Transient Global Amnesia and the Risk of Ischemic Stroke

Atul Mangla, MBBS; Babak B. Navi, MD; Kelly Layton, MPH; Hooman Kamel, MD

**Background and Purpose**—Whether transient global amnesia (TGA) represents an arterial insult that heralds ischemic stroke remains unclear. Therefore, we examined stroke risk after TGA in a population-based cohort.

**Methods**—After performing chart review at our institution to validate the International Classification of Diseases, 9th Edition, Clinical Modification diagnosis code for TGA, we used administrative claims data to identify all patients discharged from nonfederal California emergency departments or acute care hospitals between 2005 and 2010 with a primary discharge diagnosis of TGA. Patients with a primary discharge diagnosis of migraine, seizure, or transient ischemic attack were included as controls. Kaplan–Meier statistics were used to calculate rates of ischemic stroke, and Cox proportional hazards analyses were used to compare stroke risk among the 4 exposure groups while controlling for traditional stroke risk factors.

**Results**—International Classification of Diseases, 9th Edition, Clinical Modification code 437.7 had a sensitivity of 86% and a specificity of 95% for TGA. The cumulative 1-year rate of stroke was 0.54% (95% confidence interval [CI], 0.36–0.81) after TGA, 0.22% (95% CI, 0.20–0.25) after migraine, 0.90% (95% CI, 0.83–0.97) after seizure, and 4.72% (95% CI, 4.60–4.85) after transient ischemic attack. After adjustment for demographic characteristics and stroke risk factors, TGA was not associated with stroke risk when compared with migraine (hazard ratio, 0.82; 95% CI, 0.61–1.10). The likelihood of stroke after TGA was lower than after seizure (hazard ratio, 0.57; 95% CI, 0.44–0.76) or transient ischemic attack (hazard ratio, 0.27; 95% CI, 0.20–0.35).

**Conclusions**—Compared with patients diagnosed with migraine or seizure, patients diagnosed with TGA do not seem to face a heightened risk of stroke. (Stroke. 2014;45:389-393.)

Key Words: amnesia, transient global ■ stroke

Transient global amnesia (TGA) is characterized by a sudden deficit of anterograde and retrograde memory that usually lasts for a few hours and is not accompanied by other focal neurological symptoms or signs. Fisher and Adams introduced the term in 1964, although Bender and separately Guyotat and Courjon had initially described a syndrome consistent with TGA in 1956. The incidence of TGA ranges from 3 to 8 per 100,000 people per year. Most episodes (75%) occur in people between 50 and 70 years, and TGA rarely occurs in patients <40 years old. Although neurologists generally view TGA as a benign entity, its exact prognosis remains unclear. Although many studies have supported the benign nature of TGA, others have suggested that TGA may be a vascular prelude that confers the same risk of stroke as transient ischemic attack (TIA). Many of these studies were performed decades ago, and the diagnosis of vascular events such as stroke or TIA has improved since then because of better imaging and clearer definitions. Furthermore, the majority of studies on the outcome of TGA have had low statistical power, with an average of 70 patients among the studies cited above and a maximum sample size of 236. It is important to resolve clearly whether TGA is a benign entity or a stroke risk factor because uncertainty about the pathogenesis and prognosis of TGA drives potentially unnecessary stroke evaluations, especially when vascular risk factors are present. Therefore, to build on previous studies and confirm that TGA confers a low risk of subsequent stroke, we performed a retrospective cohort study using statewide administrative claims data from across California.

**Methods**

**Design**

We examined the risk of ischemic stroke after a diagnosis of TGA using statewide administrative claims data collected by the California Office for Statewide Healthcare Planning and Development. All nonfederal acute care emergency departments (EDs) and hospitals in California use standardized methods to collect and electronically submit discharge data about all patient visits and hospitalizations. These data are checked for inconsistent or invalid elements and codes before being provided in a deidentified format to the Healthcare Cost and Utilization Project of the Agency for Healthcare Research and Quality. An anonymous, patient-specific record number allows longitudinal tracking of patients across visits.

