Quantitative Intracerebral Hemorrhage Localization

John Muschelli, ScM; Natalie L. Ullman, BS; Elizabeth M. Sweeney, ScM; Ani Eloyan, PhD; Neil Martin, MD; Paul Vespa, MD; Daniel F. Hanley, MD; Ciprian M. Crainiceanu, PhD

**Background and Purpose**—The location of intracerebral hemorrhage (ICH) is currently described in a qualitative way; we provide a quantitative framework for estimating ICH engagement and its relevance to stroke outcomes.

**Methods**—We analyzed 111 patients with ICH from the Minimally Invasive Surgery Plus Recombinant-Tissue Plasminogen Activator for Intracerebral Evacuation (MISTIE) II clinical trial. We estimated ICH engagement at a population level using image registration of computed tomographic scans to a template and a previously labeled atlas. Predictive regions of National Institutes of Health Stroke Scale and Glasgow Coma Scale stroke severity scores, collected at enrollment, were estimated.

**Results**—The percent coverage of the ICH by these regions strongly outperformed the reader-labeled locations. The adjusted \( R^2 \) almost doubled from 0.129 (reader-labeled model) to 0.254 (quantitative location model) for National Institutes of Health Stroke Scale and more than tripled from 0.069 (reader-labeled model) to 0.214 (quantitative location model). A permutation test confirmed that the new predictive regions are more predictive than chance: \( P<0.001 \) for National Institutes of Health Stroke Scale and \( P<0.01 \) for Glasgow Coma Scale.

**Conclusions**—Objective measures of ICH location and engagement using advanced computed tomographic imaging processing provide finer, objective, and more quantitative anatomic information than that provided by human readers.

**Clinical Trial Registration**—URL: http://www.clinicaltrials.gov. Unique identifier: NCT00224770.

(Stroke. 2015;46:3270-3273. DOI: 10.1161/STROKEAHA.115.010369.)

**Key Words:** cerebral hemorrhage • Glasgow Coma Scale • stroke • tomography, x-ray computed

Intracerebral hemorrhage (ICH) results from a blood vessel rupturing into the brain. Quantification of hemorrhage location using X-ray computed tomography is complicated as ICH may extend into multiple brain areas, distend tissues, and break through the ventricular wall. Current practice is qualitative and may extend into multiple brain areas, distend tissues, and break through the ventricular wall. Predictive regions of National Institutes of Health Stroke Scale and Glasgow Coma Scale stroke severity scores, collected at enrollment, were estimated.

**Methods**

The population studied consists of 111 patients from the Minimally Invasive Surgery Plus Recombinant-Tissue Plasminogen Activator for Intracerebral Evacuation (MISTIE) trial recruited from 26 centers with lobar and deep ICHs ≥20 mL in volume. This sample contained 35 women and had mean (SD) age of 60.8 (11.2) years. Diagnostic computed tomographic images were acquired under a standard protocol but with differences across sites. Scans were acquired using GE (n=46), Siemens (n=27), Philips (n=20), and Toshiba (n=8) scanners had gantry tilt (n=87) and the slice thickness of the image varied within some scans (n=14). Therefore, scans had different voxel dimensions and image resolution before template registration.

ICH was manually segmented using OsiriX (v.4.1, Pixmeo; Geneva, Switzerland) by expert readers. Readers identified the anatomic location most engaged by the ICH: putamen (n=68), lobar (n=33), globus pallidus (n=6), and thalamus (n=4). The initial mean (SD) ICH volume of this sample was 37.4 (20.1) mL. The brain image was spatially registered to a computed tomographic template using the Clinical toolbox and the statistical parametric mapping software (SPM8, Wellcome Trust Center for Neuroimaging, London, United Kingdom) in MATLAB (Mathworks, Natick, MA). The binary hemorrhage mask was transformed into the template space. No scans were excluded because of inadequate registration, determined by visual inspection.

**ICH Localization and Engagement**

After registration, all scans and hemorrhage masks are in the same template space. We calculated the percentage of patients with hemorrhage at each voxel in template space, resulting in a 3D histogram of ICH presence. The Eve atlas, located in template space, labels gray matter and white matter regions. Ventricular regions were not explicitly segmented; any region not classified was labeled cerebrospinal fluid. Using Eve, we calculated the percent of the ICH engaged by region (eg, putamen engages 20% of the ICH).
Prediction of Severity Score Based on Hemorrhage Location

We investigated the prediction performance of ICH region engagement with the following stroke severity scores collected at enrollment in the trial: National Institutes of Health Stroke Scale (NIHSS) and Glasgow Coma Scale (GCS). We analyzed voxels in the template space, where at least 10 patients exhibit ICH ($V=166,202$).

