

1 **Clinical Policy: Use of Thrombolytics for the Management of Acute Ischemic Stroke in the Emergency**
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

12 Peter Panagos, MD

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)

50 John T. Finnell, MD (Board Liaison 2020-2024)

51 Travis Schulz, MLS, AHIP, Staff Liaison, Clinical Policies Committee and Writing Committee on Thrombolytics

52 Kaeli Vandertulip, MSLS, MBA, AHIP, Staff Liaison, Clinical Policies Committee
53

54 **ABSTRACT**

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

62
63 **INTRODUCTION**

64
65 Approximately 30% of all acute ischemic strokes have a large vessel occlusion (LVO), which contributes
66 to 64% of all moderate to severe disability from stroke at 3 months and over 95% of stroke deaths at 6 months.^{1,2}
67 Over the past decade, acute treatment for LVO has expanded beyond thrombolytics with evidence supporting the
68 use of endovascular therapy (EVT) such as mechanical thrombectomy.³⁻⁵

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

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

82 Studies that compared EVT alone (direct endovascular therapy or direct mechanical thrombectomy) with
83 IVT + EVT (bridging therapy) used the Modified Rankin Scale (mRS) to assess functional outcomes. The mRS
84 ranges from 0 (no neurological symptoms) to 6 (death). Good functional outcome or functional independence is
85 often defined as mRS 0 to 2, which represents patients with slight disability but who can look after their own
86 affairs without assistance. Excellent functional outcome is usually defined as mRS of 0 to 1, which represents no
87 significant disability and the ability to carry out all duties and activities.¹⁵ Although the mRS is the most common
88 tool used for evaluating disability in stroke research, there are known limitations with inter-rater reliability.¹⁶

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

94 **METHODOLOGY**

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

101 Search and Study Selection

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

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

115

116 Assessment of Risk of Bias and Determination of Classes of Evidence

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

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

135

136 Translation of Classes of Evidence to Recommendation Levels

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

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

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

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

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

156

157 Evaluation and Review of Recommendations

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

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

165

166 Application of the Policy

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

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

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

182

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

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

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

187

188 **CRITICAL QUESTION**

189

190 **In adult stroke patients who are a candidate for mechanical thrombectomy, is the use of IV thrombolysis**
191 **prior to mechanical thrombectomy (Bridge therapy) beneficial and safe versus mechanical thrombectomy**
192 **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

Potential Benefit of Implementing the Recommendations:

- 203 ● Improved functional outcomes
- 204 ● Decreased mortality

205

206

Potential Harm of Implementing the Recommendations:

- 207 ● Delays in endovascular therapy
- 208 ● Increased cost with the use of thrombolytics

209

210

211

212

213

214

215

216

217

218

219

220

221

222

223

224

225

226

227

228

Key words/phrases for literature searches: Acute Ischemic Stroke, Acute Stroke, Alteplase, Anticoagulation Bridge, Brain Ischemia, Bridge Therapy, Bridging Anticoagulation, Catheter-directed Thrombectomy, Cerebrovascular Accident, Directed, Thrombectomy, Elaxim, Emergency Department, Emergency Health Service, Emergency Medical Services, Emergency Medicine, Emergency Treatment, Emergency Ward, EMS, Endovascular Therapy, Endovascular Thrombectomy, EVT, Fibrinolytic, Fibrinolytic Agents, Guided Thrombectomy, Intravenous, Intravenous Drug Administration, Ischemic Stroke, IV, Mechanical Thrombectomy, Metalyse, Percutaneous Thrombectomy, rTPA, Stroke, Tenecteplase, Thrombectomy, Thrombolytic Therapy, Thrombolytic Treatment, Thrombolytic, Tissue Plasminogen Activator, TNKase, tPA, and variations and combinations of key words/phrases. Searches included January 2015 to search the date of April 10, 2023 (Appendix E4).

Study Selection: Five hundred fifty-seven articles were identified in the searches. Three hundred thirty-four articles were selected from the search results as candidates for further review. After grading for methodological rigor, 3 Class I studies, 7 Class II studies, and 8 Class III studies were included for this critical question (Appendix E5). Appendix E6 lists the 69 articles graded for methodological rigor but ultimately found to be fatally flawed.

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. Alteplase

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

235 All the RCTs were designed primarily to evaluate if EVT alone was non-inferior to IVT + EVT except for
236 1 trial (LeCouffe 2021) that evaluated superiority of EVT alone followed by non-inferiority of EVT alone.²² As
237 opposed to superiority studies which are designed to demonstrate better effectiveness of 1 intervention over
238 another, non-inferiority studies are powered to evaluate whether 1 intervention is potentially “less good” than
239 another intervention within a predefined range.²⁷ Non-inferiority trials are appropriate if 1 intervention has added
240 costs, risks, or limited availability that might render superiority less important.²⁸ Since intention-to-treat analysis
241 is more likely to create Type 1 error by falsely concluding non-inferiority compared with per-protocol analysis,
242 dual reporting of both analyses are preferable for non-inferiority trials.^{29,30} To achieve non-inferiority, the lower
243 limit of the confidence interval (CI) should exceed the prespecified non-inferiority margin. Each of the non-
244 inferiority RCT trials in this clinical policy used different primary end points as well as various non-inferiority
245 margins. Both per-protocol and intention-to-treat analysis were performed and remained consistent within each
246 study and is summarized in Table 1.

247 In a Class I study, the DIRECT-MT trial enrolled 654 patients from 41 academic tertiary care centers in
248 China with an internal carotid artery (ICA) or first segment middle cerebral artery (M1)/second segment middle
249 cerebral artery (M2) LVO.²¹ The primary outcome was a median 90-day mRS. Both EVT alone and IVT + EVT
250 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).
251 These results demonstrate non-inferiority as the lower limit margin was set at 0.80. There was no statistical
252 difference in sICH or death at 90 days observed between the 2 groups.

253 The DEVT trial was a Class II study that enrolled 234 patients with an ICA or M1 LVO from 33 stroke
254 centers in China.²⁴ The primary outcome was the proportion of patients achieving mRS 0 to 2 at 90 days. Results
255 from the per-protocol analysis showed an mRS 0 to 2 in 53.2% of the EVT alone group versus 46% of the IVT +
256 EVT group. The absolute difference of 7.1% (97.5% CI -5.9 to ∞) allowed them to conclude non-inferiority
257 based upon their pre-specified margin of 10%. The DEVT trial was stopped early after enrolling only 235 out of
258 the planned 970 patients because of a statistical finding of likely futility. Both groups had similar rates of sICH
259 and death at 90 days with no statistical differences observed.

260 In a Class II study, the SKIP trial enrolled 204 patients from 23 stroke centers in Japan with an ICA or
261 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
262 primary outcome was mRS 0 to 2. Results from the per-protocol analysis showed a favorable neurologic outcome
263 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
264 0.60 to ∞), which did not meet the prespecified lower margin of 0.74. The investigators were unable to conclude
265 non-inferiority. Mortality at 90 days and sICH were not observed to be statistically different between the 2
266 groups.

