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- physicians, nurses, and other health care professional and provider organizations;
- health plans, health systems, health care organizations, hospitals and integrated health care delivery systems;
- health care teaching institutions;
- health care information technology departments;
- medical specialty and professional societies;
- researchers;
- federal, state and local government health care policy makers and specialists; and
- employee benefit managers.

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Emergency Department Treatment Algorithm



Stroke Code Algorithm



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Foreword

Scope and Target Population

The scope of the following guideline is the 48 hours beginning when a patient age 18 years or older presents to a provider with symptoms of ischemic stroke or transient ischemic attack. For most stroke patients who are hospitalized, the guideline's temporal scope will expire before discharge. The guideline work group on Diagnosis and Initial Treatment of Ischemic Stroke recognizes that two time frames are critically important in the overall outcome, and fall outside the defined scope. They are prehospital care, and continuing care of stroke patients after 48 hours, which includes the development of a long-term secondary prevention strategy. While the group has not itself performed a systematic review of the primary evidence on these matters, we recommend the following guidelines from the American Heart Association/American Stroke Association.

A. Regarding prehospital care:

Acker, Joe E, et al. Implementation strategies for emergency medical services within stroke systems of care. A policy statement from the American Heart Association/American Stroke Association Expert Panel on Emergency Medical Services and the Stroke Council. *Stroke* 2007;116:3097-115.

- B. Regarding continuing care after the initial 48 hours and secondary prevention:
 - 1. Sacco RL, et al. Guidelines for prevention of stroke in patients with ischemic stroke or transient ischemic attack: a statement for healthcare professionals from the American Heart Association/ American Stroke Association Council on Stroke: co-sponsored by the Council on Cardiovascular Radiology and Intervention: the American Academy of Neurology affirms the value of this guideline. *Stroke* 2006;37:577-617.
 - 2. Adams RJ, et al. Update to the AHA/ASA recommendations for the prevention of stroke in patients with stroke and transient ischemic attack. *Stroke* 2008;39:1647-52.

Clinical Highlights and Recommendations

- Patients presenting with signs and symptoms of TIA should be evaluated for risk of immediate future events using the ABCD score. (*Annotation #23; Aim #1*)
- Patients who present in time to be candidates for treatment with intravenous tissue plasminogen activator (tPA) should be evaluated by a physician within 10 minutes, undergo a CT scan within 25 minutes of arrival in the ED, and have CT interpreted within 20 minutes of test completion. (Annotation #29; Aim #1)
- tPA, if given, should be administered within three hours (4.5 hours in selected patients; see Annotation #18, "Consider IV Tissue Plasminogen Activator [tPA]/See Stroke Code Algorithm") of stroke onset and less than 60 minutes of arrival at the ED. (*Annotations #29, 30, 33; Aim #3*)
- Patients presenting with stroke onset who are not candidates for intravenous tPA should promptly be given aspirin, after exclusion of hemorrhage on CT scan. (*Annotation #35; Aim #3*)
- Education regarding early stroke symptoms, risk factors, diagnostic procedures, and treatment options should be offered to the patient and family. This should be documented in the patient chart. (Annotation #31; Aim #6)

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- Medical management for prevention of complications within the initial 24-48 hours of diagnosis and initial treatment of ischemic stroke include: (*Annotation #38; Aim #5*)
 - continue appropriate blood pressure management;
 - continue to treat hyperthermia;
 - continue to treat hypo- or hyperglycemia;
 - continue IV fluids;
 - initiate deep vein thrombosis prophylaxis;
 - perform swallow evaluation;
 - initiate early rehabilitation; and
 - perform nutritional status assessment.

Priority Aims

- 1. Increase the percentage of patients presenting within three hours of stroke onset who are evaluated within 10 minutes of arriving in the ED. (*Annotation #29*)
- 2. Increase the percentage of patients presenting with TIA symptoms within 24 hours at high risk for stroke who are admitted to the hospital. (*Annotation #11*)
- 3. Increase the percentage of patients receiving appropriate thrombolytic and antithrombotic therapy for ischemic stroke (use of tPA and aspirin). (*Annotations #29, 30, 33, 35*)
- 4. Increase the percentage of non-tPA recipients who have hypertension appropriately managed in the first 48 hours of hospitalization or until neurologically stable. (*Annotation #37*)
- 5. Increase the percentage of patients who receive appropriate medical management for prevention of complications within the initial 24-48 hours of diagnosis: (*Annotations #31, 38*)
 - Continue to treat hypoglycemia and hyperglycemia (Annotation #31)
 - Continue to treat hyperthermia (Annotation #31)
 - Continue IV fluids (Annotation #31)
 - Continue to treat hypoxia (Annotation #31)
 - Initiate deep vein thrombosis prophylaxis (Annotation #38)
 - Perform swallow evaluation (Annotation #38)
 - Initiate early rehabilitation (early mobilization) (Annotation #38)
 - Perform nutritional status assessment (Annotation #38)
- 6. Improve patient and family education of patients with ischemic stroke in both the ED and the admitting hospital unit. (*Annotation #31*)

Key Implementation Recommendations

The following system changes were identified by the guideline work group as key strategies for health care systems to incorporate in support of the implementation of this guideline.

- 1. Hospitals should consider developing and implementing critical pathways, standing orders and a stroke process to accomplish rapid evaluation and treatment. The process should expedite the evaluation and treatment of patients who are candidates for intravenous tPA and assure uniform, guideline-driven care for all patients with respect to issues like:
 - ongoing antithrombotic therapy,
 - management of blood pressure,
 - early mobilization, and
 - use of appropriate antiembolism treatment in the paralyzed patient.
- 2. A process should be in place for the patient and family that will rapidly orient them to the suspected diagnosis, ED process, tests to be preformed, tPA treatment and its risks, and other treatment measures to be considered. This could include both caregiver face-to-face interactions with the patient and family, as well as teaching tools in written form.

System Improvement

There is evidence that benchmarking can guide and drive quality improvement. Using essentially the same quality indicators as The Joint Commission on the Accreditation of Health Care Organization (TJC) and ICSI, programs like the American Heart Association's Get With The Guidelines-Stroke (*LaBresh*, 2008 [C]; *Schwamm*, 2009b [B]) and the Paul Coverdell National Acute Stroke Registry (*Stoeckle-Roberts*, 2006 [C]) have been shown to improve the quality of stroke care.

The Joint Commission (TJC) Primary Stroke Center Certification

TJC offers certification as Primary Stroke Centers to hospitals that meet specific qualifications. The emphasis of the process is on the early recognition and management of stroke and the scope of accreditation includes integrated efforts in public awareness, emergency medical services, emergency room and hospitalization (*Alberts, 2000 [R]*). The link is: http://www.jointcommission.org/CertificationPrograms/PrimaryStrokeCenters. Beginning in October 2009 all TJC accredited hospitals will have to submit the eight National Quality Forum-endorsed stroke consensus measures. The Centers for Medicare and Medicaid Services (CMS) is also considering the reporting of stroke measures and in the near future the draft Inpatient Prospective Payment System (IPPS) Rule will be released. IPPS is the venue in which CMS communicates with hospitals and physicians about their future measurement reporting.

Among the requirements for TJC certification as a Primary Stroke Center is ongoing process improvement guided by data and benchmarking. The quality indicators chosen by TJC overlap with those developed by the ICSI Diagnosis and Initial Treatment of Ischemic Stroke guideline work group. The TJC quality indicators are:

- 1. Deep Vein Thrombosis (DVT) Prophylaxis*
- 2. Discharged on Antithrombotics*
- 3. Patients with Atrial Fibrillation Receiving Anticoagulation Therapy*
- 4. Thrombolytic Therapy Administered (in eligible patients)
- 5. Antithrombotic Therapy by End of Hospital Day Two

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- 6. Discharged on Cholesterol Reducing Medication
- 7. Dysphagia Screening
- 8. Stroke Education
- 9. Smoking Cessation/Advice Counseling
- 10. Assessed for Rehabilitation
- * Initial standard stroke measure set.

Measures 1, 4, 5, 7 and 8 are similar to or identical to those measures listed in this document and within the scope of the guideline.

Related ICSI Scientific Documents

Guidelines

- Antithrombotic Therapy Supplement
- Hypertension Diagnosis and Treatment
- Palliative Care
- Venous Thromboembolism Diagnosis and Treatment
- Venous Thromboembolism Prophylaxis

Order Sets

- Admission for Ischemic Stroke for Patients Not Receiving tPA
- Admission for Ischemic Stroke for Patients Receiving tPA

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ICSI has adopted a policy of transparency, disclosing potential conflict and competing interests of all individuals who participate in the development, revision and approval of ICSI documents (guidelines, order sets and protocols). This applies to all work groups (guidelines, order sets and protocols) and committees (Committee on Evidence-Based Practice, Cardiovascular Steering Committee, Women's Health Steering Committee, Preventive & Health Maintenance Steering Committee and Respiratory Steering Committee).

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No work group members have potential conflicts of interest to disclose.

Foreword

Introduction to ICSI Document Development

This document was developed and/or revised by a multidisciplinary work group utilizing a defined process for literature search and review, document development and revision, as well as obtaining input from and responding to ICSI members.

For a description of ICSI's development and revision process, please see the Development and Revision Process for Guidelines, Order Sets and Protocols at http://www.icsi.org.

Evidence Grading System

B.

A. Primary Reports of New Data Collection:

Class A:	Randomized, controlled trial
Class B:	Cohort study
Class C:	Non-randomized trial with concurrent or historical controls Case-control study Study of sensitivity and specificity of a diagnostic test Population-based descriptive study
Class D:	Cross-sectional study Case series Case report
Reports that	Synthesize or Reflect upon Collections of Primary Reports:
Class M:	Meta-analysis Systematic review

	Systematic review
	Decision analysis
	Cost-effectiveness analysis
Class R:	Consensus statement
	Consensus report
	Narrative review

Class X: Medical opinion

Citations are listed in the guideline utilizing the format of (*Author, YYYY [report class]*). A full explanation of ICSI's Evidence Grading System can be found at http://www.icsi.org.

Algorithm Annotations

Screening (Ambulatory) Algorithm Annotations

1. Initial Contact with Patient with Complaint of Neurological Symptoms

This contact may occur with one of several medical system personnel, including primary care physicians, other medical specialty physicians, emergency medical services, nursing staff in a clinic or urgent care setting or even non-medical triage personnel. This does not refer to the ED evaluation. This contact may be by phone or in person. Potential staff contacts should be educated in the importance of stroke symptom recognition and the appropriate triage measures that should be taken.

2. Immediate Screening for Ischemic Stroke

This should include detail as to the location, severity, duration of symptoms and any aggravating or relieving factors. Symptoms that are commonly associated with ischemic stroke or transient ischemic attack (TIA) include:*

- sudden numbness or weakness of the face, arm or leg especially on one side of the body;
- sudden mental confusion, trouble speaking or understanding;
- sudden trouble walking, dizziness, loss of balance or coordination;
- sudden trouble seeing in one or both eyes;
- sudden severe headache with no known cause.

* List from American Stroke Association for public education

The American Stroke Association, American Academy of Neurology and American College of Emergency Physicians have recently launched a public awareness campaign entitled "Give Me 5" emphasizing that stroke typically presents as problems of walking, talking, reaching, seeing and/or feeling.

Symptoms of ischemic stroke can also, of course, be represented in atypical ways.

Clinical diagnoses with neurologic symptoms that may imitate or superficially resemble ischemic stroke or TIA include the following (*Adams*, 2007 [*R*]):

• Migraine

Neurologic symptoms experienced with migraine tend to have a more gradual onset and slower development. However, the two problems may be indistinguishable.

Seizures

Although seizures typically consist of a "positive" phenomenon (jerking of a limb) rather than loss of neurologic function (weakness or paralysis of a limb), symptoms and signs during the ictus or in the postictal state may be similar to ischemic stroke (e.g., confusion or speech arrest during the ictus as in complex partial seizure, postictal confusion, postictal paralysis, and other sensory or visual phenomenon).

• Syncope

• Transient global amnesia

This is characterized by a sudden onset antegrade and retrograde memory disturbance without other neurologic symptoms. If the patient experiences symptoms of transient global amnesia, it would be inappropriate to assume the diagnosis without a complete neurologic assessment exam.

• Peripheral nerve disorders

Mononeuropathy and radiculopathy can often be distinguished from ischemic stroke by the anatomic distribution of the symptoms, and in the case of radiculopathy, by associated painful symptoms. Bell's palsy, vestibular neuritis and extraocular muscle imbalance due to cranial neuropathy may also imitate ischemic stroke; a complete history and neurologic examination are required to accurately differentiate from ischemic stroke.

• Intracranial hemorrhage

• Other intracranial masses, e.g., tumor, abscess (often differentiated by CT)

The mode of onset and early course tend to be more gradual in development but mimickry of stroke is not uncommon.

• Psychogenic presentation

Psychogenic conditions or reactions such as anxiety, panic disorder, or conversion reactions may need to be considered in some cases.

• Metabolic disorders

Hypoglycemia is the most common metabolic disorder producing neurologic symptoms that imitate stroke. A patient with known diabetes or liver disease should be screened for hypoglycemia.

This discussion is not meant to be detailed guide to discerning between ischemic stroke and other diagnoses. If there is any uncertainty as to symptom causation, the evaluation should proceed as though ischemic stroke or TIA is confirmed so as not to delay appropriate emergency treatment if needed.

4. Refer to Emergency Department (ED) or Physician's Office as Appropriate for Other Conditions

Some of the diagnoses outlined in Annotation #2, "Immedidate Screening for Ischemic Stroke," may warrant ED evaluation because of the urgency of the problem itself or the inability of the contact person to distinguish the other condition from ischemic stroke (*Adams*, 2007 [*R*]). In these uncertain cases, the contact person should continue on to box #5 in the Screening (Ambulatory) algorithm. See Appendix B, "Acute Stroke Care Networks."

5. Symptoms Present Now?

This annotation refers to ongoing symptoms suggestive of cerebral ischemia. If ischemic symptoms have resolved and were present for less than 24 hours, this is clinically defined as a TIA.

Recently, a new definition of TIA has been proposed: TIA is a transient episode of neurologic dysfunction caused by focal brain, spinal cord or retinal ischemia without acute infarction.

Applying the definition requires imaging. The pre-imaging syndrome has been designated "acute neuro-vascular syndrome."

