Clinical Policy: Use of Thrombolytics for the Management of Acute Ischemic Stroke in the Emergency 1 2 **Department** 3 This DRAFT is EMBARGOED – Not for Distribution 4 5 6 From the American College of Emergency Physicians Clinical Policies Subcommittee (Writing Committee) on 7 Thrombolytics: 8 9 Bruce M. Lo, MD, MBA, RDMS (Subcommittee Chair) 10 Christopher R. Carpenter, MD, MSc 11 Ken Milne, MD, MSc Peter Panagos, MD 12 13 Jason S. Haukoos, MD, MSc (Methodologist) 14 Deborah B. Diercks, MD, MSc (Committee Chair) 15 16 17 Members of the American College of Emergency Physicians Clinical Policies Committee (Oversight Committee): 18 19 Deborah B. Diercks, MD, MSc (Chair 2021-2024) 20 John D. Anderson, MD 21 Richard Byyny, MD, MSc (Methodologist) 22 Christopher R. Carpenter, MD, MSc 23 Benjamin W. Friedman, MD (Methodologist) 24 Seth R. Gemme, MD 25 Charles J. Gerardo, MD, MHS 26 Steven A. Godwin, MD 27 Benjamin W. Hatten, MD, MPH 28 Jason S. Haukoos, MD, MSc (Methodologist) 29 Amy Kaji, MD, MPH, PhD (Methodologist) 30 Heemun Kwok, MD, MS (Methodologist) 31 Bruce M. Lo, MD, MBA, RDMS 32 Sharon E. Mace, MD 33 Amal Mattu, MD 34 Susan B. Promes, MD, MBA 35 Kaushal H. Shah, MD 36 Richard D. Shih. MD 37 Scott M. Silvers, MD 38 Andrea Slivinski, RN, DNP (ENA Representative 2021-2024) 39 Michael D. Smith, MD, MBA 40 Molly E. W. Thiessen, MD 41 John T. Thompson, MD (EMRA Representative 2023-2024) 42 Christian A. Tomaszewski, MD, MS, MBA 43 Stacy A. Trent, MD, MPH (Methodologist) 44 Jonathan H. Valente, MD 45 Lauren M. Westafer, DO, MPH, MS 46 Stephen P. Wall, MD, MSc, MAEd (Methodologist) 47 Yanling Yu, PhD (Washington Advocates for Patient Safety) 48 Michelle P. Lin, MD, MPH, MS (Liaison with the ACEP Quality and Patient Safety Committee and E-QUAL 49 Steering Committee) John T. Finnell, MD (Board Liaison 2020-2024) 50 Travis Schulz, MLS, AHIP, Staff Liaison, Clinical Policies Committee and Writing Committee on Thrombolytics 51

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52

ABSTRACT

This clinical policy from the American College of Emergency Physicians is the revision of a clinical policy approved in 2015 addressing a critical question regarding the use of thrombolytics for the management of acute ischemic stroke. A writing committee conducted a systematic review of the literature to derive evidence-based recommendations to answer the following clinical question: In adult stroke patients who are a candidate for mechanical thrombectomy, is the use of IV thrombolysis prior to mechanical thrombectomy (Bridge therapy) beneficial and safe versus mechanical thrombectomy alone? Evidence was graded, and recommendations were made based on the strength of the available data.

INTRODUCTION

Approximately 30% of all acute ischemic strokes have a large vessel occlusion (LVO), which contributes to 64% of all moderate to severe disability from stroke at 3 months and over 95% of stroke deaths at 6 months. 1.2 Over the past decade, acute treatment for LVO has expanded beyond thrombolytics with evidence supporting the use of endovascular therapy (EVT) such as mechanical thrombectomy. 3-5

For patients who are eligible for both interventions, this has led to recent debate on the use of intravenous thrombolysis (IVT) prior to EVT in patients with an LVO. On one hand, the use of IVT may contribute to early reperfusion from an LVO and resolve residual distal thrombi after EVT.^{6,7} However, IVT alone has low recanalization rates in patients with an LVO, especially with proximal lesions, and may fragment and cause distal embolization making EVT less effective.^{8,9} IVT may also increase the risk of symptomatic intracranial hemorrhage (sICH) and delay EVT, although the outcomes of such delays in patients receiving both interventions is unclear.^{10,11}

Another challenge in determining the optimal treatment paradigm is the availability of EVT. Although approximately 90% of patients in the United States have access to a stroke center within 60 minutes, most lack timely access to an EVT-capable center, with only around 20% residing within a 15-minute and 50% within a 60-minute radius to a stroke center equipped for EVT. 12-14 This may lead to varying treatment strategies for patients with an LVO: individuals who initially present to a facility without EVT capabilities and require transfer, and those who directly present to an EVT-capable facility.

Studies that compared EVT alone (direct endovascular therapy or direct mechanical thrombectomy) with IVT + EVT (bridging therapy) used the Modified Rankin Scale (mRS) to assess functional outcomes. The mRS ranges from 0 (no neurological symptoms) to 6 (death). Good functional outcome or functional independence is often defined as mRS 0 to 2, which represents patients with slight disability but who can look after their own affairs without assistance. Excellent functional outcome is usually defined as mRS of 0 to 1, which represents no significant disability and the ability to carry out all duties and activities. Although the mRS is the most common tool used for evaluating disability in stroke research, there are known limitations with inter-rater reliability. ¹⁶

Recently, an international survey showed that 63% of stroke physicians consisting of neurologists, interventionalists, and neurosurgeons would still give IVT prior to EVT.¹⁷ However published consensus from experts have been conflicting whether to support IVT prior to EVT due to differing interpretations of the data.^{18,19} This systematic review will evaluate outcomes for patients who present with an acute stroke from an LVO and received EVT with or without IVT.

METHODOLOGY

This ACEP clinical policy was developed by emergency physicians with input from medical librarians and a patient safety advocate and is based on a systematic review and critical descriptive analysis of the medical literature and is reported in accordance with Preferred Reporting Items for Systematic reviews and Meta-Analyses (PRISMA) guidelines.²⁰

Search and Study Selection

This clinical policy is based on a systematic review with critical analysis of the medical literature meeting the inclusion criteria. Searches of PubMed, SCOPUS, Embase, Web of Science, and the Cochrane Database of Systematic Reviews were performed by a second librarian. Search terms and strategies were peer reviewed by a second librarian. All searches were limited to human studies published in English. Specific key words/phrases, years used in the searches, dates of searches, and study selection are identified under each critical question. In addition, relevant articles from the bibliographies of included studies and more recent articles identified by committee members and reviewers were included.

Using Covidence (Covidence), 2 subcommittee members independently reviewed the identified abstracts to assess for possible inclusion. Of those identified for potential inclusion, each full-length text was reviewed for eligibility. Those identified as eligible were subsequently abstracted and forwarded to the committee's methodology group (emergency physicians with specific research methodological expertise) for methodological grading using a Class of Evidence framework (Appendix E1).

Assessment of Risk of Bias and Determination of Classes of Evidence

Each study identified as eligible by the subcommittee was independently graded by 2 methodologists. Design 1 represents the strongest possible study design to answer the critical question, which relates to whether the focus was therapeutic, diagnostic, or prognostic, or a meta-analysis. Subsequent design types (ie, Design 2 and Design 3) represent respectively weaker study designs. Articles are then graded on dimensions related to the study's methodological features and execution, including but not limited to randomization processes, masking, allocation concealment, methods of data collection, outcome measures and their assessment, selection and misclassification biases, sample size, generalizability, data management, analyses, congruence of results and conclusions, and potential for conflicts of interest.

Using a predetermined process that combines the study's design, methodological quality, and applicability to the critical question, 2 methodologists independently assigned a preliminary Class of Evidence grade for each article. Articles with concordant grades from both methodologists received that grade as their final grade. Any discordance in the preliminary grades was adjudicated through discussion which involved at least 1 additional methodologist, resulting in a final Class of Evidence assignment (i.e., Class I, Class II, Class III, or Class X) (Appendix E2). Studies identified with significant methodologic limitations and/or ultimately determined to not be applicable to the critical question received a Class of Evidence grade "X" and were not used in formulating recommendations for this policy. However, content in these articles may have been used to formulate the background and to inform expert consensus in the absence of evidence. Classes of Evidence grading may be found in the Evidentiary Table included at the end of this policy.