267 The MR CLEAN-NO IV trial was a Class II study that included 539 patients from 20 hospitals in the
268 Netherlands, Belgium, and France.²² Patients had an acute ischemic stroke due to a proximal occlusion of the
269 anterior circulation. The primary outcome was median mRS at 90 days, first evaluating for superiority of EVT
270 alone over IVT + EVT. If superiority was not established, then an evaluation of non-inferiority of EVT alone
271 compared with IVT + EVT was performed. The non-inferiority margin was set at 0.8 for the adjusted common
272 OR. Median mRS favored IVT + EVT over EVT alone (2 versus 3). Results from the adjusted common OR was
273 0.84 (95% CI 0.62 to 1.15), which demonstrated neither superiority nor non-inferiority for EVT alone. No
274 statistical difference was observed between the 2 groups for sICH or death within 90 days.

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

282 In a Class III study, the DIRECT-SAFE trial enrolled 295 patients from 25 acute-care hospitals in
283 Australia, New Zealand, China, and Vietnam.²⁶ Patients needed to have an LVO in either the ICA, M1 or M2
284 segments of the middle cerebral artery (MCA), or basilar artery and were randomized with or without alteplase in
285 Asian countries (83%) and tenecteplase in non-Asian countries (17%). The primary outcome was mRS 0 to 2 at
286 90 days. Results from the per-protocol analysis showed a favorable neurologic outcome in 54% of the EVT alone

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

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

294 **Table 1.** A synthesis of the ACEP Clinical Policy Level of Evidence, direction of support for bridging therapy
 295 (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 ∞) ³	1.09 (97.5% CI 0.63 to ∞) ^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

296 ^A Adjusted common Odds Ratio

297 ^B Unadjusted difference

298 ^C Odds Ratio

299 ^D Adjusted Risk Difference

300 ^E Unadjusted Risk Difference

301

302 Systematic Reviews/Meta-Analysis

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

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

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

317 In a Class II study, Lin et al reviewed 4 RCTs (DEVT, SKIP, DIRECT-MT, and MR CLEAN NO
318 IV)^{21,22,24,25} for a total of 1,633 patients.³² Based on the literature, they assessed 5 different non-inferiority margins
319 for functional independence (mRS 0 to 2) at 90 days. There was no observed statistical heterogeneity among trials
320 ($I^2=0\%$). Although the risk difference was 1% (95% CI -4% to 5%) favoring EVT alone, the lower margin of the
321 95% CI suggests EVT alone is non-inferior to IVT + EVT except when using the most stringent of margins at
322 -1.3%. The outcome measure of mRS 0 to 1 showed a similar risk difference of 1% (95% CI -3% to 5%),
323 showing non-inferiority except when using the margin of -1.3%. SICH and mortality were not shown to be
324 different between both groups.

325 In another Class II study, Wang et al reviewed 6 RCTs (DEVT, SKIP, DIRECT-MT, DIRECT SAFE,
326 SWIFT DIRECT, and MR CLEAN NO IV)²¹⁻²⁶ for a total of 2,334 patients.¹⁰ This international workgroup
327 consisted of various stakeholders including stroke experts, pharmacists, academics, and caregivers of stroke
328 patients. The workgroup established minimally important differences through survey of their guideline panel and
329 discussion for the following outcomes: 1% for recovery with minimal disability (mRS 0 to 2), 0.8% for mortality,
330 and 1% for sICH. Pooled estimate of effect showed lack of observed statistical heterogeneity ($I^2=0\%$). They
331 concluded with low certainty of evidence that EVT alone had a smaller decrease in patients with minimal
332 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

333 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
334 certainty of evidence that EVT alone had a small decrease in sICH (RR 0.75, 95% CI 0.52 to 1.07; risk difference
335 -1.0%, 95% CI -1.8% to 0.27%).

336 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
337 observational/retrospective studies, for a total of approximately 20,000 patients.³³ A comprehensive meta-analysis
338 was performed for utilizing both RCTs and observational/retrospective studies to investigate various outcomes.
339 Functional independence was defined as mRS of 0 to 2 and excellent outcomes was defined as mRS of 0 to 1. For
340 RCTs, the IVT + EVT group reduced the risk of mortality versus EVT alone (OR 0.65, 95% CI 0.49 to 0.88,
341 $I^2=52\%$), but not functional independence (OR 1.17, 95% CI 0.99 to 1.38, $I^2=0\%$). On the other hand, the
342 observational studies showed that IVT + EVT had better outcomes for functional independence (OR 1.36, 95% CI
343 1.21 to 1.52, $I^2=48\%$), excellent outcomes (OR 1.49, 95% CI 1.26 to 1.75, $I^2=4\%$), and mortality (OR 0.73, 95%
344 CI 0.56 to 0.94, $I^2=67\%$). Neither the RCTs nor observational studies showed an increased risk in sICH.

345 In a Class II study, Ghaith et al reviewed 49 studies (4 RCTs^{21,22,24,25} and 44 observational studies) for a
346 total of 36,123 patients.³⁴ In the analysis combining both RCTs and observational studies, they demonstrated that
347 IVT + EVT had better mortality (RR 0.75, CI 95% 0.68 to 0.82, $I^2=36\%$), successful recanalization (RR 1.06,
348 95% CI 1.03 to 1.09, $I^2=50\%$), and 90-day functional independence (RR 1.21, 95% CI 1.13 to 1.29, $I^2=52\%$), but
349 no improvement in National Institutes of Health Stroke Scale (NIHSS). Subgroups were stratified accounting to
350 study design showing similar benefits with IVT + EVT for observational studies, but not for RCTs. No difference
351 was seen between the 2 groups related to sICH.

352 Lastly, in a Class III study, Katsonos et al included 6 observational studies totaling 1,723 patients.
353 Patients who received IVT at a PSC before transferring for EVT (“drip and ship” or DNS, 53% of the group) were
354 compared with those receiving EVT alone at a Comprehensive Stroke Center (CSC).³⁵ In their analysis adjusted
355 for potential confounders, “DNS patients” had higher odds of mRS 0 to 1 (adjusted OR 1.32, 95% CI 1.00 to 1.74,
356 $I^2=0\%$) and lower probability for all-cause mortality at 3-months (adjusted OR 0.50, 95% CI: 0.27 to 0.93,
357 $I^2=69\%$) compared to patients receiving EVT alone at a CSC. No differences were found between the 2 groups in
358 probability of 3-month disability, mRS 0 to 2, or sICH.

359 The majority of SRMA favored IVT + EVT. Two of the SRMA showed either improved mortality or
360 improved functional outcomes with IVT + EVT, however these results varied based on whether the analysis
361 utilized RCTs and/or observational studies.^{33,34} Of the 3 studies that looked at the RCTs alone, 1 SRMA³² showed
362 non-inferiority of EVT alone compared with IVT + EVT in various cutoffs except for the most strict cutoff for
363 functional outcomes while another SRMA¹⁰ suggested a possible small increase in mortality, a small decrease in
364 recovery with minimal disability, but moderate certainty of decreased sICH with EVT alone. The other SRMA
365 that utilized RCTs alone suggests that IVT + EVT is superior to EVT alone but is time dependent.³¹ Lastly in
366 patients who are transferred, evidence suggests patients who received IVT + EVT have better functional outcomes
367 and mortality compared with EVT alone.³⁵

368

369 Observational and Retrospective Evidence

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

375 Balodis et al reported a single-center prospective observational analysis of IVT + EVT versus EVT alone
376 for anterior cerebral artery LVO in a single Latvian university hospital.⁴² Although exclusions did not include a
377 time-of-onset for symptoms, all thrombectomy occurred within 8 hours of symptom onset and all patients
378 presenting within 4.5 hours received IVT unless contraindications were identified, or physician's preference was
379 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
380 EVT-alone group. No significant differences were observed in 90-day mortality or sICH.

