search strategy

We systematically searched several databases (August 2019 – December 2020) to identify studies examining the effect of elevating BP in AIS. The databases included MEDLINE and EPub ahead of print, In-process, Ovid Embase, Cochrane Central

Register of Controlled Trials (CENTRAL), Cochrane Database of Systematic Reviews, Web of Science and Scopus. No language restrictions were applied.

To further identify published, unpublished and ongoing trials we also searched:

• reviews of high BP in acute stroke¹⁶

• the ongoing Trials Section of Stroke and the Internet Stroke Centre Stroke Trials Registry

In addition, we also searched Google Scholar and the reference lists from published reviews and relevant trials.

The search strategy was designed and developed by the study's principal investigator (KK) with input from all collaborators. Keywords were used to search for studies testing BP elevation in AIS. An appendix detailing the search strategy is appended separately.

Inclusion criteria

We included any study which aimed to or reported on the effect of increasing BP in AIS. We included adults (aged 18 years or over) of either sex, eligible or not eligible for acute reperfusion therapy (ART) including IVT, MT, intra-arterial thrombolysis (IAT) or combined.

Exclusion criteria

We excluded studies where there was no classification or confirmation (neuroradiological or other) of AIS or where increasing BP was neither the aim nor the proposed mechanism of action.

Outcomes

The primary outcome was death or dependency assessed using the seven-level modified Rankin Scale (mRS). Secondary outcomes included stroke severity (National Institute of Health Stroke Scale-NIHSS), or any other clinical measures used to define neurological deficit, mortality and adverse events. Where available, we extracted information on systolic BP (SBP), diastolic BP (DBP) and mean arterial pressure (MAP) and heart rate (before, during and after treatment) and radiological effects of treatment (e.g. infarct growth/extension, collateral circulation).

Study extraction

Using the selection criteria, three authors (TAS, MO, CC) independently assessed published and unpublished articles from the electronic searches and with input from authors (ECS, KK) selected studies for inclusion. We resolved any disagreements by discussion.

Data extraction

For this update, data were independently extracted by three authors (TAS, MO, TJKH) and one author (KK) checked each value.

Data management

We sought information on the agent, dose, route of administration, timing, inclusion criteria, method of randomisation, allocation concealment, blinding and method of analysis (intention-to-treat).

The protocol was registered with the international prospective register of systematic reviews (CRD42021227717).

Results

The searches yielded 6995 publications. After removal of duplicates, 5972 titles were screened. Of these, eighty-four articles were selected for initial assessment and thirty-four publications were extracted for full text review. After applying the predefined selection criteria, twenty-seven studies were finally included for this review. (**FIGURE 1**)

Eleven studies were prospective of which seven were Randomised Controlled Trials (RCTs). Sixteen studies were retrospective of which ten were observational studies, two case series and four case reports.

Nine studies deliberately increased BP in patients treated with acute reperfusion therapy (ART) including IVT, MT, intra-arterial thrombolysis (IAT) or combined. Among these, six studies were during MT only, one in IVT, one in IVT and MT, and one in combination of IVT, MT and IAT.

The majority of studies (n=18) either:

- were conducted before ART
- included patients who were ineligible for ART
- did not provide information on ART

Of these, only five studies were RCTs and the other thirteen comprised observational studies and case reports.

Patients

The total number of patients in the included studies were 1886 (range: one to 256). Of these, 1551 participants received vasopressors to increase BP and 335 participants did not (controls). (**TABLE 1**)

Bias

We systematically assessed the risk of bias in the included seven RCTs using the Cochrane risk-of-bias tool. ¹⁷

Allocation

We assessed allocation concealment as 'low risk', 'high risk' or 'unclear risk' according to the Cochrane handbook for Systematic Reviews of Interventions (ref). One study of ART used computer-generated randomisation and was therefore deemed Low-Risk and the other study did not report the method of randomisation (unclear risk) (**SUPPLEMENTARY TABLE 1**).

Two trials of no ART used a computer-generated randomisation method (low risk) and the remaining three trials did not describe randomisation (unclear risk).

Blinding

In the ART studies, participants and investigators were blinded to treatments as follows:

- double-blind (n=1)
- open-label (n=1)

Of the five RCTs of no ART blinding was as follows:

- single blind (n=2)
- open-label (n=3)

Incomplete outcome data:

Both RCTs of ART were analysed by the intention-to-treat principle. Of the five RCTs of no ART, two were analysed according to the intention to treat principle and the remaining three were on a per-protocol basis.

Selective reporting

We did not see evidence of selective reporting in any of the seven RCTs.

Other sources of bias:

We did not find any other risks to the validity of the included RCTs.