<u>Translation of Classes of Evidence to Recommendation Levels</u>

Based on the strength of evidence for each critical question, the subcommittee drafted the recommendations and supporting text synthesizing the evidence using the following guidelines:

Level A recommendations. Generally accepted principles for patient care that reflect a high degree of scientific certainty (eg, based on evidence from one or more Class of Evidence I, or multiple Class of Evidence II studies that demonstrate consistent effects or estimates).

Level B recommendations. Recommendations for patient care that may identify a particular strategy or range of strategies that reflect moderate scientific certainty (e.g., based on evidence from one or more Class of Evidence II studies, or multiple Class of Evidence III studies that demonstrate consistent effects or estimates).

Level C recommendations. Recommendations for patient care that are based on evidence from Class of Evidence III studies or, in the absence of adequate published literature, based on expert consensus. In instances where consensus recommendations are made, "consensus" is placed in parentheses at the end of the recommendation.

There are certain circumstances in which the recommendations stemming from a body of evidence should not be rated as highly as the individual studies on which they are based. Factors such as consistency of results, uncertainty of effect magnitude, and publication bias, among others, might lead to a downgrading of recommendations. When possible, clinically-oriented statistics (e.g., likelihood ratios [LRs], number needed to treat) are presented to help the reader better understand how the results may be applied to the individual patient. This can assist the clinician in applying the recommendations to most patients but allow adjustment when applying to patients with extremes of risk (Appendix E3).

Evaluation and Review of Recommendations

Once drafted, the policy was distributed for internal review (by members of the entire committee) followed by external expert review and an open comment period for all ACEP membership. Comments were received during a 60-day open comment period with notices of the comment period sent electronically to ACEP members, published in *EM Today*, posted on the ACEP Web site, and sent to other pertinent physician organizations. The responses were used to further refine and enhance this clinical policy, although responses do not imply endorsement. Clinical

policies are scheduled for revision every 3 years; however, interim reviews are conducted when technology, methodology, or the practice environment changes significantly.

Application of the Policy

This policy is not intended to be a complete manual on the use of thrombolytics for the management of acute ischemic stroke but rather a focused examination of critical questions that have particular relevance to the current practice of emergency medicine. Potential benefits and harms of implementing recommendations are briefly summarized within each critical question.

It is the goal of the Clinical Policies Committee to provide evidence-based recommendations when the scientific literature provides sufficient quality information to inform recommendations for a critical question. In accordance with ACEP Resolution 56(21), ACEP clinical policies do not use race-based calculators in the formulation of the recommendations. When the medical literature does not contain adequate empirical data to inform a critical question, the members of the Clinical Policies Committee believe that it is equally important to alert emergency physicians to this fact.

This clinical policy is not intended to represent a legal standard of care for emergency physicians. Recommendations offered in this policy are not intended to represent the only diagnostic or management options available to the emergency physician. ACEP recognizes the importance of the individual physician's judgment and patient preferences. This guideline provides clinical strategies for which medical literature exists to inform the critical questions addressed in this policy. ACEP funded this clinical policy.

Scope of Application. This guideline is intended for physicians working in EDs.

Inclusion Criteria. This guideline is intended for adult patients aged 18 years and older presenting to the ED with acute ischemic stroke.

Exclusion Criteria. This guideline is not intended to be used for pediatric or pregnant patients.

CRITICAL QUESTION

 In adult stroke patients who are a candidate for mechanical thrombectomy, is the use of IV thrombolysis prior to mechanical thrombectomy (Bridge therapy) beneficial and safe versus mechanical thrombectomy alone?

193 194	Patient Management Recommendations
195	Level A recommendations.
196	Level B recommendations. In stroke patients who are candidates for both mechanical thrombectomy and
197	IV thrombolysis*, IV thrombolysis should be offered and may be given prior to mechanical thrombectomy.
198	*IV thrombolysis given within 4.5 hours from symptom onset
199	Level C recommendations. When feasible, shared decision-making between the patient (and/or their
200	surrogate) and a member of the health care team should include a discussion of potential benefits and harms prior
201	to the decision whether to administer IV thrombolytics (Consensus recommendation).
202	
203	Potential Benefit of Implementing the Recommendations:
204	 Improved functional outcomes
205	Decreased mortality
206	
207	Potential Harm of Implementing the Recommendations:
208	Delays in endovascular therapy
209	 Increased cost with the use of thrombolytics
210	
211	
212	Key words/phrases for literature searches: Acute Ischemic Stroke, Acute Stroke, Alteplase,
213	Anticoagulation Bridge, Brain Ischemia, Bridge Therapy, Bridging Anticoagulation, Catheter-directed
214	Thrombectomy, Cerebrovascular Accident, Directed, Thrombectomy, Elaxim, Emergency Department,
215	Emergency Health Service, Emergency Medical Services, Emergency Medicine, Emergency Treatment,
216217	Emergency Ward, EMS, Endovascular Therapy, Endovascular Thrombectomy, EVT, Fibrinolytic, Fibrinolytic Agents, Guided Thrombectomy, Intravenous, Intravenous Drug Administration, Ischemic Stroke, IV, Mechanical
218	Thrombectomy, Metalyse, Percutaneous Thrombectomy, rTPA, Stroke, Tenecteplase, Thrombectomy,
219	Thrombolytic Therapy, Thrombolytic Treatment, Thrombolytic, Tissue Plasminogen Activator, TNKase, tPA
220	and variations and combinations of key words/phrases. Searches included January 2015 to search the date of April
221	10, 2023 (Appendix E4).
222	10, 2023 (Appendix L4).
223	Study Selection: Five hundred fifty-seven articles were identified in the searches. Three hundred thirty-
224	four articles were selected from the search results as candidates for further review. After grading for
225	methodological rigor, 3 Class I studies, 7 Class II studies, and 8 Class III studies were included for this critical
226	question (Appendix E5). Appendix E6 lists the 69 articles graded for methodological rigor but ultimately found to
227	be fatally flawed.
228	·
229	Randomized Controlled Trials
230	Six randomized controlled trials (RCTs) were included: 1 Class I study, 4 Class II studies, 21-25 and 1 Class
231	III study. ²⁶ All included RCTs were open-labeled with masked assessment of outcomes and included only adult
232	patients who presented within 4.5 hours of symptom onset without contraindications for thrombolytics. Alterlase

at 0.9 mg/kg was used in all studies except in studies where it was noted that either a different alteplase dose was given or tenecteplase was used.

All the RCTs were designed primarily to evaluate if EVT alone was non-inferior to IVT + EVT except for 1 trial (LeCouffe 2021) that evaluated superiority of EVT alone followed by non-inferiority of EVT alone.²² As opposed to superiority studies which are designed to demonstrate better effectiveness of 1 intervention over another, non-inferiority studies are powered to evaluate whether 1 intervention is potentially "less good" than another intervention within a predefined range.²⁷ Non-inferiority trials are appropriate if 1 intervention has added costs, risks, or limited availability that might render superiority less important.²⁸ Since intention-to-treat analysis is more likely to create Type 1 error by falsely concluding non-inferiority compared with per-protocol analysis, dual reporting of both analyses are preferable for non-inferiority trials.^{29,30} To achieve non-inferiority, the lower limit of the confidence interval (CI) should exceed the prespecified non-inferiority margin. Each of the non-inferiority RCT trials in this clinical policy used different primary end points as well as various non-inferiority margins. Both per-protocol and intention-to-treat analysis were performed and remained consistent within each study and is summarized in Table 1.

In a Class I study, the DIRECT-MT trial enrolled 654 patients from 41 academic tertiary care centers in China with an internal carotid artery (ICA) or first segment middle cerebral artery (M1)/second segment middle cerebral artery (M2) LVO.²¹ The primary outcome was a median 90-day mRS. Both EVT alone and IVT + EVT had similar 90-day mRS (3 versus 3). The adjusted odds ratio (OR) for the mRS was 1.08 (95% CI 0.82 to 1.43). These results demonstrate non-inferiority as the lower limit margin was set at 0.80. There was no statistical difference in sICH or death at 90 days observed between the 2 groups.

