The Hyperdense Middle Cerebral Artery Sign in Drip-and-Ship Models of Acute Stroke Management

Lise Jodaitis\textsuperscript{a}  Noémi Ligot\textsuperscript{a}  Rudy Chapusette\textsuperscript{b}  Thomas Bonnet\textsuperscript{c}  Nicolas Gaspard\textsuperscript{a}  Gilles Naeije\textsuperscript{a}

\textsuperscript{a}Department of Neurology, Erasme Hospital, ULB, Brussels, Belgium; \textsuperscript{b}Department of Radiology, Erasme Hospital, ULB, Brussels, Belgium; \textsuperscript{c}Department of Interventional Neuroradiology, Erasme Hospital, ULB, Brussels, Belgium

Keywords
Acute stroke imaging · Hyperdense middle cerebral artery sign · Acute stroke management

Abstract

\textbf{Background:} Large vessel occlusion (LVO) leads to debilitating stroke and responds modestly to recombinant tissue plasminogen activator (rt-TPA). Early thrombectomy improves functional outcomes in selected patients with proximal occlusion but it is not available in all medical facilities. The best imaging modality for triage in an acute stroke setting in drip-and-ship models is still the subject of debate. \textbf{Objectives:} We aimed to assess the diagnostic value of millimeter-sliced noncontrast computed tomography (NCCT) hyperdense middle cerebral artery sign (HMCAS) in itself or associated with clinical data for early detection of LVO in drip-and-ship models of acute stroke management. \textbf{Methods:} NCCT of patients admitted to the Erasme Hospital, ULB, Brussels, Belgium, for suspicion of acute ischemic stroke between January 1 and July 31, 2017, were collected. Patients with brain hemorrhages were excluded, leading to 122 cases. The presence of HMCAS on NCCT was determined via visual assessment by 6 raters blinded to all other data. An independent rater assessed the presence of LVO on digital subtraction angiography imaging or contrast-enhanced CT angiography (CTA). The sensitivity, false-positive rate (FPR), and accuracy of HMCAS and the dot sign to detect LVO were calculated. The interobserver agreement of HMCAS was assessed using Gwet’s AC1 coefficient. Then, on a separate occasion, the first 2 observers rereviewed all NCCT provided with clinical clues. The sensitivity, FPR, and accuracy of HMCAS were recalculated. \textbf{Results:} HMCAS was found in 21% of the cases and a dot sign was found in 9%. The mean HMCAS sensitivity was 62% (95% CI 45–79%) and its accuracy was 86% (95% CI 79–92%) for detecting...
LVO. The interobserver reliability coefficient was 80% for HMCAS. Combined with clinical information, HMCAS sensitivity increased to 81% (95% CI 68–94; \( p = 0.041 \)) and accuracy increased to 91% (95% CI 86–96%). **Conclusion:** When clinical data are provided, detection of HMCAS on thinly sliced NCCT could be enough to decide on transfer for thrombectomy in drip-and-ship models of acute stroke management, especially in situations where CTA is less available and referral centers for thrombectomy fewer and further apart.

## Introduction

Stroke is the second cause of death and the main cause of acquired disability [1]. Most debilitating strokes are due to proximal arterial occlusion (i.e., internal carotid artery, M1/M2 segment of the middle cerebral artery, A1 segment of the anterior cerebral artery, and basilar and intracranial vertebral arteries), which accounts for 40% of all ischemic strokes [2]. Thrombectomy improves functional outcomes in selected patients with large vessel occlusion (LVO) [3–7] but is not available in all medical facilities. To meet that issue, acute stroke management led to intra and extra hospital re-organization. In extra hospital management, the development of “hub-and-spoke” stroke networks, where hospitals able to provide recombinant tissue plasminogen activator (rt-TPA) are organized around a referral center that performs thrombectomy, increased the number of stroke interventions [8, 9]. However, despite widening of the time window therapy to 16–24 h [10], many patients are still excluded from recanalization therapy due to a lack of viable brain tissue to salvage. One of the most common causes is the time lost in hospital-to-hospital transfer [11].

In this work, we aim to assess the place of a simple and widely available noncontrast computed tomography (NCCT) indirect sign of LVO, i.e., the hyperdense middle cerebral artery sign (HMCAS), to identify earlier a subset of patients with LVO and gain time for treatment in drip-and-ship models of acute stroke management. We thus determined the diagnostic value and the interrater reliability of the HMCAS and the dot sign (which is the equivalent of the HMCAS within the Sylvian fissure) using millimeter-sliced NCCT in isolation or in combination with clinical information for early detection of LVO strokes.