Outcomes

Eighteen studies reported on functional outcome after BP elevation: nine used mRS, five reported NIHSS and four studies assessed both. (**TABLE 1**)

Two studies reported on language comprehension using lexical semantics.^{18, 19} One study mainly assessed safety²⁰ and two others on morbidity, symptomatic intracerebral haemorrhage, Glasgow Coma scale and death.^{21, 22} Two studies reported neurological deficits using non-standard scales.^{23, 24} One study used day 90 mRS as primary outcome, but also examined revascularisation using the Thrombolysis in Cerebral Infarction (TICI) score and haemodynamics.²⁵

One small prospective study reported on the effect of BP elevation on collateral circulation.²⁶

Two studies reported on volume of cerebral infarct after BP elevation.^{27, 28}

The primary outcome in one study was the need for haemodynamic intervention to maintain SBP target 140-180mm Hg.²⁹

Intervention

Vasopressors were used as a single agent or in combination in the included twentyseven studies. Overall, the most commonly used drug to increase BP was the selective- α 1 agonist phenylephrine.

In the ART studies, phenylephrine was used in four studies combined with other vasopressors^{25, 29-31} and in one study as a single agent.³² Norepinephrine was used thrice as a single agent ^{26, 33, 34} and once in combination.³⁰ Ephedrine was used three

times combined with other vasopressors.^{25, 29, 31} Metaraminol was used as a single agent in one study.³⁵

Of the studies of no ART, phenylephrine was used in eleven studies as a single agent to elevate BP. Dopamine was used once as single agent³⁶ and once in combination with other vasopressors.²⁰ Norepinephrine was used once alone²¹ and in one study combined with esmolol and sodium nitroprusside.³⁷ Dobutamine³⁸ and Epinephrine²⁴ were as used single agents in one study each. Saxena et al tested Diaspirin Cross-Linked Hemoglobin (a haemoglobin based oxygen carrying solution) to increase BP for three days in 85 patients within 18 hours of stroke onset.³⁹ In one retrospective study, the sympathomimetic drug midodrine was tested in combination with phenylephrine, dopamine, and fludrocortisone in 46 patients.²⁰ Another study reported using vasopressor therapy in patients but did not mention which agent was used.²³

The effects of vasopressors on receptors and pharmacodynamics are listed in **SUPPLEMENTARY TABLE 2**.

Haemodynamics

The baseline BP, definition of target achieved and duration of treatment varied between the studies (**TABLE 2**). The measures of BP were defined as: SBP (n=15 studies), Diastolic BP (DBP) (n=4), Mean Arterial Pressure (MAP) (n=11) or combined (n=5).

Target blood pressure

The target BP was predefined in twenty-one studies, either as a percentage increase from baseline (n=8), or absolute SBP, DBP, MAP or as combined (n=13).

Of the nine ART studies which assessed increasing BP, six studies aimed for a predefined SBP target which ranged between: 140-180 mm Hg $(n=2)^{29, 32}$; 140-160 mm Hg $(n=2)^{26, 34}$; 160-180 mm Hg $(n=1)^{.35}$ One study aimed for a target as SBP \ge 140 mm Hg and MAP \ge 70 mm Hg.²⁵ In one study, the target BP was set at 10-20% higher from baseline.³³

In the eighteen studies where participants did not receive ART or were not eligible, fifteen reported a predefined BP target. In eight studies, BP target was set as 10%, 20% or a 10-20% increase from baseline with seven assessing MAP and one using SBP. In the remaining seven studies, the BP target varied from SBP 140-220 mm Hg, 90-130 MAP and one study used DBP 85-100 mm Hg.

In one prospective RCT of no ART (n=218), patients were randomised to three systolic BP (SBP) treatment groups: 140-160mm Hg, 161-180mm Hg and 181-200mm Hg.³⁷

Duration of treatment

The duration of BP elevation was reported in only one of the ART studies and the median was one hundred and five minutes.³⁴ From the published data, we were unable to determine treatment duration in other ART studies.

Of the eighteen studies of no ART, thirteen reported on duration of BP. In these studies the duration of BP elevation ranged from:

- 5 minutes 1 hours (n=3)
- 1 hour 24 hours (n=2)
- 24 hours 72 hours (n=3)
- 72 hours 168 hours (n=1)

• > 168 hours (n=1). In this study, the threshold SBP for neurological deterioration (defined as SBP below which a sustained neurological decline occurred at least twice, and above which the decline was rapidly reversed with increase in BP) was observed in 10 of 33 patients treated with phenylephrine. Patients in this study were treated for up to 24 days.

In one study, phenylephrine was used in 6 patients with ischaemic stroke (within 1 week of onset) with diffusion and perfusion-weighted imaging mismatch (DWI/PWI) mismatch on magnetic resonance imaging (MRI) until functional improvement or target MAP of 130 mm Hg was achieved.¹⁹ In two other studies, patients had their infusion continued and maintained until the motor score on the National Institutes of Health Stroke Scale (NIHSS) remained stable for 24 hours.^{40, 41} (**TABLE 2**)

Blood pressure target achieved

In the ART studies, only one RCT achieved a predefined SBP target (160-180 mm Hg).³⁵ In this study, patients randomised to treatment with metaraminol had a mean SBP of 167 mm Hg throughout the duration of MT. There was insufficient information to make a judgment whether the pre-defined target was achieved in the other ART studies.