At each voxel, we tested if the mean score (NIHSS or GCS) was different in patients with ICH at each voxel, giving a voxel-wise $P$ value. We selected voxels, referred as highest predictive regions (HPR), based on the 1000, 2000, or 3000 smallest $P$ values or 3 different $P$ value thresholds: 0.05, 0.01, and 0.001. Selected voxels are referred as HPR.

For each HPR, we calculated the overlap of the patient-level ICH and the HPR, which we will call HPR coverage. If ICH in a scan covers the entire HPR, coverage is 100%; 0% coverage indicates no overlap. For each HPR coverage and severity score, we fit the following linear model:

$$\text{Stroke severity}_i = \beta_0 + \beta_1 \text{HPR coverage}_i + \beta_2 \text{Age}_i + \beta_3 \text{Sex}_i + \beta_4 \text{TotalVol}_i + \epsilon_i$$

where $i=$scan. We fit the same model replacing HPR coverage, with a categorical indicator for each reader-labeled ICH location: thalamus, globus pallidus, putamen, or lobar.

For each model, we estimated the adjusted $R^2$. The HPR with the highest adjusted $R^2$ was selected and compared with the adjusted $R^2$ for the reader-labeled model. We calculated engagement of these HPR with the Eve atlas–labeled neuroanatomic regions.

To determine whether these regions were predictive above chance, a permutation test was conducted: the severity score was randomly permuted, an HPR was estimated using the permuted severity scores, and the adjusted $R^2$ was calculated for each permutation. Comparing the permutation distribution of the adjusted $R^2$ with the adjusted $R^2$ obtained from the original data provided the $P$ value.

Results

ICH Localization and Engagement

Figure 1 represents the 3D histogram of hemorrhage prevalence—colors represent the percentage of patients with ICH engagement at that area. ICH is distributed medially in the brain, with a lower concentration at the cortical surface and higher on the left side. Most voxels have a low prevalence of ICH engagement; the median number of patients with ICH at a given voxel is 3 (3%), although a small group of voxels ($V=5685$) has a high prevalence (>40%).

Prediction of Severity Score Based on Hemorrhage Location

Figure 2 displays the HPRs for $P$ values <0.01 for NIHSS (Figure 2A) and the smallest 1000 $P$ values for GCS (Figure 2B). Figure 2C and 2D displays the NIHSS and GCS relationship with HPR coverage. The larger the HPR coverage, the higher (more severe) the NIHSS score, and the lower (deeper unconsciousness) the GCS score.

Adjusted $R^2$ model estimates indicated that HPR coverage models strongly outperform reader-labeled location models: it almost doubled from 0.129 (reader-labeled model) to 0.254 (best coverage model) for NIHSS and more than tripled from 0.069 (reader-labeled model) to 0.214 (best coverage model) for GCS.

The permutation test $P$ values for HPR coverage were <0.001 for NIHSS and <0.01 for GCS; HPRs were more predictive than HPRs obtained by chance.

ICH Localization and Engagement

Table represents the 5 most-engaged regions for the population 3D histogram and the HPRs from the GCS and NIHSS.
The engagement represents the percent engagement of a specific area compared with all areas engaged. The 3D histogram of ICH engages the insular, putamen, and primarily the cerebrospinal fluid, especially the ventricles. The NIHSS HPR engages primarily areas of the internal capsule, thalamus, superior corona radiata, and cerebrospinal fluid regions. The GCS HPR engages primarily the thalamus and superior corona radiata.

