The association of insular stroke with lesion volume

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Abstract

The insula has been implicated in many sequelae of stroke. It is the area most commonly infarcted in people with post-stroke arrhythmias, loss of thermal sensation, hospital acquired pneumonia, and apraxia of speech. We hypothesized that some of these results reflect the fact that: (1) ischemic strokes that involve the insula are larger than strokes that exclude the insula (and therefore are associated with more common and persistent deficits); and (2) insular involvement is a marker of middle cerebral artery (MCA) occlusion. We analyzed MRI scans of 861 patients with acute ischemic hemispheric strokes unselected for functional deficits, and compared infarcts involving the insula to infarcts not involving the insula using t-tests for continuous variables and chi square tests for dichotomous variables. Mean infarct volume was larger for infarcts including the insula (n = 232) versus excluding the insula (n = 629): 65.8 ± 78.8 versus 10.2 ± 15.9 cm³ (p < 0.00001). Even when we removed lacunar infarcts, mean volume of non-lacunar infarcts that included insula (n = 775) were larger than non-lacunar infarcts (n = 227) that excluded insula: 67.0 cm³ ± 79.2 versus 11.5 cm³ ± 16.7 (p < 0.00001). Of infarcts in the 90th percentile for volume, 87% included the insula (χ² = 181.8; p < 0.00001). Furthermore, 79.0% infarcts due to MCA occlusion included the insula; 78.5% of infarcts without MCA occlusion excluded the insula (χ² = 93.1; p < 0.0001). The association between insular damage and acute or chronic sequelae likely often reflects the fact that: (1) ischemic strokes that involve the insula are larger (and therefore are less likely to recover from large volume strokes (and are therefore less likely to recover from post-stroke sequelae that have been attributed to the insula can be explained by the fact that patients with strokes involving the insula have larger volume strokes (and are therefore less likely to recover from

1. Introduction

The insula, also known as the insular cortex, is the small part of the brain that lies deep within the lateral sulcus. It is a triangular region, covered by the fronto-parietal and temporal opercula (Varnavas and Grand, 1999). The insula has bidirectional connections with most of the brain, including the frontal, temporal, parietal, cingulate, subcortical structures (basal ganglia and thalamus), and limbic structures (amygdala, periamygdaloid areas, and entorhinal cortex) (Augustine, 1996). The arterial supply of the insula is complex: arteries supplying the insula arise from the middle cerebral artery (MCA), with predominance from the M2 segment and a small contribution from the insular branches of the M1 segment (Türe et al., 2000). Because it receives blood supply from both M1 and M2, complete MCA occlusion may be necessary to produce a substantial infarct.

The insular cortex has been implicated in visceral sensory, visceromotor, and interoceptive functions (Augustine, 1985), and also in complex processes such as emotions, music, and language (Shura et al., 2014). It has been reported that stroke-induced insular damage is associated with a variety of deficits, including disrupted autonomic nervous system (Augustine, 1985; De Raedt et al., 2015; Inamasu et al., 2013), emotion (Calder, 1996; Ibanez et al., 2010), pain functions (Cattaneo et al., 2007; Cereda et al., 2002), apraxia of speech (Dronkers, 1996), other aspects of speech production such as respiration (Ackermann and Riecker, 2010), auditory processing (Bamio et al., 2003), gustatory functions (Flynn, 1999; Small, 2010), and somatosensory function (Stephani et al., 2011). Insular stroke has been associated with cardiac arrhythmias and cardiac death after stroke (Oppenheimer, 2006; Seifert et al., 2015; Tokgozoglu et al., 1999) as well as hospital acquired pneumonia (Kemmling et al., 2013). It is possible that the frequent association between insular damage and at least some of the post-stroke sequelae that have been attributed to the insula can be explained by the fact that patients with strokes involving the insula have larger volume strokes (and are therefore less likely to recover from...
any sequelae), compared to patients with stroke with no insular involvement (Hillis et al., 2004). Previous studies have shown that the insula is frequently involved in any large MCA territory infarct (Nakano et al., 2001; Truwit et al., 1990). However, it has not been shown that strokes involving the insula are larger than other hemispheric strokes. The goal of this study is to test the hypotheses that: (1) ischemic strokes including the insula are larger in volume than other hemispheric strokes; and (2) insular infarct is a marker of MCA occlusion. We tested these hypotheses in a series of 861 patients with acute ischemic hemispheric stroke, unselected for clinical deficits. This study was a retrospective analysis of prospectively collected data.