In the seventeen studies where participants did not receive ART, ten reported reaching the predefined BP target. The number of patients who reached the predefined target varied between studies and the timing within target also varied (**TABLE 2**).

In one retrospective case analysis of 100 patients, 46 received various agents to maintain 20% increase from baseline BP. Of these, only 16 patients reached the predefined target.²⁰ In a case series of acute lacunar ischaemic strokes, phenylephrine was successfully used to increase SBP to 20% from admission in 42% of patients.⁴⁰ In four small studies, the predefined BP target was attained in all patients.^{18, 19, 27, 42}

Collateral circulation

Only one study of MT reported the effect of collateral status (assessed using Computed Tomography angiography) on functional outcome following treatment with nor-epinephrine to increase BP.²⁶ This study reported a favourable outcome (mRS \leq 2) in 60% of patients with moderate or good collateral circulation compared to only 28% of patients with poor collaterals.²⁶

Amongst the studies where patients did not receive ART, none reported on collateral circulation before, during or after treatment.

Effect of blood pressure elevation on outcomes

Of the ART studies, three studies found no significant association between increasing BP during MT and mRS.^{25, 34, 35} Similar results were observed in one study using a combination of MT, IAT and IVT.³¹ In three studies of MT, worse outcomes (mRS >2) were associated with haemodynamic fluctuations in MAP^{30, 32} or SBP.²⁶

One study of MT reported a non-significant improvement in NIHSS compared to baseline after vasopressor therapy.³⁴

In one study where the primary outcome was haemodynamic intervention during MT, intraprocedural hypotension (SBP < 140 mm Hg) was higher among patients who received conscious sedation compared to no sedation (80% vs 50%).²⁹ The majority were self-limiting, but 15% of all patients required treatment with phenylephrine and ephedrine to increase BP.

Of the studies of no ART, three studies reported significant improvement in mRS at discharge^{40, 41} or at day 90^{41, 43} in non-cardioembolic stroke with vasopressor therapy.

Following treatment, changes in NIHSS were reported in eight studies. In each study, patients who had BP elevation had lower NIHSS scores compared to baseline, of which five were statistically significant ($p \le 0.05$) (TABLE 2).

One study reported improvement in participant language assessed using the lexical semantics with increase in MAP.¹⁹ The same group reported that in a single patient with large fronto-temporal stroke, language comprehension improved when perfusion of the Wernicke's area increased with a parallel increase in MAP.¹⁸

Three studies assessed safety as a key endpoint in patients (n=95) assigned to vasopressors and each reported that BP elevation was safe and feasible.²⁰⁻²² The investigators concluded that there was an acceptable rate of complications, considering the severity of stroke and the relative aggressive nature of intervention.

In the study by Saxena et al, functional outcome was significantly worse in the group treated with Diaspirin cross-linked Hemoglobin (mRS 3 to 6 at 3 months: 85% in treated patients compared to 51% in the non-treated group).³⁹ The authors concluded that differences in baseline stroke severity and dose-dependent increase in endothelin levels could explain the results.

Using perfusion-weighted MRI, one study reported significant reduction in volume of hypoperfused brain after infusion of phenylephrine.²⁷ This corresponded to a significant improvement in mean NIHSS from 9.3 on admission to 4.8 at day 3. There were no adverse events in the group treated with phenylephrine.

Another group reported a case of a 76 year-old man with fluctuating right-hand clumsiness and mild right leg weakness and MRI confirmed a left anterior cerebral artery infarction with DWI/PWI mismatch.²⁸ After treatment with phenylephrine, a repeat MRI showed that the perfusion deficit had decreased by 40%.

The authors of three studies (2 case series and 1 prospective RCT) concluded that AIS patients with atherosclerosis were more likely to have post stroke hypotension and therefore vasopressors were indicated.^{20, 41, 43}

Adverse effects

Of the studies with ART, only one study reported adverse events.

Marzan et al reported a single episode of paroxysmal atrial fibrillation followed by ventricular tachycardia with norepinephrine in one patient, so treatment was discontinued.³³ In this study, 4 patients died; however it is not clear if these patients received ART.

Eight studies of no ART reported adverse effects with vasopressor therapy. Of these, the most common was intracerebral haemorrhage (n=44). Other complications included haemorrhagic transformation of infarct, brain herniation and worsening of cerebral oedema (n=8). Pulmonary oedema and chest tightness were also reported, but less frequent (n=3 each) with treatment.

Thirty-two patients who received vasopressor therapy and no ART died^{21, 23, 24, 37} but it is unclear whether this was directly related to treatment, duration or other complications. Other studies reported on mortality, but did not differentiate between the intervention group versus controls.

