Adjunctive Efficacy of Intra-Arterial Conebeam CT Angiography Relative to DSA in the Diagnosis and Surgical Planning of Micro-Arteriovenous Malformations

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ABSTRACT

BACKGROUND AND PURPOSE: Micro-arteriovenous malformations are an underrecognized etiology of intracranial hemorrhage. Our study aimed to assess the adjunctive efficacy of intra-arterial conebeam CTA relative to DSA in the diagnosis and surgical planning of intracranial micro-AVMs.

MATERIALS AND METHODS: We performed a retrospective study of all micro-AVMs (≤1-cm nidus) at our institution. Blinded neuroradiologists qualitatively graded DSA and intra-arterial conebeam CTA images for the detection of specific micro-AVM anatomic parameters (arterial feeder, micronidus, and venous drainer) and defined an overall diagnostic value. Statistical and absolute differences in the overall diagnostic values defined the relative intra-arterial conebeam CTA diagnostic values, respectively. Blinded neurosurgeons reported their treatment approach after DSA and graded the adjunctive value of intra-arterial conebeam CTA to improve or modify treatment. Intra-arterial conebeam CTA efficacy was defined as interobserver agreement in the relative intra-arterial conebeam CTA diagnostic and/or treatment-planning value scores.

RESULTS: Ten patients with micro-AVMs presented with neurologic deficits and/or intracranial hemorrhages. Both neuroradiologists assigned a higher overall intra-arterial conebeam CTA diagnostic value (P = .05), secondary to improved evaluation of both arterial feeders and the micronidus, with good interobserver agreement (τ = 0.66, P = .018) in the relative intra-arterial conebeam CTA diagnostic value. Both neurosurgeons reported that integrating the intra-arterial conebeam CTA data into their treatment plan would allow more confident localization for surgical/radiation treatment (8/10; altering the treatment plan in 1 patient), with good interobserver agreement in the relative intra-arterial conebeam CTA treatment planning value (τ = 0.73, P = .025).

CONCLUSIONS: Adjunctive intra-arterial conebeam CTA techniques are more effective in the diagnostic identification and anatomic delineation of micro-AVMs, relative to DSA alone, with the potential to improve microsurgical or radiosurgery treatment planning.

ABBREVIATIONS: mAVM = micro-AVM; IA-CBCTA = intra-arterial conebeam CTA; 3DRA = 3D rotational angiography

Cerebral micro-arteriovenous malformations (mAVMs) are a rare subgroup, accounting for ~8% of intracranial AVMs in the surgical series by Stiver and Ogilvy and may represent 21% of AVMs presenting with intracranial hemorrhage in young adults. In 2013, Alén et al reported approximately 87% of patients with mAVMs presenting with spontaneous hemorrhage, while previous studies reported a 100% incidence. Yasargil defined mAVMs as a particular entity of pial AVMs characterized by an occult nidus of ≤1 cm and differentiated this subgroup, which could not be visualized on angiography or gross pathology specimens but could be identified on histopathology if the hematoma was carefully removed.

Micro-AVMs may be suspected on cerebral DSA as an abnormal draining vein with variable detection of a feeding artery and/or micronidus, often only suggested by small coalescing arterioles. Subtle angiographic findings make the imaging diagnosis challenging and broaden the differential diagnosis to include the possibility of a dural or pial arteriovenous fistula. Additionally, the small mAVM size and associated hemorrhage add difficulty in lesion localization for treatment planning with either microsurgical resection or stereotactic radiation therapy.

Digital flat panel detector CT technology has improved the imaging capabilities of modern angiographic equipment with the
feasibility of routinely performing 3D rotational angiography (3DRA) and intra-arterial conebeam CTA (IA-CBCTA). IA-CBCTA incorporates the high spatial vascular resolution of 3DRA with CT postprocessing techniques for increased contrast resolution to visualize the peripheral osseous and soft tissues. High spatial resolution is maintained and displayed in submillimeter multiplanar reconstructions for improved diagnostic sensitivity and cross-sectional mapping. Prior studies have described the efficacy of IA-CBCTA for the anatomic localization of intracranial and spinal dural AVFs. In this study, we hypothesized that the efficacy of IA-CBCTA relative to DSA would improve the diagnostic identification, localization, and treatment planning of mAVMs.