**Validation of the TGA Diagnosis Code**

Because the accuracy of the International Classification of Diseases, 9th Edition, Clinical Modification (ICD-9-CM) code for TGA (437.7) has not been validated, we first performed a retrospective chart review at our institution to determine its sensitivity and specificity. To do so, we identified 25 ED visits or hospitalizations from 2008 through...
2012 with a primary ICD-9-CM discharge diagnosis of 437.7, as well as 25 discharges with other primary cerebrovascular diagnosis codes (eg, TIA) and no TGA code. A single board–certified neurologist (A. Mangla) reviewed the medical records from these 50 encounters while blinded to their ICD-9-CM diagnoses, and adjudicated cases of TGA based on its standard definition as the sudden onset of anterograde and retrograde memory lasting <24 hours and unaccompanied by other neurological deficits. Using his clinical determination of the primary diagnosis as the gold standard, we calculated the sensitivity and specificity of ICD-9-CM code 437.7 for TGA.

**Patient Population**

After validating the TGA diagnosis code, we used it to identify a cohort of all patients discharged from a nonfederal California ED or acute care hospital between 2005 and 2010 with a primary discharge diagnosis of TGA.

As control groups, we identified all patients discharged from a nonfederal California ED or acute care hospital from 2005 through 2010 with a primary discharge diagnosis of migraine (ICD-9-CM code 346.x), seizure (345.x), or TIA (435.x). We selected migraine and seizure as negative controls given that these are paroxysmal neurological complaints referable to the central nervous system and are not typically treated as stroke heralds, and we chose TIA as a positive control given its well-established risk of subsequent stroke.27

In all 4 groups, we excluded patients with any concurrent secondary diagnoses of ischemic stroke (ICD-9-CM code 433.x1, 434.x1, or 436), intracerebral hemorrhage (431), or subarachnoid hemorrhage (430) at baseline. Furthermore, to focus our analysis on incident cases of stroke, we excluded patients with any diagnoses of ischemic stroke at visits before baseline. Finally, to maximize our ability to ascertain follow-up events, we excluded patients who were not California residents at baseline.

**Measurements**

The primary outcome was any hospital discharge diagnosis of ischemic stroke (ICD-9-CM code 433.x, 434.x, or 436) without concurrent codes for rehabilitation (V57), trauma (800–804 or 850–854), intracerebral hemorrhage (431), or subarachnoid hemorrhage (430) at baseline. Furthermore, to focus our analysis on incident cases of stroke, we excluded patients with any diagnoses of ischemic stroke at visits before baseline. Finally, to maximize our ability to ascertain follow-up events, we excluded patients who were not California residents at baseline.

**Statistical Analysis**

We used standard descriptive statistics to report and compare proportions with exact binomial confidence intervals (CI) and means with SD. Kaplan–Meier survival statistics were used to calculate cumulative rates of stroke. Patients entered observation at the first visit that resulted in a discharge diagnosis of TGA, migraine, seizure, or TIA, and were censored at the time of stroke, in-hospital death, or December 31, 2011. Patients were allowed multiple concurrent exposures (eg, a patient discharged with TGA might at a later encounter also receive a diagnosis of migraine), except that because TIA is a well-established herald of stroke,27 patients with TGA, migraine, and seizure had these time-varying covariates set to zero if they developed a TIA. Because we only considered exposures occurring through the end of 2010, the data from 2011 ensured ≥1 year of follow-up for all patients. Multivariable Cox proportional hazards models were used to examine the association between our exposures and outcome while controlling for potential confounders. Because our goal was to isolate the relationship between our exposures and outcome, all covariates were left in the model regardless of statistical significance. We performed tests of interaction to explore the effects of concurrent exposures (eg, patients with both TGA and migraine).

To assess the robustness of our results in the face of changes to the structure of our baseline analyses, we additionally performed stratified analyses that defined exposures based on ED discharges only versus hospital discharges only. Furthermore, we performed sensitivity analyses using a more sensitive definition of ischemic stroke that did not exclude patients with concurrent rehabilitation, trauma, or hemorrhage codes.