381 Broocks et al retrospectively analyzed a cohort of acute ischemic stroke patients treated at 1 of 2 high-
382 volume tertiary stroke centers in Germany and the United States for ICA or MCA LVO.⁴³ The Alberta Stroke
383 Program Early CT Score (ASPECTS) was determined on pre-treatment non-contrast head CT by 1 neuro-
384 radiologist.⁴⁴ Most had ASPECTS >5 (86%). Overall, those receiving IVT + EVT had better NIHSS at 24 hours
385 (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

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

388 Casetta et al reviewed the Italian Registry of Endovascular Stroke Treatments prospective observational
389 data from 13 hospitals which included 1,148 patients with either an ICA or M1/M2 LVO who were eligible for
390 IVT.⁴⁵ EVT was performed within 6 hours of symptom onset and decisions about IVT were left to the discretion
391 of the treating neurology team. Although the median time from symptom onset to hospital arrival was similar
392 between the 2 groups (95 minutes for IVT + EVT versus 96 minutes for EVT alone patients), the symptom onset
393 to groin puncture was significantly prolonged in the IVT + EVT subset (230 minutes versus 210 minutes in EVT).
394 Multivariate analysis for stroke patients surviving with mRS 0 to 3 demonstrated a significant benefit favoring
395 IVT + EVT (adjusted OR 1.42; 95% CI 1.04 to 1.95) and a significantly lower risk of death or unfavorable
396 outcome in that same group (adjusted OR 0.62; 95% CI 0.45 to 0.84). No differences were found regarding sICH.

397 Di Maria et al retrospectively evaluated acute ischemic stroke patients involving the proximal or distal
398 MCA or ICA within 6-hours of symptoms.⁴⁶ A stroke neurologist decided whether or not to treat with IVT. IVT +
399 EVT patients were matched with patients treated with EVT alone using a propensity score. An mRS 0 to 2 was
400 more likely with IVT + EVT (OR 1.31; 95% CI 1.02 to 1.68). All-cause mortality and sICH did not differ
401 between groups. Only ASPECTS \geq 7 demonstrate the benefit of IVT + EVT compared with EVT alone (OR 1.48,
402 95% CI 1.10 to 2.0).

403 Zha et al reported a post-hoc analysis of a prospective study across 16 Chinese stroke centers.⁴⁷ The
404 primary outcome of mRS 0 to 2 at 90 days. In a multivariable analysis, IVT + EVT more frequently demonstrated
405 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
406 0 to 2. The 90-day mortality rate was significantly lower in the IVT + EVT cohort (13.9% versus 27.7%).

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

411

412 Summary

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

418 From a safety standpoint, although some studies showed a decrease in mortality with IVT + EVT, most
419 studies showed no difference. Lastly, although there have been concerns about the increased risk of sICH with
420 the addition of IVT before EVT, no study included in our review showed an increased risk of sICH. However,
421 safety data from these studies may have also been under-reported.^{48,49} It is important that with any intervention,
422 shared decision making is made when feasible with the patient and/or family.

423

424 Future Research

425 Existing research predominantly employed alteplase as the primary thrombolytic agent. Subsequent
426 investigations should explore alternative thrombolytics, such as tenecteplase.⁵⁰ Future studies should also look at
427 timing of thrombolytics prior to EVT with patient outcomes. In addition, the role of ASPECTS score and other
428 tools in identifying individuals unlikely to benefit from the addition of IVT prior to EVT should be explored
429 prospectively.⁴³ Furthermore, future studies ought to consider larger sample sizes, utilizing more stringent non-
430 inferiority margins or ideally conducting superiority studies, as well as evaluating the cost-effectiveness of
431 different treatment strategies.⁵¹

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

437

438 ***Relevant industry relationships: There were no relevant industry relationships disclosed by the***
439 ***subcommittee members for this topic.***

440 *Relevant industry relationships are those relationships with companies associated with products or*
441 *services that significantly influence the specific aspect of disease addressed in the critical question.*
442

DRAFT

443 **REFERENCES**

- 444
- 445 1. Lakomkin N, Dhamoon M, Carroll K, et al. Prevalence of large vessel occlusion in patients presenting with
- 446 acute ischemic stroke: a 10-year systematic review of the literature. *J Neurointerv Surg.* 2019;3:241-245.
- 447
- 448 2. Malhotra K, Gornbein J, Saver JL. Ischemic Strokes Due to Large-Vessel Occlusions Contribute
- 449 Disproportionately to Stroke-Related Dependence and Death: A Review. *Front Neurol.* 2017;8:651.
- 450
- 451 3. Powers WJ, Rabinstein AA, Ackerson T, et al. Guidelines for the Early Management of Patients With Acute
- 452 Ischemic Stroke: 2019 Update to the 2018 Guidelines for the Early Management of Acute Ischemic Stroke: A
- 453 Guideline for Healthcare Professionals From the American Heart Association/American Stroke Association.
- 454 *Stroke.* 2019;50:e344-e418.
- 455
- 456 4. Heran M, Lindsay P, Gubitz G, et al. Canadian Stroke Best Practice Recommendations: Acute Stroke
- 457 Management, 7th Edition Practice Guidelines Update, 2022. *Can J Neurol Sci.* 2024;51:1-31.
- 458
- 459 5. Turc G, Bhogal P, Fischer U, Khatri P, Lobotesis K, Mazighi M, Schellinger PD, Toni D, de Vries J, White P,
- 460 Fiehler J. European Stroke Organisation (ESO) - European Society for Minimally Invasive Neurological
- 461 Therapy (ESMINT) Guidelines on Mechanical Thrombectomy in Acute Ischemic Stroke. *J Neurointerv Surg.*
- 462 2023;15:e8.
- 463
- 464 6. Desilles JP, Loyau S, Syvannarath V, et al. Alteplase Reduces Downstream Microvascular Thrombosis and
- 465 Improves the Benefit of Large Artery Recanalization in Stroke. *Stroke.* 2015;46:3241-3248.
- 466
- 467 7. Seners P, Turc G, Maier B, et al. Incidence and Predictors of Early Recanalization After Intravenous
- 468 Thrombolysis: A Systematic Review and Meta-Analysis. *Stroke.* 2016;47:2409-2412.
- 469
- 470 8. Tsivgoulis G, Katsanos AH, Schellinger PD, et al. Successful Reperfusion With Intravenous Thrombolysis
- 471 Preceding Mechanical Thrombectomy in Large-Vessel Occlusions. *Stroke.* 2018;49:232-235.
- 472
- 473 9. Ohara T, Menon BK, Al-Ajlan FS, et al. Thrombus Migration and Fragmentation After Intravenous Alteplase
- 474 Treatment: The INTERSeCT Study. *Stroke.* 2021;52:203-212.
- 475
- 476 10. Wang X, Ye Z, Busse JW, et al. Endovascular thrombectomy with or without intravenous alteplase for acute
- 477 ischemic stroke due to large vessel occlusion: a systematic review and meta-analysis of randomized trials.
- 478 *Stroke Vasc Neurol.* 2022;7:510-517.
- 479
- 480 11. Atchaneeyasakul K, Desai S, Malhotra K, et al. Intravenous alteplase delays door-to-puncture time in acute
- 481 ischemic stroke with large vessel occlusion. *J Stroke Cerebrovasc Dis.* 2021;30:105732.
- 482
- 483 12. Zachrison KS, Cash RE, Adeoye O, et al. Estimated Population Access to Acute Stroke and Telestroke
- 484 Centers in the US, 2019. *JAMA Netw Open.* 2022;2:e2145824.
- 485
- 486 13. Sarraj A, Savitz S, Pujara D, et al. Endovascular Thrombectomy for Acute Ischemic Strokes: Current US
- 487 Access Paradigms and Optimization Methodology. *Stroke.* 2020;51:1207-1217.
- 488
- 489 14. Aldstadt J, Waqas M, Yasumiishi M, et al. Mapping access to endovascular stroke care in the USA and
- 490 implications for transport models. *J Neurointerv Surg.* 2022;14:neurintsurg-2020-016942.
- 491
- 492 15. van Swieten JC, Koudstaal PJ, Visser MC, et al. Interobserver agreement for the assessment of handicap in
- 493 stroke patients. *Stroke.* 1988;19:604-607.
- 494