## Materials and Methods

### Study Design and Sample

Patients who were admitted to the emergency department of a tertiary academic medical center (Hôpital Erasme) with a suspected acute ischemic stroke and who benefited from an NCCT between January 1 and July 31, 2017, were retrospectively included. Spontaneous intracranial hemorrhage visible on the initial NCCT and patients without proper brain and cervical vessel injection imaging were excluded. The remaining NCCT were then anonymized by removing the patient’s name and image markings and they were given a randomly assigned number. Demographic and clinical data such as the National Institutes of Health Stroke Scale (NIHSS) at admission and the final diagnosis were retrieved from the patients’ medical records.

### Standard of Care

At our institution, all patients presenting to the emergency department with acute (<7.5 h or of uncertain timing) neurologic symptoms benefit from an immediate bedside
evaluation, including neurological assessment with the NIHSS and NCCT imaging. In the absence of clinical or radiological contraindications, patients admitted within 4.5 h of onset are given IV rtTPA in the imaging suite; they then undergo further imaging with a contrast-enhanced CT angiography (CTA) to assess extra- and intracranial arteries. When iodine contrast injection is contraindicated due to an iodine intolerance or renal failure, subjects benefit from magnetic resonance angiography (MRA). Patients with LVO identified on CTA who are admitted within 7.5 h of onset and have a premorbid modified Rankin scale score < 2 are eligible for thrombectomy. Patients with wake-up strokes or strokes admitted > 4.5 h after onset receive a perfusion CT before thrombolysis or thrombectomy is performed, based on mismatch criteria.

Imaging Parameters and Review

The imaging parameters were: continuous axial slice acquisition from the vertex to the skull base, a slice thickness of 1.5 mm, an increment thickness of 1.0 mm, a tube current of 165 mAS, a tube voltage of 120 kv, and 64 × 0.6 mm collimation.

First, in order to determine HMCAS and dot sign reliability, anonymized NCCT were retrospectively presented and independently reviewed by 6 raters blinded to all clinical and other imaging data, i.e., 4 neurologists (N.G., L.J., N.L., and G.N.), 1 neuroradiologist (R.C.), and 1 interventional neuroradiologist (T.B.). The presence or absence of HMCA and dot signs was determined by visual assessment and, respectively, defined as a relative hyperdensity of an MCA in comparison to its contralateral counterpart and as the hyperdensity of an arterial structure (seen as a dot on axial slices) in the Sylvian fissure relative to the contralateral side or to other vessels within the Sylvian fissure [12].

Then, to determine the yield of the HMCAS in a more practical real-life situation, 2 of the initial raters (2 clinical neurologists, i.e., L.J. and G.N.) reassessed the presence of the HMCAS and the dot sign after gaining access to the patient’s clinical details.

Statistical Analysis

Positive HMCA and dot signs detected by clinicians were considered true positives if a proximal MCA (M1, M2) occlusion was confirmed on arterial CTA, MRA, or digital subtraction angiography (DSA). Means and 95% CI for sensitivity, the false-positive rate (FPR), and the accuracy of both signs, and of the presence of any of the two, were calculated. The interrater reliability for the HMCAS and dot sign was assessed using Gwet's AC1 coefficient, a chance-adjusted index for the reliability of categorical measurements that has been shown to provide a more stable interrater reliability coefficient than Cohen's \( \kappa \) [13]. Means and 95% CI for Gwet’s AC1 were calculated using leave-N-out jackknife resampling with 100 iterations, leaving 50% of the sample out at each iteration. Then, the sensitivity, the FPR and the accuracy of both signs, and of the presence of any of the 2, were reassessed by 2 of the initial raters who had access on that occasion to the laterality and extent of the neurological findings on examination. Data are presented as means (±SD), means (95% CI), or numbers (%). Differences in test performance with and without clinical information were assessed using log-linear analysis in a model containing 4 variables (sign [present or absent], proximal occlusion [present or absent], rater [1 vs. 2], and clinical information [available vs. not available]). Statistics were performed in R (The R Foundation, Vienna, Austria).

Results

Patients

The patients demographic and clinical features summarized in Table 1. One hundred forty-nine potential subjects were admitted during the study period. We excluded 27 of them,
i.e., 15 due to acute intracerebral hemorrhage and 12 due to a lack of adequate imaging performed at our institution (7 had NCCT in an outside hospital prior to admission to our institution, 1 had a brain MRI without a previous NCCT, and 4 did not get the angiographic sequences [CTA, MRA, or DSA] within the desired time window). Thus, 122 patients were included in this study.