The DEVT trial was a Class II study that enrolled 234 patients with an ICA or M1 LVO from 33 stroke centers in China. He primary outcome was the proportion of patients achieving mRS 0 to 2 at 90 days. Results from the per-protocol analysis showed an mRS 0 to 2 in 53.2% of the EVT alone group versus 46% of the IVT + EVT group. The absolute difference of 7.1% (97.5% CI –5.9 to ω) allowed them to conclude non-inferiority based upon their pre-specified margin of 10%. The DEVT trial was stopped early after enrolling only 235 out of the planned 970 patients because of a statistical finding of likely futility. Both groups had similar rates of sICH and death at 90 days with no statistical differences observed.

In a Class II study, the SKIP trial enrolled 204 patients from 23 stroke centers in Japan with an ICA or M1 LVO.²⁵ Whereas 0.9 mg/kg of alteplase was used in other trials, this trial used 0.6 mg/kg of alteplase. The primary outcome was mRS 0 to 2. Results from the per-protocol analysis showed a favorable neurologic outcome in 60.8% of the EVT alone group versus 58.8% of the IVT + EVT group and an OR of 1.06 (1-sided 97.5% CI 0.60 to ω), which did not meet the prespecified lower margin of 0.74. The investigators were unable to conclude non-inferiority. Mortality at 90 days and sICH were not observed to be statistically different between the 2 groups.

The MR CLEAN-NO IV trial was a Class II study that included 539 patients from 20 hospitals in the Netherlands, Belgium, and France.²² Patients had an acute ischemic stroke due to a proximal occlusion of the anterior circulation. The primary outcome was median mRS at 90 days, first evaluating for superiority of EVT alone over IVT + EVT. If superiority was not established, then an evaluation of non-inferiority of EVT alone compared with IVT + EVT was performed. The non-inferiority margin was set at 0.8 for the adjusted common OR. Median mRS favored IVT + EVT over EVT alone (2 versus 3). Results from the adjusted common OR was 0.84 (95% CI 0.62 to 1.15), which demonstrated neither superiority nor non-inferiority for EVT alone. No statistical difference was observed between the 2 groups for sICH or death within 90 days.

The SWIFT DIRECT was a Class II trial that enrolled 408 patients with anterior strokes from 48 EVT-capable centers in Europe and Canada.²³ The primary outcome was mRS 0 to 2 at 90 days. Results from the perprotocol analysis showed favorable neurologic outcomes in 57% of the EVT alone group versus 64% of the IVT + EVT group. Absolute risk difference was -4.6% (95% CI -14.8 to 5.8%), with the lower limit of 1-sided 95% CI of -13.2%. The lower limit exceeded the prespecified 12% and non-inferiority of EVT alone could not be concluded in the overall study population or in any of the pre-specified subgroups. There was no statistical difference in sICH or mortality by 90 days between both groups.

In a Class III study, the DIRECT-SAFE trial enrolled 295 patients from 25 acute-care hospitals in Australia, New Zealand, China, and Vietnam.²⁶ Patients needed to have an LVO in either the ICA, M1 or M2 segments of the middle cerebral artery (MCA), or basilar artery and were randomized with or without alteplase in Asian countries (83%) and tenecteplase in non-Asian countries (17%). The primary outcome was mRS 0 to 2 at 90 days. Results from the per-protocol analysis showed a favorable neurologic outcome in 54% of the EVT alone

group vs 62% of the IVT + EVT group. Risk difference was -0.062 (95% CI -0.173 to 0.049). The lower end of the 95% CI exceeded -0.1 prespecified threshold and therefore non-inferiority of EVT alone was not demonstrated. Safety outcomes were not statistically different with 1% sICH in both groups and a similar number of deaths at 90 days.

Of the 6 RCTs, 4 did not show non-inferiority of EVT alone compared with IVT + EVT, thus supporting the use of IVT in this patient population.^{22,23,25,26} In all RCT studies, sICH and death was not statistically significant between the 2 groups, although the studies were not all powered for safety.²¹⁻²⁶

Table 1. A synthesis of the ACEP Clinical Policy Level of Evidence, direction of support for bridging therapy (BT), original investigator's non-inferiority (NI) margin, and Per Protocol and Intention to Treat analysis.

RCT	Level of Evidence	Study Size	NI Margin	Per Protocol	Intention to Treat	Support BT?
DIRECT MT ²¹	I	654	0.8	1.08 (95% CI 0.82 to 1.43) ¹	1.07 (95% CI 0.81 to 1.40) ^A	No
DEVT ²⁴	II	234	-10%	7.1% (97.5% CI -5.9 to ∞) ²	7.7% (97.5% CI -5.1% to ∞) ^B	No
SKIP ²⁵	II	204	0.74	1.06 (97.5% CI 0.60 to ω) ³	1.09 (97.5% CI 0.63 to ω) ^c	Yes
MR CLEAN NO IV ²²	II	539	0.8	0.84 (95% CI 0.61 to 1.16) ¹	0.84 (95% CI 0.62 to 1.15) ^A	Yes
SWIFT DIRECT ²³	II	408	-12%	-4.6% (95% CI -14.8 to 5.8%) ⁴	-7.3% (95% CI -16.6 to 2.1) ^D	Yes
DIRECT SAFE ²⁶	III	295	-0.1	-0.062 (95% CI -0.173 to 0.049) ⁴	-0.051 (95% CI -0.160 to 0.059) ^E	Yes

A Adjusted common Odds Ratio

Systematic Reviews/Meta-Analysis

Six systematic reviews/meta-analysis (SRMA) were included in this guideline. Three SRMAs included RCTs only, which were included in this review. 10,31,32 Two other SRMAs included both RCTs and observational studies, including studies that were eliminated during the critical appraisal (grading) process. 33,34 Lastly, 1 SRMA

²⁹⁷ B Unadjusted difference

^{298 &}lt;sup>C</sup> Odds Ratio

²⁹⁹ DAdjusted Risk Difference

³⁰⁰ E Unadjusted Risk Difference

compared patients who were transferred from a primary stroke center (PSC) with IVT compared with patients who arrived at an EVT-capable center who did not receive IVT, but did not include any RCTs.³⁵

In a Class I study, Kaesmacher et al included 6 randomized clinical trials (DEVT, SKIP, DIRECT-MT, DIRECT-SAFE, SWIFT-DIRECT, and MR CLEAN NO IV)²¹⁻²⁶ totaling 2,023 patients comparing EVT alone versus IVT + EVT for patients with anterior circulation LVO only.³¹ The primary outcome was time from symptom onset to expected administration of IVT plus thrombectomy versus thrombectomy alone with a minimal clinically important difference for the rate of mRS 0 to 2 of 1.3% at 90 days. There was a statistically significant interaction between time from symptoms onset to expected administration of IVT and the association of allocated treatment with functional outcomes (adjusted OR per 1-hour delay, 0.84; 95% CI 0.72 to 0.97). The benefit of IVT + EVT decreased with longer times from symptom onset to IVT administration and the benefit was not statistically significant after 2 hours 20 minutes.

In a Class II study, Lin et al reviewed 4 RCTs (DEVT, SKIP, DIRECT-MT, and MR CLEAN NO IV)^{21,22,24,25} for a total of 1,633 patients.³² Based on the literature, they assessed 5 different non-inferiority margins for functional independence (mRS 0 to 2) at 90 days. There was no observed statistical heterogeneity among trials (I^2 =0%). Although the risk difference was 1% (95% CI –4% to 5%) favoring EVT alone, the lower margin of the 95% CI suggests EVT alone is non-inferior to IVT + EVT except when using the most stringent of margins at –1.3%. The outcome measure of mRS 0 to 1 showed a similar risk difference of 1% (95% CI –3% to 5%), showing non-inferiority except when using the margin of –1.3%. SICH and mortality were not shown to be different between both groups.