- 495 16. Pożarowszczyk N, Kurkowska-Jastrzębska I, Sarzyńska-Długosz I, et al. Reliability of the modified Rankin
496 Scale in clinical practice of stroke units and rehabilitation wards. *Front Neurol.* 2023;14:1064642.
497
- 498 17. Singh N, Kashani N, Ganesh A, et al. Understanding physician and patient preferences for thrombolysis in
499 ischemic stroke eligible for endovascular thrombectomy. *Stroke Vasc Interv Neurol.* 2022;2:e000218.
500
- 501 18. Masoud HE, de Havenon A, Castonguay AC, et al. 2022 Brief practice update on intravenous thrombolysis
502 before thrombectomy in patients with large vessel occlusion acute ischemic stroke: a statement from Society
503 of Vascular and Interventional Neurology Guidelines and Practice Standards (GAPS) Committee. *Stroke Vasc*
504 *Interv Neurol.* 2022;2:1-10.
505
- 506 19. Ye Z, Busse JW, Hill MD, et al. Endovascular thrombectomy and intravenous alteplase in patients with acute
507 ischemic stroke due to large vessel occlusion: A clinical practice guideline. *J Evid Based Med.* 2022;15:263-
508 271.
509
- 510 20. Page MJ, McKenzie JE, Bossuyt PM, et al. The PRISMA 2020 statement: an updated guideline for reporting
511 systematic reviews. *BMJ.* 2021;372:n71.
512
- 513 21. Yang P, Zhang Y, Zhang L, et al. Endovascular Thrombectomy with or without Intravenous Alteplase in
514 Acute Stroke. *N Engl J Med.* 2020;382:1981-1993.
515
- 516 22. LeCouffe NE, Kappelhof M, Treurniet KM, et al. A Randomized Trial of Intravenous Alteplase before
517 Endovascular Treatment for Stroke. *N Engl J Med.* 2021;385:1833-1844.
518
- 519 23. Fischer U, Kaesmacher J, Strbian D, et al. Thrombectomy alone versus intravenous alteplase plus
520 thrombectomy in patients with stroke: an open-label, blinded-outcome, randomised non-inferiority trial.
521 *Lancet.* 2022;400(10346):104-115.
522
- 523 24. Zi W, Qiu Z, Li F, et al. Effect of Endovascular Treatment Alone vs Intravenous Alteplase Plus Endovascular
524 Treatment on Functional Independence in Patients With Acute Ischemic Stroke: The DEVT Randomized
525 Clinical Trial. *JAMA.* 2021;325:234-243.
526
- 527 25. Suzuki K, Matsumaru Y, Takeuchi M, et al. Effect of Mechanical Thrombectomy Without vs With
528 Intravenous Thrombolysis on Functional Outcome Among Patients With Acute Ischemic Stroke: The SKIP
529 Randomized Clinical Trial. *JAMA.* 2021;325:244-253.
530
- 531 26. Mitchell PJ, Yan B, Churilov L, et al. Endovascular thrombectomy versus standard bridging thrombolytic
532 with endovascular thrombectomy within 4·5 h of stroke onset: an open-label, blinded-endpoint, randomised
533 non-inferiority trial. *Lancet.* 2022;400(10346):116-125.
534
- 535 27. Kaul S, Diamond GA. Good enough: a primer on the analysis and interpretation of noninferiority trials. *Ann*
536 *Intern Med.* 2006;145:62-69.
537
- 538 28. Al Deeb M, Azad A, Barbic D. Critically appraising noninferiority randomized controlled trials: a primer for
539 emergency physicians. *CJEM.* 2015;17:231-236.
540
- 541 29. Piaggio G, Elbourne DR, Pocock SJ, et al. Reporting of noninferiority and equivalence randomized trials:
542 extension of the CONSORT 2010 statement. *JAMA.* 2012;308:2594-2604.
543
- 544 30. Wiens BL, Zhao W. The role of intention to treat in analysis of noninferiority studies. *Clin Trials.*
545 2007;4:286-291.
546
547

