Thrombolytic Therapy for Patients Who Wake-Up With Stroke

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Background and Purpose—Approximately 25% of ischemic stroke patients awaken with their deficits. The last-seen-normal time is defined as the time the patient went to sleep, which places these patients outside the window for thrombolysis. The purpose of this study was to describe our center’s experience with off-label, compassionate thrombolysis for wake-up stroke (WUS) patients.

Methods—A retrospective review of our database identified 3 groups of ischemic stroke patients: (1) WUS treated with thrombolysis; (2) nontreated WUS; and (3) 0- to 3-hour intravenous tissue plasminogen activator-treated patients. Safety and clinical outcome measures were symptomatic intracerebral hemorrhage, excellent outcome (discharge modified Rankin score, 0–1), favorable outcome (modified Rankin score, 0–2), and mortality. Outcome measures were controlled for baseline NIHSS using logistic regression.

Results—Forty-six thrombolysed and 34 nonthrombolysed WUS patients were identified. Sixty-one percent (28/46) of the treated WUS patients underwent intravenous thrombolysis alone whereas 30% (14/46) were given only intra-arterial thrombolysis. Four patients received both intravenous and intra-arterial thrombolysis (9%). Two symptomatic intracerebral hemorrhages occurred in treated WUS (4.3%). Controlling for NIHSS imbalance, treated WUS had higher rates of excellent (14% vs 6%; \(P=0.06\)) and favorable outcome (28% vs 13%; \(P=0.006\)), but higher mortality (15% vs 0%) compared to nontreated WUS. A second comparison controlling for baseline NIHSS between treated WUS and 174 intravenous tissue plasminogen activator patients treated within 3 hours of symptoms showed no significant differences in safety and clinical outcomes.

Conclusion—Thrombolysis may be safe in WUS patients. Our center’s experience supports considering a prospective, randomized trial to assess the safety and outcome of thrombolysis for this specific patient population. (Stroke. 2009;40:827-832.)

Key Words: awakening ■ ischemic ■ sleep ■ stroke ■ thrombolysis

The only Food and Drug Administration-approved therapy for acute ischemic stroke, intravenous tissue plasminogen activator (IV tPA), is limited to patients who present with a known symptom onset of \(\leq 3\) hours.\(^1\) Approximately 16% to 28% of ischemic stroke patients awaken with their deficits.\(^2-7\) In these wake-up strokes (WUS), the onset of symptoms is defined as the last-seen-normal (LSN) time. Because this is the time the patient went to sleep, unfortunately it usually places these patients outside the window for thrombolysis or entry into reperfusion clinical trials. Numerous case series and clinical trials have demonstrated an early-morning peak occurrence of ischemic strokes.\(^2,4-11\) Similar to acute myocardial infarction and sudden death,\(^12,13\) the predominant stroke onset time has been shown to occur during the waking hours. In a meta-analysis of 31 publications reporting the circadian timing of 11 816 strokes (8250 ischemic), the onset of symptoms was 55% more likely to occur between 6:00 AM and noon.\(^14\) If a portion of the morning-onset strokes occur near the time of awakening, reperfusion therapies may still be of potential benefit, provided patients present and are triaged emergently.

Some authors\(^3,6,15\) have reported similar clinical and radiographic features between WUS and known-onset ischemic strokes. Fink et al\(^1\) reported similar rates of diffusion-weighted imaging/perfusion-weighted imaging mismatch in both WUS and strokes of known onset. Another group found no significant differences in early ischemic changes in blinded assessment of hyperacute noncontrast CT scans between stroke of known onset and stroke at awakening.\(^15\) Nadeau et al\(^16\) recently published their center’s experience with WUS. Although their data were derived from an incomplete stroke registry (missing 60% of their stroke admissions),

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patients who awoke with their stroke seemed to have worse outcomes. However, after excluding subarachnoid cases and focusing only on ischemic WUS, there was no significant difference between WUS and non-WUS in terms of poor outcome. These studies lead to the hypothesis that some WUS patients may be candidates for thrombolysis. However, there are no published studies to our knowledge on thrombolytic therapy for this patient population.