In another Class II study, Wang et al reviewed 6 RCTs (DEVT, SKIP, DIRECT-MT, DIRECT SAFE, SWIFT DIRECT, and MR CLEAN NO IV)²¹⁻²⁶ for a total of 2,334 patients. ¹⁰ This international workgroup consisted of various stakeholders including stroke experts, pharmacists, academics, and caregivers of stroke patients. The workgroup established minimally important differences through survey of their guideline panel and discussion for the following outcomes: 1% for recovery with minimal disability (mRS 0 to 2), 0.8% for mortality, and 1% for sICH. Pooled estimate of effect showed lack of observed statistical heterogeneity (I^2 =0%). They concluded with low certainty of evidence that EVT alone had a smaller decrease in patients with minimal disability (risk ratio (RR) 0.97, 95% CI 0.89 to 1.05; risk difference –1.5%; 95% CI –5.4% to 2.5%) and a small

increase in mortality (RR 1.07, 95% CI 0.88 to 1.29; risk difference 1.2%, 95% CI –2.0% to 4.9%), but moderate certainty of evidence that EVT alone had a small decrease in sICH (RR 0.75, 95% CI 0.52 to 1.07; risk difference –1.0%, 95% CI –1.8% to 0.27%).

In a Class I study, Zheng et al reviewed a total of 55 studies that included 9 RCTs^{21,22,24,25,36-40} and 46 observational/retrospective studies, for a total of approximately 20,000 patients.³³ A comprehensive meta-analysis was performed for utilizing both RCTs and observational/retrospective studies to investigate various outcomes. Functional independence was defined as mRS of 0 to 2 and excellent outcomes was defined as mRS of 0 to 1. For RCTs, the IVT + EVT group reduced the risk of mortality versus EVT alone (OR 0.65, 95% CI 0.49 to 0.88, P=52%), but not functional independence (OR 1.17, 95% CI 0.99 to 1.38, P=0%). On the other hand, the observational studies showed that IVT + EVT had better outcomes for functional independence (OR 1.36, 95% CI 1.21 to 1.52, P=48%), excellent outcomes (OR 1.49, 95% CI 1.26 to 1.75, P=4%), and mortality (OR 0.73, 95% CI 0.56 to 0.94, P=67%). Neither the RCTs nor observational studies showed an increased risk in sICH.

In a Class II study, Ghaith et al reviewed 49 studies (4 RCTs^{21,22,24,25} and 44 observational studies) for a total of 36,123 patients.³⁴ In the analysis combining both RCTs and observational studies, they demonstrated that IVT + EVT had better mortality (RR 0.75, CI 95% 0.68 to 0.82, P=36%), successful recanalization (RR 1.06, 95% CI 1.03 to 1.09, P=50%), and 90-day functional independence (RR 1.21, 95% CI 1.13 to 1.29, P=52%), but no improvement in National Institutes of Health Stroke Scale (NIHSS). Subgroups were stratified accounting to study design showing similar benefits with IVT + EVT for observational studies, but not for RCTs. No difference was seen between the 2 groups related to sICH.

Lastly, in a Class III study, Katsonos et al included 6 observational studies totaling 1,723 patients. Patients who received IVT at a PSC before transferring for EVT ("drip and ship" or DNS, 53% of the group) were compared with those receiving EVT alone at a Comprehensive Stroke Center (CSC). In their analysis adjusted for potential confounders, "DNS patients" had higher odds of mRS 0 to 1 (adjusted OR 1.32, 95% CI 1.00 to 1.74, I^2 =0%) and lower probability for all-cause mortality at 3-months (adjusted OR 0.50, 95% CI: 0.27 to 0.93, I^2 =69%) compared to patients receiving EVT alone at a CSC. No differences were found between the 2 groups in probability of 3-month disability, mRS 0 to 2, or sICH.

The majority of SRMA favored IVT + EVT. Two of the SRMA showed either improved mortality or improved functional outcomes with IVT + EVT, however these results varied based on whether the analysis utilized RCTs and/or observational studies. 33,34 Of the 3 studies that looked at the RCTs alone, 1 SRMA showed non-inferiority of EVT alone compared with IVT + EVT in various cutoffs except for the most strict cutoff for functional outcomes while another SRMA suggested a possible small increase in mortality, a small decrease in recovery with minimal disability, but moderate certainty of decreased sICH with EVT alone. The other SRMA that utilized RCTs alone suggests that IVT + EVT is superior to EVT alone but is time dependent. Lastly in patients who are transferred, evidence suggests patients who received IVT + EVT have better functional outcomes and mortality compared with EVT alone. 35

Observational and Retrospective Evidence

Multiple non-randomized Class III studies have also explored the role of thrombolysis with thrombectomy. Abilleira et al analyzed Spanish stroke registry data from Catalonia to compare EVT alone with IVT + EVT.⁴¹ After adjusting for higher proportion of patients with heart failure, atrial fibrillation, oral anticoagulation, and previous stroke among patients receiving EVT alone, no differences in 90-day mortality, symptomatic bleeding at 24 to 36 hours, or mRS 0 to 2 were noted between the 2 treatment groups.

Balodis et al reported a single-center prospective observational analysis of IVT + EVT versus EVT alone for anterior cerebral artery LVO in a single Latvian university hospital.⁴² Although exclusions did not include a time-of-onset for symptoms, all thrombectomy occurred within 8 hours of symptom onset and all patients presenting within 4.5 hours received IVT unless contraindications were identified, or physician's preference was not to provide IVT. A 90-days mRS of 0 to 2 was observed in 44% of the IVT + EVT group versus 42% in the EVT-alone group. No significant differences were observed in 90-day mortality or sICH.

Broocks et al retrospectively analyzed a cohort of acute ischemic stroke patients treated at 1 of 2 high-volume tertiary stroke centers in Germany and the United States for ICA or MCA LVO.⁴³ The Alberta Stroke Program Early CT Score (ASPECTS) was determined on pre-treatment non-contrast head CT by 1 neuro-radiologist.⁴⁴ Most had ASPECTS >5 (86%). Overall, those receiving IVT + EVT had better NIHSS at 24 hours (11 versus 13) and mRS at 90 days (3 versus 4). More patients in the IVT + EVT cohort had an mRS 0 to 2 at 90

days (43% versus 32%). Among the 14% with ASPECTS <6, no difference was seen for mRS 0 to 2. ASPECTS was the only variable demonstrating a significant interaction with IVT.

Casetta et al reviewed the Italian Registry of Endovascular Stroke Treatments prospective observational data from 13 hospitals which included 1,148 patients with either an ICA or MI/M2 LVO who were eligible for IVT.⁴⁵ EVT was performed within 6 hours of symptom onset and decisions about IVT were left to the discretion of the treating neurology team. Although the median time from symptom onset to hospital arrival was similar between the 2 groups (95 minutes for IVT + EVT versus 96 minutes for EVT alone patients), the symptom onset to groin puncture was significantly prolonged in the IVT + EVT subset (230 minutes versus 210 minutes in EVT). Multivariate analysis for stroke patients surviving with mRS 0 to 3 demonstrated a significant benefit favoring IVT + EVT (adjusted OR 1.42; 95% CI 1.04 to 1.95) and a significantly lower risk of death or unfavorable outcome in that same group (adjusted OR 0.62; 95% CI 0.45 to 0.84). No differences were found regarding sICH.

Di Maria et al retrospectively evaluated acute ischemic stroke patients involving the proximal or distal MCA or ICA within 6-hours of symptoms.⁴⁶ A stroke neurologist decided whether or not to treat with IVT. IVT + EVT patients were matched with patients treated with EVT alone using a propensity score. An mRS 0 to 2 was more likely with IVT + EVT (OR 1.31; 95% CI 1.02 to 1.68). All-cause mortality and sICH did not differ between groups. Only ASPECTS ≥7 demonstrate the benefit of IVT + EVT compared with EVT alone (OR 1.48, 95% CI 1.10 to 2.0).

Zha et al reported a post-hoc analysis of a prospective study across 16 Chinese stroke centers.⁴⁷ The primary outcome of mRS 0 to 2 at 90 days. In a multivariable analysis, IVT + EVT more frequently demonstrated a higher mRS 0 to 1 at 90-days (adjusted OR 2.731; 95% CI 1.238 to 6.023), but not the primary outcome of mRS 0 to 2. The 90-day mortality rate was significantly lower in the IVT + EVT cohort (13.9% versus 27.7%).