- 548 31. Kaesmacher J, Cavalcante F, Kappelhof M, et al. Time to Treatment With Intravenous Thrombolysis Before
549 32. Thrombectomy and Functional Outcomes in Acute Ischemic Stroke: A Meta-Analysis. *JAMA*.
550 2024;331:764-777.
551
- 552 32. Lin CH, Saver JL, Ovbiagele B, et al. Endovascular thrombectomy without versus with intravenous
553 thrombolysis in acute ischemic stroke: a non-inferiority meta-analysis of randomized clinical trials. *J*
554 *Neurointerv Surg*. 2022;14:227-232.
555
- 556 33. Zheng M, Li L, Chen L, et al. Mechanical thrombectomy combined with intravenous thrombolysis for acute
557 ischemic stroke: a systematic review and meta-analyses. *Sci Rep*. 2023;13:8597.
558
- 559 34. Ghaith HS, Elfil M, Gabra MD, et al. Intravenous thrombolysis before mechanical thrombectomy for acute
560 ischemic stroke due to large vessel occlusion; should we cross that bridge? A systematic review and meta-
561 analysis of 36,123 patients. *Neurol Sci*. 2022;43:6243-6269.
562
- 563 35. Katsanos AH, Sarraj A, Froehler M, et al. IV Thrombolysis Initiated Before Transfer for Endovascular Stroke
564 Thrombectomy: A Systematic Review and Meta-analysis. *Neurology*. 2023;100:e1436-e1443.
565
- 566 36. Coutinho JM, Liebeskind DS, Slater LA, et al. Combined intravenous thrombolysis and thrombectomy vs
567 thrombectomy alone for acute ischemic stroke: A pooled analysis of the SWIFT and STAR studies. *JAMA*
568 *Neurol*. 2017;74:268–274.
569
- 570 37. Gariel F, Lapergue B, Bourcier R, et al. Mechanical thrombectomy outcomes with or without intravenous
571 thrombolysis. *Stroke*. 2018;49:2383–2390.
572
- 573 38. Chalos V, LeCouffe NE, Uyttenboogaart M, et al. Endovascular treatment with or without prior intravenous
574 alteplase for acute ischemic stroke. *J Am Heart Assoc*. 2019;8:e011592.
575
- 576 39. Huu An N, Dan Luu V, Duy Ton M, et al. Thrombectomy alone versus bridging therapy in acute ischemic
577 stroke: Preliminary results of an experimental trial. *Clin Ter*. 2022;173:107–114.
578
- 579 40. Sakai N, Takeuchi M, Imamura H, et al. Safety, pharmacokinetics and pharmacodynamics of DS-1040, in
580 combination with thrombectomy, in Japanese Patients with acute ischemic stroke. *Clin Drug Investig*.
581 2022;42:137–149.
582
- 583 41. Abilleira S, Ribera A, Cardona P, et al. Outcomes After Direct Thrombectomy or Combined Intravenous and
584 Endovascular Treatment Are Not Different. *Stroke*. 2017;48:375-378.
585
- 586 42. Balodis A, Radzina M, Miglane E, et al. Endovascular thrombectomy in anterior circulation stroke and
587 clinical value of bridging with intravenous thrombolysis. *Acta Radiol*. 2019;60:308-314.
588
- 589 43. Broocks G, Heit JJ, Kuraitis GM, et al. Benefit of Intravenous Alteplase before Thrombectomy Depends on
590 ASPECTS. *Ann Neurol*. 2022;92:588-595.
591
- 592 44. Barber PA, Demchuk AM, Zhang J, Buchan AM. Validity and reliability of a quantitative computed
593 tomography score in predicting outcome of hyperacute stroke before thrombolytic therapy. ASPECTS Study
594 Group. Alberta Stroke Programme Early CT Score [published correction appears in *Lancet*
595 2000;355(9221):2170]. *Lancet*. 2000;355(9216):1670-1674.
596
- 597 45. Casetta I, Pracucci G, Saletti A, et al. Combined intravenous and endovascular treatment versus primary
598 mechanical thrombectomy. The Italian Registry of Endovascular Treatment in Acute Stroke. *Int J Stroke*.
599 2019;14:898-907.
600

- 601 46. Di Maria F, Mazighi M, Kyheng M, et al. Intravenous Thrombolysis Prior to Mechanical Thrombectomy in
602 Acute Ischemic Stroke: Silver Bullet or Useless Bystander?. *J Stroke*. 2018;20:385-393.
603
- 604 47. Zha M, Huang K, Yang D, et al. Bridge mechanical thrombectomy may be a better choice for acute large
605 vessel occlusions. *J Thromb Thrombolysis*. 2021;52:291-300.
606
- 607 48. Hodkinson A, Kirkham JJ, Tudur-Smith C, et al. Reporting of harms data in RCTs: a systematic review of
608 empirical assessments against the CONSORT harms extension. *BMJ Open*. 2013;3:e003436.
609
- 610 49. Zorzela L, Golder S, Liu Y, et al. Quality of reporting in systematic reviews of adverse events: systematic
611 review. *BMJ*. 2014;348:f7668.
612
- 613 50. Lo BM, Carpenter CR, Ducey S, et al. Clinical Policy: Critical Issues in the Management of Adult Patients
614 Presenting to the Emergency Department With Acute Ischemic Stroke. *Ann Emerg Med*. 2023;82:e17-e64.
615
- 616 51. Kaul S. Understanding the Merits and Drawbacks of Noninferiority Trials in Cardiovascular Medicine. *Can J*
617 *Cardiol*. 2021;37:1378-1393.

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

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

620 †Objective is to measure therapeutic efficacy comparing interventions.

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

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

623

624 **Appendix E2.** Approach to downgrading strength of evidence.

625

626

627

628

629

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

630

631

632

633

634

635

636

637

638

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

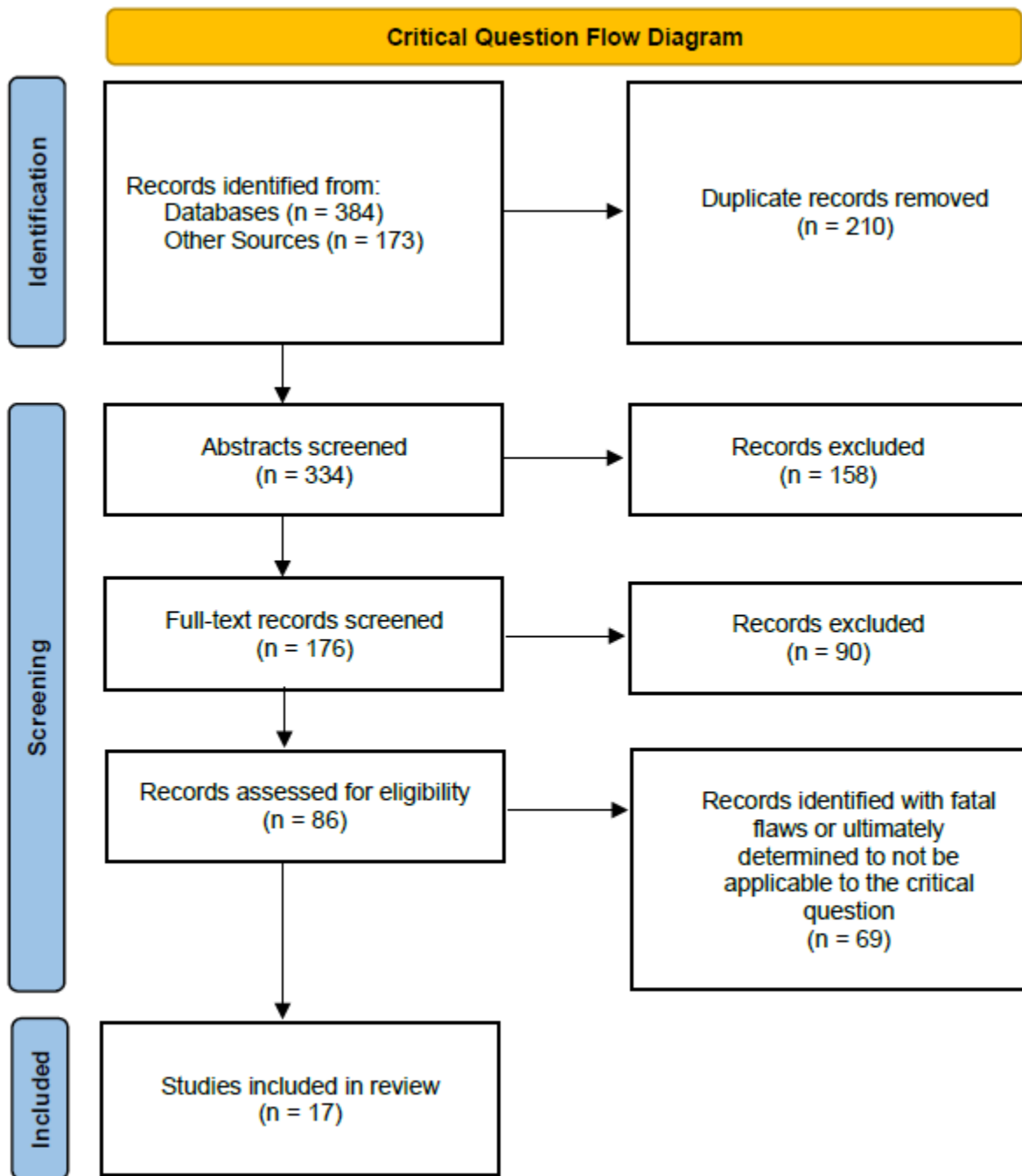
LR (+)	LR (-)	
1.0	1.0	Does not change pretest probability
1–5	0.5–1	Minimally changes pretest probability
10	0.1	May be diagnostic if the result is concordant with pretest probability
20	0.05	Usually diagnostic
100	0.01	Almost always diagnostic even in the setting of low or high pretest probability

639 *LR*, likelihood ratio.