The objective of this study was to collect all WUS patients treated with off-label thrombolysis from our institutional stroke database and report the baseline patient demographics, safety, and early clinical outcomes from this cohort. In addition, records of consecutive nontreated WUS patients and standard-of-care 0- to 3-hour IV tPA patients were collected for comparison.

Patients and Methods

Patient Selection

We retrospectively identified 3 historical groups of ischemic stroke patients from our database between March 2003 and January 2008: (1) WUS patients treated with off-label thrombolysis; (2) WUS patients not treated with thrombolysis; and (3) 0- to 3-hour standard IV-tPA patients. All patients were assessed by staff stroke neurologists or stroke fellows at a single academic emergency department. WUS patients all met the following criteria: (1) patients were neurologically normal when LSN before going to sleep and were either witnessed with deficits on awakening or could report that they awoke with stroke symptoms; (2) patients had a disabling neurological deficit that would usually be treated with tPA in 0- to 3-hour onset patients; and (3) hypodensity in one-third middle cerebral artery (MCA) territory on noncontrast cranial CT scan.

In situations in which the onset time was unclear (eg, vague or intermittent symptoms), inclusion was adjudicated among our stroke faculty. If a consensus was reached among stroke faculty that the onset of symptoms was at awakening (and not the day before), then those patients were included in the cohort. WUS patients whose LSN on the previous evening.

Outcome Assessment

Safety was assessed by the development of symptomatic intracerebral hemorrhage (sICH), defined as any intracerebral hemorrhage associated with a ≥4-point increase in the NIHSS. Clinical outcomes were defined as excellent outcome (discharge modified Rankin score, 0–1), favorable outcome (discharge modified Rankin score, 0–2), and mortality. Data were abstracted from charts by 1 author (A.D.B.).

Statistical Analysis

Categorical variables were analyzed using *χ*² and Fisher exact test when appropriate. Continuous variables were analyzed using independent samples *t* test or Mann–Whitney *U* when appropriate. Treated WUS were compared to standard 0- to 3-hour IV tPA patients and nontreated WUS. Outcome measures were controlled for baseline NIHSS using multivariate logistic regression. The analysis was performed using SPSS for Windows, version 15 (SPSS Inc).

Results

During the study timeframe, a total of 1253 ischemic stroke patients were admitted to our center and 338 received IV tPA alone (27%). This does not include patients treated with IAT therapy or combination IV plus IAT. After removing patients treated within research protocols and off-label treatments, 174 remained who were treated within 0 to 3 hours. This group served as the standard-of-care 0- to 3-hour IV tPA arm in the study.

A total of 80 consecutive WUS patients were identified from our database (6.6% of all ischemic strokes). Of these, 46 received off-label thrombolysis (treated WUS) and 34 did not receive any intravenous or endovascular treatments (nontreated WUS). In treated WUS patients, 28 received full-dose (0.9 mg/kg, maximum 90 mg) IV tPA alone; 14 received IAT alone; and 4 received combination full-dose IV tPA plus IA thrombolysis. Only 2 patients had preceding vague symptoms requiring adjudication for inclusion into the thrombolysed WUS group. Of the 46 WUS patients, 7 woke-up with stroke symptoms but lacked an exact LSN time. Their chart documentation noted a LSN time of the previous evening.

For comparison, 34 nontreated WUS patients were identified. Half (17/34) of the nontreated WUS patients had an unknown exact LSN time but were known to have been neurologically normal before going to sleep the previous evening.
In the thrombolysed WUS patients, 16 underwent pretreatment multimodal neuroimaging either with CT perfusion (n=10) or with MRI diffusion-weighted imaging/perfusion-weighted imaging (n=6). Two CT perfusion studies were nondiagnostic because of poor contrast bolus and 1 CT perfusion study was normal. The remaining patients were treated using noncontrast CT alone to exclude hemorrhage or hypodensity > one-third MCA territory. Emergent vascular imaging was performed before treatment in 30 of the 46 thrombolysed WUS patients. This included 14 CT angiograms, 10 transcranial Doppler ultrasounds, and 6 MR angiograms. Eighty-six percent (26/30) of patients who had vascular imaging had documented occlusions of intracranial arteries.