The work group will use "clinical TIA" in this document in lieu of acute neurovascular syndrome (*Easton*, 2009 [*R*]).

6. Possible Ischemic Stroke – Symptoms Onset within 24 Hours? Key Point:

• The onset of symptoms should be defined as the last time the patient was known to be normal or at previous prestroke baseline.

If the symptoms resolve completely and then recur, for the purposes of determining whether thrombolysis can be considered for stroke, the time of onset would be the last time the patient was normal (just prior to the onset of the second set of symptoms.) Patients may be unable to give this information if they have an aphasia or mental confusion. Family members or other witnesses may need to give this information. If the patient was sleeping and awakened with the problem, the time of onset would be the moment the patient was last known to be normal just before falling asleep.

7. Ischemic Stroke Symptoms Present for > 24 Hours/Symptoms Mild and Stable

Patients with stable mild deficits present longer than 24 hours may be transported to the ED for evaluation and treatment by means other than 911. As a rule, they should be admitted to the hospital to assure thorough and expeditious evaluation and treatment. Outpatient evaluation and treatment is an acceptable alternative if it can be done as quickly as it could be done inpatient and if all goals of inpatient assessment (diagnosis of mechanism, initiation of appropriate secondary prevention, prevention of complications, early assessment for and deployment of rehabilitative services) can be successfully addressed.

9. Clinical TIA – Symptoms within Two Hours?

Patients presenting with history of clinical TIA may have neurological deficits of which they are not aware. To avoid missing the thrombolytic treatment window, patients with clinical TIAs presenting within two hours of symptom onset should be triaged like patients with stroke, i.e., call 911 (*Adams*, 2007 [*R*]).

11. Transport to Emergency Department

Patients should be taken to the ED expeditiously; use of 911 Emergency is at the provider's discretion. Alternatively, if such a program were available, the patient may be assessed in a specialized clinic or other program in which the evaluation can be carried out as quickly and treatment initiated as definitively as if the patient were admitted to the hospital. This work group otherwise recommends that the physician strongly consider hospitalization for clinical TIA patients who appear within 24 hours of the event to expedite workup and possibly administer tPA if the deficit recurs.

13. Rapid Outpatient Evaluation or Admit to Hospital

Patients should receive rapid outpatient evaluation (TIA clinic or other program) or be admitted to the hospital as soon as possible (*Johnston*, 2006 [R]). In addition to a risk assessment for stroke, the patient should be diagnostically evaluated for:

- brain imaging: MRI (preferred) or CT
- vascular imaging: ultrasound (if symptoms suggest ischemia in the carotid distribution), computed tomography angiography (CTA) or magnetic resonance angiography (MRA)

- cardiac rhythm assessment; (monitor rhythm if admitted)
- echocardiogram (if suspect cardioembolic source)

Emergency Department Treatment Algorithm Annotations

18. Consider IV Tissue Plasminogen Activator (tPA)/See Stroke Code Algorithm

Key Points:

- Treatment with IV tPA should begin within three hours (180 minutes) (270 minutes in selected patients; see Annotation #18, "Consider IV Tissue Plasminogen Activator [tPA]/See Stroke Code Algorithm") of symptom onset.
- Patients with persisting symptoms presenting to the ED within 150 minutes of symptom onset should be evaluated rapidly for treatment with IV tPA.
- Occasionally, patients may be able to receive IV tPA even if they present later than 150 minutes, (240 minutes in selected patients) (4.5 hours in selected patients; see Annotation #18, "Consider IV Tissue Plasminogen Activator [tPA]/See Stroke Code Algorithm") if their workup, such as laboratory evaluation, has been completed and they have IV access in place.
- Intra-arterial thrombolysis may be an option for treatment of selected patients who have a major stroke with symptoms onset less than six hours previous due to occlusion of the MCA or basilar artery or who are not otherwise candidates for intravenous tPA if the patient can be treated in a timely manner at an experienced stroke center with immediate access to cerebral angiography and qualified interventionalists. (A protocol for intra-arterial thrombolysis will be institutional specific and is out of the scope of this guideline.)

Patients presenting to the ED soon after the onset of symptoms may be candidates for treatment with intravenous (IV) tissue plasminogen activator (tPA) and will therefore require a rapid evaluation and treatment initiation (*Albers*, 2004 [*R*]). (See Appendix B, "Acute Stroke Care Networks.") Although the time window from onset of symptoms to treatment can be up to 180 minutes (to 270 minutes in selected patients) (4.5 hours in selected patients; see Annotation #18, "Consider IV Tissue Plasminogen Activator [tPA]/See Stroke Code Algorithm"), the evaluation in the ED will require at least 30 minutes in most cases (CT scan of head, laboratory tests performed and results have returned, IV access obtained, and neurological exam and history) (*Adams*, 2007 [*R*]). We have therefore chosen 150 minutes (240 minutes in selected patients; see Annotation #18, "Consider IV Tissue Plasminogen Activator [tPA]/See Stroke Code Algorithm") as a practical cutoff time for this triage decision.

There are important exceptions to this time limitation guideline for triage of the patients into the "stroke code" process. In certain instances, the time required for evaluation may be shorter, and "stroke code" may be feasible for patients presenting as late as 165 or 170 (255-260 minutes in selected patients; see Annotation #18, "Consider IV Tissue Plasminogen Activator [tPA]/See Stroke Code Algorithm") minutes after onset. One example would be the patient who is already in the hospital and has undergone the appropriate laboratory evaluation, has an IV access in place, and much of the history is already known. In that case, a

Algorithm Annotations

brief neurologic exam and rapid evaluation with CT may be the only items required prior to treatment and could theoretically be performed in 10-15 minutes.

Initial Thrombolytic Trials

Thrombolytic therapy for ischemic stroke using intravenous tissue plasminogen activator (tPA) has now been tested in several large, randomized, placebo-controlled clinical trials. The National Institute of Neurological Disorders and Stroke (NINDS) stroke trial (actually a combination of two trials, one with 24-hour and the other with a 90-day outcome measure) compared placebo with tPA at a dose of 0.9 mg/kg given within three hours of symptom onset in 624 patients (*National Institute of Neurological Disorders and Stroke tPA Study Group, 1995 [A]*). The time of stroke onset was strictly defined, blood pressure was maintained within a specified range, and other anticoagulant and antiplatelet drugs were avoided within the first 24 hours after treatment. The predetermined threshold for a clinically important difference at 24 hours was not met. However, at three months and one year (*Kwiatkowski, 1999 [A]*), there were significantly increased percentage of patients (11%-13%) with favorable outcomes in the tPA group compared with controls. Results were consistent across all four of the standard outcome measures that were assessed. Treatment with tPA resulted in a significantly increased risk of symptomatic intracerebral hemorrhage (6.4% tPA treated vs. 0.6% in placebo group, p < 0.001). Mortality was lower at three months in those treated with tPA (17% vs. 21% in placebo treated), but this did not reach statistical significance. On the basis of the favorable results from these combined trials, the FDA approved tPA for use in the United States in 1996.

The European Cooperative Acute Stroke Study (ECASS) performed concurrent with the NINDS trial also compared intravenous tPA to placebo in a randomized, placebo-controlled trial in 620 patients. However, the study design was different in a number of respects, including longer time window to treatment (six hours), higher dose of tPA (1.1 mg/kg), and lack of strict blood pressure control (*Hacke, 1995 [A]*). Over 80% of the patients were treated between three and six hours after symptoms began. The intention-to-treat analysis did not demonstrate significant improvement in the primary outcome measure (combination of Barthel Index and modified Rankin Scale at three months). However, there was a high rate of major protocol violation (109 patients). In a secondary analysis including only the target population, there was a significant difference in favor of tPA treatment, but the margin was of questionable clinical significance.

In 1998, a third large tPA study was completed. ECASS II had a nearly identical protocol to the original ECASS study except that the tPA dose was lowered to 0.9 mg/kg (*Hacke*, 1998 [A]). A total of 800 patients were enrolled, and approximately 80% were treated three to six hours after stroke onset. The results indicated that there was no significant difference in neurological function between tPA and placebo patients. There was a trend in favor of treatment that did not reach statistical significance.

The Thrombolytic Therapy in Acute Ischemic Stroke Study sponsored by Genentech also used a six-hour time window, but was similar in other respects to the NINDS tPA trial (*Clark, 2000 [A]*). Only 15% of patients were enrolled within three hours. Although a significantly higher percentage of tPA treated patients showed early improvement at 24 hours measured with the National Institutes of Health Stroke Scale (NIHSS), these findings were reversed at one month with the placebo group having a statistically higher percentage of patients showing improvement on this scale. Symptomatic intracerebral hemorrhage was significantly increased compared to placebo (11% vs. 0%), with the greatest risk of hemorrhage in patients treated between five and six hours.

Three similar large-scale clinical trials utilizing intravenous streptokinase compared to placebo in randomized, placebo-controlled trial design were performed concurrent with the NINDS tPA and ECASS studies (*Multicenter Acute Stroke Trial (MAST), 1996 [A]; Multicenter Acute Stroke Trial (MAST), 1995 [A]).* All three studies were terminated before completion because of safety concerns with excessive rates of intracerebral hemorrhagic complications and higher mortality in the treated groups. These early studies support a limited use of intravenous tPA at a dose of 0.9 mg/kg with appropriate precautions and treatment beginning within 3 hours of symptom onset (*Ingall, 2004 [A]*). Following FDA approval of tPA for stroke, several reports of community experience with this treatment appeared in the literature (*Tanne, 1999 [D]; Buchan, 2000 [A]; Hanson, 2000 [D]; Charipar, 2008 [D]; Chiu, 1998 [D]; Wang, 2000a [D]*). Although clinical results for the most part have been concordant with those in the NINDS study, two of these reports document an increase in intracerebral hemorrhagic complications when patients are treated outside the NINDS protocol further supporting the importance of following the NINDS tPA study protocol.

The significance of early time to treatment was further emphasized by a secondary analysis of the NINDS tPA study population showing gradual decline in measured efficacy even within the three-hour time window (*Marler, 2000 [C]*). American Heart Association (AHA) consensus guidelines for the use of tPA have been published, and treating physicians are encouraged to evaluate patients for this treatment and initiate treatment with urgency (*Adams, 2007 [D]*). Another analysis using pooled data from all the early tPA trials suggested efficacy might be achieved even after the three-hour window (*ATLANTIS, ECASS, and NINDS tPA Study Group Investigators, 2004 [M]*).

Recent Thrombolytic Trials

ECASS III (*Hacke*, 2008 [A]) was a multicenter trial conducted in Europe to evaluate intravenous recombinant tissue plasminogen activator (tPA or alteplase) versus placebo administered between 3 and 4.5 hours after onset of ischemic stroke symptoms. A total of 821 patients were enrolled, nearly one-third more than in NINDS, the U.S. trial that led to the adoption of tPA as the gold standard for the treatment of acute ischemic stroke within the first three hours after symptom onset. Both arms were acceptably well balanced, although initial stroke severity and previous history of stroke were greater in the placebo group.

Treatment with tPA was associated with a significant improvement in the rate of favorable functional outcome as defined by a modified Rankin of 0 or 1 (i.e., mild or no deficits). These rates were 52% for the tPA group and 45% for the placebo group (OR 1.34; 95% CI 1.02-1.76) in the intention-to-treat analysis. The benefit was more pronounced when only patients treated with tPA according to the protocol were analyzed (OR 1.47; 95% CI 1.10-1.97) and it was also present when a global outcome measure also including the NIHSS, the Barthel index and the Glasgow Outcome scale was used as the endpoint. Overall, the chances to regain full independence were 28% higher among patients treated with tPA, and 14 patients had to be treated for one additional patient to achieve a favorable outcome.

Mortality was not significantly different between the groups, but slightly higher in the placebo arm. The rate of symptomatic intracranial hemorrhage as defined by the NINDS criteria was 7.9% in the tPA group (versus 6.4% in the NINDS trial). Furthermore, only 2.4% of patients were thought to have worsened because of intracranial hemorrhage in the active treatment group.

The positive results of ECASS III are similar to these of a pooled analysis of previous tPA trials (*ATLANTIS*, 2004 [M]). Yet, some have expressed skepticism, pointing out the discrepancy with previous negative results of ATLANTIS, a U.S. trial evaluating intravenous tPA versus placebo within five hours of stroke onset (*Clark, 1999* [A]). ECASS III was different from the NINDS and ATLANTIS trials. The main difference with NINDS was the lower stroke severity of enrolled patients, which explains the much higher rate of complete or near complete recovery at 90 days in ECASS III (modified Rankin 0-1; 52% in ECASS III vs. 39% in NINDS in the tPA arm; and 45% in ECASS III vs. 26% in NINDS in the placebo arm). The main difference with ATLANTIS was the shorter time from symptom onset to enrollment in ECASS III (76% treated with tPA after 4 hours in ATLANTIS vs. 37% in ECASS III).

While attention to detail and critical analysis of the data are essential before applying trial results to change clinical practice, the results of ECASS III provide high-quality evidence supporting the use of intravenous tPA in acute stroke patients up to 4.5 hours after symptom onset. However, extending the therapeutic window for intravenous thrombolysis should not allow for complacency. It is very clear that thrombolytic treatment should be started as soon as possible. Thrombolysis is most effective when initiated within 90 minutes of

symptom onset (*ATLANTIS*, 2004 [*M*]), and any delay decreases the benefit. Also, it will be prudent to be selective when extending the window to 4.5 hours. The criteria used for patient selection in ECASS III should be replicated when considering thrombolytic treatment between 3 and 4.5 hours after symptom onset in clinical practice. Age greater than 80, combined history of previous stroke and diabetes mellitus, use of anticoagulation regardless of INR and very high initial stroke severity (NIHSS > 25 or radiological evidence of infarction involving more than one-third of the middle cerebral artery territory) were exclusion criteria in ECASS III. Therefore, intravenous tPA in patients with these characteristics should continue to be used only within 3 hours of symptom onset. Finally, obtaining written informed consent before administration of intravenous tPA between 3 and 4.5 hours of symptom onset may be advisable. NOTE: Extending the treatment window for tPA to 4.5 hours has not been approved by the FDA.

(del Zoppo, 2009 [R])

Acute imaging with diffusion-perfusion MRI and other modalities is being evaluated to expand the therapeutic window for intravenous thrombolysis (*Davis*, 2008 [A]; Furlan, 2006; [A] Hacke, 2005 [A]). See Annotation #30, under "Evaluation (Should Occur Concurrently with Intervention) – Emerging Technologies," for more discussion.