Of the 6 studies, 4 showed an improvement in functional outcomes with IVT + EVT compared with EVT alone. 43,45-47 In several studies, the use of ASPECTS further defined which patients benefited from IVT prior to EVT. 43,46 In 2 studies, mortality was decreased with IVT + EVT, but no difference in the others. 45,47 Lastly, there was no increase in sICH with IVT + EVT compared with EVT alone in any of the studies.

Summary

The majority of published research favored the use of IVT + EVT over EVT alone. This includes RCTs where the majority of trials failed to show non-inferiority with EVT alone, despite using wide non-inferiority thresholds. However, there are a number of limitations to these trials including different outcome measures and different non-inferiority thresholds. Among systematic reviews, inclusion of observational studies increased observed statistical heterogeneity.

From a safety standpoint, although some studies showed a decrease in mortality with IVT + EVT, most studies showed no difference. Lastly, although there have been concerns about the increased risk of sICH with the addition of IVT before EVT, no study included in our review showed an increased risk of sICH. However, safety data from these studies may have also been under-reported. It is important that with any intervention, shared decision making is made when feasible with the patient and/or family.

Future Research

Existing research predominantly employed alteplase as the primary thrombolytic agent. Subsequent investigations should explore alternative thrombolytics, such as tenecteplase. Future studies should also look at timing of thrombolytics prior to EVT with patient outcomes. In addition, the role of ASPECTS score and other tools in identifying individuals unlikely to benefit from the addition of IVT prior to EVT should be explored prospectively. Furthermore, future studies ought to consider larger sample sizes, utilizing more stringent non-inferiority margins or ideally conducting superiority studies, as well as evaluating the cost-effectiveness of different treatment strategies. Figure 1.

Since the majority of the literature has focused on anterior strokes, future studies should also evaluate the role of IVT before EVT in posterior circulation strokes. Finally, more studies evaluating the role of thrombolytics in patients with an LVO who are candidates for EVT but need to be transferred are needed. This includes patients who are considered for prehospital diversion to EVT-capable centers and the use of mobile stroke units to triage potential patients for EVT.

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618 Appendix E1. Literature classification schema.*

Design/ Class	Therapy [†]	Diagnosis [‡]	Prognosis [§]
1	Randomized, controlled trial or meta-analysis of randomized trials	Prospective cohort using a criterion standard or meta-analysis of prospective studies	Population prospective cohort or meta-analysis of prospective studies
2	Nonrandomized trial	Retrospective observational	Retrospective cohort Case control
3	Case series	Case series	Case series

^{*}Some designs (eg, surveys) will not fit this schema and should be assessed individually.

Appendix E2. Approach to downgrading strength of evidence.

	I	Design/Class	gn/Class	
Downgrading	1	2	3	
None	I	II	III	
1 level	II	III	X	
2 levels	III	X	X	
Fatally flawed	X	X	X	

Appendix E3. Likelihood ratios and number needed to treat.*

LR (+)	LR (-)	
1.0	1.0	Does not change pretest probability
1-5	0.5-1	Minimally changes pretest probability
10	0.1	May be diagnostic if the result is concordant with pretest probability
20	0.05	Usually diagnostic
100	0.01	Almost always diagnostic even in the setting of low or high pretest probability

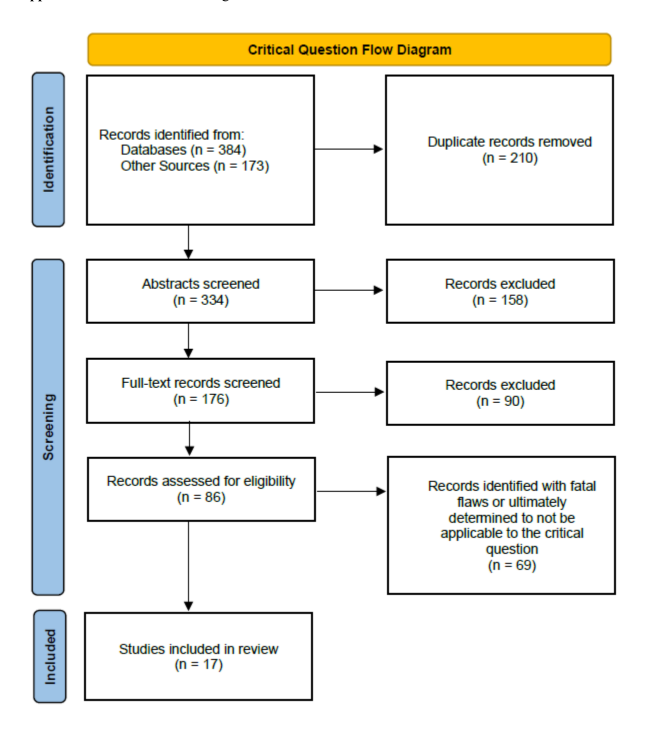
LR, likelihood ratio.

[†]Objective is to measure therapeutic efficacy comparing interventions.

[‡]Objective is to determine the sensitivity and specificity of diagnostic tests.

[§]Objective is to predict outcome, including mortality and morbidity.

^{*}Number needed to treat (NNT): number of patients who need to be treated to achieve 1 additional good outcome; NNT=1/absolute risk reduction×100, where absolute risk reduction is the risk difference between 2 event rates (ie, experimental and control groups).



Appendix E5. Literature Searches

Search Date	Database	Search Strings	Filters
4/10/2023	PubMed	((Mechanical Thrombectomy[tiab]) OR (Bridge Therapy[tiab]) OR (Percutaneous Thrombectomy[tiab]) OR (Endovascular Therapy[tiab]) OR (EVT[tiab]) OR (Endovascular Thrombectomy[tiab]) OR (Guided Thrombectomy[tiab]) OR (Catheter-directed Thrombectomy[tiab]) OR ("Thrombectomy"[mh]) OR ("Bridge Therapy"[Mesh])) AND ((Tissue Plasminogen Activator[tiab]) OR (Alteplase[tiab]) OR (tPA[tiab]) OR (rTPA) OR (Tenecteplase[tiab]) OR (Thrombolytic*[tiab]) OR (Fibrinolytic*[tiab]) OR ("Tissue Plasminogen Activator"[mh]) OR ("Tenecteplase"[mh]) OR ("Fibrinolytic Agents" [Pharmacological Action]) OR ("Thrombolytic Therapy"[mh])) AND ((Intravenous[tiab]) OR (IV[tiab]) OR ("Administration, Intravenous"[mh])) AND((Acute Stroke[tiab]) OR (Acute Ischemic Stroke[tiab]) OR (Brain Ischemia[tiab]) OR ("Stroke"[mh]) OR ("Ischemic Stroke"[mh]) OR ("Brain Ischemia"[mh])) AND ((Emergency Medicine[tiab]) OR (Emergency Treatment[tiab]) OR (Emergency Medicine"[mh]) OR ("Emergency Medicin	2015- Current
4/10/2023	Scopus	TITLE-ABS-KEY("Mechanical Thrombectomy" OR "Bridge Therapy" OR "Anticoagulation Bridge" OR "Percutaneous Thrombectomy" OR "Endovascular Therapy" OR "EVT" OR "Endovascular Thrombectomy" OR "Guided Thrombectomy" OR "Directed Thrombectomy" OR "Catheter-directed Thrombectomy") AND TITLE-ABS-KEY("Tissue Plasminogen Activator" OR "Alteplase" OR "tPA" OR "rTPA" OR "Tenecteplase" OR "Metalyse" OR "TNKase" OR "Elaxim" OR "Thrombolytic*" OR "Fibrinolytic*") AND TITLE-ABS-KEY("Intravenous" OR "IV") AND TITLE-ABS-KEY("Stroke" OR "Acute Stroke" OR "Acute Ischemic Stroke" OR "Brain Ischemia") AND TITLE-ABS-KEY("Emergency Medicine" OR "Emergency Treatment" OR "Emergency Department" OR "Emergency Medical Service*")	2015- Current
4/10/2023	Embase	('Mechanical Thrombectomy':de,ti,ab,kw OR 'Bridge Therapy':ti,ab,kw OR 'Bridging Anticoagulation':de OR 'Percutaneous Thrombectomy':de,ti,ab,kw OR 'Endovascular Therapy':ti,ab,kw OR 'EVT':ti,ab,kw OR 'Endovascular Thrombectomy':ti,ab,kw OR 'Guided Thrombectomy':ti,ab,kw OR 'Directed Thrombectomy':ti,ab,kw OR 'Catheter-directed Thrombectomy':ti,ab,kw OR ('Tissue Plasminogen Activator':de,ti,ab,kw OR 'Alteplase':de,ti,ab,kw OR 'tPA':ti,ab,kw OR 'rTPA':ti,ab,kw OR "Tenecteplase':de,ti,ab,kw OR 'Metalyse':ti,ab,kw OR 'TNKase':ti,ab,kw OR 'Elaxim':ti,ab,kw OR 'Thrombolytic*':ti,ab,kw OR 'Thrombolytic Therapy':de,ti,ab,kw OR 'Thrombolytic treatment':de,ti,ab,kw OR 'Fibrinolytic':de,ti,ab,kw) AND ('Intravenous':ti,ab,kw OR 'Intravenous Drug Administration':de,ti,ab,kw OR 'IV':ti,ab,kw) AND ('Stroke':ti,ab,kw OR 'Cerebrovascular Accident':de OR 'Acute Stroke':ti,ab,kw OR 'Acute Ischemic Stroke':de,ti,ab,kw OR 'Brain Ischemia':de,ti,ab,kw) AND ('Emergency Medicine':de,ti,ab,kw OR 'Emergency Treatment':de,ti,ab,kw OR 'Emergency Department':ti,ab,kw OR 'Emergency Ward':de,ti,ab,kw OR 'Emergency Medical Service*':ti,ab,kw OR 'Emergency Health Service':de,ti,ab,kw)	2015- Current