Baseline demographic and radiological data for group comparisons are displayed in Tables 1 and 2. All 18 patients who received IAT were given thrombolitics (11 reteplase, 6 urokinase, and 1 tenecteplase). Five IAT patients underwent embolectomy with the MERCI retriever system (Concentric Medical, Inc). Two IAT patients received balloon angioplasty and intracranial stent placement.

The timing parameters (eg, door-to-needle time), incidence of sICH, and clinical outcomes are displayed in Figure 1. Seven of the 46 WUS patients died: 3 deaths were the result of malignant MCA syndrome, 1 was from a basilar thrombosis that led to massive brain stem infarction, and 1 patient was given comfort care per family wishes after large right hemispheric infarction unresponsive to tPA and no hemorrhage. The remaining 2 patients died from sICH.

**Thrombolysed vs Nonthrombolysed WUS**

After controlling for the baseline NIHSS imbalance, thrombolysed WUS patients had higher rates of excellent (OR, 9.0; 95% CI, 0.9–90; P=0.06) and favorable outcomes (OR, 9.2; 95% CI, 0.1.9–45; P=0.006). In unadjusted (χ²) analysis, thrombolysed WUS had a nonsignificant increase in sICH (P=0.64), but a higher rate of death (P=0.02). A regression to control for the NIHSS imbalance was not able to be performed because the nonthrombolysed WUS patients experienced zero deaths.

Twenty-three of the 46 thrombolysed WUS (50%) were treated within 3 hours of awakening. There were no significant differences in outcomes between those thrombolysed within 3 hours of awakening with their deficits compared to those treated after 3 hours (sICH: P=0.76; excellent outcome: P=0.31; favorable outcome: P=0.54; death: P=0.5). Eleven patients had mismatch on multimodal imaging, 4 of whom had an excellent or favorable clinical outcome. There were no significant differences between treated WUS with mismatch compared to all other treated WUS patients in any of the outcome measures (sICH: P=0.44; excellent outcome: P=0.61; favorable outcome: P=0.62; death: P=0.57). Similarly, the presence of hypodensity on CT scan did not influence the study outcome measures (sICH: P=0.36; ex-
excellent outcome: \(P=0.32\); favorable outcome: \(P=0.56\); death: \(P=0.61\).

Two patients (both received IV tPA alone) experienced sICH: 1 patient presented with a left MCA occlusion and baseline NIHSS of 20 and the other presented with a right MCA occlusion and NIHSS of 14. No sICH occurred in IAT-treated patients.

Thrombolysed WUS vs 0- to 3-Hour IV tPA Patients
After controlling for the higher median NIHSS scores in the thrombolysed WUS patients, treated WUS experienced similar rates of excellent outcomes (14% vs 32%; OR, 0.48; 95% CI, 0.18–1.27; \(P=0.14\)) and favorable outcomes (28% vs 48%; OR, 0.64; 95% CI, 0.3–1.38; \(P=0.64\)) compared to the standard-of-care 0- to 3-hour IV tPA-treated patients.

WUS patients who received only IV tPA (\(n=28\)) were compared to the 0- to 3-hour standard of care group (Table 3). IV tPA-treated WUS patients had longer onset-to-needle and door-to-needle times, but after controlling for baseline NIHSS, still experienced similar rates of excellent outcomes (19% vs 32%; OR, 0.57; 95% CI, 0.19–1.68; \(P=0.31\)) and favorable outcomes (33% vs 48%; OR, 0.63; 95% CI, 0.25–1.57; \(P=0.32\); Figure 2).