21. Emergency Department Diagnostic Evaluation

Patients with a history of clinical TIA should be evaluated promptly (*Adams*, 2007 [*R*]). The following diagnostic evaluations should typically be performed (*Albers*, 2002 [*R*]; *Calvet*, 2007 [*D*]; *Coutts*, 2005 [*C*]; *Johnston*, 2002 [*R*]). The speed and venue of the assessment described below will depend on the currency of the symptoms and the physician's assessment of risk of early recurrence of clinical TIA or the development of stroke. The work group recommends that patients presenting less than 24 hours since initial clinical TIA with high risk symptoms (see Annotation #23, "High Risk for Stroke?") generally not leave the ED until the following are completed or scheduled within the next few hours on an inpatient basis.

- Laboratory tests
 - Complete blood count
 - Electrolytes (sodium, potassium, chloride, CO₂), BUN, creatinine, glucose
 - Prothrombin time / international normalized ratio (INR)
 - Activated partial thromboplastin time (aPTT)
 - Cardiac biomarkers (troponin)
- Electrocardiogram
- Brain and vascular imaging (see below)
 - Magnetic resonance imaging MRI (preferred)/magnetic resonance angiography (MRA)
 - Computed tomography (CT)/computed tomography angiography (CTA)
 - CT/carotid ultrasound, if symptoms referable to carotid distribution

Brain Imaging

If the patient is not having symptoms at the time of presentation, a diffusion-weighted MRI (DW-MRI) is preferred, if available. Restricted diffusion in the setting of a clinical transient ischemic attack identifies higher risk of stroke. At this time, an MRA of the carotids and intracranial artery can be performed.

If MRA is not available, a CT of the head would be indicated and, if feasible, a CTA of the head and neck can also be performed.

(Boulanger, 2007 [B]; Douglas, 2003 [D])

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23. High Risk for Stroke?

Key Points:

- Risk of stroke is greatest in the immediate aftermath of clinical TIA or minor stroke.
- Features of presentation define those at highest risk.
- Hospitalization should be strongly considered for those at highest risk.

The major issues in dealing with clinical TIA patients is making the best decisions about the speed of workup, the appropriate evaluation to guide preventive therapy, and the most efficacious therapies to avert stroke. To make the best decisions the provider must know what the early risk of stroke is for the given patient, whether speed of workup and treatment matter, and, if so, what treatments should be deployed. Information about these points is just becoming available. That a clinical TIA is a risk factor for stroke is not new news. The traditional wisdom is that a patient has a 30%-40% risk of having a stroke in the five years following a clinical TIA. The more salient question is about the short-term risk.

Several studies have identified factors in patients presenting with clinical transient ischemic attacks (TIAs) that predict progression to ischemic stroke. Older studies using a clinical definition of TIA examined factors predisposing to ischemic stroke within a time frame of months or years after the clinical TIA, but provide limited information as to acute risk. Examples of risk factors identified in these older studies include advanced age, presenting with more than four clinical TIAs in the two weeks prior to the index clinical TIA, and the comorbidities of hypertension, myocardial infarction, cardiac arrhythmia, and diabetes mellitus (*Dennis*, 1990 [B]; Friday, 1997 [B]; Hankey, 1992 [B]; Kernan, 1991 [B]; Streifler, 1995 [D]). The presentation of transient monocular blindness or amaurosis fugax generally confers a more benign prognosis (*Dennis*, 1989 [D]; Evans, 1994 [B]; Wilterdink, 1992 [M]). Brown, et al., proposed an algorithm for triage and evaluation of patients with clinical TIA and minor ischemic stroke, citing these data to estimate specific risk and make timing decisions (*Brown*, 1994 [R]; Flemming, 2004 [R]).

Data from recent studies also using a clincial definition of TIA are more relevant to the issue at hand in the ED, i.e., the risk over the next few days and weeks. A cohort study described the early (one-week and three-month) risk of ischemic stroke, cardiovascular events, and death in 1,707 patients presenting with a diagnosis of clinical TIA to emergency departments (ED) within the Kaiser-Permanente system in northern California (*Johnston*, 2000 [B]). Fifteen percent of the patients were admitted for further monitoring and the rest were discharged from the emergency department. The risk of stroke or admission for other cardiovascular events (myocardial infarction, unstable angina, cardiac arrhythmia, congestive heart failure) were reported as follows:

Event	7 days	3 months
Stroke	6%*	10.5%
TIA		13%
CV events		2.7%
Death		2.6%

* Over half occurred within two days.

Taken in total, 26.2% of patients returned to the hospital within three months with some other cerebrovascular or cardiovascular event.

In multivariate analysis, five factors independently predicted higher risk:

- Age greater than 60 years
- Diabetes mellitus
- TIA lasting longer than 10 minutes

- TIA including weakness as a symptom
- TIA including abnormal speech as a symptom

The features comprise a risk stratification scheme known as the "California Score."

The same issue of early risk was examined in the population-based Oxford Vascular Study (*Coull*, 2004 [B]). The short-term fate of 87 consecutive Oxfordshire residents with clinical TIA or minor stroke was examined. Risk of stroke was 8% at one week, 11.5% at one month and 17.3% at three months. The risks were slightly higher after minor stroke (11.5%, 15% and 18.5%, respectively) at the three time intervals.

Yet another study examined patients with carotid stenosis randomized to medical therapy after first TIA in the North American Symptomatic Carotid Endarterectomy Trial. The study found stroke risk of 5.5% at two days and 20.1% at 90 days following the qualifying TIA, emphasizing that early risk is substantial after TIA in this setting (*Eliasziw*, 2004 [B]). The study also showed that benefit of carotid endarterectomy in such patients drops significantly as time to the procedure exceeds two weeks from the ischemic event, arguing that speed of assessment and treatment are all-important (*Rothwell*, 2004 [M]).

Giles, in a meta-analysis of studies of stroke risk on day two and seven post-TIA, reported that variability in reports of rates of stroke may relate to the setting in which the patients are seen for evaluation and whether treatments are offered (*Giles*, 2007 [M]).

From these studies it is clear that in general TIA represents a potent short-term risk factor for stroke. From the Kaiser Permanente study, it is also suggested that risk is heterogeneous, i.e., some patients are at higher risk than others. If true, it might then be possible to identify those at highest short-term risk prospectively and reliably. They could be triaged to an expedited management track.

Confirmation information has now been accumulated. Analysis of the Oxfordshire population-based sample of TIA episodes (n=209) yielded an "ABCD" score identifying those at high risk of stroke (*Rothwell*, 2005 [B]).

Table 1.

A – for age	Over the age of 60 years	1 point
B – for blood pressure	A systolic greater than 140 mmHg or diastolic greater than 90 mmHg	1 point
C – for clinical features	Unilateral weakness Speech disturbance without weakness Other clinical features	2 points 1 point 0 points
D – for duration of symptoms	Symptoms lasting greater than 60 minutes Symptoms lasting 10-59 minutes Symptoms lasting less than 10 minutes	2 points 1 point 0 points

The elements of the scale from this derivation sample are:

(Rothwell, 2005 [B])

The ABCD score was subsequently validated in a second population-based sample of TIA episodes (n=190). The seven-day risks of stroke in the combined derivation and validation samples (n=299) were:

0-4 points (73% of combined samples): 0.4% (95% CI 0-1.1%)

5 points (18% of combined samples): 12.1% (4.2%-20.0%)

6 points (9% of combined samples): 31.4% (16.0%-46.8%)

Note the similarity of the ABCD score features, derived and validated in Great Britain, to those of the California Score described above. Next challenge was to demonstrate the generalizability of these approaches, i.e., to show the reliability of the prediction models in all patient groups and settings.

The ABCD scheme was applied in additional cohorts. One (*Cucchiara*, 2006 [C]) found the scheme not as sensitive for high risk as originally reported, but the study used a different outcome set than the original report. The other (*Tsivgoulis*, 2006 [C]) found the scheme very reliable in identifying high-risk patients. Both cohorts were already-hospitalized patients under care of neurologists. It might be argued that a more relevant setting to study the validity of the scheme would be in a community-based sample of patients seen by non-neurologists. Finally, retrospective analysis (*Bray*, 2007 [C]) concluded that the ABCD score was highly predictive in identifying patients with TIA at a high short-term risk of stroke.

Recently, the group from Kaiser Permanente (California Score) and Oxford (ABCD Score) together, validated the two similar prognostic scores in four independent groups of patients and generated a new unified score (the ABCD² Score) to predict the risk of stroke in the two days following a TIA (*Johnston*, 2007 [C]). This new score was derived and validated in patients seen in emergency departments and outpatient clinics and is a more accurate predictor than either of the two previous scores (California Score and ABCD Score) in the derivation and validation groups. This score also predicted the risk of stroke within two days, which is more useful in the outpatient setting. Data from the validation groups included 4,799 patients.

Table 2.	
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A – for age	60 years or older	1 point
B – for blood pressure	a systolic 140 mmHg or greater or diastolic of 90 mmHg or greater	1 point
C – for clinical features	Unilateral weakness Speech disturbance without weakness	2 points 1 point
D – for duration of symptoms	Symptoms lasting greater than 60 minutes Symptoms lasting 10-59 minutes Symptoms lasting less than 10 minutes	2 points 1 point 0 points
D – Diabetes		1 point

Based on these results, the authors suggest admitting patients who present with a TIA and have a ABCD² score of 4 or greater. Risk of stroke at two days:

Low risk (0-3 points): 1.0%

Moderate risk (4-5 points): 4.1%

High risk (6-7 points): 8.1%

These reports highlighted the frequent early occurrence of stroke and other cardiovascular events and the validity of risk stratification schemes. It has not been clear whether hospitalization or expedited outpatient management would mitigate high risk. Very recently, it has been shown that deployment of streamlined systems that address TIAs very quickly (e.g., within 24-48 hours) with definitive diagnostic testing and initiation of secondary prevention are associated with reducing the rate of early stroke (EXPRESS, SOS-TIA). These studies used historic cohorts as controls, randomization of rapid vs. slow assessment being ethically impossible. The EXPRESS Trial (*Rothwell*, 2007 [C]), for example, compared stroke rate at 90 days in TIA patients treated in an expedited process with that of an historical control group in the same medical system. The expedited process reduced time from TIA to initial assessment by stroke specialists from 3 days to

less than 1 day and the time to initiation of secondary prevention from 20 days to 1 day. The 90-day risk fell from 10.3% to 2.1%. The reported systems have been expedited outpatient assessments (EXPRESS, SOS-TIA) or in hospital protocols (FASTER) (*Kennedy*, 2007 [A]; Lavallee, 2007 [D]; Rothwell, 2007 [C]). Interestingly, these studies did not use stratification to select patients at higher risk. Even so, the studies showed value in expedited evaluation. Not clear at this point is what role stratification should play. It is also not clear what interventions made the apparent difference.

What these data show is that not only are TIAs dangerous and amenable to risk stratification, but early medical treatment reduces risk. Based on what is known and acknowledging the continuing areas of uncertainty, the work group recommends that patients seen within 24-48 hours of initial TIA be admitted to hospital or to a program of expedited outpatient assessment. The factors outlined above that predict high risk of recurrence might influence decision-making in this patient group. Also, detection of ischemic infarct by diffusion-weighted MRI would confer risk similar to an ABCD² score of 4 or greater. Although the data available cannot define an appropriate triage decision for all patients, this information should serve as a guide for rational triage of the patient with TIA. Certain diagnostic entities, if suspected, may require hospitalization for specific management, even with presentation later than 24-48 hours from TIA occurrence or lower ABCD² score (e.g., carotid or vertebral artery dissection, carotid stenosis, specific coagulopathy or arteriopathy, cerebral venous thrombosis). Not settled is whether the assessment of those at low risk can be safely pursued at a more leisurely pace or foregone altogether. At present, the work group is not prepared to recommend that patients be selected for hospitalization based solely on the ABCD² scheme. It recognizes that it may be being used in that way in some hospitals in the region and encourages that the effectiveness of the approach be monitored in those hospitals.

In summary, the work group recommends consideration of hospitalization for patients with first TIA within the past 24-48 hours to facilitate early deployment of lytic therapy, if necessary, and to expedite institution of definitive secondary prevention. For others, the risk stratification data described above might also justify hospitalization rather than expedited ambulatory management. Whatever the strategy, speed is key. Patients managed in the outpatient setting should be fully educated about the need to return immediately if symptoms recur, to allow use of lytic therapy.

(Goldstein, 2006; [R]; Johnston, 2006 [R]; Purroy, 2004 [R])

24. Admit to Monitored Unit (Intensive Care or Telemetry)

Patients with TIA symptoms within 24-48 hours and at high risk for stroke (see Annotation #23, "High Risk for Stroke?") should be admitted to a monitored unit (ideally telemetry) for observation and further evaluations. Admitting patients expedites diagnostic evaluation, allows for ready access to fibrinolysis should the patient have an acute stroke, facilitates early carotid revascularization if indicated, and offers greater opportunity for risk factor modification for secondary stroke prevention. Again, expedited outpatient programs may be equivalent (See Annotation #26, "Rapid Outpatient Evaluation or Admit to Hospital").

The following diagnostic evaluations should be performed for inpatients (*Adams*, 2007 [*R*]; *Albers*, 2002 [*R*]; *Johnston*, 2000 [*R*]):

Laboratory Tests

- If not already completed (see Annotation #21, "Emergency Department Diagnostic Evaluation"), test for complete blood count, electrolytes, BUN, creatinine, glucose, PT, aPTT, cardiac biomarkers
- Fasting lipid profile
- Fasting glucose

- Serial cardiac biomarkers if suspect acute coronary syndrome
- Consider A1c if suspect diabetes
- Echocardiogram
- Brain and vascular imaging (Douglas, 2003 [D])
 - Magnetic resonance imaging (MRI)/magnetic resonance angiography (MRA)
 - Computed tomography (CT)/computed tomography angiography (CTA)
 - CT/carotid ultrasound if symptoms referable to carotid distribution
- Risk factor assessment and counseling

26. Rapid Outpatient Evaluation or Admit to Hospital

Patients with TIA symptoms that occurred more than 24 hours ago but within the last seven days should be evaluated as soon as possible (*Albers 2002 [R]*; *Johnston*, 2002 [*R]*). Organizations have started TIA clinics for the rapid evaluation of patients in the outpatient setting. Patients who cannot be evaluated rapidly as an outpatient should be admitted to the hospital. The following diagnostic evaluations should be performed within 48 hours:

- Laboratory Tests
 - Complete blood count
 - Electrolytes (sodium, potassium, chloride, CO₂)
 - Sedimentation rate (ESR)
 - BUN
 - Creatinine
 - Fasting lipid profile
 - Fasting glucose
 - Consider A1c if suspect diabetes

All patients with clinical TIA will either be admitted or followed up in an outpatient clinic (either TIA clinic or primary care). Risk factor assessment and modification should take place in the outpatient (or inpatient) setting.