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

643

644



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"[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 Department[tiab] OR Emergency Medical Service*[tiab] OR EMS[tiab] OR ("Emergency Medicine"[mh] OR ("Emergency Service, Hospital"[mh] OR ("Emergency Treatment"[mh] OR ("Emergency Medical Services"[mh]))	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) AND ('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 Therapy" OR "EVT" OR "Endovascular Thrombectomy" OR "Guided Thrombectomy" OR "Directed 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 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/in-person interview (intention-to-treat [ITT] analysis)	N=656; 327 EVT alone; 329 IVT+EVT; EVT alone noninferior aOR 1.07 (95% CI 0.81 to 1.40, $P=.04$), but was associated with lower percentage with successful reperfusion before thrombectomy (2.4% vs 7%) and overall successful reperfusion (79.4% vs 84.5%) and 90 d mortality 17.7% in EVT only vs 18.8% in IVT+EVT	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

Evidentiary Table (continued).

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

Graded Randomized Controlled Trials

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

658 **Evidentiary Table (continued).****Graded Randomized Controlled Trials**

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

659

Graded Systematic Reviews/Meta-Analysis

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

661 **Evidentiary Table (continued).**

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

662

663 **Evidentiary Table (continued).**

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

664

665 **Evidentiary Table (continued).**

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

666

DRAFT

Observational and Retrospective Evidence

Author & Year Published	Class of Evidence	Setting & Study Design	Methods & Outcome Measures	Results	Limitations & Comments
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	N=1,166; 599 received EVT only and 567 IVT followed by EVT; OR for mRS 0 to 2 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	N=146; 84 received bridging thrombolysis followed by thrombectomy, 62 received thrombectomy alone; mRS 0 to 2: 44% in bridging group vs 42% in thrombectomy only group, OR 0.48 (95% CI 0.22 to 1.07), $P=.14$; mortality: 17% in bridging group vs 21% in thrombectomy only group, $P=.57$; symptomatic hemorrhage: 12% in bridging group vs 10% in thrombectomy only group, $P=.79$	Single center; non-randomized; limited adjustment, including for treatment by indication; unclear outcome assessment blinding

Observational and Retrospective Evidence

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

671 **Evidentiary Table (continued).****Observational and Retrospective Evidence**

Author & Year Published	Class of Evidence	Setting & Study Design	Methods & Outcome Measures	Results	Limitations & Comments
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 imaging 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 2.09, <i>P</i> =.5)	Propensity score methods, including matching and adjustment; residual confounding still possible; no apparent blinding for outcome assessment

672

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, $P=.46$; propensity score matched mRS 0 to 1: 43% in bridging thrombolysis group vs 25% in thrombectomy only group, $P=.023$; propensity score matched sICH: 11% in bridging thrombolysis group vs 9% in thrombectomy alone group, $P=1.0$; propensity score matched mortality: 15% in bridging thrombolysis group vs 25% in thrombectomy alone group, $P=.31$	Non-randomized limited power\ limited detail regarding use of propensity score methods and thus concern related to remaining imbalances between groups

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

677 **Appendix E6.** Articles graded for methodological rigor but ultimately found to be fatally flawed.
678

679 Abilleira, S., et al., *Outcomes of a contemporary cohort of 536 consecutive patients with acute ischemic stroke treated*
680 *with endovascular therapy.* Stroke, 2014. 45(4): p. 1046-52.

681 Al-Khaled, M., et al., *Comparing outcome and recanalization results in patients with anterior circulation stroke following*
682 *endovascular treatment with and without a treatment with rt-PA: A single-center study.* Brain Behav, 2018. 8(5): p.
683 e00974.

684 Alonso de Leciana, M., et al., *Mechanical thrombectomy in patients with medical contraindications for intravenous*
685 *thrombolysis: a prospective observational study.* J Neurointerv Surg, 2017. 9(11): p. 1041-1046.

686 Anadani, M., et al., *Endovascular therapy with or without intravenous thrombolysis in acute stroke with tandem*
687 *occlusion.* J Neurointerv Surg, 2022. 14(4): p. 314-320.

688 Bellwald, S., et al., *Direct Mechanical Intervention Versus Bridging Therapy in Stroke Patients Eligible for Intravenous*
689 *Thrombolysis: A Pooled Analysis of 2 Registries.* Stroke, 2017. 48(12): p. 3282-3288.

690 Berkhemer, O.A., et al., *A randomized trial of intraarterial treatment for acute ischemic stroke.* N Engl J Med, 2015.
691 372(1): p. 11-20.

692 Bourcier, R., et al., *Is bridging therapy still required in stroke due to carotid artery terminus occlusions?* J Neurointerv
693 Surg, 2018. 10(7): p. 625-628.

694 Broeg-Morvay, A., et al., *Direct Mechanical Intervention Versus Combined Intravenous and Mechanical Intervention in*
695 *Large Artery Anterior Circulation Stroke: A Matched-Pairs Analysis.* Stroke, 2016. 47(4): p. 1037-44.

696 Brooks, G., et al., *Impact of intravenous alteplase on sub-angiographic emboli in high-resolution diffusion-weighted*
697 *imaging following successful thrombectomy.* Eur Radiol, 2021. 31(11): p. 8228-8235.

698 Chalos, V., et al., *Endovascular Treatment With or Without Prior Intravenous Alteplase for Acute Ischemic Stroke.* J Am
699 Heart Assoc, 2019. 8(11): p. e011592.

700 Chang, A., et al., *Intravenous Tissue Plasminogen Activator in Combination With Mechanical Thrombectomy: Clot*
701 *Migration, Intracranial Bleeding, and the Impact of "Drip and Ship" on Effectiveness and Outcomes.* Front Neurol, 2020.
702 11: p. 585929.

703 Choi, J.H., et al., *Comparison of Outcomes After Mechanical Thrombectomy Alone or Combined with Intravenous*
704 *Thrombolysis and Mechanical Thrombectomy for Patients with Acute Ischemic Stroke due to Large Vessel Occlusion.*
705 World Neurosurg, 2018. 114: p. e165-e172.