Discussion

This review found several retrospective observational studies evaluating BP elevation in AIS and seven small prospective RCTs. Although the studies in this review varied in inclusion/exclusion criteria and methodology, the results show that treatment was feasible. Although no formal meta-analysis was possible, this review found that patients who may be more likely to receive vasopressor therapy included:

- stroke from atherosclerosis
- large vessel occlusion in the anterior circulation including patients treated with MT, or IAT or combined
- ischaemic stroke presenting outside time-windows for IVT
- patients with fluctuating or worsening neurological symptoms including lacunar syndrome

A recent trial showed that increasing BP was significantly associated with early neurological improvement and functional independence day 90 (mRS 0-2) in patients with acute lacunar ischaemic stroke.⁴³ One explanation for the observed results is that increasing BP directly increases blood volume to an end-artery in the brain and improves microcirculation.⁴¹

The relationship between increasing BP in acute stroke and collateral circulation was unclear but two studies reported that a modestly higher target (170-190 mm Hg) was significantly associated with improved collaterals including patients treated with IVT.^{44, 45} This suggests that there is no 'ceiling effect 'for BP on collateral status, however the effects when SBP exceeds 180 mm Hg need further exploration. The results are relevant in AIS, where cerebral autoregulation is impaired and blood flow is directly dependent on systemic BP. The development of collateral circulation is dynamic and failure is associated with increasing infarct growth and poor outcomes.⁴⁶ Nevertheless, these data suggest the prognostic importance of hypotension in AIS and a potential therapeutic role for vasopressors.

In this review, it was unclear whether the rates of mortality were in relation to treatment and adverse events were not consistently reported. The studies varied in the target BP for vasopressor therapy; earlier studies tested a SBP

target of 150-175 mm Hg and one trial tested different BP thresholds.³⁷ In this trial, higher rates of symptomatic intracerebral haemorrhage (sICH) were seen in the patient group having a SBP target 180-200mm Hg. This could be relevant as the association between BP and outcomes after MT indicate a J-shaped or U-shaped curve suggesting a narrow window of therapeutic target and response.⁴⁷

Amongst various vasopressors, this review found that phenylephrine was the most commonly used agent with few significant neurological adverse events. Phenylephrine has selective α 1-agonist properties and likely to increase BP without substantial cerebral vasoconstriction; the effects are attributed to the low density of α 1-receptors in the cerebral vessels. Additionally phenylephrine has the advantage that it can be given as a bolus and infused though a peripheral line unlike other agents, which generally require a central venous catheter. In addition to phenylephrine, norepinephrine was commonly tested to augment cerebral blood flow and preserve the ischaemic penumbra. The study by Schwartz et al and the results of a retrospective study³³ indicate that norepinephrine is effective in elevating BP but is more likely to cause adverse cardiac effects due to its β -1 receptor action. Atrial fibrillation was reported in one study with phenylephrine and norepinephrine but one case-note review reported no significant differences in mortality.²² In one study, cumulative dose of norepinephrine to maintain SBP at 140-160 mm Hg during MT was associated with worse functional outcome (mRS >2) at day $90.^{26}$ This effect was independent of the duration of MT indicating that norepinephrine could have an adverse effect on collateral circulation. This finding is relevant as norepinephrine is widely used to stabilise BP during interventions.²⁶

Other vasoactive agents including epinephrine, dobutamine, dopamine and Diaspirin cross-linked Haemoglobin have been tested but the present results indicate less convincing evidence for increasing BP in AIS. Moreover, patients with history of recent congestive cardiac failure, myocardial infarction, unstable angina, bradycardia, uncontrolled hypertension or significant peripheral arterial disease, and treated with digoxin, mono-amine oxidase inhibitors or tricyclic antidepressants will not be eligible for treatment with these agents.

An important issue with the studies was that none clearly defined post-stroke hypotension. Moreover, there was very little information on whether patients had a history of high BP and whether they were on antihypertensives. Furthermore, it was unclear how BP was measured before, during and after treatment. As a result it is difficult to assess the quality of the reported readings.

The studies in this review varied by examining SBP, MAP and/or DBP as the therapeutic physiological target during pressor therapy. This reflects the uncertainty on which measure best describes regional cerebral blood flow in AIS or may be more than one parameter is needed. MAP is dependent on cardiac output and peripheral vascular resistance and indirectly associated with Cerebral Perfusion Pressure, (ie Cerebral Perfusion Pressure= MAP-Intracranial Pressure). It is commonly used in critical care management of patients with subarachnoid haemorrhage or severe traumatic brain injury. ⁴⁸ The role of MAP in AIS is unclear ⁴⁹ and the interaction with pressor therapy and effects on outcome needs to be tested in well-designed prospective studies.

Considering the present results, it may be postulated that patients with AIS who might benefit from vasopressor therapy include those:

- with sustained SBP <130 to 150 mm Hg or those with significant drop in BP (>20 mm Hg). This could include those treated with IVT, LVO and low BP
- ineligible for IVT to maintain cerebral perfusion until development of collateral circulation or recanalisation of the occluded vessel, but with levels of BP still below those where guidelines recommend antihypertensive treatment.^{14, 15}
- severe ipsilateral extracranial or intracranial large vessel occlusion or stenosis
- eligible for MT for large vessel occlusion as 'bridging therapy ' until collateral circulation is restored. This could include those presenting in later-time windows (12-24 hours) with a salvageable ischaemic penumbra.
- patients with unstable collateral circulation including those fluctuant or progressive neurological symptoms
- MT during GA when periprocedural BP is known to be labile⁵⁰

It is essential that future trials report on how BP is measured by including the following information:

- equipment including manufacturer, model, method of measurement (invasive or non-invasive) and validity of the equipment used
- who measured BP and were they trained, assessed or reassessed
- measurements: the number of readings at each time point and site of reading

Future studies evaluating the efficacy of manipulating BP in acute stroke should also account for the volume of the ischaemic core at presentation (marker of salvageable brain and a risk factor for sICH) and the presence or absence of LVO (marker for the potential for BP manipulation to sustain collateral circulation or to maintain brain perfusion where normal autoregulatory mechanisms are impaired).