2. Methods
2.1. Participants

Patients were identified from a prospective database of patients who presented with acute ischemic hemispheric stroke to the Johns Hopkins Medical Institutions stroke service (East Baltimore campus or Johns Hopkins Bayview Medical Center) and consented for a study of language or cognitive deficits associated with areas of acute ischemia. All participants gave informed consent (if they demonstrated intact comprehension), or their closest relative or legal representative consented (if the patient had impaired comprehension) to the study according to the Declaration of Helsinki. The study was approved by the Johns Hopkins Medicine Institutional Review Board.

Inclusion criteria were: (1) symptoms of acute ischemic hemispheric stroke; (2) age 18 or older; (3) able to complete testing within 48 h from onset of symptoms. Exclusion criteria included: (1) contra-indication for MRI; (2) premorbid dementia or other neurological disease; (3) impaired level of consciousness or need for ongoing sedation; (3) hemorrhage; and (4) stroke limited to the brainstem or cerebellum. For this study we also excluded patients who were found to have: (1) chronic stroke; (2) no lesion on diffusion-weighted image (DWI); (3) other (non-stroke) lesions on MRI, such as tumor; and (4) poor quality DWI, preventing reliable measurement of lesion volume. From the database we identified 861 patients with acute strokes unselected for functional deficits. The mean age of participants was 61.1 years (SD of 15.5 years).

2.2. Imaging

MRI scans were obtained within 48 h of symptom onset. Participants underwent the following imaging: T2, fluid attenuation inversion recovery (FLAIR; to evaluate for old lesions), susceptibility weighted images (to evaluate for hemorrhage), PWI (to evaluate for areas of hypoperfusion), and DWI (to evaluate for acute ischemia). DWI scans were 5 mm in thickness and provided whole-brain coverage. MR/CT angiography (to evaluate for stenosis or occlusion) was obtained for 620 patients. Acute DWI scans were evaluated by a technician blinded to clinical data (including MRA/CTA) for presence or absence of involvement of the insula. A neurologist, also blinded to clinical data, evaluated a subset of 620 participants with magnetic resonance angiography (MRA) or computed tomography angiogram (CTA) revealed that 62 (10.0%) patients had MCA occlusion and 63 (10.2%) had MCA stenosis. Among the 62 patients with occlusion of the MCA, 49 (79%) had insular infarcts and 13 (21%) had infarcts not involving the insula ($\chi^2 = 93.1$, df1, $p < 0.0001$; Table 2). Among the 63 patients with MCA stenosis, 29 (46.0%) had infarcts involving the insula and 34 (54.0%) had infarcts not involving the insula. Insular cortex involvement was strongly associated with the presence of either MCA occlusion or stenosis ($\chi^2 = 114.7$, df2, $p < 0.0001$) (Table 3, Fig. 1).

Interestingly, the volume of infarcts was larger for patients with MCA occlusion compared to those without MCA occlusion, but only in patients with insular infarcts (Table 3). This finding should be considered preliminarily because there were only a small number of patients without insular infarcts who had MCA occlusion. But there was no trend for their infarcts to be larger than those of other patients without insular involvement. We can speculate that cases of MCA occlusion without acute insular stroke are those cases in which the occlusion has been gradual, and thus does not cause a large stroke.

We also evaluated whether or not there was any difference in the relationship between volume of infarct and MCA occlusion in left versus right hemisphere stroke, because earlier research has demonstrated a higher frequency of left MCA occlusion relative to right MCA occlusion (Hedna et al., 2013). Among left hemisphere strokes, 26/308 (8.4%)...
had MCA occlusion; not surprisingly, those with MCA occlusion had larger infarcts than those without ($86.8 \pm 107.2$ vs $15.8 \pm 24.4$ cm$^3$; $t = -6.6$; $p < 0.00001$). Among right hemisphere strokes, $32/286$ (11.1%) had MCA occlusion; those with MCA occlusion had larger infarcts than those without ($78.4 \pm 85.4$ vs $19.6 \pm 40.2$ cm$^3$; $t = -6.6$; $p < 0.00001$). Even in those with bilateral strokes, the $4/26$ with MCA occlusion on one side had larger strokes than those without occlusion ($195.1 \pm 2.2$ vs. $11.3 \pm 20.2$; $t = -4.3$; $p = 0.0002$). There was not a significant difference between the frequency of insular involvement between left and right hemisphere stroke (36.0 vs. 37.6%). The association between MCA occlusion and insular involvement was strong for both left hemisphere stroke ($\chi^2 = 24.1$; $p < 0.0001$) and right hemisphere stroke ($\chi^2 = 68.5$; $p < 0.0001$).