706 Ciccone, A., E. Berge, and U. Fischer, *Systematic review of organizational models for intra-arterial treatment of acute*
707 *ischemic stroke.* Int J Stroke, 2019. 14(1): p. 12-22.
708
709
710
711
712
713
714
715
716
717
718
719
720

- 721 Coutinho, J.M., et al., *Combined Intravenous Thrombolysis and Thrombectomy vs Thrombectomy Alone for Acute*
722 *Ischemic Stroke: A Pooled Analysis of the SWIFT and STAR Studies*. JAMA Neurol, 2017. 74(3): p. 268-274.
- 723
- 724 D'Anna, L., et al., *Endovascular Thrombectomy with or without Intravenous Thrombolysis for Anterior Circulation Large*
725 *Vessel Occlusion in the Imperial College London Thrombectomy Registry*. J Clin Med, 2023. 12(3).
- 726
- 727 Dávalos, A., et al., *Retrospective multicenter study of Solitaire FR for revascularization in the treatment of acute ischemic*
728 *stroke*. Stroke, 2012. 43(10): p. 2699-705.
- 729
- 730 Del Toro-Pérez, C., et al., *Direct Mechanical Thrombectomy vs. Bridging Therapy in Stroke Patients in A "Stroke Belt"*
731 *Region of Southern Europe*. J Pers Med, 2023. 13(3).
- 732
- 733 Dicipinigitis, A.J., et al., *Endovascular thrombectomy with and without preceding intravenous thrombolysis for treatment*
734 *of large vessel anterior circulation stroke: A cross-sectional analysis of 50,000 patients*. J Neurol Sci, 2022. 434: p.
735 120168.
- 736
- 737 Faizy, T.D., et al., *Association Between Intravenous Thrombolysis and Clinical Outcomes Among Patients With Ischemic*
738 *Stroke and Unsuccessful Mechanical Reperfusion*. JAMA Netw Open, 2023. 6(5): p. e2310213.
- 739
- 740 Ferrigno, M., et al., *Intravenous Recombinant Tissue-Type Plasminogen Activator: Influence on Outcome in Anterior*
741 *Circulation Ischemic Stroke Treated by Mechanical Thrombectomy*. Stroke, 2018. 49(6): p. 1377-1385.
- 742
- 743 Gamba, M., et al., *Intravenous fibrinolysis plus endovascular thrombectomy versus direct endovascular thrombectomy for*
744 *anterior circulation acute ischemic stroke: clinical and infarct volume results*. BMC Neurol, 2019. 19(1): p. 103.
- 745
- 746 Gariel, F., et al., *Mechanical Thrombectomy Outcomes With or Without Intravenous Thrombolysis*. Stroke, 2018. 49(10):
747 p. 2383-2390.
- 748
- 749 Gong, L., et al., *Bridging Therapy Versus Direct Mechanical Thrombectomy in Patients with Acute Ischemic Stroke due to*
750 *Middle Cerebral Artery Occlusion: A Clinical- Histological Analysis of Retrieved Thrombi*. Cell Transplant, 2019. 28(6):
751 p. 684-690.
- 752
- 753 Goyal, N., et al., *Comparative safety and efficacy of combined IVT and MT with direct MT in large vessel occlusion*.
754 *Neurology*, 2018. 90(15): p. e1274-e1282.
- 755
- 756 Goyal, N., et al., *Impact of pretreatment with intravenous thrombolysis on reperfusion status in acute strokes treated with*
757 *mechanical thrombectomy*. J Neurointerv Surg, 2019. 11(11): p. 1073-1079.
- 758
- 759 Guedin, P., et al., *Prior IV Thrombolysis Facilitates Mechanical Thrombectomy in Acute Ischemic Stroke*. J Stroke
760 *Cerebrovasc Dis*, 2015. 24(5): p. 952-7.
- 761
- 762 Guimarães Rocha, M., et al., *Primary Thrombectomy Versus Combined Mechanical Thrombectomy and Intravenous*
763 *Thrombolysis in Large Vessel Occlusion Acute Ischemic Stroke*. J Stroke Cerebrovasc Dis, 2019. 28(3): p. 627-631.
- 764

765 Hassan, A.E., et al., *Pre-thrombectomy intravenous thrombolytics are associated with increased hospital bills without*
766 *improved outcomes compared with mechanical thrombectomy alone.* J Neurointerv Surg, 2019. 11(12): p. 1187-1190.
767

768 Heinrichs, A., et al., *Relevance of standard intravenous thrombolysis in endovascular stroke therapy of a tertiary stroke*
769 *center.* Acta Neurol Belg, 2018. 118(1): p. 105-111.
770

771 Hinsenveld, W.H., et al., *Intravenous Thrombolysis Is Not Associated with Increased Time to Endovascular Treatment.*
772 Cerebrovasc Dis, 2020. 49(3): p. 321-327.
773

774 Huu An, N., et al., *Thrombectomy Alone versus Bridging Therapy in Acute Ischemic Stroke: Preliminary Results of an*
775 *Experimental Trial.* Clin Ter, 2022. 173(2): p. 107-114.
776

777 Imbarrato, G., J. Bentley, and A. Gordhan, *Clinical Outcomes of Endovascular Thrombectomy in Tissue Plasminogen*
778 *Activator versus Non-Tissue Plasminogen Activator Patients at Primary Stroke Care Centers.* J Neurosci Rural Pract,
779 2018. 9(2): p. 240-244.
780

781 Jian, Y., et al., *Direct versus Bridging Mechanical Thrombectomy in Elderly Patients with Acute Large Vessel Occlusion:*
782 *A Multicenter Cohort Study.* Clin Interv Aging, 2021. 16: p. 1265-1274.
783

784 Kaesmacher, J. and J.F. Kleine, *Bridging Therapy with i. v. rtPA in MCA Occlusion Prior to Endovascular*
785 *Thrombectomy: a Double-Edged Sword?* Clin Neuroradiol, 2018. 28(1): p. 81-89.
786

787 Kandregula, S., et al., *Direct thrombectomy versus bridging thrombolysis with mechanical thrombectomy in middle*
788 *cerebral artery stroke: a real-world analysis through National Inpatient Sample data.* Neurosurg Focus, 2021. 51(1): p.
789 E4.
790

791 Kass-Hout, T., et al., *Is bridging with intravenous thrombolysis of any benefit in endovascular therapy for acute ischemic*
792 *stroke?* World Neurosurg, 2014. 82(3-4): p. e453-8.
793

794 Leker, R.R., et al., *Direct Thrombectomy versus Bridging for Patients with Emergent Large-Vessel Occlusions.* Interv
795 Neurol, 2018. 7(6): p. 403-412.
796

797 Leker, R.R., et al., *Is Bridging Necessary? A Pilot Study of Bridging versus Primary Stentriever-Based Endovascular*
798 *Reperfusion in Large Anterior Circulation Strokes.* J Stroke Cerebrovasc Dis, 2015. 24(6): p. 1163-7.
799

800 Lin, L., et al., *Bridging Thrombolysis Before Endovascular Therapy in Stroke Patients With Faster Core Growth.*
801 Neurology, 2023. 100(20): p. e2083-e2092.
802

803 Machado, M., et al., *Functional Outcome After Mechanical Thrombectomy with or without Previous Thrombolysis.* J
804 Stroke Cerebrovasc Dis, 2021. 30(2): p. 105495.
805

806 Maier, I.L., et al., *Bridging-therapy with intravenous recombinant tissue plasminogen activator improves functional*
807 *outcome in patients with endovascular treatment in acute stroke.* J Neurol Sci, 2017. 372: p. 300-304.
808

809 Maingard, J., et al., *Outcomes of endovascular thrombectomy with and without bridging thrombolysis for acute large*
810 *vessel occlusion ischaemic stroke*. Intern Med J, 2019. 49(3): p. 345-351.