Conclusions

This updated review found that increasing BP is feasible in AIS (with close monitoring) and to date very few RCTs have been performed. The interpretation of the results of the published studies was complicated by differences in study methods, participants, measures of BP, duration of treatment and outcome measures. The small sample sizes of what were mostly retrospective studies, limit any reliable conclusion of the effects of BP elevation in AIS.

This review found that phenylephrine was the most common drug to increase BP in acute stroke. However, treatment should only be considered after possible causes of low BP are investigated and treated. This review suggests that vasopressor therapy should be ideally restricted to the acute phase of stroke when potential viable ischaemic penumbra exists. Investigations to demonstrate this viable penumbra should be performed before treating patients and beyond the acute phase of stroke.

This review indicates that the risk-benefit of BP elevation (SBP, DBP, MAP individually or combined) in patients with AIS needs to be tested in well-designed prospective RCTs. These studies should assess which measures of BP, (SBP, DBP, MAP) either alone, or combined), which targets best predict the effects of treatment and which clinical scenarios (such as confirmed LVO) are most likely to benefit from pressor therapy.

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DISCLOSURES

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FIGURE 1. Flow chart of searches

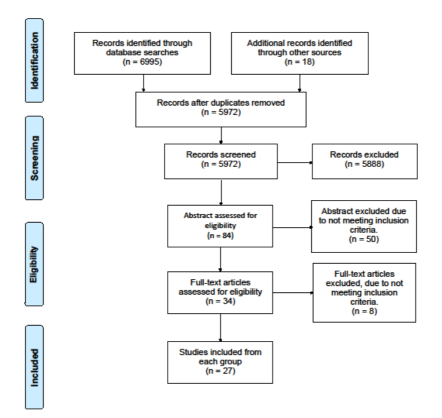


TABLE 1. Characteristics of included studies

Study	Design & Main inclusion criteria	Intervention & Duration of intervention	Outcomes	Results	Adverse effects	Modality of reperfusion
	·	S	tudies of ART			
Alcaraz 2019	Retrospective N=126 EVT anterior circulation	PhE., Ephedrine -	Haemodyna mic intervention	24 received vasopressor	-	EVT
Deng 2019	RCT N=51 A/C: 25/26 Anterior LVO mRS <3	Metaraminol -	90 days mRS	No significant change in mRS	C: 1 Perforated vessel + 1 ICH	EVT
Rasmussen 2018	RCT N=128 A/C: 100/28 NIHSS ≥ 10 Infarct <70 ml, Independently living Groin puncture <6h OTT Anterior circulation LVO	Ephedrine, PhE -	90 days mRS TICI score Haemodyna mic parameters	No significant change in mRS at 90 days	None	EVT
Schönenberg er 2018	Retrospective N=150 (GA/CS: 73/77) AIS, ICA occlusion and/or MCA. NIHSS > 10. Planned EVT	NEp Median 105 min (IQR: 80–150)	24h NIHSS 90 days mRS	No significant association between SBP, DBP and MAP and change in 24h NIHSS. No change in 90 day mRS	-	150 EVT 96 IVT + EVT

Treurniet 2018	Subgroup analysis of MR CLEAN N=60 BP data available.	NEp., Ep., PhE -	90 days mRS	 > change in MAP was associated with worse mRS (acOR) 0.95 per mmHg, (95% CI 0.92- 0.99) decrease. MAP during GA 10 mmHg lower than baseline MAP constituted a 1.67 times lower odds of a shift towards good outcome on the mRS. 	None	EVT
Whalin 2017	Retrospective N=256 EVT for AIS	PhE -	90 days mRS ≤ 2	No difference in BP between 2 groups. Low MAP before reperfusion was associated with mRS >2 (P=0.04)	19 ICH	EVT
Mundiyanapu rath 2016	Prospective study N=64 EVT of ICA or MCA in AIS NIHSS>10	NEp. -	Favorable outcome: mRS ≤ 2 Collateral status	27 with mRS ≤2 37 with mRS >2 mRS >2 was associated with higher cumulative dose NEp (OR 0.142, CI: 0.038- 0.525, p=0.003) Median mRS at 90 days was 3	None	EVT
Sivasankar 2016	Retrospective N=84 EVT of anterior circulation and/or treated with IA-tPA.	PhE., Ep., Ephedrine During EVT	mRS	Change in mRS was not associated with the use of vasopressor.	None	77 EVT (47 MT alone, 6 IA-tPA