All analyses were performed using Stata MP (version 12, College Station, TX). The threshold of statistical significance was \(\alpha=0.05\). Our institutional review board certified our analysis of the publicly available and deidentified California data set as exempt from review, and separately approved the validation of ICD-9-CM codes via retrospective chart review at our institution.

**Results**

Validation of ICD-9-CM code 437.7 via medical record review indicated that this code has a sensitivity of 86% (95% CI, 67–96) and a specificity of 95% (95% CI, 77–100) for TGA. Across California between 2005 and 2010, we identified 4299 eligible patients with TGA, 170,000 with migraine, 71,087 with seizure, and 115,105 eligible patients with TIA. At baseline, patients with TGA were older than those with migraine or seizure and younger than those with TIA (Table 1). Their burden of cardiovascular comorbidities and risk factors was much lower than those with TIA, generally similar to those with seizure, and much higher than those with migraine (Table 1).

The cumulative 1-year rate of ischemic stroke after TGA was 0.54% (95% CI, 0.36–0.81) compared with 0.22% (95% CI, 0.20–0.25) after migraine, 0.90% (95% CI, 0.83–0.97) after seizure, and 4.72% (95% CI, 4.60–4.85) after TIA (Figure). Cumulative 5-year rates of ischemic stroke were 2.44% (95% CI, 1.86–3.21) after TGA, 0.86% (95% CI, 0.80–0.91) after migraine, 2.90% (95% CI, 2.72–3.10) after seizure, and 12.23% (95% CI, 12.00–12.46) after TIA.

After adjustment for demographic characteristics and traditional stroke risk factors (Table 2), TGA was not associated with stroke when compared with migraine (hazard ratio, 0.82; 95% CI, 0.61–1.10). The likelihood of stroke after TGA was lower than after seizure (hazard ratio, 0.57; 95% CI, 0.44–0.76) or TIA (hazard ratio, 0.27; 95% CI, 0.20–0.35). We found no significant evidence of interaction between TGA and the control exposures.

The results of our multivariable analyses did not substantially change when we stratified exposures by ED visits only or hospitalizations only, or when we performed sensitivity analyses using a more sensitive definition of ischemic stroke.

**Discussion**

In a large and demographically heterogeneous population, we found that patients discharged from an ED or hospital with a primary diagnosis of TGA faced a low risk of subsequent ischemic stroke. Their risk was similar to the risk of stroke after a diagnosis of migraine, lower than the risk after presentation with seizure, and much lower than the risk after TIA.

Our results are consistent with several smaller studies that suggest that TGA is benign in terms of future vascular
and our results do not support other studies that suggest that TGA heralds stroke in a similar fashion as TIA. Furthermore, our findings do not support the proposed pathogenetic explanation of TGA as a form of arterial ischemia. Our findings are not inconsistent with alternative and still unproven explanations of TGA as a manifestation of venous insufficiency or migrainous phenomena, but our epidemiological analysis cannot further inform the debate about the exact pathogenesis of TGA. From a practical standpoint, our study suggests that patients with classical presentations of TGA do not require a vascular evaluation or any specific intensification of stroke risk factor treatment, even if they have vascular comorbidities.

The results of our study should be interpreted in light of several limitations. First, our analysis relied on administrative data and ICD-9-CM diagnosis codes without detailed clinical data, and therefore may be susceptible to incorrect or biased ascertainment of TGA cases; we also lacked access to ICD-10 codes, which are likely to offer advantages such as greater specificity. This concern is somewhat mitigated by our validation of the TGA ICD-9-CM code via detailed chart review, whose results indicate that cases identified on the basis of this diagnosis code are likely to be representative of patients evaluated in an ED or hospital for TGA. Second, our use of administrative data may have led to underascertainment of strokes. However, the rate of stroke after TIA in our sample is broadly consistent with rates in recent population-based prospective studies, suggesting adequate capture of incident strokes in our study. Furthermore, even if our overall ascertainment of strokes were insensitive, there is no obvious reason to suspect differential misclassification of strokes in patients with TGA versus migraine or seizure, which supports our finding that TGA is not a risk factor for stroke compared with these negative controls. Third, this was a hospital-based study in that we lacked access to