811
812 Masoud HE, de Havenon A, Castonguay AC, et al. 2022 Brief Practice Update on Intravenous Thrombolysis Before
813 Thrombectomy in Patients With Large Vessel Occlusion Acute Ischemic Stroke: A Statement from Society of Vascular
814 and Interventional Neurology Guidelines and Practice Standards (GAPS) Committee. Stroke Vasc Interv Neurol.
815 2022;2:1-10.

816
817 Merlino, G., et al., *Short and long-term outcomes after combined intravenous thrombolysis and mechanical thrombectomy*
818 *versus direct mechanical thrombectomy: a prospective single-center study*. J Thromb Thrombolysis, 2017. 44(2): p. 203-
819 209.

820
821 Minnerup, J., et al., *Outcome After Thrombectomy and Intravenous Thrombolysis in Patients With Acute Ischemic Stroke:*
822 *A Prospective Observational Study*. Stroke, 2016. 47(6): p. 1584-92.

823
824 Mokin, M., et al., *Recent Endovascular Stroke Trials and Their Impact on Stroke Systems of Care*. J Am Coll Cardiol,
825 2016. 67(22): p. 2645-55.

826
827 Mokin, M., et al., *Intravenous alteplase has different effects on the efficacy of aspiration and stent retriever*
828 *thrombectomy: analysis of the COMPASS trial*. J Neurointerv Surg, 2022. 14(10): p. 992-996.

829
830 Park, H.K., et al., *Preceding Intravenous Thrombolysis in Patients Receiving Endovascular Therapy*. Cerebrovasc Dis,
831 2017. 44(1-2): p. 51-58.

832
833 Pfefferkorn, T., et al., *Preceding intravenous thrombolysis facilitates endovascular mechanical recanalization in large*
834 *intracranial artery occlusion*. Int J Stroke, 2012. 7(1): p. 14-8.

835
836 Pienimäki, J.P., et al., *In-Hospital Intravenous Thrombolysis Offers No Benefit in Mechanical Thrombectomy in*
837 *Optimized Tertiary Stroke Center Setting*. Cardiovasc Intervent Radiol, 2021. 44(4): p. 580-586.

838
839 Platko, S., et al., *Intravenous thrombolysis prior to mechanical thrombectomy does not affect clinical or procedural*
840 *outcomes in patients with large vessel occlusion acute ischemic stroke*. J Clin Neurosci, 2022. 100: p. 120-123.

841
842 Purruicker, J.C., et al., *Efficacy and safety of bridging thrombolysis initiated before transfer in a drip-and-ship stroke*
843 *service*. Stroke Vasc Neurol, 2022. 7(1): p. 22-28.

844
845 Rai, A.T., et al., *Intravenous thrombolysis before endovascular therapy for large vessel strokes can lead to significantly*
846 *higher hospital costs without improving outcomes*. J Neurointerv Surg, 2018. 10(1): p. 17-21.

847
848 Regenhardt, R.W., et al., *'Drip-and-ship' intravenous thrombolysis and outcomes for large vessel occlusion thrombectomy*
849 *candidates in a hub-and-spoke telestroke model*. J Neurointerv Surg, 2022. 14(7): p. 650-653.

850
851 Reiff, T., et al., *Safety of Mechanical Thrombectomy with Combined Intravenous Thrombolysis in Stroke Treatment 4.5 to*
852 *9 Hours from Symptom Onset*. J Stroke Cerebrovasc Dis, 2020. 29(11): p. 105204.

- 854 Rossi, R., et al., *The administration of rtPA before mechanical thrombectomy in acute ischemic stroke patients is*
855 *associated with a significant reduction of the retrieved clot area but it does not influence revascularization outcome.* J
856 Thromb Thrombolysis, 2021. 51(2): p. 545-551.
- 857
- 858 Sakai, N., et al., *Safety, Pharmacokinetics and Pharmacodynamics of DS-1040, in Combination with Thrombectomy, in*
859 *Japanese Patients with Acute Ischemic Stroke.* Clin Drug Investig, 2022. 42(2): p. 137-149.
- 860
- 861 Sallustio, F., et al., *Effect of mechanical thrombectomy alone or in combination with intravenous thrombolysis for acute*
862 *ischemic stroke.* J Neurol, 2018. 265(12): p. 2875-2880.
- 863
- 864 Sarraj, A., et al., *Clinical and Neuroimaging Outcomes of Direct Thrombectomy vs Bridging Therapy in Large Vessel*
865 *Occlusion: Analysis of the SELECT Cohort Study.* Neurology, 2021. 96(23): p. e2839-e2853.
- 866
- 867 Smith, E.E., et al., *Outcomes After Endovascular Thrombectomy With or Without Alteplase in Routine Clinical Practice.*
868 JAMA Neurol, 2022. 79(8): p. 768-776.
- 869
- 870 Tajima, Y., et al., *Effectiveness of Low-Dose Intravenous Tissue Plasminogen Activator before Stent Retriever or*
871 *Aspiration Mechanical Thrombectomy.* J Vasc Interv Radiol, 2019. 30(2): p. 134-140.
- 872
- 873 Tong, X., et al., *Thrombectomy Versus Combined Thrombolysis and Thrombectomy in Patients With Acute Stroke: A*
874 *Matched-Control Study.* Stroke, 2021. 52(5): p. 1589-1600.
- 875
- 876 Tu, W.J., et al., *Endovascular thrombectomy or bridging therapy in minor ischemic stroke with large vessel occlusion.*
877 Thromb Res, 2022. 219: p. 150-154.
- 878
- 879 Wang, H., et al., *Direct endovascular treatment: an alternative for bridging therapy in anterior circulation large-vessel*
880 *occlusion stroke.* Eur J Neurol, 2017. 24(7): p. 935-943.
- 881
- 882 Weber, R., et al., *Comparison of outcome and interventional complication rate in patients with acute stroke treated with*
883 *mechanical thrombectomy with and without bridging thrombolysis.* J Neurointerv Surg, 2017. 9(3): p. 229-233.
- 884
- 885 Wee, C.K., et al., *Outcomes of Endovascular Thrombectomy with and without Thrombolysis for Acute Large Artery*
886 *Ischaemic Stroke at a Tertiary Stroke Centre.* Cerebrovasc Dis Extra, 2017. 7(2): p. 95-102.
- 887
- 888 Wei, D., et al., *Mobile Interventional Stroke Teams Lead to Faster Treatment Times for Thrombectomy in Large Vessel*
889 *Occlusion.* Stroke, 2017. 48(12): p. 3295-3300.
- 890
- 891 Ye, Z., et al., *Cost-effectiveness of endovascular thrombectomy with alteplase versus endovascular thrombectomy alone*
892 *for acute ischemic stroke secondary to large vessel occlusion.* CMAJ Open, 2023. 11(3): p. E443-e450.
- 893
- 894 Yi, H.J., J.H. Sung, and D.H. Lee, *Bridging Intravenous Thrombolysis Before Mechanical Thrombectomy for Large*
895 *Artery Occlusion May be Detrimental with Thrombus Fragmentation.* Curr Neurovasc Res, 2020. 17(1): p. 18-26.
- 896