Table 1. Patient demographics and clinical features

| Age, years   | 71±17             |
| Sex, male   | 70 (57)           |
| Confirmed ischemic stroke | 84 (69) |

Table 2. Interobserver agreement and accuracy of the HMCAS and the dot sign when blinded to clinical data

|                      | Gwet’s AC1 PA | Se FPR | Mean accuracy |
|----------------------|---------------|--------|---------------|
| HMCAS                | 80 (73–87)    | 70 (62–78) | 62 (45–79)   | 5 (1–10)   | 86 (79–92) |
| Dot sign             | 87 (82–91)    | 75 (68–83) | 19 (6–32)    | 6 (1–10)   | 73 (65–81) |
| HMCAS or dot sign    | 73 (65–80)    | 63 (54–71) | 68 (51–84)   | 11 (5–17)  | 83 (77–90) |

Data are presented as mean percents (95% CI). PA, percent agreement; Se, sensitivity.

HMCAS on NCCT when Blinded to Clinical Data

The HMCAS was identified in 21% (95% CI 18–26) and the dot sign in 9% (95% CI 5–11) of the cases. The interrater reliability for the HMCAS, the dot sign, or the presence of any of the 2 signs was substantial (Gwet’s AC1 = 80%, 95% CI 73–87), almost perfect (Gwet’s AC1 = 87%, 95% CI 82–91), and substantial (Gwet’s AC1 = 73%, 95% CI 65–87), respectively. The sensitivity and FPR of the HMCAS were 62% (95% CI 45–79) and 5% (95% CI 1–10), respectively, yielding a very good test accuracy of 86% (95% CI 79–92). The dot sign had a much lower sensitivity (19%, 95% CI 6–32) and a similar FPR (6%, 95% CI 1–10). The presence of
either the HMCAS or the dot sign only marginally improved the sensitivity (68%, 95% CI 51–84) compared to the HMCAS but with a higher FPR (11%, 95% CI 5–17), resulting in a marginally lower accuracy (83%, 95% CI 77–90). See Table 2.

### HMCAS on NCCT Combined with Clinical Data

Disclosure of clinical information to the raters significantly improved the sensitivity and accuracy of the HMCAS \((p = 0.04\) for the effect of clinical information). We found no significant difference between the 2 raters. See Table 3.

### Discussion/Conclusion

The HMCAS was found in 21% of cases suspicious for acute stroke, with good interobserver agreement. When clinical data were provided, as in real-life situations, the HMCAS on millimeter-sliced NCCT led to 81% sensitivity and 91% accuracy for MCA proximal occlusion detection, which could help to identify earlier a subset of patients with LVO in drip-and-ship models of acute stroke management.

Despite the study’s retrospective design and the sample size, the population studied mirrors the baseline characteristics of the patients included in the major trials on thrombectomy for proximal occlusion in terms of age and NIHSS [3, 6]. Similarly, the proportion of proximal occlusions within our cohort of ischemic strokes (40.5%) parallels previous prospective reports [2], as does the average NIHSS score of our patients with LVO [14]. Finally, the stroke etiology distribution in our population was comparable to that in a recent study that analyzed the associations between HMCAS and stroke etiology [15]. So, the evidence provided by our study is likely to be transposable to most acute stroke settings.

The HMCAS was found in 17–50% of acute MCA occlusions and has a specificity of 95%, with a lower sensitivity of 52% according to a recent meta-analysis that reviewed studies from 1990 to 2013 [16]. In our study, its sensitivity was of 62% for raters blinded to clinical information and 77–82% when raters were aware of clinical details, which was thus higher than expected. This can be explained by both technical and practical differences. First, previous studies have demonstrated that using thinner (<3 mm) NCCT slices increases the sensitivity of the HMCAS for detecting LVO, as the average diameter of the intracranial artery diameter is <3 mm [17]. In our acute stroke management, we systematically use 1.5-mm-slice section CT, whereas most prior studies have been conducted with ≥5 mm slices. Second, the raters in our study are clinicians and radiologists routinely involved in acute stroke management, which probably led to the high interrater reliability coefficient of 0.80 and the 81% sensitivity for LVO when clinical data were added to the model. This is a message but

| HMCAS reader 1 without clinical information | 62 (45–79) | 2 (0–5) | 88 (82–93) |
| HMCAS reader 2 without clinical information | 62 (44–80) | 10 (4–17) | 82 (75–89) |
| HMCAS reader 1 with clinical information | 77 (62–91) | 3 (0–7) | 91 (86–96) |
| HMCAS reader 2 with clinical information | 85 (73–97) | 6 (1–11) | 92 (87–97) |

Data are presented as mean percents (95% CI). Se, sensitivity.
also a limitation of our study, because our results may not be generalizable to settings where NNCT are not interpreted by stroke specialists.