Appendix E5. Literature Searches (continued)

Search Date	Database	Search Strings	Filters
8/24/2022	Web of Science	TS=("Mechanical Thrombectomy" OR "Bridge Therapy" OR "Anticoagulation Bridge" OR "Percutaneous Thrombectomy" OR "Endovascular Thrombectomy" OR "Guided Thrombectomy" OR "Directed Thrombectomy" OR "Catheter-directed Thrombectomy") AND TS=("Tissue Plasminogen Activator" OR "Alteplase" OR "tPA" OR "rTPA" OR "Tenecteplase" OR "Metalyse" OR "TNKase" OR "Elaxim" OR "Thrombolytic*" OR "Fibrinolytic*") AND TS=("Intravenous" OR "IV") AND TS=("Stroke" OR "Acute Stroke" OR "Acute Ischemic Stroke" OR "Brain Ischemia") AND TS=("Emergency Medicine" OR "Emergency Treatment" OR "Emergency Department" OR "Emergency Medical Services")	2011- Current
8/24/2022	Cochrane Library	("Mechanical Thrombectomy":ti,ab,kw OR "Bridge Therapy":ti,ab,kw OR "Bridging Anticoagulation":ti,ab,kw OR "Percutaneous Thrombectomy":ti,ab,kw OR "Endovascular Therapy":ti,ab,kw OR "EVT":ti,ab,kw OR "Endovascular Thrombectomy":ti,ab,kw OR "Guided Thrombectomy":ti,ab,kw OR "Directed Thrombectomy":ti,ab,kw OR "Catheter-directed Thrombectomy":ti,ab,kw) AND ("Tissue Plasminogen Activator":ti,ab,kw OR "Alteplase":ti,ab,kw OR "tPA":ti,ab,kw OR "rTPA":ti,ab,kw OR "Tenecteplase":ti,ab,kw OR "Metalyse":ti,ab,kw OR "TNKase":ti,ab,kw OR "Elaxim":ti,ab,kw OR "Thrombolytic*":ti,ab,kw OR "Thrombolytic Therapy":ti,ab,kw OR "Thrombolytic treatment":ti,ab,kw OR "Fibrinolytic":ti,ab,kw) AND ("Intravenous":ti,ab,kw OR "Intravenous Drug Administration":ti,ab,kw OR "IV":ti,ab,kw) AND ("Stroke":ti,ab,kw OR "Acute Ischemic Stroke":ti,ab,kw OR "Brain Ischemia":ti,ab,kw) AND ("Emergency Medicine":ti,ab,kw OR "Emergency Treatment":ti,ab,kw OR "Emergency Department":ti,ab,kw OR "Emergency Ward":ti,ab,kw OR "Emergency Medical Service*":ti,ab,kw OR "Emergency Health Service":ti,ab,kw)	2011- Current

Evidentiary Table.

	Graded Randomized Controlled Trials							
Author & Year Published	Class of Evidence	Setting & Study Design	Methods & Outcome Measures	Results	Limitations & Comments			
Yang et al (2020) ²¹	I	Multi-center (Chinese tertiary care centers); prospective randomized open-label, non- inferiority trial w/blinded outcome assessments	Adults ≥18 y, AIS of ICA or first segment middle cerebral artery (M1)/second segment middle cerebral artery (M2) or both by computed tomography angiography (CTA) that could be treated <4.5 h after symptom onset and NIHSS ≥2; 2 arms: EVT alone vs IVT+EVT in patients with AIS with LVO; primary outcome: 90 d mRS for noninferiority (logistic regression – ordinal) margin of 0.8 via telephone/inperson interview (intention-to-treat [ITT] analysis)	thrombectomy (2.4% vs 7%) and overall successful reperfusion (79.4% vs	Open label, not generalizable outside China, excluded those with missing outcomes, no adjustment for multiple comparisons, and this is a non-inferiority trial, whereas the Clinical Policies Committee question is for superiority			

	Graded Randomized Controlled Trials								
Author & Year	Class of	Setting & Study	Methods & Outcome	Results	Limitations & Comments				
Published	Evidence	Design	Measures						
LeCouffe et al	II	Multicenter,	Adult patients with	N=539; median mRS 3 for	Open label, unblinded to				
$(2021)^{22}$		randomized,	AIS randomly	thrombectomy alone group	treatment although blinded				
		open label,	assigned to either	vs mRS 2 for bridge	outcome assessment				
		clinical trial	endovascular treatment	thrombolysis plus					
		from 20	or IVT followed by	thrombectomy, OR 0.84					
		hospitals in	endovascular	(95% CI 0.62 to 1.15,					
		Europe	treatment; outcomes:	<i>P</i> =.28); mortality: 21% for					
			mRS at 90 d; sICH;	thrombectomy alone group					
			mortality	vs 16% for bridge					
				thrombolysis plus					
				thrombectomy, OR 1.39					
				(95% CI 0.84 to 2.30); sICH:					
				6% for thrombectomy alone					
				group vs 5% for bridge					
				thrombolysis plus					
				thrombectomy group, OR					
				1.30 (95% CI 0.60 to 2.81)					
Fischer et al	II	Multicenter,	Adults with acute	N=408: thrombectomy alone	Open label design could result in				
$(2022)^{23}$		academic centers	AIS+LVO, onset <4.5	(N=201) vs thrombectomy +	differential treatment bias; pre-				
		in Europe and	h; thrombectomy alone	IV alteplase (N=207); mRS 0	specified non-inferiority				
		Canada; non-	vs thrombectomy + IV	to 2: thrombectomy alone	margin=12%				
		inferiority,	alteplase; efficacy	57% vs thrombectomy + IV					
		randomized	outcome: mRS 0 to 2	alteplase 65%; adjusted risk					
		clinical trial	at 90 d; safety	difference –7.3, one-sided					
			outcome: ICH	(95% CI –16.6 to 2.1); ICH:					
				thrombectomy alone 2% vs					
				thrombectomy + IV alteplase					
				3%, risk difference –1.0%					
				(95% CI –4.8 to 2.7)					