Table 1. Baseline Characteristics of Patients with TGA Compared With Negative Controls (Migraine and Seizure) and Positive Controls (TIA)*

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>TGA (N=4299)</th>
<th>Migraine (N=170 400)</th>
<th>Seizure (N=71 087)</th>
<th>TIA (N=115 105)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, mean (SD), y</td>
<td>65 (13)</td>
<td>38 (12)</td>
<td>44 (16)</td>
<td>71 (15)</td>
</tr>
<tr>
<td>Female</td>
<td>2239 (52.1)</td>
<td>140120 (82.2)</td>
<td>32695 (46.0)</td>
<td>65067 (56.5)</td>
</tr>
<tr>
<td>Race</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>3202 (79.4)</td>
<td>92705 (58.0)</td>
<td>37328 (55.5)</td>
<td>73869 (67.8)</td>
</tr>
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<td>Black</td>
<td>94 (2.3)</td>
<td>16643 (10.4)</td>
<td>10011 (14.9)</td>
<td>8113 (7.4)</td>
</tr>
<tr>
<td>Hispanic</td>
<td>367 (9.1)</td>
<td>41047 (25.7)</td>
<td>16016 (23.8)</td>
<td>18664 (17.1)</td>
</tr>
<tr>
<td>Other</td>
<td>368 (9.1)</td>
<td>9315 (5.8)</td>
<td>3946 (5.9)</td>
<td>8265 (7.6)</td>
</tr>
<tr>
<td>Payment source</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Medicare</td>
<td>1752 (40.8)</td>
<td>13905 (8.2)</td>
<td>20559 (28.9)</td>
<td>73924 (64.2)</td>
</tr>
<tr>
<td>Medicaid</td>
<td>81 (1.9)</td>
<td>35371 (20.8)</td>
<td>16571 (23.3)</td>
<td>7066 (6.1)</td>
</tr>
<tr>
<td>Private</td>
<td>2184 (50.8)</td>
<td>89105 (52.3)</td>
<td>18414 (25.9)</td>
<td>27581 (24.0)</td>
</tr>
<tr>
<td>Self-pay</td>
<td>184 (4.3)</td>
<td>23036 (13.5)</td>
<td>10934 (15.4)</td>
<td>3571 (3.1)</td>
</tr>
<tr>
<td>Other</td>
<td>97 (2.3)</td>
<td>8929 (5.2)</td>
<td>4597 (6.5)</td>
<td>2938 (2.6)</td>
</tr>
<tr>
<td>Hypertension</td>
<td>1792 (41.7)</td>
<td>16740 (9.8)</td>
<td>15424 (21.7)</td>
<td>72206 (62.7)</td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>380 (8.8)</td>
<td>6243 (3.7)</td>
<td>7473 (10.5)</td>
<td>28160 (24.5)</td>
</tr>
<tr>
<td>CHD</td>
<td>314 (7.3)</td>
<td>1227 (0.7)</td>
<td>3279 (4.6)</td>
<td>21242 (18.5)</td>
</tr>
<tr>
<td>Atrial fibrillation</td>
<td>172 (4.0)</td>
<td>376 (0.2)</td>
<td>1599 (2.2)</td>
<td>12301 (10.7)</td>
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<tr>
<td>CHF</td>
<td>53 (1.2)</td>
<td>379 (0.2)</td>
<td>1581 (2.2)</td>
<td>7514 (6.5)</td>
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<tr>
<td>CKD</td>
<td>71 (1.7)</td>
<td>609 (0.4)</td>
<td>2304 (3.2)</td>
<td>7004 (6.1)</td>
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<tr>
<td>COPD</td>
<td>73 (1.7)</td>
<td>1158 (0.7)</td>
<td>2376 (3.3)</td>
<td>7534 (6.5)</td>
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<tr>
<td>PVD</td>
<td>56 (1.3)</td>
<td>181 (0.1)</td>
<td>664 (0.9)</td>
<td>4619 (4.0)</td>
</tr>
</tbody>
</table>

CHD indicates coronary heart disease; CHF, congestive heart failure; CKD, chronic kidney disease; COPD, chronic obstructive pulmonary disease; PVD, peripheral vascular disease; SD, standard deviation; TGA, transient global amnesia; and TIA, transient ischemic attack.