- Echocardiogram
- Brain and vascular imaging (Douglas, 2003 [D])
 - Magnetic resonance imaging (MRI)/magnetic resonance angiography (MRA)
 - Computed tomography (CT)/computed tomography angiography (CTA)
 - CT/carotid ultrasound, if symptoms referable to carotid distribution
- Risk factor assessment and counseling factor

Stroke Code Algorithm Annotations

29. Admit and Begin Stroke Code

Key Points:

- The "door to first physician contact" goal is within 10 minutes.
- The "door to initiation of CT scan" goal is within 25 minutes.
- The "door to drug" goal for thrombolytic treatment is within 60 minutes.

The goal of the stroke code is to rapidly administer tPA in appropriately screened candidates. The onset of symptoms to treatment can be up to 180 minutes (or 270 minutes in selected patients), but the NIH recommendation of "door to drug" is within 60 minutes (*Adams*, 2007 [*R*]).

This committee uses the term "stroke code" to refer to a process in the ED for the rapid evaluation and treatment of patients who have presented in a time frame qualifying them for thrombolytic therapy. This process may take many forms. It might include a formal "stroke team" that is called whenever a possible candidate for tPA has presented, or it may include the ED staff who have been trained in the rapid evaluation and treatment of stroke patients. The general concept is one that includes the following.

- Rapid triage of patients as soon as they arrive in the ED
- Immediate initiation of phlebotomy for appropriate blood tests, followed by CT scan or other equivalent imaging
- First physician contact for history and exam occurring early in the ED visit. The NIH recommendation for timing of "door to first physician contact" for thrombolytic candidates is within 10 minutes
- Rapid access to the best neurologic and radiologic expertise for evaluation of the patient and interpretation of the CT scan prior to treatment

This may include a neurologist and neuroradiologist present at the time of treatment. Alternatively, it may be a primary care physician with expertise in stroke diagnosis and administration of tPA and a general radiologist with expertise in reviewing head CT scans. The NIH recommendation for the timing of "door to initiation of CT scan" for thrombolytic candidates is within 25 minutes.

• The goal of the stroke code should be to administer tPA in appropriately screened candidates. The NIH recommendation for the timing of "door to drug" for thrombolytic treatment is within 60 minutes (*Adams*, 2007 [R]); Bock, 1999 [NA])

30. Evaluation (Should Occur Concurrently with Intervention)

Key Points:

- Apart from history and examination (NIHSS) relevant to thrombolytic therapy, CT scan and glucose, other tests are not necessary before administering IV tPA in most patients. Obtaining them should not delay treatment.
- Review tPA indications/contraindications and document as to whether patient is eligible.
- Perform baseline National Institutes of Health Stroke Scale (NIHSS).

• Perform non-contrast head CT to exclude hemorrhage.

Review History and tPA Treatment Indications and Contraindications, and Baseline NIHSS

Take a complete patient history, including a review of indications and contraindications for treatment with tPA (*Adams*, 2007 [*R*]).

Indications for tPA

- Acute onset of focal neurological symptoms, consistent with ischemic stroke in patients 18 years of age and older.
- Clearly defined onset of stroke less than 3 hours (4.5 hours in selected patients; see Annotation #18, "Consider IV Tissue Plasminogen Activator [tPA]/See Stroke Code Algorithm") prior to planned start of treatment; if the patient awakens with symptoms, onset is defined as the time of the last known baseline neurological status prior to retiring.
- CT scan does not show evidence of intracranial hemorrhage, non-vascular lesions (e.g., brain tumor, abscess) or signs of advanced cerebral infarction such as sulcal edema, hemispheric swelling, or large areas of low attenuation consistent with extensive volume of infarcted tissue.
- A patient with a seizure at the time of onset of stroke may be eligible for treatment as long as the physician is convinced that residual impairments are secondary to stroke and not a postictal phenomenon (*Adams*, 2007 [*R*]).

Contraindications for tPA

The clinical history, laboratory and radiological contraindications for thrombolytic therapy (tPA) that are listed below should be considered relative contraindications. Clinical judgement should weigh the patient's risk for receiving tPA compared with the benefits of thrombolytic therapy.

Clinical contraindications

- Clearly defined onset of stroke greater than 3 hours prior to planned start of treatment; if the patient awakens with symptoms, onset is defined as the time of the last known baseline neurological status prior to retiring.
- Rapidly improving symptoms
- Mild stroke symptoms/signs (NIHSS less than four)
 - Sensory symptoms only
 - Ataxia without other deficits
 - Dysarthria without other deficits
 - Mild motor signs (non-disabling)
 - Visual field defect without other deficits
- In the setting of middle cerebral artery (MCA) stroke, an obtunded or comatose state may be a relative contraindication.
- Clinical presentation suggestive of subarachnoid hemorrhage, regardless of CT result
- Hypertension systolic blood pressure (SBP) greater than 185 mmHg or diastolic blood pressure (DBP) greater than 110 mmHg

Patients with a systolic blood pressure greater than 185 mmHg or diastolic blood pressure greater than 110 mmHg are excluded only if the blood pressure remains elevated on consecutive measurements, and if aggressive treatment is required to lower the blood pressure into an appropriate range.

Throughout this guideline, the work group frequently refers to blood pressure limits that are represented as systolic/diastolic. These ranges are intended to show the blood pressure limits as exceeding as either a given systolic level **OR** a given diastolic level.

History contraindications

- Minor ischemic stroke within the last month
- Major ischemic stroke or head trauma within the last three months
- History of intracerebral or subarachnoid hemorrhage if recurrence risk is substantial
- Untreated cerebral aneurysm, arteriovenous malformation (AVM) or brain tumor
- Gastrointestinal or genitourinary hemorrhage within the last 21 days
- Arterial puncture at a non-compressible site within the last seven days or lumbar puncture within the last three days
- Major surgery or major trauma within the last 14 days
- Clinical presentation suggestive of acute myocardial infarction (MI) or post-MI pericarditis
- Patient taking oral anticoagulants and INR greater than 1.7
- Patient receiving heparin within the last 48 hours and has an elevated aPTT
- Patient receiving low-molecular-weight heparin within the last 24 hours
- Pregnant, or anticipated pregnant, female
- Known hereditary or acquired hemorrhagic diathesis or unsupported coagulation factor deficiency

Laboratory contraindications

Glucose should always be measured prior to giving tPA; other parameters should be checked before treatment if there is reason to believe they may be abnormal (e.g., INR and aPTT should be checked if patient has been exposed recently to warfarin or heparin or if there is history of liver disease).

- Glucose less than 50 or greater than 400 mg/dL
- Platelet count less than 100,000 mm³
- INR greater than 1.7
- Elevated aPTT
- Positive pregnancy test

Radiology contraindications

- Intracranial hemorrhage
- Large area of low attenuation consistent with an infarcted brain

Early changes of this type suggest that onset of symptoms occurred earlier than the history first indicated. Recheck patient history and time of symptom onset.

• Intracranial tumor, aneurysm, arteriovenous malformation (AVM) or other space-occupying lesion

Once indications and contraindications have been reviewed, the patient should be appropriately managed and documentation of why tPA was given or not given must occur.

Baseline NIHSS

A history and neurological examination must be performed to assess whether the presentation is consistent with a stroke diagnosis and to estimate the severity of the deficit (*Adams*, 2007 [*R*]). Use of the NIHSS by physicians and nursing staff is encouraged, as the scale provides a uniform method of evaluation to facilitate comparison between examiners during the early hours of the stroke care. We encourage use of the NIHSS as an initial evaluation tool and after treatment to assess for change.

The NIHSS is a quantitative measure of neurologic deficit in stroke patients that covers the key aspects of the neurological exam, including level of consciousness and orientation, eye movements, visual fields, facial weakness, motor strength in limbs, coordination, sensation, language and comprehension of language, articulation, and neglect. It can be performed in rapid fashion (five to eight minutes), which is an important feature in this clinical setting (*Adams*, 2007 [*R*]; Brott, 1992 [*R*]).

The NIHSS has been demonstrated in several evaluations to have both validity and reliability as follows:

- Content validity
 - The items contained in the NIHSS were selected on the basis of expert opinion and literature review, thus satisfying the requirements for content validity (*Boysen*, 1992 [R]).
- Concurrent criterion validity
 - The NIHSS correlates with lesion volume on CT scan (Brott, 1989 [D]).
 - The NIHSS correlates with other measures of neurological outcome (Duncan, 1992 [C]).
- Construct validity
 - Factor analysis reported by Lyden et al. defined two constructs relating to right and left hemisphere function confirming construct validity of this scale (i.e., it is measuring what it was designed to measure). This final factor structure remained consistent in both tPA-treated and placebo patients over time after ischemic stroke treatment (*Lyden*, 1999 [C]).
- Predictive validity
 - The NIHSS predicts three-month outcome (Adams, 1999 [B]; Muir, 1996 [C]).
- Interrater and intrarater reliability
 - The NIHSS has been shown to be a reproducible measure, both comparing different examiners and comparing repeated evaluations by the same examiner. Reliability has been demonstrated for neurologists, other physicians and nursing caregivers (*Dewey*, 1999 [C]; Goldstein, 1989 [C]; Goldstein, 1997 [C]). Although the NIHSS was originally designed as a research tool, it has proven to be an excellent measure of neurologic status and can be an important tool for the standardization and communication of clinical information between nurse caregivers and between nurse caregivers and other health care professionals (*Spilker*, 1997 [C]).

Perform Vital Signs Every 15 Minutes with Neurological Checks (not NIHSS)

It is the standard of practice to perform a baseline NIHSS neurological assessment (*Adams*, 2007 [*R*]). For subsequent neuro checks, a less extensive tool is appropriate. Performing a full NIHSS assessment every 15 minutes is often not feasible and may not be a good use of time. There is not evidence showing that performing a full NIHSS assessment every 15 minutes improves patient outcomes or improves the assessment and early detection of changes in patient condition. Unfortunately, there is not a standard validated non-NIHSS neurological assessment that is utilized by health care providers or that has been studied.

The work group has gathered the abbreviated neurological assessments used by several organizations and proposes the following non-NIHSS neuro check as an option.

Level of Consciousness - measures the level of alertness and cognition of the patient

- Is the patient alert, alert with stimulation or requires repeated stimulation to remain alert, or comatose?
- Is the patient able to correctly mouth his/her name and age?
- Is the patient able to correctly follow simple commands of opening and closing his/her eyes?

Motor Functions - measures the motor functions and patient's ability to follow commands

- Is the patient able to perform a series of arm movements?
- Is the patient able to perform a series of leg movements?

Language Skills – measures the amount of aphasia and dysarthria in response to asking patients to describe an item or read several sentences

See Appendix C for examples of non-NIHSS neuro check forms.

The work group would like to encourage organizations to measure the use of non-NIHSS assessment tools to grow the evidence in this area.

Record Weight (estimate if needed)

Draw Blood for Lab Tests

Necessary/critical laboratory tests (results must be available before treatment in all cases):

- Glucose
- PT/INR (if patient on warfarin)

Recommended laboratory tests (results must be available before treatment if physical exam and/or patient history indicates the possibility of abnormal results):

- Complete blood count (CBC) with platelet count
- Electrolytes, BUN, creatinine
- PT/INR, aPTT

Others to consider:

- Troponin
- AST

These tests are used to evaluate for dehydration, metabolic disorders that might influence neurologic status (especially hypoglycemia and hyperglycemia), hematologic disorders such as polycythemia that may affect cerebral perfusion, or coagulopathies that could affect the treatment decision (*Adams*, 2007 [*R*]). Prior to administration of tPA, the glucose level should be reviewed. If the patient is known to be on warfarin or has received heparin within the last 24 hours, the prothrombin time and partial thromboplastin time must be reviewed prior to treatment. A urine or serum pregnancy test should be obtained in women of childbearing potential if there is substantial reason to believe the patient may be pregnant.

Perform Electrocardiography

An EKG should be performed for the purpose of screening for concomitant cardiac disease, either acute or chronic, that may impact immediate treatment decisions.

Perform CT Head without Contrast (or Other Equivalent Imaging)

A CT scan without contrast must be performed prior to treatment with tPA, primarily for the purpose of excluding hemorrhage. Early signs of infarct should also be sought as this finding confers greater risk of symptomatic intracerebral hemorrhage with tPA treatment (*Adams, 2007 [R]*). It has been recently shown that MRI scans of the brain with diffusion- and susceptibility-weighted (gradient echo) sequences are much more sensitive than CT in detecting new infarction and chronic hemorrhage as well as of equal sensitivity for acute hemorrhage (*Chalela, 2007 [C]; Fiebach, 2004 [C]*). Consequently, when it is possible to perform MRI as quickly as CT with equally expert and timely interpretation, MRI may be used in this situation. Whichever is used, it is recommended that the greatest level of radiologic expertise possible be obtained for interpretation, with the caveat that this CT reading should not create excessive delays in the evaluation and treatment process. A procedure for rapid teleradiography CT readings should be organized and in place if needed to provide this expertise quickly.

Other Cardiac Assessment as Appropriate (Telemetry)

Consider If Intra-Arterial Recanalization Candidate

Intra-arterial thrombolytic therapy may be a treatment option for selected patients presenting in an early time frame but beyond the 3-hour (or 4.5 hours in selected patients; see Annotation #18, "Consider IV Tissue Plasminogen Activator [tPA]/See Stroke Code Algorithm") time window for intravenous tPA (*Adams*, 2007 [*R*]).

The availability of this option will be institution dependent, and patients must be highly selected. If considering this treatment option for a patient, a physician must explain to the patient and family that this is not standard of usual care and has substantial risk. Despite the limitations of available study data, in cases of more severe presentation with basilar artery or middle cerebral artery or basilar artery occlusion, intra-arterial thrombolytic treatment may be appropriate because the prognosis without treatment is poor.