	Graded Randomized Controlled Trials							
Author & Year	Class of	Setting & Study	Methods & Outcome	Results	Limitations & Comments			
Published	Evidence	Design	Measures					
Zi et al	II	Multicenter	Adults ≥18 y, AIS of	N=234, 116 EVT, 118 in	Infused whole dose of tPA			
$(2021)^{24}$		(China)	proximal circulation	IVT+EVT	despite achieving reperfusion			
		noninferiority	occlusion strokes that		earlier which might pose a			
		study, 4-block	could be treated <4.5 h	Primary Outcome: median	bleeding risk; within-site			
		randomized 1:1	after symptom onset;	mRS EVT alone was 2, 1 to	correlations analysis was post-			
			2 arms: EVT alone vs	4, and IVT+EVT was 3, 1 to	hoc and successful reperfusion			
			IVT+EVT in patients	4, and unadjusted difference	before EVT; study was powered			
			with AIS;	was 0, -1 to 0, aOR is 1.13	for noninferiority, rather than			
			outcomes: proportion	(95% CI 0.71 to 1.79) and no	whether IVT+EVT was			
			of patients with mRS 0	difference in secondary	"beneficial" (Clinical Policies			
			to 2 at 90 d (assessors	outcomes	Committee question)			
			were blinded					
			neurologists) vs	Safety Outcomes: 90 d				
			telephone call or video call with non-	mortality was 17.2% in EVT				
				only vs 17.8% in IVT+EVT				
			inferiority margin of - 10%; safety outcomes	-0.5, -10.3 to 9.2%) and sICH difference was 6.1% vs				
			were sICH within 48 h	6.8%, difference -0.8%,				
			and 90 d mortality	(95% CI –7.1 to 5.6);				
			and 90 d mortanty	asymptomatic hemorrhage				
				was 15.7% vs 25.6%, 10%				
				difference, 95% CI –20.3 to				
				0.3%, clot migration				
				occurred in 113 (17.7%) vs				
				28 of 117 (23.9%) in				
				IVT+EVT group with no				
				differences in serious adverse				
				events				

Evidentiary rable (C	Graded Randomized Controlled Trials							
Author & Year	Class of	Setting & Study	Methods & Outcome	Results	Limitations & Comments			
Published	Evidence	Design	Measures					
Suzuki et al	II	Multicenter,	Adult patients	N=204; mRS 0 to 2; 59% in	Open label, unblinded			
$(2021)^{25}$		randomized,	randomly assigned to	MT group vs 57% in bridge				
		open label,	MT alone or IVT+MT;	thrombolysis plus				
		noninferiority	outcomes: mRS 0 to 2	thrombectomy, <i>P</i> =.18;				
		clinical trial	at 90 d; mortality;	among 7 secondary efficacy				
		from 23 centers	sICH	endpoints and 4 safety				
		in Japan		endpoints, 10 were not				
				different, including mortality				
				(8% vs 9%, P=1.0) and sICH				
				(6% vs 8%, <i>P</i> =.78)				
Mitchell et al	III	Multicenter,	Adult patients with	N=295; 148 assigned to	Open label, unblinded to			
$(2022)^{26}$		randomized,	AIS eligible for	direct thrombectomy and 147	treatment although blinded			
		open label,	thrombolysis,	assigned to bridge therapy;	outcome assessment; trial			
		noninferiority	allocated 1:1 to either	mRS 0 to 2: 55% for	terminated early; some			
		clinical trial	direct thrombectomy	thrombectomy group vs 61%	imbalances in baseline			
		from 25 acute-	or IVT plus	for bridge thrombolysis plus	characteristics			
		care hospitals in	thrombectomy;	thrombectomy, OR 0.75				
		Australia, New	outcomes: mRS 0 to 2	(95% CI 0.45 to 1.24, <i>P</i> =.19)				
		Zealand, China,	at 90 d; mRS 0 to 1 at	for noninferiority, <i>P</i> =.26 for				
		and Vietnam	90 d; sICH; mortality	superiority; sICH: 1% vs 2%,				
				OR 1.70 (95% CI 0.22 to				
	(13.04, $P=0.61$); mortality:				
				15% vs 16%, OR 0.92 (95%				
				CI 0.46 to 1.84, <i>P</i> =.82)				

	Graded Systematic Reviews/Meta-Analysis								
Author & Year	Class of	Setting & Study	Methods & Outcome	Results	Limitations & Comments				
Published	Evidence	Design	Measures						
Kaesmacher et al	I	Individual	Systematic review and	6 randomized clinical trials;	Trials performed at				
$(2024)^{31}$		participant data	meta-analysis to	N=2,313, 1,160 IVT +	thrombectomy-capable stroke				
		meta-analysis	estimate the	thrombectomy, 1,153	centers; only patients with				
		from 6	association of	thrombectomy alone;	anterior circulation large-vessel				
		randomized	treatment with IVT	median time from symptom	occlusion were included; nearly				
		clinical trials	plus thrombectomy vs	onset to IVT administration	all patients in the IVT +				
		190 sites across	thrombectomy alone	was 2 h 28 min (inter quartile	thrombectomy group were				
		15 countries	and better outcomes	range [IQR] 1 h 46 min to 3	treated with alteplase; thus,				
			was modified by the	h 17 min);	results may not be generalizable				
			time from stroke	statistically significant	to those treated with tenecteplase				
			symptom onset to	interaction between time					
			treatment; primary	from symptom onset to					
			outcome: disability at	administration of IVT and					
			90 d using the mRS	functional outcome (aOR per					
				1-h delay 0.84 (95% CI 0.72					
				to 0.97), P =.02 for interaction); after 2 h 20 min,					
				the benefit associated with					
				IVT + thrombectomy was					
				not significant and the point					
				estimate crossed the null					
				association at 3 h 14 min					
Lin et al	II	Meta-analysis of	Trials comparing	N=4 trials with 1,633	Included studies with different				
$(2022)^{32}$	11	randomized	thrombectomy along	participants; 817 assigned to	noninferiority margins				
(= = =)		clinical trials	vs IVT plus	thrombectomy alone vs 816	ine initial entropy in the game				
			thrombectomy among	to bridge thrombolysis plus					
			adults with AIS-LVO;	thrombectomy; pooled					
			Primary outcome:	difference with risk					
			functional	difference of 1% for good					
			independence (mRS 0	functional outcomes (95% CI					
			to 2) at 90 d	-4% to 5%); pooled					
				difference in sICH was also					
				1%, 95% CI –1% to 3%					

Graded Systematic Reviews/Meta-Analysis					
Author & Year	Class of	Setting & Study	Methods & Outcome	Results	Limitations & Comments
Published	Evidence	Design	Measures		
Wang et al	II	Meta-analysis of	Trials of adult patients	N=6 trials with 2,334	Only used fixed effects
$(2022)^{10}$		randomized	with AIS comparing	participants; mRS 0 to 2:	modeling; limited
		clinical trials	thrombectomy alone	pooled RR 0.97 (95% CI	subgroup/sensitivity analyses
			vs IVT plus	0.89 to 1.05); sICH: pooled	
			thrombectomy;	RR 0.75 (95% CI 0.52 to	
			outcomes: mRS 0 to 2;	1.07); mortality: 1.07 (95%	
			sICH; mortality	CI 0.88 to 1.29)	
Zheng et al	I	Meta-analysis	RCTs of MT alone vs	mRS 0 to 2: 6 studies. aOR	Heterogeneity is less of a factor
$(2023)^{33}$			MT+IVT for patients	1.17 (95% CI 0.99 to 1.38);	in the adjusted analysis. Data
			with AIS secondary to	sICH: 6 studies; aOR: 1.07	reported here are from RCTs
			anterior circulation	(95% CI 0.79 to 1.46);	although the published
			large vessel occlusion;	mortality: 6 studies; aOR	manuscript also includes data
			outcomes: 3 mo mRS	0.65 (95% CI 0.49 to 0.88)	from observational studies
			score 0 to 2; sICH at	favoring IVT+EVT	
			24 h or 36 h; mortality	mRS score 0 to 1: 4 studies;	
			at discharge or 3 mo; 3	aOR: 1.11 (95% CI 0.90 to	
			mo mRS 0 to 1	1.38)	