Marzan 2004	Retrospective N=34	NEp.	mRS	mRS at discharge 3±1	N=4 Died	alone, 24 MT + IA- tPA) - EVT + IV- tPA 8 IVT
	NIHSS ≥ 5, SBP < 140 mmHg, No evidence for hypovolemia	14-96 h			1 Ventricular tachycardia	26 None
		Studie	s of no ART	'	,	
Bang 2019	RCT N=153 A:76/C: 77 NoncardioembolicAIS NIHSS 4–18, ineligible EVT or IVT or with progressive stroke*.	PhE Mean 4.9 ±1.9 days	7-day NIHSS 90 days mRS	A: NIHSS improvement in 44/76 (58%) Favourable day 7 NIHSS OR 2.49 (95% CI 1.25–4.96, (p=0.010) OR 90 day mRS (0-2): 2.97 (95% CI 1.32–6.68, (p=0.009). C: NIHSS improvement in 24/77 (31.2%)	sICH: 1A/0C (p=0.313) HTI: 5A/0C (p=0.022)	*
Nasi 2019	RCT N=218 A1: 140-160mmHg n= 77 A2: 161-180mmHg n= 75 A3: 181-200mmHg n= 66	NEp, Esmolol, Ntr.pru 24 h	90 days mRS	No change in mRS at 90 days	sICH was more frequent in higher SBP range (p=0.048). A1: 1, A2: 2,	*

	AIS ≤ 12 OTT ineligible for IVT				A3: 6 mRS 6: N=30 (12 received vasopressor)	
Kang 2017	Retrospective N=66 A/C: 25/41 Lacunar infarct diameter on MRI <20mm Motor progression	PhE Until 24h stable NIHSS score + 24h tapering dose.	NIHSS at discharge mRS ≤ 2 at discharge and 90 days	A vs C: NIHSS: 4.4±2.5 vs 6.0±3.7, (p=0.036) mRS ≤ 2 at discharge: 21 vs 20 (p=0.004) 90 days mRS ≤2: 18 vs 15 (p=0.0011)	None	*
Lim 2011	Retrospective N= 82 A/C: 52/30 Lacunar syndrome with motor weakness. DWI confirmed infarct <20 mm in diameter	PhE Until 24h stable NIHSS score + 24h tapering dose.	NIHSS mRS Hospitalisati on days	A vs C NIHSS motor score at discharge 1.01 \pm 1.47 vs 1.86 \pm 1.92 (p=0.042) mRS \leq 2: 32 vs 15 (p=0.044) Hospitalisation days: 13.1 \pm 7 vs 18.6 \pm 17.3 (p=0.047)	Chest tightness 4 (3A/ 1C) Dysuria (n=2)	*
Koenig 2006	Retrospective N=100 Symptoms of AIS, MRI <7 OTT MRI suggestive of AIS	PhE., Dpm., Mddr., Fl.cort. -	Safety NIHSS	No significant change in NIHSS	C:2 NSTEMI 2 PE. A:1 sICH, 1CE, 1PE	*

Chalela 2005	Case report N=1 A2 occlusion. perfusion/diffusion mismatch	PhE 5 min	MRI Perfusion	Perfusion deficit decreased 40% and hypoperfusion improved. CBV did not change.	None	*
Hillis 2004	RCT N=15 A/C: 10/5 Quantifiable, aphasia, hemi-spatial neglect, and/or hemiparesis < 7 days OTT >20% + >30 ml MRI mismatch	PhE 24-72 hours	PWI volume at baseline vs follow up Mean NIHSS at baseline vs day 3	PWI volume: A: 147ml to 58.3ml (p=<0.001) C: 106ml to 94ml (p= 0.19) Mean NIHSS A: 9.3 to 4.8 (p<0.001) C: 12 to 11.8 (p=0.5)	HTI: C:1	*
Hillis 2003	RCT N=15 A/C: 9/6 <7 days from AIS. Perfusion/diffusion mismatch Neurological deficits	PhE <72 h	NIHSS & cognition	A: Mean NIHSS from day 1-3: 10.2-5.6 (p=0.002) C: No significant difference in NIHSS between day 1-3 A vs C: Mean NIHSS: day 3: 5.6 vs 12.3 (p=0.01) 6-8 weeks: mean 2.8 vs 9.7 (p=0.04) A: cognitive battery improved 58.7 \pm 36% to 27.9 \pm 32% (p <0.002) C: No significant change in cognitive battery (64.3 \pm 31 vs. 67.3 +/- 34% errors)	None	*