### 4. Discussion

In a relatively large cohort of patients with acute ischemic hemispheric stroke, we demonstrate that the mean volume of stroke with insular involvement is significantly larger than the mean volume of strokes without insular involvement. Moreover, the largest strokes, and those due to MCA occlusion, usually involved the insula. The large lesion volume in strokes affecting the insula could explain poor clinical outcomes and persistent deficits compared to strokes sparing the insula. Previous studies that indicate acute infarction measuring $>70$ cm$^3$ predicts poor outcome in acute stroke setting (Yoo et al., 2009; Yoo et al., 2010).

These data illustrate a pitfall in lesion-overlap studies of patients with post-stroke deficits; the area of greatest overlap in any deficit due to MCA stroke is likely to be in the insula. Simply overlapping lesions to determine if the same structure is involved in all the strokes in patients with a particular deficit is not a valid method of determining structure–function relationships (Rorden and Karnath, 2004). Furthermore, these data also call for caution in interpreting other lesion-deficit association studies of acute or chronic stroke (e.g. using voxel-symptom lesion mapping), because there will nearly always be greatest power to detect an association between the deficit and the lesion in the insula. For example, one study found that hospital acquired pneumonia (HAP) was associated with damage to the insula (Kemmling et al., 2013); this result may reflect the strong association between damage to the insula (large MCA volume strokes) and persistent deficits in mobility and other known risk factors for HAP. Patients with and without HAP in that study were matched for NIHSS scores, but there was a significantly greater percentage of left hemisphere strokes among those without HAP; the equivalent NIHSS score is associated with smaller volume of lesion in left hemisphere compared to right hemisphere stroke (Fink et al., 2002; Gottesman et al., 2009; Menezes et al., 2007; Woo et al., 1999). This bias can account for the fact that only the right insula (included in the largest strokes) was associated with HAP.

### Tables

| Table 1 | Demographic information (mean age 61.1(±SD 15.5)). |
|---------|---------------------------------------------------|
|          | Sex          | Male | Female | Race          | African Americans | Caucasians | Other | Hemisphere involved | Left | Right | Bilateral |
| All infarcts |              | 49.6%| 50.4% |              |                |             |      |                   |      |      | 3.3%      |
| Lacunar infarcts |        | 56.5%| 43.5% |              |                |             |      |                   |      |      | 6.7%      |

| Table 2 | MCA occlusion and insular infarct ($\chi^2 = 93.1, df_1, p < 0.0001$). |
|---------|-----------------------------------------------------------------------|
|          | MCA patent | MCA occluded | Difference in volume between occluded vs non-occluded MCA | Total         |
| No insular infarct | N = 438 | 9.1 ± 14.6 cm$^3$ | ns | 451 9.1 ± 14.5 cm$^3$ |
| Insular infarct  | N = 120 | 47.3 ± 54.8 cm$^3$ | N = 49 110.6 ± 111.2 cm$^3$ | t = −4.93 $p = 0.00001$ | 169 65.8 ± 80.5 cm$^3$ |
| Difference in volume for insular vs no insular infarct | t = −8.71 $p < 0.00001$ | t = −3.24 $p = 0.0019$ | 620 |

| Table 3 | MCA occlusion/stenosis and insular infarct ($\chi^2 = 114.7, df_2, p < 0.0001$). |
|---------|---------------------------------|
|          | MCA patent | MCA occluded | MCA stenosed | Total |
| No insular infarct | 404 | 13 | 34 | 451 |
| Insular infarct  | 91 | 49 | 29 | 169 |
| 495 | 62 | 63 | 620 |
infarct has also been found to be associated with a wide range of chronic motor, sensory and cognitive sequelae (Ibanez et al., 2010). But in studies that show the overlap of lesions of all participants, the insula is often the area with most lesions (e.g. (Hope et al., 2013; Wu et al., 2015), and thus the greatest power to detect the association. Therefore, it is essential to control for volume of infarct as well as demographic factors that may influence the outcome (e.g. (Hope et al., 2013; Wu et al., 2015).