	Graded Systematic Reviews/Meta-Analysis					
Author & Year	Class of	Setting & Study	Methods & Outcome	Results	Limitations & Comments	
Published	Evidence	Design	Measures			
Ghaith et al	II	Meta-analysis	Included studies on	N=49 studies; pooled RR for	Subgroup analysis by study	
$(2022)^{34}$			patients with AIS-	favorable neurological	design demonstrated significant	
			LVO,	outcome, 45% for bridge	differences in reported efficacy;	
			exposed/experimental	thrombolysis plus	heterogeneity among studies,	
			group received	thrombectomy group vs 39%	although random effects	
			IVT+MT and	for thrombectomy alone, RR	modeling used to mitigate	
			comparison group only	1.21 (95% CI 1.13 to 1.29,		
			MT; outcomes:	<i>P</i> <.0001); subgroup analyses		
			favorable neurological	by study design showed		
			function based on	favorable outcomes for		
			mRS; mortality,	bridge thrombolysis among		
			successful	observational studies (RR		
			recanalization,	1.25, 95% CI 1.17 to 1.34)		
			complications;	but not for experimental		
			comparative studies	studies (RR 0.99, 95% CI		
			designs including both	0.89 to 1.09); sICH: RR 0.88		
			experimental and	(95% CI 0.70 to 1.10, <i>P</i> =.27)		
			quasi-experimental, or			
			observational designs			

	Graded Systematic Reviews/Meta-Analysis						
Author & Year	Class of	Setting & Study	Methods & Outcome	Results	Limitations & Comments		
Published	Evidence	Design	Measures				
Katsanos et al	III	Meta-analysis	Observational studies	mRS 0 or 1:5 studies, 1,518	Included primarily lower quality		
$(2023)^{35}$			of patients with LVO	participants; aOR 1.32 (95%	studies which studies patients		
			receiving IVT at a	CI 1.00 to 1.74) favoring	who received thrombectomy		
			primary stroke center	IVT+EVT mRS 0 to 2: 5	rather than patients who were		
			before transfer for	studies, 1,518 participants;	eligible for thrombectomy		
			EVT vs transfer for	aOR 1.22 (95% CI 0.95 to			
			EVT alone; outcomes:	1.58); symptomatic ICH: 5			
			3 mo mRS of 0 to 1; 3	studies, 1,535 participants;			
			mo mRS scores of 0 to	aOR 0.72 (95% CI 0.42 to			
			2; sICH within 48 h; 3	1.25); mortality: 5 studies;			
			mo all-cause mortality	1,549 participants; aOR: 0.50			
				(95% CI 0.27 to 0.93)			
				favoring IVT+EVT			

Observational and Retrospective Evidence					
Author & Year	Class of	Setting & Study	Methods & Outcome	Results	Limitations & Comments
Published	Evidence	Design	Measures		
Abilleira et al (2017) ⁴¹	III	Regional registry retrospective cohort from Catalonia, Spain	Patients with anterior circulation stroke caused by large vessel occlusion; EVT vs bridging thrombolysis prior to EVT; outcomes: mRS 0 to 2 at 3 mo; death; symptomatic bleeding 24 h to 36 h	at 90 d: 0.97 (95% CI 0.74 to 1.27); OR for death: 1.07 (95% CI 0.74 to 1.54); OR for symptomatic bleeding: 0.56 (95% CI 0.25 to 1.27)	Discrepancies in important baseline features is accounted for by using propensity score to stratify subjects into blocks; outcome assessments are unblinded; study population included only patients who received thrombectomy rather than those who were eligible for thrombectomy
Balodis et al (2019) ⁴²	III	Prospective single-center study from Latvia	Patients with acute stroke and eligible for endovascular treatment; EVT vs bridging thrombolysis prior to EVT; outcomes: mRS 0 to 2 at discharge and 90 d; symptomatic and asymptomatic intracranial hemorrhage; mortality	thrombectomy, 62 received thrombectomy alone; mRS 0 to 2: 44% in bridging group vs 42% in thrombectomy only group, OR 0.48 (95%	Single center; non-randomized; limited adjustment, including for treatment by indication; unclear outcome assessment blinding

	Observational and Retrospective Evidence						
Author & Year	Class of	Setting & Study	Methods & Outcome	Results	Limitations & Comments		
Published	Evidence	Design	Measures				
Broocks et al	III	Multicenter,	Adults with AIS+LVO	N=720, IVT (N=366) vs no	Multivariable regression analysis		
$(2022)^{43}$		academic center	who received EVT,	IVT (N=354); proportions	with propensity weighting, but		
		in Germany and	with or without IVT,	with favorable outcome: IVT	residual confounding due to		
		the United	2013 to 2021;	(43%) vs none (32%); aOR	treatment indication may bias		
		States;	outcome: functional	1.57 (95% CI 1.16 to 2.14)	estimates		
		retrospective	independence (mRS 0	for functional independence,			
		cohort	to 2) at 90 d	favoring IVT			
Casetta et al	III	Regional	All patients who	N=1,148, 635 with IV	Propensity score methods,		
$(2019)^{45}$		registry,	underwent	thrombolytics plus	including use of IPTW; residual		
		multicenter	endovascular	thrombectomy, 513 with	confounding still possible;		
		prospective	treatment, either	thrombectomy only; IPTW	unclear blinding outcome		
		enrollment from	thrombectomy only vs	mRS 0 to 2: OR 1.3 (95% CI	assessment		
		an Italian	IV thrombolytics plus	0.98 to 1.75); IPTW sICH:			
		registry; 13	thrombectomy for	OR 2.1 (95% CI 0.93 to			
		centers	anterior circulation	1.62)			
			stroke; outcomes:				
			mRS at 90 d; sICH				

· ·	Observational and Retrospective Evidence						
Author & Year	Class of	Setting & Study	Methods & Outcome	Results	Limitations & Comments		
Published	Evidence	Design	Measures				
Di Maria et al (2018) ⁴⁶	III	Retrospective registry cohort from 3 stroke centers located in France	Adult patients with AIS within 6 h of onset with imagining evidence of anterior circulation occlusion; outcomes: mRS 0 to 2 at 90 d; sICH	N=1,507; of the 1,507, 65% received IV thrombolytics; 407 propensity score matched patients and use of multiple imputation to account for missing data; propensity matched mRS 0 to 2: 49% in the thrombolytics plus thrombectomy group vs 45% in the thrombectomy only group, OR 1.21 (95% CI 0.90 to 1.63), <i>P</i> =.21; sICH: 9% for the thrombolytic plus thrombectomy vs 7% for the thrombectomy only group, OR 1.21 (95% CI 0.70 to 0.95% CI 0.70 to 0.95% CI 0.70 to 0.95% CI 0.70 to	Propensity score methods, including matching and adjustment; residual confounding still possible; no apparent blinding for outcome assessment		
				0.90 to 1.63), <i>P</i> =.21; sICH: 9% for the thrombolytic plus thrombectomy vs 7% for the			

•	Observational and Retrospective Evidence					
Author & Year Published	Class of Evidence	Setting & Study Design	Methods & Outcome Measures	Results	Limitations & Comments	
Zha et al (2021) ⁴⁷	III	Post-hoc analysis of a multicenter, prospective cohort study from China	Adult, AIS with baseline mRS <2 who received thrombectomy within 8 h or bridge thrombolysis (within 4.5 h) plus thrombectomy; outcomes: mRS 0 to 2 at 90 d and successful recanalization; sICH; mortality	N=245; propensity score matching with use of multiple imputation for missing values, resulting in 65 pairs; propensity score matched mRS 0 to 2: 49% in bridging thrombolysis group vs 42% in thrombectomy only group, <i>P</i> =.46; propensity score matched mRS 0 to 1: 43% in bridging thrombolysis group vs 25% in thrombectomy only group, <i>P</i> =.023; propensity score matched sICH: 11% in bridging thrombolysis group vs 9% in thrombectomy alone group, <i>P</i> =1.0; propensity score matched mortality: 15% in bridging thrombolysis group vs 25% in thrombectomy alone group, <i>P</i> =.31	Non-randomized limited power\ limited detail regarding use of propensity score methods and thus concern related to remaining imbalances between groups	

AIS, acute ischemic stroke; aOR, adjusted odds ratio; CI, confidence interval; d, day; EVT, endovascular thrombectomy; h, hour; ICH, intercranial hemorrhage; IPTW, inverse probability of treatment weighting; IQR, interquartile range; IVT, intravenous thrombolysis; LVO, large vessel occlusion; min, minutes; mo, month; MT, mechanical thrombectomy; OR, odds ratio; RR, risk ratio; sICH, symptomatic intracranial hemorrhage; vs, versus; y, year.

- Appendix E6. Articles graded for methodological rigor but ultimately found to be fatally flawed.
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