Oliviera-Filho 2002	Case report N=1	Dpm. Circa 48 h	NIHSS	Improved according to graph in the publication but no further details	-	*
Schwarz 2002	Prospective N=19 Acute MCA territory stroke	NEp. -	sICH, ICH GCS & Death	15 severely disabled at dis3)4 died	scharge (GCS	*
Hillis 2001	Case series N=6 <7 days from AIS. Aphasia PWI/DWI mismatch	PhE Until function improved and/or MAP 130 or adverse effects	Lexical semantics	6 improvement in lexical semantics when MAP was elevated	-	*
Hillis 2001	Case report N=1	PhE 12 h	Language comprehensi on	Improvement in parallel with MAP Improved perfusion on MRI	-	*
Rordorf 2001	Prospective N=13 NIHSS>4, No ICH, ≤ 12 hours from onset	PhE ≥30 min	NIHSS	7 improved NIHSS	None	-
Saxena 1999	RCT N=85 A/C: 40/45 AIS in the anterior circulation ≤ 18 hours from onset	DCLHb. Every 6h for 72h	Daily physical examination while admitted and at 90 days. mRS at 90 days.	Adverse events: A: Patient 1: scleral icterus hypertension, PE, CE, dea Patient 2: pancreatic insuf nausea, anaemia Outcome at 90 days was worse in treatment group. Treatment with DCLHb (p 3.9; CI, 1.4 to 12.0) were predictors of a worse outc	ath. ificiency significantly (p=0.002). =0.015; OR, independent	*

				C: 51% 90 days mRS 3-6 Reperfusion of the hemisp	5 90 days mRS 3-6 (23% death) 6 90 days mRS 3-6 (9% death) fusion of the hemisphere through ral flow with patients symptoms ed		
Duke 1998	Case report N=1 Embolic occlusion of the middle cerebral artery	Dobutamine -	Clinical	Reperfusion of the hemisp collateral flow with patients resolved	U	-	
Rordorf 1997	Retrospective N=63 A/C: 30/33 Cerebral ischaemia	PhE Mean 110h (range, 7 to 576 h).	Morbidity and mortality	No significant difference in outcome between groups.	CE A:2, C:6. Radiological abnormalitie s: A:8 C: 12, Arrhythmia A:1	*	
Meier 1991	Prospective N=81 A/C=37/44 <6 h from stroke onset Acute blood flow defect in carotid system	Ep 5 min x3	Consciousne ss and severity of paresis Mortality	A: Improved Consciousness 22%, Improvement in Severity of paresis 24%, Improved paresis and consciousness 16%. No effect 38%, A: Mortality 21 days	None	*	

				14/37 C: Mortality 21 days 28/44		
Wise 1972	Case series N=13 Nonhypertensive with focal brain ischaemia without change BP≤4 hours of onset	Vasopressor** -	Neurological function	N=5/13 improvement of brain function immediately	-	*

Patients under 18 years of age were not included.

* Progressive stroke was defined as symptom worsening (i.e., a ≥2-point increase in NIHSS score, including an increase in the motor score for the affected upper and lower limbs) during hospitalisation and the presence of new lesions or infarct growth on diffusion-weight imaging magnetic resonance scan performed within 24 hours of treatment.

** Not specified further in the publication

A = Active group; AIS = acute ischaemic stroke; ART: acute reperfusion therapy; BP = blood pressure; C = Control group; CBV = cerebral blood volume

CE = Cerebral oedema; CS = conscious sedation; DBP = diastolic blood pressure; DCLHb = Diaspirin cross-linked hemoglobin

Dpm = Dopamine; Ep = Epinephrine; EVT = endovascular treatment; Fl.cort = Fludrocortisone; GA = general anaesthesia

GCS = Glasgow coma scale; HTI = Haemorrhagic transformation of infarct; IA-tPA = Intra-arterial tissue plasminogen activator

ICA = Internal carotid artery; ICH: intracerebral haemorrhage; IVT = intravenous thrombolysis; MAP = mean arterial pressure; MCA = middle

cerebral artery Mddr = Midodrine; mRS = modified Rankin scale; MT = Mechanical thrombectomy; NEp = norepinephrine

NIHSS = National Institutes of Health Stroke Scale; NSTEMI = non ST elevation myocardial infarction; Ntr.pru.= Nitroprusside

OTT = onset to treatment; PE = Pulmonary oedema; PhE = Phenylepinephrine; SBP = systolic blood pressure; sICH = symptomatic intracranial hemorrhage; - = Not reported; *= indicates no ART/conducted before ART