Isolated infarcts of the insula generally produce minor and transient deficits in somatosensory, gustatory, vestibular, or cardiovascular functions (Boucher et al., 2015; Cereda et al., 2002; Frontzek et al., 2014; von Brevern et al., 2014). Transient neuropsychological disorders, including aphasia (left posterior insula), dysarthria, and somatoparaphrenia have also been reported in acute insular stroke (Cereda et al., 2002). However, some reported deficits observed in association with acute insular infarct may be due to hypoperfusion of the MCA territory, as illustrated in Fig. 2. Further investigation of deficits associated with isolated insular lesions (without cortical hypoperfusion or damage to surrounding areas) is needed to determine if the insula itself is essential for many of the functions that have been found to be impaired when it is damaged. Outcome in strokes restricted to the insula appears to be uniformly good, given the transient nature of the reported deficits; but long term outcome studies of patients with such lesions (particularly with detailed evaluation of sensory, emotional, vestibulatory, and gustatory functions) are lacking. Given that the insula is highly connected to many vestibular, somatosensory, viscerosensory, gustatory areas, as well as the limbic system and other networks critical for complex social, emotional, and cognitive functions (involving amygdala, temporopolar, parahippocampal, orbitofrontal, and anterior cingulate cortices) (see (Shura et al., 2014) for review) it may indeed serve as a critical node for a variety of disparate functions, including at least some those in which it has been implicated in previous studies.

We did not replicate the interesting finding of a previous study (Hedna et al., 2013) that left MCA occlusion is more common than right MCA occlusion, and associated with more severe stroke. The lack of a hemispheric difference in our study might be due to an ascertainment bias in our sample. We included only patients who gave consent for participation in a study of stroke recovery or whose identified decision-maker provided informed consent. Patients who are severely aphasic (often due to left MCA occlusion) cannot provide informed consent; and it is often difficult to obtain timely consent from a legal decision-maker. Therefore, it is possible that we enrolled disproportionately fewer patients with left MCA occlusion. Nevertheless, there was a very strong association between insular damage and MCA occlusion in both left and right hemisphere stroke.

In summary, some deficits that have been ascribed to insular lesions might be better explained as a consequence of large lesions or hypoperfusion due to MCA occlusion or stenosis (especially in the acute stage). That is not to say that the insula is not critical for some of the functions for which it has been implicated. Indeed, there is independent evidence that it plays a role in corticolimbic networks underlying social, emotional, and autonomic functions. Functional neuroimaging studies show activation of the insula associated with many tasks (e.g. (Brown et al., 2011; Gu et al., 2012; Jabbi et al., 2007; Lamm et al., 2011; Moser et al., 2009; Phillips et al., 1997); but functional imaging can only show that a region is engaged in a task, not that it is essential for the task (Fellows et al., 2005; Müller and Knight, 2006). Neuroimaging studies also show that the insula is functionally connected, as well as structurally connected, with many regions (Cerliani et al., 2012; Cloutman et al., 2012; Dennis et al., 2014). Lesions to the insula, or to its connections, could plausibly disrupt a variety of functions. Lesion-deficit studies complement functional imaging studies, but need to adequately control for volume of lesion and consider the potential influence of hypoperfusion beyond the infarct.

Previous studies have shown a frequent involvement of the insula in MCA territory infarction (Fink et al., 2005; Nakano et al., 2001; Truwit et al., 1990), but ours may show the clearest association with infarct volume. Fink et al. (2005) reported that insular infarct was associated with lenticulostriate territory infarction, infarct of more than one-third of MCA territory infarction; higher NIHSS score, and proximal vascular occlusion. They also showed that anterior insula infarct was associated with infarct in the superior division MCA territory, while posterior insula infarct was associated with infarct in the inferior division. Our results confirm that involvement of the insula on early DWI is a marker of probable MCA occlusion. Furthermore, in our study, only patients with insular stroke showed larger strokes in association with MCA occlusion. These findings have important clinical significance because the presence of insular involvement on early DWI could signal the need for urgent vessel imaging to consider rapid intervention such as thrombolysis or embolectomy to prevent progression to large volume stroke.

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