If the patient is an appropriate candidate for this treatment, consideration should be given to immediate transfer to an institution offering this intervention. If an endovascular interventionist skilled in this technique is available to the hospital, the patient should be mobilized quickly. See also Appendix B, "Acute Stroke Care Networks."

Middle cerebral artery occlusion

Criteria for consideration of angiographic evaluation for intra-arterial treatment:

- Middle cerebral artery occlusion defined by:
 - Symptom complex consistent with this vascular distribution:
 - Contralateral hemiplegia and face weakness
 - Contralateral hemisensory loss
 - Aphasia if ischemia is on left, "neglect" if on right
 - Commonly, contralateral homonymous visual field deficit, reduced level of arousal, eye deviation toward side of brain ischemia (away from side of weakness)
 - MCA "clot sign" on baseline pretreatment CT scan with appropriate clinical presentation

- CT angiogram, MRA or transcranial Doppler (TCD) demonstration of the occlusion with appropriate clinical presentation

Treatment should begin greater than three hours but less than six hours from onset of symptoms.

In the case of middle cerebral artery occlusion, the estimated degree of benefit may seem to be less dramatic than that with basilar occlusion, but the supporting studies offer a superior level of evidence. In the first of two PROACT studies (Prolyse in Acute Cerebral Thromboembolism Study), an improved rate of recanalization was established when comparing use of intra-arterial recombinant pro-urokinase (r-proUK) plus IV heparin to intra-arterial infusion of placebo plus IV heparin (57.7% versus 14.3%, 2p=0.017) (del Zoppo, 1998 [A]). This was a small phase II trial (n=40, 26 received r-proUK and 14 received placebo). Clinical efficacy was not a primary endpoint and it was not established in this study. In PROACT II intra-arterial r-proUK plus IV heparin (n=121) was compared to IV heparin alone (n=59) (Furlan, 1999 [A]). Significant clinical benefit with treatment was established, showing a 15% absolute increase in the number of patients with good outcome at three months (primary outcome measure was modified Rankin score of two or less). The complication of symptomatic intracerebral hemorrhage was higher than that seen in the NINDS IV tPA study (10.2% vs. 6.4% respectively) (National Institute of Neurological Disorders and Stroke tPA Stroke Study Group, 1995 [R]). However, the pretreatment severity of stroke in PROACT II was also higher than that in the NINDS study, probably accounting for this excess of hemorrhagic complications. Despite these promising results, r-proUK was not approved by the FDA for the indication of ischemic stroke, and r-proUK has never been available in the U.S. for any indication. The results of PROACT II, however, served as a proof of principle for the efficacy of an intra-arterial lytic approach to proximal MCA (M, or M, segment) occlusion in the three- to six-hour time frame. It should be recalled that the available lytic agent, tPA, has not been examined in a randomized trial of the intra-arterial route.

Basilar artery occlusion

- Basilar artery occlusion defined by the following.
 - Symptom complex consistent with this vascular distribution:
 - Quadriparesis, sometimes with posturing bulbar dysfunction (dysarthia, dysphagia, dysphonia)
 - Typically dysconjugate eye movement deficits
 - Commonly, depressed level of arousal, respiratory abnormalities
 - Hyperdense "clot sign" in basilar artery on baseline non-contrast CT scan with appropriate clinical presentation
 - CT angiogram, MRA or TCD demonstration of the occlusion with appropriate clinical presentation

Treatment should begin greater than three hours but less than 12 hours from onset of symptoms.

The occurrence of acute basilar artery occlusion with bilateral brainstem symptoms is typically a catastrophic neurological event portending a poor prognosis if reperfusion does not occur, with estimations of over 75% mortality and severe disability in survivors (*Archer, 1977 [D]; Caplan, 1983 [D]; Kubik, 1946 [D]*). Several investigators have reported their results in series of treated patients with basilar thrombosis using intra-arterial urokinase or tPA, showing recanalization rates between 40% and 78% and good outcome by various measures in 20% to 50% (*Becker, 1996 [D]; Brandt, 1996 [D]; Cross, 1997 [D]; Gonner, 1998 [D]; Hacke, 1988 [C]; Wijdicks, 1997 [D]; Zeumer, 1993 [D])*. These are dramatic results when compared to the natural history of this disease reported in the literature.

Algorithm Annotations

There are no randomized, controlled trials of cases of basilar occlusion comparing intravenous tPA to intra-arterial thrombolysis within 3 hours of symptom onset or intra-arterial therapy to placebo controls in any time window, but the limited number of patients presenting with this specific entity would make this a difficult undertaking. In patients presenting within a 3-hour time window, (4.5 hours in select patients; see Annotation #18, "Consider IV Tissue Plasminogen Activator [tPA]/See Stroke Code algorithm") the work group suggests that IV tPA be administered. Subsequent intra-arterial treatment may be considered in some centers.

Newer approaches

Other approaches to acute reperfusion are under study. One has been to combine the speed of the intravenous therapy with the superior recanalization effect of intra-arterial administration. The Emergency Management of Stroke (EMS) Bridging Trial was a small study (n=35) comparing combined use of intravenous and intra-arterial tPA in patients presenting within the three-hour time window (*Lewandowski*, 1999 [A]). This study demonstrated the feasibility of the combined intravenous/intra-arterial approach showing better recanalization rates when compared to intra-arterial treatment alone. The study was too small to adequately assess efficacy and safety. A larger study (n=80) by the Interventional Management of Stroke (IMS) Investigators compared efficacy and safety of the approach using matched historical controls from the NINDS IV tPA trial (*IMS Study Investigators*, 2004 [A]). Efficacy was slightly greater and safety similar to the historical IV-tPA-treated patients. A randomized trial is ongoing. Another approach uses mechanical rather than chemical clot removal. The Mechanical Embolus Removal in Cerebral Ischemia phase I study (MERCI I) showed that recanalization using a corkscrew-shaped retrieval device could be safely performed with promising efficacy results in patients presenting up to eight hours after occlusion of a proximal artery, e.g., internal carotid, MCA, BA, vertebral artery (*Gobin*, 2004 [D]). A phase 2 trial found similar recanalization rates but less favorable outcomes. Nevertheless, FDA approval was established.

At this point in time, there are no studies comparing intravenous to intra-arterial therapy within the 3-hour window. Intravenous treatment with tPA is proven effective in this time frame. Intra-arterial thrombolysis with the agents currently available for use has theoretic advantages in certain stroke types (demonstrated large vessel occlusion of the internal carotid, middle cerebral or basilar arteries), but its superiority in producing improved clinical outcomes remains unproved. Also, there are logistic difficulties with intra-arterial catheter technique that may delay the time to intervention, thus limiting the benefit for these patients. For these reasons, treatment within the 3-hour time window with intra-arterial instead of intravenous thrombolysis cannot be recommended for these large vessel occlusion cases. Centers with expertise in use of this technique or mechanical clot removal should be encouraged to continue utilizing intra-arterial thrombolysis in appropriate candidates presenting beyond the 3-hour time window (presentation within a 3 to 6 hour time window for MCA occlusion [M₁ or M₂ segment], for intra-arterial tPA, 3 to 8 hours for mechanical clot removal and a 3- to 12-hour time window for tPA, 3 to 8 hours for mechanical removal of basilar occlusion) while collecting outcome data and reporting their experience to the medical community. Even more desirable, these same centers should be participating in randomized, controlled trials so that the efficacy of this therapy can be fully established and its role in the acute ischemic stroke treatment armamentarium can be clarified for all.

Emerging Technologies

Several groups have reported use of imaging to support decisions about reperfusion therapies (*Albers*, 2006 [*C*]; *Furlan*, 2006 [*A*]; *Hacke*, 2005 [*A*]). MR and CT technologies can provide information about the status of the vascular supply and parenchyma of acutely ischemic brain. Vascular studies (MRA, CTA) demonstrate presence and location of occlusive thrombus, as well as collateral channels. This knowledge enables the offending thrombus to be targeted more precisely and may expand reperfusion options to intra-arterial chemical or mechanical means. Special CT and MR imaging protocols measure cerebral transit time, blood flow, blood volume and in the case of MR, presence or absence of cytotoxic edema characteristic of

infarcting or infarcted brain by detecting restricted proton diffusion. From these parenchymal data, presence and extent of penumbra vs. infarct can be inferred.

It has been argued that information available from these technologies may be of equal or greater importance as time elapsed since symptom onset in deciding whether and how to undertake a reperfusion therapy. The hypothetical ideal reperfusion scenario is when a vascular occlusion is identified and the parenchymal signature is that of penumbra with little or no infarct. On theoretical grounds, treating in such a scenario might be defended even if the time limit were exceeded. A small randomized trial recently supported this idea by showing that intravenous tPA may be beneficial beyond 3 hours compared with placebo in patients with perfusion-diffusion "mismatch" (i.e., a larger volume of brain with reduced flow than that of already infarcted tissue) using perfusion- and diffusion-weighted MRI (*Davis*, 2008 [A]). Management protocols using such approaches are being assessed in several hospitals in this region. Their use to supersede time-based reperfusion policies, though supported by emerging information, must still be considered experimental outside of standard care.

31. Intervention (Should Occur Concurrently with Evaluation)

Key Points:

- Apart from treating elevated blood pressure in preparation for thrombolytic therapy, none of these treatments is necessary before administering tPA. They should not delay therapy.
- Education on the suspected diagnosis of ischemic stroke and the possible treatment plans should occur with the patient and family/caregiver.
- Patients with a systolic blood pressure greater than 185 mmHg or diastolic blood pressure greater than 110 mmHg are excluded only if the blood pressure remains elevated on consecutive measurements despite treatment with approved measures (e.g., labetolol, nitro paste) and if aggressive treatment is required to lower the blood pressure into an appropriate range.
- Prevent dehydration in patients by maintaining euvolemia with isotonic fluids. Hypotonic fluids should be avoided because they promote brain swelling.
- Initiate treatment if necessary to correct hyperthermia, hypo- or hyperglycemia, and hypoxia.

Educate Patient and Family

A process should be in place for the patient and family that will rapidly orient them to the suspected diagnosis, ED process, tests to be performed, tPA treatment and its risks, and other treatment measures to be considered. This could include caregiver face-to-face interaction with the patient and family, as well as teaching tools in written form. Education should be documented in the medical record.

Treat Hypertension If Greater than 185 Systolic or 110 Diastolic

Patients with a systolic blood pressure greater than 185 mmHg or diastolic blood pressure greater than 110 mmHg are excluded only if the blood pressure remains elevated on consecutive measurements (*Adams*, 2007 [R]), and if aggressive treatment is required to lower the blood pressure into an appropriate range (e.g., if more than a few doses of any medication is required or if nitroprusside drip is required).

Guidelines for blood pressure management in this setting have been slowly evolving (*Adams*, 2007 [*R*]; *International Society of Hypertension Group*, 2003 [*R*]; *Powers*, 1993 [*R*]; *Stead*, 2004 [*M*]; *Strandgaard*, 1996 [*R*]). Although empirically many have considered hypertension in the acute stroke setting to be potentially injurious, it is not clear now that this is not the case.

A full understanding of this issue requires understanding of the physiology. Cerebral blood flow (CBF) is regulated by the relationship between cerebral perfusion pressure (CPP) and cerebrovascular resistance (CVR) (CBF=CPP/CVR). CPP represents the difference between arterial blood pressure forcing the blood into the cerebral circulation and the venous back pressure. Under normal circumstances, the venous back pressure is negligible and CPP is equal to arterial blood pressure. Normally, changes in blood pressure (or CPP) over a wide range have little effect on CBF. This phenomenon, termed autoregulation, is mediated via changes in the CVR. An increase in CPP (or arterial blood pressure) produces vasoconstriction and a decrease produces vasodilatation. This autoregulation keeps the cerebral blood flow at a steady level over a range of 60-150 mmHg mean arterial pressure. In individuals with chronic hypertension, the range for autoregulation is shifted upwards so that they may be more tolerant of higher blood pressure and less tolerant of lower blood pressure (decreased cerebral blood flow).

Acute ischemic stroke will cause a change in autoregulation in the ischemic zone by two mechanisms:

First, when an artery is occluded, a central core of severe ischemia is produced. This is surrounded by a zone with less reduction in blood flow where perfusion is maintained by collateral circulation, termed the penumbra. These blood vessels in the penumbra are maximally dilated, and for that reason blood flow through them may be completely dependent on blood pressure.

Second, during the acute period, the phenomenon of autoregulation even outside of the penumbra can be impaired in patients both with and without persistent arterial occlusion, changing the autoregulation curve so that maintenance of blood flow is completely dependent on the blood pressure.

These abnormalities in autoregulation may persist for days or weeks. There is evidence to suggest that there is slow improvement in this phenomenon in the acute period. But early on, lowering the blood pressure may reduce blood flow to critical levels in the ischemic region, potentially extending the area of infarct. This is supported by data from both animal and human studies (*Christensen, 2002 [D]; Powers, 1993 [R]*).

Although the potential dangers of lowering arterial blood pressure in patients with acute ischemic stroke are accepted theory influencing practice, documentation of actual risk is based on a few published case reports (*Britton*, 1980 [D]; Grossman, 1996 [R]; Lavin, 1986 [D]). The theoretical adverse effects are substantial. Whether carefully controlled treatment of hypertension in acute stroke might be beneficial has not been adequately studied.

A Cochrane review (2003) consisting of 34 randomized controlled trials and 5,368 patients examined the effect of various drugs on blood pressure (BP) during the first 72 hours of acute ischemic stroke (AIS). Drugs shown to actually reduce BP included oral and IV calcium channel blockers, oral beta-blockers, Glyceryl Trinitrate, ACE inhibitors, Prostacyclin (PGI2), and Streptokinase. The effect of blood pressure reduction was not clear, likely due to the significant imbalances in baseline blood pressure between treatment and control groups. Outcomes sought included early death and overall case fatality. The review concluded that there is insufficient evidence to evaluate the effect of altering blood pressure on outcome after acute ischemic stroke. Another systematic review demonstrated increased mortality, early deterioration, and dependency associated with higher blood pressure in the acute stroke setting (*Willmot, 2003 [M]*).

The above review includes the IST trial (*Leonardi-Bee*, 2002 [A]), which demonstrated a U-shaped curve when BP was plotted against survival (i.e., increased mortality at lowest and highest pressure with lowest mortality at systolic pressure around 150 mmHg). Other investigators have also confirmed this (*Castillo*, 2004 [D]), providing grounds for the current consensus-based guidelines to treat BP (*Adams*, 2007 [R]).