Study	Baseline BP (mmHg)	Target BP (mm Hg or % as reported)	BP during/end of treatment (mm Hg)	Change in BP (mm Hg or % as reported)	Attained target
		Studies	of ART		
Alcaraz 2019	Mean SBP±SD: 143.63±24.8 Mean DBP±SD: 79.34±13.2	SBP 140-180	-	-	-
Deng 2019	-	160–180	Mean SBP: A/C: 167 (IQR 150–175 [113– 188]) / 139 (IQR 135–143 [115– 154])	diff in median SBP = 28	100%
Rasmussen 2018	GA mean SBP±SD 143 <u>+</u> 15, MAP 95 <u>+</u> 8 CS mean SBP±SD 155 <u>+</u> 20, MAP 101+12	SBP≥140 & MAP≥70	-	-	-
Schönenberge r 2018	Mean SBP±SD: 165.3±26.3 Mean DBP±SD: 93.9±17.2 Mean MAP±SD: 117.7±17.8	SBP : 140 to 160	Mean SBP±SD: 142.2±20.5 No control group	-	-
Treurniet 2018	Median (IQR): SBP: 140 (126–155) DBP: 80 (70–90) MAP: 100 (92–110)	No target	median (IQR) during GA: SBP: 119 (106–130) DBP: 64 (59-71) MAP: 81 (77-91)	-	-
Whalin 2017	Mean SBP±SD:157.6 ± 29.3 Mean MAP±SD :108.5 ± 18.6	SBP 140-180	-	-	-
Mundiyanapur ath 2016	Median SBP: 150.0 (IQR 140.0– 170.0)	SBP 140-160	Median 146.0 (IQR 131.0–159.0) after extubation	-	-
Sivasankar 2016	Baseline MAP±SD: MAC 103.3±28.1; TIVA	No target	Mean MAP±SD during procedure: MAC 95.1±16.2; TIVA 89.0±10.8;	-	-

	102.0±25.5; Vol 109.5±17.7; Comb 104.8±13.7		Vol 95.8±12.2; C	Comb 94.3±11.0		
Marzan 2004	Mean SBP <u>+</u> SD 127 <u>+</u> 14, Mean DBP <u>+</u> SD 65 <u>+</u> 10	10-20% > baseline SBP	Mean SBP±SD: Mean DBP±SD:		SBP: 17 <u>+</u> 10% DBP: 18 <u>+</u> 18%	¶
		Studies o	f no ART		I	
Bang 2019	Mean SBP±SD: A/C: 144.2±18.5/146.5±16.8	>20% from baseline	Mean SPB±SD A/C: 178.6 ± 18.	7/146.5±16.8	-	86%
Nasi 2019	Median SBP: A1:166 (144-185) A2:163 (140-189) A3:169 (151-203)	A1: 140-160 A2: 161-180 A3: 181-200	Median SBP: A1: 153 (IQR 147-160) A2: 163 (IQR 151-170) A3: 178 (IQR 167-184)		-	A1: 70%, A2: 61%; A3: 60%
Kang 2017	Mean SBP±SD: A/C: 139.1±18.5/150.4±21.5	>20% from baseline SBP	Mean SBP±SD A/C: 175.4±21.2/138.41±17.9 (p= <0.0001)		-	80%
Lim 2011	Mean SBP±SD A/C:161.7±24.8/168.0±31.9	>20% from baseline	Mean SBP±SD A/C:168.7±13.6/	′158.2±17.9	-	42%
Koenig 2006	Mean MAP*: A/C: 106/100	MAP 10-20% > baseline	Mean MAP A/C: 103±14/96:	±13(p=0.002)	-	35%
Chalela 2005	MAP 91	-	MAP 142		MAP increase 51	-
Hillis 2004	Mean MAP±SD A/C: 97.9±15	10–20% MAP increment, max 130	-		-	100%
Hillis 2003	Mean MAP: A/C: 94.2±15/103.5±15	MAP 90-130	Mean MAP A/C: 115.8±16/1	08.2±14	-	100%
Olivia-Filho 2002	SBP 120	160 SBP	-	-	-	-
Schwarz 2002	Mean MAP <u>+</u> SD: 83.6 <u>+</u> 1.6	> 10 MAP	Mean MAP±SD:	108.9±2.0	MAP increase 25.3	-

Hillis 2001	-	Increments of 10% MAP ≤130	-	-	-	-
Hillis 2001	MAP: 72-88	MAP 90-100	-		-	100%
Rordorf 2001	Mean SBP <u>+</u> SD Responders: 140±13 Not responders: 141±23	SBP 160 or > 20% from baseline	174±15	D Responders: lot responders: -	-	100%
Saxena 1999	Mean MAP <u>+</u> SDA/C: 113 <u>+</u> 14/-	-	Mean MAP±S 134±20/109±2		-	-
Duke 1998	SBP 110-120	-	SBP 110-120		-	-
Rordorf 1997	Mean SBP 152**	-	Mean SBP 15	6	-	-
Meier 1991	-	210-220	-		-	-
Wise 1972	-	SBP: 150-170 DBP: 85-100	-		-	38%

Heart rate was mentioned in Schönenberger, Marzena and Saxena. * Estimated from fig. 3 in the study publication. ** Calculated from table 1 in original article.

¶ Systolic blood pressure values varied within 5% of the target value in 16 (47%), within 10% of the target value in 14 (41%), and within 15% of the target value in the remaining 4 (12%) patients.

A: active; ART: acute reperfusion therapy; C: Control; Comb = Combined intravenous and volatile anaesthesia; CS = Conscious sedation; DBP = Diastolic blood pressure; GA = General anaesthesia; HR = Heart rate; MAC = Monitored anaesthesia care; MAP = Mean arterial pressure; PAWP = Pulmonary arterial wedge pressure; SBP = Systolic blood pressure; TIVA = Total intravenous anaesthesia; Vol = Volatile; - = Not reported