The advantage of adding clinical data has already been highlighted by Lim et al. [18] in a study that assessed the sensitivity and specificity of the HMCAS on thin-slice (1.5 mm) NCCT combined with the NIHSS score and a brief clinical history and found up to 79% sensitivity for LVO in patients with an NIHSS > 10 [18]. It also parallels results from a recent study that demonstrated that combining the NIHSS score and the absolute attenuation value of the middle cerebral artery on NCCT showed 85% sensitivity for LVO [19].

The good accuracy of the HMCAS on NCCT for LVO when assessed by stroke clinicians is an important element in the global context of acute stroke management choice of settings. Indeed, the American Stroke Association recommends: “transport patients rapidly to the closest available appropriate institution to provide initial emergency care, including administration of intravenous rt-TPA and if needed, to arrange transfer to centers capable of performing endovascular stroke treatment with comprehensive per-procedural care” [20]. This drip-and-ship model effectively increases the rate of stroke interventions [8] but is also associated with longer transfer times that may delay endovascular treatments [11, 21]. Several approaches have been tried to prevent interhospital delays between primary and secondary stroke centers in suspected LVO, i.e., by-passing time-consuming imaging in patients arriving soon after symptom onset (like perfusion CT or MRI that can induce almost a 20-min delay in acute ischemic stroke treatment [22]), relying on telemedicine, angiography suites, and mobile strokes. In that context, our findings support that HMCAS on thin slices NCCT could efficiently identify 8 out of 10 patients with acute MCA LVO and allow earlier transfers for thrombectomy in a substantial subset of patients. However, these findings do not argue for the replacement of CTA by NCCT in drip-and-ship models of acute stroke management as 20% of LVO would be missed. Indeed, CTA provides almost 100% accuracy for LVO [23] and important information on stroke etiology and should remain the first imaging for triage in acute stroke management. However, it is not realized around the clock in many primary stroke centers and may not be feasible for technical problems or contraindications. Furthermore, the yield and prognosis value of CTA are higher when CTA is realized close to the thrombectomy procedure, even in the angiography suite [24–26].

So, shipping suspected LVO based on identification of the HMCAS on thin-slice NCCT in order to perform CTA in the thrombectomy stroke center may both be efficient and adequate in selected situations. Finally, many middle-income countries that now account for a large part of stroke mortality and related disability [27] are likely to develop stroke networks based on the drip-and-ship model for economic reasons. So, the presence of HMCAS on 1.5-mm NCCT could be enough to take the “drip-and-shift” decision in LVO and gain precious time for endovascular treatment.

Another limitation of this study is the focus on MCA occlusions that prevents our results from being applied to other sites of proximal occlusion. However, carotid and MCA thrombi represent the vast majority of LVO (respectively, 24% within the carotid artery and 52% within the proximal MCA) while ACA and posterior circulation occlusions occur only in a minority of cases (1 and 16%, respectively) [28]. However, NCCT arterial hyperdensity signs in studies that looked at both anterior and posterior circulation LVO yielded sensitivity results similar to those for MCA occlusions [18]. So, it is likely that our findings may still be valid for proximal posterior occlusions.

In conclusion, in drip-and-ship models of acute stroke management, detection of the HMCAS on millimeter-sliced NCCT could identify earlier a subset of patients with LVO and hurry the transfer for thrombectomy in primary stroke centers, especially in situations where CTA is less available and referral centers are fewer and further apart. However, even combined with clinical data, the HMCAS on NCCT still misses 20% of LVO, which warrants the use of CTA when feasible.
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Statement of Ethics

The study protocol was approved by the research institute’s committee on human research. The authors have no ethical conflicts to disclose.

Disclosure Statement

The authors have no conflict of interests to declare.

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Author Contributions

Lise Jodaitis: main writer. Gilles Naeije: substantial contributions to the conception and design of this work, acquisition of data, critical revision of this work, and final approval of the version for publication. Noémie Ligot, Rudy Chapusette, and Thomas Bonnet: substantial contributions to acquisition of data, critical revision of this work, and final approval. Nicolas Gaspard: substantial contributions to acquisition, analysis, and interpretation of data for this work, critical revision of this work, final approval.

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