Taking the above studies into consideration, the AHA issued a revised 2007 edition of "Guidelines for the Early Management of Patients with Acute Ischemic Stroke" (*Adams*, 2007 [*R*]). In the absence of unambiguous data, these consensus-based guidelines recommend the following measures for treatment of BP in patients with AIS.

Table 3.

Approach to Elevated Blood Pressure in Acute Ischemic Stroke				
Blood Pressure Level (mmHg)	Treatment			
A. Not eligible for thrombolytic therapy Systolic < 220 OR diastolic < 120	Observe unless other end-organ involvement, e.g., aortic dissection, acute myocardial infarction, pulmonary edema, hypertensive encephalopathy.			
	Treat other symptoms of stroke such as headache, pain, agitation, nausea and vomiting.			
	Treat other acute complications of stroke, including hypoxia, increased intracranial pressure, seizures or hypoglycemia.			
Systolic > 220 OR diastolic > 120	Labetalol 10-20 mg IV over 1-2 min. May repeat or double every 10 min. (maximum dose 300 mg) or nicardipine 5 mg/hr IV infusion as initial dose; titrate to desired effect by increasing 2.5 mg/hr. every 5 min. to maximum of 15 mg/hr. Aim for a 15% reduction of blood pressure.			
Diastolic > 140	Nitroprusside 0.5 microgm/kg/min. IV infusion as initial dose with continuous blood pressure monitoring. Aim for a 10% to 15% reduction of blood pressure.			
B. Eligible for thrombolytic therapy				
Systolic >185 OR diastolic >110 mmHg	Labetalol 10-20 mg IV over 1-2 min. May repeat x 1 OR nitropaste 1-2 inches; OR nicardipine infusion, 5 mg/hour, titrate up by 2.5 mg/hour at 5- to 15-minute intervals, maximum dose 15 mg/hour; when desired blood pressure attained, reduce to 3 mg/hour.			
	If blood pressure does not decline and remains > 185/110 mmHg, do not administer rtPA.			
During and after treatment				
Monitor BP Blood pressure level	Check BP every 15 min. during treatment and then for another 2 hours, then every 30 minutes for 6 hours, then every hour for 16 hours.			
Diastolic 105 to 120 mmHg of	Labetalol 10 mg IV over 1-2 minutes, may repeat every 10 to 20 minutes, maximum dose of 300 mg, OR			
	Labetalol 10 mg IV followed by an infusion at 2 to 8 mg/minute			
Systolic > 230 mmHg or Diastolic 121-140 mmHg	Labetalol 10 mg IV over 1 to 2 minutes, may repeat every 10 to 20 minutes, maximum dose of 300 mg OR			
	Labetalol 10 mg IV followed by an at 2 to 8 mg/minute OR Nicardipine infusion, 5 mg/hour, titrate up to desired effect by increasing 2.5 mg/hour every 5 minutes to a maximum of 15 mg/hour. If blood pressure not controlled, consider sodium nitroprusside 0.5 mcg/kg/minute.			

Adapted with permission from the American Heart Association. Adams Jr HP, del Zoppo G, Alberts MJ, et al. Guidelines for the early management of adults with acute ischemic stroke: a guideline from the American heart association/American stroke association stroke council, clinical cardiology council, cardiovascular radiology and intervention council, and the atherosclerotic peripheral vascular disease and quality of care outcomes in research interdisciplinary working groups: the American academy of neurology affirms the value of this guideline as an educational tool for neurologists. *Stroke* 2007;38:1655-1711. (Class R)

Initiate Two Intravenous Lines

Two intravenous (IV) lines should be started so that tPA may have a dedicated line.

Start Intravenous Fluids

Treatment with a 0.9% normal saline at a rate of 75-125 cc/hr or 2-3 L/day should be administered to avoid dehydration (*Adams*, 2007 [*R*]). The rate may be adjusted for febrile patients. IV fluids are particularly important, of course, for patients in whom oral intake is prevented or limited by swallowing problems. Dehydration is fairly common on admission in stroke patients.

Hemorrheologic disturbances may be a factor in limiting cerebral blood flow in the setting of ischemic stroke. Attempts to affect blood viscosity by lowering hematocrit to increase blood flow and oxygen delivery have suggested the possibility of a useful therapeutic intervention (*Thomas, 1977 [C]; Wade, 1983 [D]*). Results have been mixed in studies of hemodilution techniques that attempt to decrease blood viscosity utilizing phlebotomy and volume expansion with dextran or pentastarch. Although there were promising results in small clinical trials, when subjected to more rigorous study with large controlled trials, this treatment was unsuccessful (*Italian Acute Stroke Study Group, 1988 [A]; Scandinavian Stroke Study Group, 1987 [A]*). Although proponents of this treatment have argued that results would be improved with earlier time-to-treatment, a more individualized approach with treatment decisions, or a more aggressive hypervolemic hemodilution approach, additional large-scale trials have not been undertaken. In fact, use of a hypervolemic approach in order to further raise cardiac output by volume expansion was complicated in some cases by cerebral edema and increased mortality, raising questions regarding the safety of this treatment (*Hemodilution in Stroke Study Group, 1989 [A]*). Therefore, hemodilution therapy is not recommended since the clinical benefit has not been established and the possibility of risk due to the development of cerebral edema has been suggested. Also, there is a risk of heart failure.

However, in the general medical management of patients with stroke, it is important to administer adequate fluids to avoid the development of dehydration or to treat it when present since dehydration with hemoconcentration may impair cerebral blood flow (*Thomas*, 1977 [C]). Dehydration with hemoconcentration may also increase the risk of thrombus formation and recurrent embolization in cardiogenic stroke (*Arboix*, 1998 [B]; Yasaka, 1993 [R]; Yasaka, 1990 [C]). Therefore, it is suggested that isotonic intravenous fluids be administered to not only those admitted with dehydration or at risk for dehydration due to problems with swallowing, but to all stroke patients. Hypotonic fluids should be avoided because they promote brain swelling.

Treat Hyperthermia

The acutely injured brain, whether due to trauma or ischemia, is inordinately susceptible to the damaging effects of brain temperature elevation. This fact is well supported by both animal and human studies (*Ginsberg*, 1998 [R]; *Terént*, 1981 [B]).

Interventions for patients with temperatures of greater than 99.5°F (37.5°C) include appropriate dosing of acetaminophen (1 gram orally or 650 mg rectally every four to six hours, not to exceed 4-6 grams in 24 hours) and regular monitoring of temperature status (every four hours). For those patients with extreme hyperthermia, greater than 103°F (39.4°C), aggressive interventions, including cooling blankets and ice packs, are encouraged. Causes for temperature elevation should be sought and treated.

In human studies, early hyperthermia in acute stroke is associated with increased risk of poor outcome, higher mortality and increased infarct volume (*Azzimondi*, 1995 [B]; Castillo, 1998 [D]; Hajat, 2000 [M]; Reith, 1996 [B]). The causality and the relationship of temperature elevation to these poor outcomes are not fully understood. Whether intervention with cooling methods will result in improved outcomes is unknown.

Treat Hyperglycemia

Hyperglycemia may adversely influence clinical outcome.

- Early identification of patients with hyperglycemia in the setting of acute ischemic stroke or in those at risk for cerebral ischemia (ED evaluation of glucose level) is recommended (*Leigh*, 2004 [C]; *Ribo*, 2005 [C]).
- Avoid any agents or factors that might induce hyperglycemia.
 - Eliminate glucose from any IV solutions used. (Recommend use of normal saline.)
 - Avoid use of corticosteroids, even in those patients with cerebral edema, as they are not helpful and may be harmful. Separate recommendations are needed for those on maintenance corticosteroids for concurrent conditions, and treatment decisions are left to the discretion of the physician.
- Use appropriate measures to maintain euglycemia, carefully avoiding hypoglycemia.
- Continue to monitor glucose with bedside testing in those receiving treatment in order to maintain euglycemia.

Most observational studies document either increased mortality or decreased functional outcome, or both, with higher glucose. Some have speculated that early hyperglycemia in the setting of acute stroke is simply a marker of physiologic stress and an epiphenomenon in those who have suffered severe stroke (Bruno, 1999 [B]; Jorgensen, 1994 [B]; Kiers, 1992 [B]; Woo, 1990 [B]). Others have documented that it is an independent predictor of poor outcome and propose that it has a causative role (Baird, 2003 [D]; Lindsberg, 2004 [R]; Parson, 2002 [D]). Despite the extensive body of literature describing this relationship a definite, clinical trial aimed at intervention to improve outcome is still lacking. A study utilizing a continuous infusion of insulin, glucose and potassium in the setting of acute ischemic stroke was recently completed (Gray, 2007 [A]). The trial was discontinued due to low enrollment. When stopped, it showed no benefit by the primary (mortality) or secondary (death or disability) endpoints. Significant but modest reductions in glucose level as well as blood pressure were seen in those randomized to the active treatment. The study was underpowered and had other limitations, making its negative results not definitive. It remains unclear whether early hyperglycemia in the setting of acute stroke is a marker of physiologic stress or an independent predictor of poor outcome. Usual management of hyperglycemia (glucose levels greater than 140 mg/dL) with gentle dosing of subcutaneous insulin, avoiding hypoglycemia, in a timely manner during acute ischemia would seem prudent until ongoing clinical trials address the appropriateness of more aggressive treatment measures (Adams, 2007 [R]).

33. Initiate tPA

Treatment should consist of tPA 0.9 mg/kg intravenously to a maximum dose of 90 mg. Ten percent of this dose should be given as a bolus over one to two minutes and the remainder infused over one hour (*Adams*, 2007 [*R*]). This dosing may be based upon actual or estimated weight.

35. Initiate Aspirin Unless Contraindicated

Key Points:

• Aspirin should be given orally, rectally or via nasogastric tube promptly in patients who are not tPA candidates unless contraindicated (aspirin allergy, GI bleeding). (See "Perform Swallow Evaluation," Annotation #38, if considering oral administration.)

- There is no evidence to support therapeutic anticoagulation with unfractionated heparin, LMW heparin or heparinoids. There is, as yet, insufficient evidence to decide whether specific subgroups of ischemic stroke (e.g., dissection, cardio-embolism with intra-cardiac clot) will benefit from therapeutic anticoagulation.
- If a decision is made to use continuous heparin infusion, boluses should be avoided, and aPTT should be maintained in the 1.5-2 times baseline range.
- Low-dose prophylactic parenteral anticoagulation (e.g., enoxaparin, 40 mg subcutaneously daily) is beneficial for prevention of DVT or PE in stroke patients with limited mobility.

Aspirin

Patients who are not candidates for tPA should be given aspirin promptly in a dose of 325 mg (*Adams*, 2007 [R]) orally, rectally or by nasogastric tube and should be continued on a similar daily dose (*Albers*, 2004 [R]). Exceptions to this approach would be justified in those with contraindications to aspirin therapy (e.g., aspirin allergy, gastrointestinal hemorrhage). For patients with an aspirin allergy, 75 mg of clopidogrel may be reasonable. Intravenous loading with 150-600 mg of clopidogrel establishes antiplatelet effect more rapidly; however, efficacy in this setting is unproven.

Initiation of aspirin therapy should be withheld for 24 hours for patients who have received tPA.

Although the benefits of aspirin therapy for long-term preventive therapy for stroke are well established, the use of aspirin to improve outcome in the acute treatment setting has also been demonstrated. Large randomized controlled trials have identified a small but measurable benefit with use of aspirin in the first 48 hours following ischemic stroke onset (*Bath*, 2001b [A]; Chinese Acute Stroke Trial Collaborative Group, 1997 [A]; International Stroke Trial Collaborative Group, 1997 [A]; Sandercock, 1993 [M]).

The studies together demonstrate benefit of small magnitude, but with statistical significance in the following outcome measures:

- Preventing early recurrent ischemic stroke 7 fewer per 1,000 treated (p< 0.0001) (comparable to the preventive effects of aspirin for one year)
- Decreasing death from any cause 5 fewer per 1,000 treated (p=0.05)
- Decreasing death or early recurrence of nonfatal stroke 9 fewer per 1,000 treated (p=0.001)
- Decreasing death or dependency at discharge or six months 13 fewer per 1,000 treated (p=0.007)

Also, the measured hazard appears to be small and statistically insignificant:

• Hemorrhagic stroke or transformation – 2 more per 1,000 in ASA treated (p=0.06)

Considerations with Heparin Use

In contrast, to the proof of efficacy for aspirin, results from the International Stroke Trial provide powerful evidence against the routine use in patients with acute ischemic stroke of any heparin regimen as intensive as the moderate-dose subcutaneous regimen utilized in this very large clinical trial (unfractionated heparin – 12,500 units subcutaneous twice daily) (*Publications Committee for the Trial of ORG 10172 in Acute Ischemic Stroke*, 1998 [A]).

This would include the adjusted-dose, continuous infusion of unfractionated heparin. The commonly cited indications of vertebrobasilar distribution ischemia or ischemic stroke in the setting of atrial fibrillation were

analyzed separately and there was no measurable benefit in these specific subgroups. Similarly, the weight of available data regarding use of full dose low-molecular-weight heparin for the acute treatment of stroke do not support their routine use for limiting disability or decreasing mortality in this setting.

In summary, the routine use of acute anticoagulation treatment with unfractionated heparin, low-molecularweight heparin, or heparinoid in acute ischemic stroke is not supported by the available evidence evidence (*International Stroke Trial Collaborative Group*, 1997 [A]). This treatment does not appear to improve clinical outcome from the index stroke. There may be subgroups that benefit, but further studies of this problem are required for confirmation.

Despite these discouraging results, the use of continuous heparin infusion in acute stroke has continued to be common in clinical practice (*Albers*, 2004 [*R*]; *Berge*, 2000 [*A*]; *Coull*, 2002 [*M*]; *Diener*, 2001 [*A*]).

Given these data, if the decision is made to use full-dose continuous heparin infusion for a specific indication (e.g., large vessel atherothrombosis or dissection), physicians are strongly encouraged to discuss with their patients the lack of proof for this therapy and to detail the potential hazards.

Heparin use for VTE prophylaxis

For patients at high risk for VTE where pharmacologic prophylaxis is contraindicated, elastic stockings are recommended, and intermittent pneumatic compression (IPC) should be used if the patient is confined to bed. See Annotation #38, "Other Post-ED Medical Management (first 24-48 hours)" (*Kamphnisen, 2005 [R]*).