Discussion
Our study details 1 center’s experience with off-label thrombolysis in patients with WUS. Although these patients have historically been denied thrombolysis and have limited treatment options, some WUS patients may be candidates for treatment. In fact, other stroke centers have shown similar rates of poor outcome between WUS and non-WUS patients and suggest that WUS patients should be a target for further therapeutic considerations in the future.\(^{16}\) Whereas other investigators have described clinical and radiographic similarities between WUS and known-onset stroke patients, our study is the first to our knowledge to present the outcomes of a cohort of WUS patients treated with thrombolysis. In our study, WUS patients had more severe strokes compared with the NINDS tPA arm (median NIHSS, 16 vs 14) and our cohort of IV 0- to 3-hour tPA-treated patients (16 vs 11). The etiology in our WUS patients was predominantly cardioembolic (43%). This is in contrast to other published reports\(^{2,4}\) that noted only 17% to 19% cardioembolic strokes in WUS. Our incidence of cardioembolic stroke may be higher because of the small numbers of patients in the study, and because compassionate thrombolysis was offered to patients who have moderate to severe strokes without treatment options.

Because previous studies have reported an early-morning predominance of stroke and similar rates of MRI mismatch compared to those of strokes of known onset, there is evidence to suggest that WUS may occur on awakening. Therefore, rapid institution of thrombolysis for WUS may be a consideration similar to strokes of known onset. In our small cohort, the incidence of sICH was within the range of previous IV tPA studies.\(^{1,18–20}\) Importantly, this incidence takes into account both IV and endovascular approaches.

Twenty-six of the 28 (93%) IV tPA-alone WUS patients received treatment within 4.5 hours of awakening. Door-to-needle times were prolonged in our cohort when compared with current treatment guidelines that mandate <60 minutes.\(^{21}\) Delays may have been attributable to many factors, including slower response of emergency personnel/stroke team activation after patients were confirmed to have an unknown onset time, inability to consent aphasic patients, difficulty reaching next-of-kin, or obtaining supplemental neuroimaging. If thrombolysis of WUS is to be prospectively studied, techniques to reduce these delays need to be under-
taken. Nevertheless, it is possible that patients may still derive benefit from IV tPA when treated within 4.5 hours of symptom onset.22

Despite significantly higher NIHSS scores, treated WUS patients as a group experienced better clinical outcomes compared to nontreated WUS patients; however, these results must be interpreted with caution because of the low numbers of patients and retrospective nature of this study. Higher

decision-making in the absence of data from randomized studies, the decisions to pursue such approaches rests with the individual physician who has a duty to explain to the patient or the patient’s legal decision-maker about the potential benefits vs the risks of invasive diagnostic studies such as cerebral angiogram and experimental treatments.

Some investigators may feel that the pursuit of invasive angiography and intra-arterial recanalization techniques in stroke patients with unknown onset time raises ethical considera-
tions. In the absence of data from randomized studies, the decisions to pursue such approaches rests with the individual physician who has a duty to explain to the patient or the patient’s legal decision-maker about the potential benefits vs the risks of invasive diagnostic studies such as cerebral angiogram and experimental treatments.

This study has many limitations, including its retrospective nature and small numbers. The nonrandomized nature of our study limits the general application of our findings to all WUS patients. Furthermore, lack of long-term outcome data (eg, 90 days) limits comparison with other clinical trial results. The higher incidence of cardioembolic strokes in WUS patients compared to that in other studies indicates that there was a selection bias in our cohort. This bias is a major limitation to our study. Finally, treatment occurred at an experienced academic center and may not be applicable to the community setting.

In conclusion, although thrombolysis may be safe in WUS patients, future, well-designed, prospective studies with pre-specified inclusion and exclusion criteria and treatment strategy are needed to further assess the safety of thrombolysis in WUS patients. For research purposes, those interested in offering tPA to WUS patients should proceed only with an

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<th>Table 3. Baseline Clinical and Radiographic Characteristics of IV tPA-Treated Alone WUS vs 0- to 3-Hour IV tPA-Treated Ischemic Strokes</th>
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<td>Early changes on baseline CT (%)</td>
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<td>Hypodensity on baseline CT (%)</td>
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Figure 2. Timing parameters and clinical outcomes of IV-tPA treated WUS and 0- to 3-hour IV-tPA treated ischemic strokes.

* Controlled for baseline NIHSS
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Disclosures
None.

References
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