Table. Distribution of the Top 5 Areas of Engagement

<table>
<thead>
<tr>
<th>Area</th>
<th>Population Prevalence</th>
<th>NIHSS HPR</th>
<th>GCS HPR</th>
</tr>
</thead>
<tbody>
<tr>
<td>CSF (ventricular and subarachnoid spaces)</td>
<td>7.9</td>
<td>10.9</td>
<td>...</td>
</tr>
<tr>
<td>Insular</td>
<td>7.6</td>
<td>...</td>
<td>...</td>
</tr>
<tr>
<td>Superior temporal gyrus</td>
<td>5.5</td>
<td>...</td>
<td>...</td>
</tr>
<tr>
<td>Putamen</td>
<td>4.8</td>
<td>...</td>
<td>...</td>
</tr>
<tr>
<td>External capsule</td>
<td>3.9</td>
<td>...</td>
<td>...</td>
</tr>
<tr>
<td>Posterior limb of internal capsule</td>
<td>...</td>
<td>12.0</td>
<td>...</td>
</tr>
<tr>
<td>Superior corona radiate</td>
<td>...</td>
<td>11.0</td>
<td>27.9</td>
</tr>
<tr>
<td>Thalamus</td>
<td>...</td>
<td>10.1</td>
<td>33.9</td>
</tr>
<tr>
<td>Caudate nucleus</td>
<td>...</td>
<td>8.4</td>
<td>9.6</td>
</tr>
<tr>
<td>Postcentral WM</td>
<td>...</td>
<td>...</td>
<td>6.7</td>
</tr>
<tr>
<td>Superior longitudinal fasciculus</td>
<td>...</td>
<td>...</td>
<td>5.9</td>
</tr>
</tbody>
</table>

CSF indicates cerebrospinal fluid; GCS, Glasgow Coma Scale; HPR, highest predictive regions; NIHSS, National Institutes of Health Stroke Scale; and WM, white matter.
Discussion
We have characterized the localization of ICH in a population from prospective clinical trial images using a 3D histogram. These 3D histograms provide detailed visualization and allow a finer comparison of ICH location across groups.

We also demonstrate how labeled atlases can automatically describe ICH engagement by neuroanatomic regions at a patient or population level. These measures are more interpretable for clinical relevance and may translate to better determination of disability. The HPR for NIHSS engages primarily areas of the internal capsule, thalamus, superior corona radiata, and cerebellopontine angle. Venkatasubramanian et al\(^7\) found degeneration of the corticospinal tract correlates with motor skill in chronic stroke patients. The corticospinal tract Wallerian degeneration in intracerebral hemorrhage evacuation (MISTIE) clinical trial. Acta Neurochir Suppl. 2008;104:147–151.

The strength of our study is that the MISTIE trial is prospective with standardized protocols. This analysis is limited by the inclusion criteria of the MISTIE trial population: ICH ≥ 20 mL. We have performed permutation testing on these pilot results, but a larger cohort is needed to validate the predictive models and ICH location distribution.

Summary
The summary of Eve atlas ICH engagement provides a more refined description of location than that provided by expert human readers. We have shown that using image-based measurements better predicts initial severity scores compared with using clinical location.

Sources of Funding
The project described was supported by the grant R01EB012547 (National Institutes of Health), T32AG000247 (National Investigation Agency), R01NS046309, R01NS060910, R01NS085211, R01NS046309, U01NS080824, and U01NS062851 (National Institute of Neurological Disorders and Stroke), and R01MH095836 (National Institute of Mental Health).

Disclosures
Johns Hopkins University holds a use patent for intraventricular tissue plasminogen activator. Dr Hanley received significant (over 5%) support on National Institutes of Health Grant R01NS046309. The other authors report no conflicts.

References
Quantitative Intracerebral Hemorrhage Localization
John Muschelli, Natalie L. Ullman, Elizabeth M. Sweeney, Ani Eloyan, Neil Martin, Paul
Vespa, Daniel F. Hanley and Ciprian M. Crainiceanu

Stroke. 2015;46:3270-3273; originally published online October 8, 2015;
doi: 10.1161/STROKEAHA.115.010369
Stroke is published by the American Heart Association, 7272 Greenville Avenue, Dallas, TX 75231
Copyright © 2015 American Heart Association, Inc. All rights reserved.
Print ISSN: 0039-2499. Online ISSN: 1524-4628

The online version of this article, along with updated information and services, is located on the
World Wide Web at:
http://stroke.ahajournals.org/content/46/11/3270

Permissions: Requests for permissions to reproduce figures, tables, or portions of articles originally published
in Stroke can be obtained via RightsLink, a service of the Copyright Clearance Center, not the Editorial Office.
Once the online version of the published article for which permission is being requested is located, click
Request Permissions in the middle column of the Web page under Services. Further information about this
process is available in the Permissions and Rights Question and Answer document.

Reprints: Information about reprints can be found online at:
http://www.lww.com/reprints

Subscriptions: Information about subscribing to Stroke is online at:
http://stroke.ahajournals.org/subscriptions/