See the ICSI Venous Thromboembolism Prophylaxis guideline for more information.

36. Post-Emergency Department Medical Management (Postthrombolysis)

- Admit to intensive care unit or acute stroke care unit/cardiac monitoring.
- **Perform vital signs and neurological checks (not NIHSS)** every 15 minutes for two hours, then every 30 minutes for six hours, then every 60 minutes for 24 hours (recommend use of an abbreviated NIHSS for neurological checks). (See Appendix C, "Non-NIHSS Neuro Check.")
- Treat BP if greater than 180/105
 - First 24 hours: Treat if SBP greater than 180 mmHg or DBP greater than 105 mmHg.
 - Monitor BP and any corresponding neurological changes in the ED and first few days of hospitalization.
- Initiate bleeding precautions:
 - Avoid placement of central venous access or arterial puncture for the first 24 hours.
 - Placement of an indwelling bladder catheter should be avoided during drug infusion and for at least 30 minutes after infusion ends.
 - Insertion of a nasogastric tube should be avoided, if possible, during the first 24 hours.
 - Avoid use of anticoagulant, antiplatelet, or non-steroidal anti-inflammatory agents for the first 24 hours.
 - Monitor for CNS hemorrhage.

Algorithm Annotations

- If any signs of CNS hemorrhage (e.g., neurological deterioration, development of severe headache, sudden severe elevation of BP, or new nausea or vomiting) or signs of major systemic hemorrhage, institute the following measures:
 - Discontinue infusion of thrombolytic drug.
 - Obtain hemoglobin, hematocrit, partial thromboplastin time, prothrombin time/INR, platelet count, fibrinogen (also type and cross match if transfusions will be needed).
 - Obtain surgical consultation if necessary.
 - Obtain emergent CT head without contrast if CNS hemorrhage suspected.
- **Initiate antithrombotic therapy** 24 hours after tPA administration (antiplatelet agent or anticoagulant as appropriate).

37. Post-Emergency Department Medical Management (Not a Thrombolysis Candidate)

Treat BP If Greater than 220/120 mmHg or MAP Greater than 130 mmHg.

Recommendations – ischemic stroke, **not** a tPA candidate:

- Treat BP only if systolic blood pressure (SBP) is greater than 220 mmHg, diastolic blood pressure (DBP) is greater than 120 mmHg, and/or mean arterial pressure (MAP) is greater than 130 mmHg.
- Use easily titrated agents, choosing those with the least effect on cerebrovasculature (labetolol, nitropaste or nicardipine). American Heart Association (AHA) recommendations support oral dosing, but if swallowing is affected, intravenous agents should be used.

Dosing examples:

labetalol oral	100-200 mg by mouth initially and every two hours as needed, up to 800 mg total in 24 hours
	or
labetalol IV	10 mg IV over 1-2 min., repeat or double dose every 10-20 min
nitropaste	1 to 2 inches
nitroprusside	0.3 mcg/kg/min IV; titrate for BP control as needed up to 10 mcg/kg/min
nicardipine	5 mg/hr IV infusion, titrate for BP control, increasing 2.5 mg/hr every 5 min. to maximum of 15 mg/hr

- Avoid agents that tend to cause precipitous drops in BP (e.g., sublingual calcium channel blockers).
- Treat hypotension (IV fluids; treat congestive heart failure or arrhythmia and consider pressors).
- Monitor BP and any corresponding neurological changes in the ED and first few days of hospitalization. Avoid overtreating BP.

In patients with markedly increased blood pressure on presentation with acute stroke, measured reduction (e.g., 15% reduction targeted for the first 24 hours) is reasonable. The threshold for initiating such treatment remains 220 mmHg systolic and/or 120 mmHg diastolic. This is despite preliminary evidence that initiating treatment at a lower level may be safe and beneficial (*CHHIPS*, 2008 [*NA*]). In patients who are on an anti-

hypertensive medication program at the time of the ischemic stroke, these medications should generally be withheld for the initial 24 hours. They should be reinstated after 24 hours, assuming that oral or tube administration is possible and hypotension is not present (Adams, 2007 [R]). Many potential reasons for deviating from this general principle exist. For example, suspension of a beta-blocker in a patient with coronary heart disease may be dangerous, and discontinuation of clonidine may cause rebound hypertension.

38. Other Post-Emergency Department Medical Management (First 24-48 Hours)

Continue to Treat Hyperthermia, Hyperglycemia or Hypoglycemia

Refer to Annotation #31, "Intervention (Should Occur Concurrently with Evaluation)."

Initiate Deep Vein Thrombosis (DVT) Prophylaxis

Consider DVT prophylaxis in any patient admitted to the hospital with lower extremity weakness related to an ischemic stroke. The risk of DVT is high (25%-50%), and prophylaxis with parenteral anticoagulant decreases the incidence (10% to 20%). The risk of pulmonary embolism appears to be decreased, as well, although numbers have been small and statistical significance not achieved (*Counsell*, 2001 [M]).

All patients should receive patient education that includes signs and symptoms of VTE and therapy options and encouraged to ambulate early and perform flexion/extension exercises (*Geerts*, 2004 [R]). Elastic stockings should be used for patients at high risk for VTE. Intermittent pneumatic compression should be considered for patients at high risk for VTE who have contraindications to pharmacologic prophylaxis.

The PREVAIL Trial compared the low-molecular-weight enoxaparin (40 mg/day) with unfractionated heparin (5,000 units twice daily) for 10 days after stroke preventing walking. There was a 43% reduction in the incidence of venous thromboembolism in the enoxaparin group (10%) compared with the unfractionated heparin group (18%). Overall bleeding rates were similar. Based on this trial, low-molecular-weight heparin is superior to unfractionated heparin in prevention of venous thromboembolism after stroke with inability to ambulate (*Sherman*, 2007 [A]).

Low-molecular-weight heparin is renally cleared. For patients with a CrCl less than 30 mL/min, use unfractionated heparin. The patient should be monitored for the possible development of heparin-induced thrombocytopenia (HIT) and bleeding. Obtain a platelet count and hemoglobin every other day, beginning on the second day of heparin therapy.

See the ICSI Antithrombotic Therapy Supplement and the Venous Thromboembolism Prophylaxis guideline.

Perform Swallow Evaluation

Pneumonia is a common finding among patients with acute strokes, its incidence ranging from 6% to 32% (*Perry, 2001 [M]*) and is associated with stroke-related dysphagia symptoms. Implementation of a coordinated swallow evaluation on all acute stroke patients has been shown to significantly decrease the incidence of pneumonia among patients with acute stroke (*Odderson, 1995 [D]*). This study used a screening tool consisting of three components: 1) the patient is alert, follows simple requests, has a clear, strong voice, and can produce a strong cough; 2) the patient can handle his/her own secretions without difficulty and can swallow ice chips and sips of ice water briskly; and 3) The larynx elevates completely at the time of swallowing, the voice remains clear after swallow and there is no coughing afterward. (See Appendix D, "Stroke Dysphagia Screen.")

The work group recommends that a bedside swallow test be performed prior to the patient's ingestion of anything by mouth (including oral aspirin or other medications). This screen may be performed by a nurse

and should include pre-specified screening questions identifying patients at high risk for aspiration. If result of screening tool is negative, bedside swallow evaluation shall be performed using 2-3 ounces of water. If no clinical signs of aspiration occur, patient may receive medications, including aspirin, by mouth. If result of screening tool is positive or if bedside swallow evaluation reveals clinical signs of aspiration, the patient shall be given nothing by mouth, referred for formal swallow evaluation to be performed by a speech language pathologist, and aspirin administered via nasogastric tube or per rectum. If this swallow screen is not to be performed in the emergency department, aspirin should be administered rectally or via nasogastric tube.

Bedside swallow assessment or more formal swallow evaluation, and dietary adjustments based on this information, have not been adequately evaluated in sufficiently powered randomized clinical trials. Because these interventions are safe and have a reasonable probability of improving care by decreasing complications, it is reasonable to advocate their use in this setting despite absence of proof of efficacy. Several previously published guidelines advocate these practices (*Bath*, 2001a [M]).

Initiate Rehabilitation Early

Early mobilization within 48 hours of admission, in the form of early initiation of appropriate rehabilitation swivels or other nursing intervention, is advocated for the purpose of preventing complications related to immobility, including deep vein thrombosis, contractures, joint disorders, and pressure sores/decubitus ulcers (*Adams*, 1994 [*R*]; *Helgason*, 1997 [*R*]). This recommendation is not based on existing randomized trial data, and it is unlikely that such a trial will be carried out in the future.

Perform Nutritional Status Assessment

Assessment of the patient's baseline nutritional status and the implementation of treatments to correct any major nutritional problems are recommended (*Adams*, 2007 [*R*]). Poor nutritional status in patients admitted for stroke is associated with increased morbidity and mortality (*FOOD Trial Collaboration*, 2003 [*B*]). However, a trial did not find benefit in administering nutritional supplementation (*Food Trial Collaboration*, 2005 [*A*]).

Early Treatment of Ischemic Brain Edema

Although ischemic brain swelling typically peaks between three and five days after stroke onset, marked early swelling (in the first 24-48 hours) causing mass effect and tissue shift can occur in the most severe cases ("malignant" ischemic brain edema). Low attenuation changes exceeding two-thirds of the MCA territory and large areas of hypoperfusion on perfusion scans (CT perfusion or MR perfusion) on initial radiological evaluation are associated with high risk of developing malignant brain edema. Patients with these features should be strictly monitored with serial neurological examinations, ideally in a stroke unit. Repeating CT scan of the brain to evaluate for progression of regional mass effect is indicated if the patient develops any signs of neurological deterioration. The value of serial CT scans of the brain in the absence of clinical changes remains to be established.

Decompressive hemicraniectomy with durotomy improves survival and functional outcome (*Vahedy*, 2007 [M]). The optimal timing of the procedure is not well established, but most experts recommend early intervention. Improvement in functional outcome has only been shown for patients 60 years old or younger.

Osmotherapy (mannitol 20% or hypertonic saline) may be used to treat ischemic brain edema, but there is very limited data supporting its value (*Bardutzky*, 2007 [*R*]). Mannitol 20% is usually administered as a bolus of 1-2 g/kg of body weight, followed by repeated boluses as needed for neurological decline or scheduled doses of 0.25 to 0.5 g/kg every four to six hours. In patients with established signs of herniation, a rescue dose of 23.4% of saline solution (30 cc) may be useful (*Koenig*, 2008 [D]).

Hyperventilation should be avoided except for mild to moderate hyperventilation (target pCO_2 30-34 mmHg) for brief periods of time because of the risk of exacerbating ischemia by causing vasoconstriction.

Appendix A – Glossary of Abbreviations

ACCESS	Acute Candesartan Cilexetil	MRA	magnetic resonance angiogram			
	Evaluation in Stroke Survivors	MRI	magnetic resonance imaging			
AHA	American Heart Association	NIH	National Institutes of Health			
aPTT	activated partial thromboplastin time	NIHSS	National Institutes of Health Stroke Scale			
AVM	arteriovenous malformation	NINDS	National Institute of Neurological Disorders			
BID	twice daily		and Stroke			
BP	blood pressure	PROACT	Prolyse in Acute Cerebral			
BUN	blood urea nitrogen	DT	Inromboembolism			
CBF	cerebral blood flow	PT	prothrombin time			
CNS	central nervous system	r-pro UK	recombinant prourokinase			
CPP	cerebral perfusion pressure	tPA	tissue plasminogen activator			
СТ	computed tomography	SBP	systolic blood pressure			
СТА	computed tomograph angiography	SPECT	single proton emission tomography			
CV	cardiovascular	TCD	transcranial Doppler			
CVR	cerebrovascular resistance	TIA	transient ischemic attack			
DBP	diastolic blood pressure	TOAST	Trial of ORG 10172 in Acute Ischemic Stroke Trial			
DVT	deep vein thrombosis					
ECASS	European Cooperative Acute Stroke Study					
ED	emergency department					
EKG	electrocardiogram					
EMS	emergency medical system					
FDA	Food and Drug Administration					
ICSI	Institute for Clinical Systems Improvement					
ICU	intensive care unit					
INR	international normalized ratio					
IV	intravenous					
MAP	mean arterial pressure					
MCA	middle cerebral artery					

myocardial infarction

MI

Appendix B – Acute Stroke Care Networks

Two advances in ischemic stroke care, IV tPA and coordinated inpatient care processes (often considered together as "stroke unit care") are known to improve outcomes. The urgency, complexity and potential risks of the former and training required for the latter challenge the resources of emergency rooms and hospitals that may care for only a few ischemic stroke patients each year. At the same time, hospitals with large volumes of such patients have developed "frontline" capability driven by local competition and supported by quality improvement programs offered through The Joint Commission for the Accreditation of Health Care Organization (TJC), the American Heart Association, National Stroke Association, CDC, and others. Currently there exist significant care inequities across the region.

In contrast to those who arrive at frontline stroke hospitals, stroke patients in remote/rural areas in our state and region are likely not to have access to advantages of informed, urgently deployed reperfusion techniques (especially IV tPA but also intra-arterial chemical and mechanical reperfusion therapies for selected cases) and stroke unit care during their hospitalizations. It is likely that outcomes are impacted by low rates of IV tPA use, suboptimal patient selection for IV tPA, and hospital processes that are not guideline based. In fact, one might calculate from number-needed-to-treat metrics that many poor outcomes each year in Minnesota might result from disparities in care based on geography.