*Data are presented as number (%) of participants unless otherwise specified.

![Figure](http://stroke.ahajournals.org/ Downloaded from)
ambulatory care visits, and therefore our results cannot be
generalized to patients presenting with TGA in such settings.
However, this would have biased our study in a conservative
direction because patients diagnosed with TGA in the ambu-
latory setting would most likely have been less severely ill
than those who presented to the hospital, and therefore
be unlikely to face a higher risk of stroke.

In summary, patients with TGA do not seem to face a
heightened risk of ischemic stroke compared with patients
diagnosed with migraine or seizure. Our findings may be
helpful in reassuring patients with this dramatic and dis-
tressing syndrome that their prognosis is good, and in reas-
suring clinicians that these patients do not need extensive
testing or specific therapy if the clinical evaluation is con-
sistent with TGA.

Sources of Funding
B.B. Navi receives research funding from the National Institutes
of Health (NIH) (KL2TR000458, U01NS062835). H. Kamel receives re-
search funding from the NIH (K23NS082367).

Disclosures
None.

Table 2. Baseline Characteristics of Patients, Stratified by

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Stroke (N=11674)</th>
<th>No Stroke (N=349217)</th>
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</thead>
<tbody>
<tr>
<td>Age, mean (SD), y</td>
<td>71.4 (14.7)</td>
<td>50.1 (20.2)</td>
</tr>
<tr>
<td>Female</td>
<td>6699 (57.4)</td>
<td>233422 (66.8)</td>
</tr>
<tr>
<td>Race</td>
<td></td>
<td></td>
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<tr>
<td>White</td>
<td>7185 (65.0)</td>
<td>199919 (60.8)</td>
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<tr>
<td>Black</td>
<td>1225 (11.1)</td>
<td>33636 (10.2)</td>
</tr>
<tr>
<td>Hispanic</td>
<td>1986 (17.2)</td>
<td>74198 (22.6)</td>
</tr>
<tr>
<td>Other</td>
<td>741 (6.7)</td>
<td>21153 (6.4)</td>
</tr>
<tr>
<td>Payment source</td>
<td></td>
<td></td>
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<tr>
<td>Medicare</td>
<td>7865 (67.4)</td>
<td>102275 (29.3)</td>
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<tr>
<td>Medicaid</td>
<td>1045 (8.9)</td>
<td>58044 (16.6)</td>
</tr>
<tr>
<td>Private</td>
<td>2049 (17.6)</td>
<td>135235 (38.7)</td>
</tr>
<tr>
<td>Self-pay</td>
<td>419 (3.6)</td>
<td>37306 (10.7)</td>
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<tr>
<td>Other</td>
<td>293 (2.5)</td>
<td>16268 (4.7)</td>
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<tr>
<td>Hypertension</td>
<td>7231 (61.9)</td>
<td>98931 (28.3)</td>
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<td>Diabetes mellitus</td>
<td>3250 (27.8)</td>
<td>39006 (11.2)</td>
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<td>CHD</td>
<td>2305 (19.7)</td>
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<td>Atrial fibrillation</td>
<td>1518 (13.0)</td>
<td>12930 (3.7)</td>
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<td>921 (7.9)</td>
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<td>800 (6.9)</td>
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<td>COPD</td>
<td>759 (6.5)</td>
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<tr>
<td>PVD</td>
<td>475 (4.1)</td>
<td>5045 (1.4)</td>
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</tbody>
</table>

CHD indicates coronary heart disease; CHF, congestive heart failure; CKD, chronic kidney disease; COPD, chronic obstructive pulmonary disease; PVD, peripheral vascular disease; and SD, standard deviation.

*Data are presented as number (%) of participants unless otherwise specified. P<0.001 for all comparisons.

References


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*Stroke*. 2014;45:389-393; originally published online December 5, 2013;
doi: 10.1161/STRK.113.003916

*Stroke* is published by the American Heart Association, 7272 Greenville Avenue, Dallas, TX 75231
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Print ISSN: 0039-2499. Online ISSN: 1524-4628

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