Ad hoc systems have been developed in Minnesota and elsewhere to eliminate the disparities in care resulting from geographic exigencies (*Frey*, 2005 [*D*]; *Hoody*, 2008 [*D*]; *Rymer*, 2003 [*D*]; *Silliman*, 2003 [*D*]; *Silverman*, 2005 [*D*]; *Switzer*, 2008 [*R*]; *Switzer*, 2009 [*R*]; *Vaishnav*, 2008 [*D*]; *Wang*, 2000b [*D*]; *Wang*, 20004 [*D*]). Typically the systems that have evolved provide support for remote/rural emergency rooms and hospitals by neurologists or other stroke experts that are physically located elsewhere, usually at a frontline hospital. A single frontline hospital (a "hub") may provide support for several remote/ rural emergency rooms and hospitals ("spokes"). Several models of support have evolved. In some models, the stroke cases are transferred to the frontline stroke center. Models include:

- Field to comprehensive stroke center transport For example, a ground ambulance or helicopter is deployed by regional EMS to a farmhouse and the patient is transport to a frontline stroke center without intermediate hospitalization, i.e., "spokeless." This model has not proliferated beyond a few areas.
- "Trip to drip" A stroke expert or team of experts travels quickly to the remote/rural emergency room or hospital to provide or supervise acute care. This model is limited by distances and availability of mobile stroke experts/teams.
- "Ship and drip" The stroke patient is transferred prior to initiation of reperfusion therapies from remote/rural emergency room or hospital to a frontline stroke center.
- "Drip and ship" After assessment of eligibility with input of supportive expert, IV tPA is started at the remote/rural hospital, and the patient is immediately transported to the front line stroke center.
- "Drip and keep" IV tPA is given with support as above, and the patient remains at remote/rural hospital for acute hospitalization.

There is considerable latitude for individual variation within these models. For example, the formality of expectations between the hub and spoke varies from casual (e.g., a phone call to a hospital neurology consulting line) to explicit, including expectation for 24/7 availability of a stroke expert on the hub side and for formal training, ongoing education, joint protocol development, and joint post-stroke care planning on the spoke side. The form of communication from hub to spoke varies from simple phone call, phone call/ teleradiography, to telemedicine/teleradiography.

Literature supports the feasibility and safety of administering IV tPA in all of these models. There is level I evidence showing that compared with phone call alone, telemedicine avoids protocol deviations in administration of IV tPA (*Meyer*, 2008 [A]). The trial was not powered to analyze actual outcomes.

Working relationships between individual stroke centers and remote/rural hospitals and emergency rooms will necessarily and appropriately be customized according to the unique circumstances and goals of the entities. It seems prudent, however, that these evolving relationships be guided by basic principles. The work group makes the following recommendations:

- Agreements between hub and spokes should be formal as opposed to ad hoc.
- Hub should provide 24/7 availability of support for reperfusion therapies by one or several of the following:
 - telephone +/- teleradiography
 - telemedicine +/- teleradiography
- Protocols defining care processes between hub and spoke should be jointly developed and support the agreed-upon model, e.g., 3, 4 or 5.
- Initial and ongoing education of spoke personnel relevant for the care model should be provided.
- Initial and ongoing training of spoke personnel (e.g., NIHSS performance and interpretation, conducting an informed consent discussion regarding IV tPA) relevant to the care model should be provided.
- There should be joint planning for stroke unit care at hub (models 3 and 4) or spoke (model 5) and case follow-up. Planning for optimal care of non-reperfusion cases should also be provided.

(Schwamm, 2009a [R]; Schwamm, 2009c [R])

Appendix C – Non-NIHSS Neuro Check

Function & Measurement Format	Scores:			
Level of Consciousness: 0=Alert 1=Not alert, but arousable with minimum stimulation 2=Not alert, requires repeated stimulation to attend 3=Coma				
LOC Questions: Ask Patient the Month and His/Her Age 0=Answers both correctly 1=Obeys one correctly 2=Both incorrect				
LOC Commands: Ask Patient to Open and Close Eyes 0=Opens both correctly 1=Obeys one correctly 2=Both incorrect				
 Motor Functions: Arms 0=Normal extension arms 90° or 45° for 10 seconds without drift 1=Drift 2=Some effort against gravity 3=No effort against gravity 4=No movement 9=Untestable – Joint fused or limb amputated (Do not include this in total score) 				
Motor Functions: Legs 0=Normal – hold leg 30° position for 5 seconds 1=Drift 2=Some effort against gravity 3=No effort against gravity 4=No movement 9=Untestable – Joint fused or limb amputated				
Best Language 0=No aphasia 1=Mild to moderate 2=Severe aphasia 3=Mute				
Dysarthria 0=Normal articulation 1=Mild to moderate slurring of words 2=Near unintelligible or unable to speak 3=Mute				

Appendix D – Stroke Dysphagia Screen

Who should be assessed?

Patients who present with TIA, stroke or stroke symptoms.

How do you assess?

Use this algorithm for a quick three-step process!



Provided by HealthPartners Medical Group and Regions Hospital.



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Brief Description of Evidence Grading

Individual research reports are assigned a letter indicating the class of report based on design type: A, B, C, D, M, R, X.

A full explanation of these designators is found in the Foreword of the guideline.

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This section provides resources, strategies and measurement specifications for use in closing the gap between current clinical practice and the recommendations set forth in the guideline.

The subdivisions of this section are:

- Priority Aims and Suggested Measures
- Key Implementation Recommendations
- Knowledge Resources
- Resources Available

Priority Aims and Suggested Measures

1. Increase the percentage of patients presenting within three hours of stroke onset who are evaluated within 10 minutes of arriving in the ED.

Possible measure for accomplishing this aim:

- a. Percentage of patients presenting within three hours of stroke onset who are evaluated by a physician within 10 minutes of arriving in the ED.
- 2. Increase the percentage of patients presenting with TIA symptoms within 24 hours at high risk for stroke who are admitted to the hospital.

Possible measure for accomplishing this aim:

- a. Percentage of patients admitted who have documentation of TIA symptoms within the last 24 hours.
- 3. Increase the percentage of patients receiving appropriate thrombolytic and antithrombotic therapy for ischemic stroke (use of tPA and aspirin).

Possible measures for accomplishing this aim:

- a. Percentage of eligible patients presenting with ischemic stroke treated with tPA. (This is similar to TJC process measure.)
- b. Percentage of patients who are not candidates for tPA treatment who receive aspirin within 24 hours of hospitalization, after a negative head CT, unless contraindicated.
- c. Percentage of patients receiving tPA according to guideline. (Refer to Annotations #29 and 30). (This is similar to TJC process measure.)
- d. Percentage of patients who are candidates for tPA with a "door to drug" time (time of arrival to time of drug administration) of less than 60 minutes.
- e. Percentage of patients who undergo a CT scan within 25 minutes of arrival in the ED.
- 4. Increase the percentage of non-tPA recipients who have hypertension appropriately managed in the first 48 hours of hospitalization or until neurologically stable.

Possible measure for accomplishing this aim:

- a. Percentage of non-tPA recipients who have hypertension appropriately managed according to the guideline. (Refer to Annotation #31).
- 5. Increase the percentage of patients who receive appropriate medical management for prevention of complications within the initial 24-48 hours of diagnosis:
 - Continue to treat hypoglycemia and hyperglycemia
 - Continue to treat hyperthermia
 - Continue IV fluids
 - Continue to treat hypoxia
 - Initiate deep vein thrombosis prophylaxis
 - Perform swallow evaluation

- Initiate early rehabilitation (early mobilization)
- Perform nutritional status assessment

Possible measures for accomplishing this aim:

- a. Percentage of patients who receive appropriate intervention for hypoglycemia and hyperglycemia.
- b. Percentage of patients who receive appropriate intervention for hyperthermia.
- c. Percentage of patients who receive IV fluids.
- d. Percentage of patients who receive appropriate treatment for hypoxia.
- e. Percentage of patients with ischemic stroke with paralysis or other reason for immobility receiving appropriate prevention for venous thromboembolism (subcutaneous heparin or pneumatic compression device). (This is similar to TJC process measure.)
- f. Percentage of patients who are at risk for aspiration who receive an early swallow evaluation. (This is similar to TJC process measure.)
- g. Percentage of patients mobilized from bed within 48 hours of admission.
- 6. Improve patient and family education of patients with ischemic stroke in both the ED and the admitting hospital unit.

Possible measures for accomplishing this aim:

- a. Percentage of patients presenting in the ED with ischemic stroke for whom patient/family education is documented in the medical record.
- b. Percentage of patients admitted to a hospital unit with ischemic stroke for whom patient/family education is documented in the medical record.

Measurement Specification

Possible Success Measure #5f

Patients with ischemic stroke who undergo evidence-based bedside testing protocol approved by the hospital before being given any food, fluids or medication by mouth.

Population Definition

Adults patients (18 years and older) initially presenting with acute symptoms of ischemic stroke.

Data of Interest

Patients who are already identified as nothing by mouth upon presentation of acute ischemic symptoms.

Numerator/Denominator Definitions

Numerator: # of patients who were screened for dysphagia before taking any food, fluids or medication (including aspirin) by mouth.

Denominator: # of all patients screened for acute ischemic stroke.

Method/Source of Data Collection

Concurrent and retrospective data collection through administrative data/claims data, and medical record.

Time Frame Pertaining to Data Collection

Data may be collected monthly or quarterly. For TJC primary stroke center certification, data reporting is quarterly with monthly data points.

Key Implementation Recommendations

The following system changes were identified by the guideline work group as key strategies for health care systems to incorporate in support of the implementation of this guideline.

- 1. Hospitals should consider developing and implementing critical pathways, standing orders and a stroke process to accomplish rapid evaluation and treatment. The process should expedite the evaluation and treatment of patients who are candidates for intravenous tPA and assure uniform, guideline-driven care for all patients with respect to issues like:
 - ongoing antithrombotic therapy,
 - management of blood pressure,
 - early mobilization, and
 - use of appropriate antiembolism treatment in the paralyzed patient.
- 2. A process should be in place for the patient and family that will rapidly orient them to the suspected diagnosis, ED process, tests to be preformed, tPA treatment and its risks, and other treatment measures to be considered. This could include both caregiver face-to-face interactions with the patient and family, as well as teaching tools in written form.

System Improvement

There is evidence that benchmarking can guide and drive quality improvement. Using essentially the same quality indicators as The Joint Commission for the Accreditation of Health Care Organization (TJC) and ICSI, programs like the American Heart Association's Get With The Guidelines-Stroke (*LaBresh*, 2008 [C]; *Schwamm*, 2009b [B]) and the Paul Coverdell National Acute Stroke Registry (*Stoeckle-Roberts*, 2006 [C]) have been shown to improve the quality of stroke care.

The Joint Commission (TJC) Primary Stroke Center Certification

TJC offers certification as Primary Stroke Centers to hospitals that meet specific qualifications. The emphasis of the process is on the early recognition and management of stroke and the scope of accreditation includes integrated efforts in public awareness, emergency medical services, emergency room and hospitalization (*Alberts, 2000 [R]*). The link is: http://www.jointcommission.org/CertificationPrograms/PrimaryStrokeCenters. Beginning in October 2009 all TJC accredited hospitals will have to submit the eight National Quality Forum-endorsed stroke consensus measures. The Centers for Medicare and Medicaid Services (CMS) is also considering the reporting of stroke measures and in the near future the draft Inpatient Prospective Payment System (IPPS) Rule will be released. IPPS is the venue in which CMS communicates with hospitals and physicians about their future measurement reporting.

Among the requirements for TJC certification as a Primary Stroke Center is ongoing process improvement guided by data and benchmarking. The quality indicators chosen by TJC overlap with those developed by the ICSI Diagnosis and Initial Treatment of Ischemic Stroke guideline work group. The TJC quality indicators are:

- 1. Deep Vein Thrombosis (DVT) Prophylaxis*
- 2. Discharged on Antithrombotics*
- 3. Patients with Atrial Fibrillation Receiving Anticoagulation Therapy*
- 4. Thrombolytic Therapy Administered (in eligible patients)

- 5. Antithrombotic Therapy by End of Hospital Day Two
- 6. Discharged on Cholesterol Reducing Medication
- 7. Dysphagia Screening
- 8. Stroke Education
- 9. Smoking Cessation/Advice Counseling
- 10. Assessed for Rehabilitation
- * Initial standard stroke measure set.

Measures 1, 4, 5, 7 and 8 are similar to or identical to those measures listed in this document and within the scope of the guideline.

Knowledge Products and Resources

Criteria for Selecting Resources

The following resources were selected by the Diagnosis and Treatment of Ischemic Stroke guideline work group as additional resources for providers and/or patients. The following criteria were considered in selecting these resources.

- The site contains information specific to the topic of the guideline.
- The content is supported by evidence-based research.
- The content includes the source/author and contact information.
- The content clearly states revision dates or the date the information was published.
- The content is clear about potential biases, noting conflict of interest and/or disclaimers as appropriate.

Resources Available to ICSI Members Only

ICSI has a wide variety of knowledge resources that are *only* available to ICSI members (these are indicated with an asterisk in far left-hand column of the Resources Available table). In addition to the resources listed in the table, ICSI members have access to a broad range of materials including tool kits on CQI processes and Rapid Cycling that can be helpful. To obtain copies of these or other Knowledge Resources, go to http://www.icsi.org/improvement_resources. To access these materials on the Web site, you must be logged in as an ICSI member.

The resources in the table on the next page that are not reserved for ICSI members are available to the public free-of-charge.

Resources Available

*	Author/Organization	Title/Description	Audience	Web Sites/Order Information
	AHA/ASA	Top Ten Things to Know: Use of Telemedicine within Stroke Systems of Care	Health Care Providers	http://www.american- heart.org/presenter. jhtml?identifier=3066480
	AHA/ASA	Top Ten Things to Know: Recom- mendations for Implementation of Telemedicine within Stroke Systems of Care	Health Care Providers	http://www.american- heart.org/downloadable/ heart/124172827688420090507_ topTenTelemedicinePolicy.pdf
	ASA (American Stroke Association)	Comprehensive Web sitePatient education resources	Health Care Providers; Patients and Families	http://www.strokeassociation.org
	Association of Black Cardiologists	Patient education resources	Health Care Providers; Patients and Families	http://www.abcardio.org
	GLRSN (Great Lakes Regional Stroke Net- work)	Comprehensive Web sitePatient education resources	Health Care Providers; Patients and Families	http://tigger.uic.edu/depts/ glstrknet/
	Minnesota Stroke Association	Patient education resources	Patients and Families	http://www.strokemn.org/
	NSA (National Stroke Association)	 Comprehensive Web site Patient education resources Links to survivor/caregiver products and services and additional related Web sites 	Health Care Providers; Patients and Families	http://www.stroke.org
	NINDS (National Institute of Neurological Disorders and Stroke)	 Links to clinical trials Vontains entire discussion and guidelines for system change to address stroke treatment 	Health Care Providers; Patients and Families	http://www.ninds.nih.gov/
	The Brain Attack Coalition	 Contains tools for health care professionals developing systems to enable the rapid diagnosis and treatment of acute stroke Patient education resources 	Health Care Providers; Patients and Families	http://www.stroke-site.org/

* Available to ICSI members only.