MATERIALS AND METHODS

Patient Study

Institutional review board approval was obtained for a retrospective study of patients presenting with intracranial mAVMs (≤1 cm) between January 2010 and February 2017. All patients were identified through the neuroangiographic database of our institution, and at least 1 pretreatment DSA study with adjunctive IA-CBCTA imaging was required for study inclusion. The clinical and radiologic findings of these patients and mAVM lesions were reviewed, including patient demographics, presentations, intracranial hemorrhage locations and volumes (gross manual calculations on presenting CT/MR imaging), mAVM classification according to the Spetzler-Martin grading system, and subsequent management/treatment.

Image Acquisition

Biplane DSA imaging studies were acquired at 2 tertiary academic institutions (Northwestern Memorial Hospital and Lurie Children’s Hospital affiliated with Northwestern University, Chicago, Illinois) with identical biplane angiography suites (Artis Zee/Zeego flat detector biplane angiosuite; Siemens, Erlangen, Germany) and acquisition protocols. 3DRA or IA-CBCTA scanning was performed with a single flat panel detector rotational fluoroscopy and angiography unit (2k detector, amorphous silicon cesium iodide scintillator; 1920 × 2480 pixel resolution, 3.25 lp/mm; size, 30 × 40 cm). The motorized frontal C-arm was used to acquire 496 projection frames over a 200° arc with 5- to 8-second rotation times at 80 kV and 260 mA and a radiation dose of 1.2 μGy/frame. Although acquisition parameters remained constant, IA-CBCTA techniques varied slightly with respect to intra-arterial contrast injection rates dependent on vessel injection sites, including the internal carotid artery (4 mL/s) and vertebral artery (3–4 mL/s), with total volumes of 15–32 mL. Shorter acquisition times (5–8 seconds) and nondilute iodinated contrast assisted with opacification of subtle microvascular anatomy and abnormal early venous drainage without venous contamination. IA-CBCTA acquisition data were transferred to an independent 3D postprocessing workstation (Leonardo; Siemens). Multiplanar, subtracted, and unsubtracted CT reconstructions (sagittal, coronal, and axial planes with overlapping 0.5-mm slice thickness) were generated and transferred to a PACS for direct visualization on the PACS workstation.

Image Analysis

Images were retrospectively and independently reviewed by 2 interventional neuroradiologists and 2 vascular neurosurgeons on a de-identified PACS system (including the postprocessed multiplanar IA-CBCTA reconstructions). Both neuroradiologists graded the DSA (biplane, magnified, oblique views) and 3DRA images without access to the IA-CBCTA dataset. Subsequent IA-CBCTA images (subtracted/unsubtracted multiplanar reconstructions) acquired from the same intra-arterial injection site were scored by the observers. Qualitative image analyses were performed on high-definition liquid crystal display monitors routinely used for diagnostic reporting and, based on the level of image quality, anatomic and angiographic characterization on a scale of 0–2 (2, excellent/good; 1, relevant visibility with restrictions; 0, poor, nondiagnostic). The reviewers scored the following parameters: 1) arterial feeders (anatomic localization, origin/course, single versus multiple); 2) venous drains (anatomic localization, course/destination, single versus multiple); and 3) nidus site (anatomic localization, size, and differentiation from arteriovenous shunting). The total score for each technique was defined as the overall diagnostic value for interpretation.

For evaluation of treatment-planning efficacy, the 2 vascular neurosurgeons reported their presumed treatment strategy and approach (if applicable) after evaluating DSA and 3DRA imaging and subsequently studied the IA-CBCTA multiplanar reconstructions. Both neurosurgeons then graded the additive or relative treatment-planning values of IA-CBCTA according to a scale of 0–2 (2, altered treatment plan; 1, more confident treatment plan; or 0, no value). They were required to provide a reason if they determined that IA-CBCTA improved or altered the treatment plan (grade 1 or 2) in comparison with DSA imaging alone.

Statistical Analysis

For each observer, the Wilcoxon test was used to assess statistical differences in DSA-versus-IA-CBCTA scoring for each of the 3 mAVM anatomic parameters and the overall diagnostic values (sum of all 3 parameter scores) between the 2 modalities. Statistical and absolute differences between the overall IA-CBCTA and DSA diagnostic value defined the relative IA-CBCTA diagnostic value/scores, respectively. A P value <.05 was considered a statistically significant difference.

Interobserver agreement for DSA and IA-CBCTA scoring of each mAVM parameter and the overall diagnostic values were assessed using the Kendall τ coefficient. Interobserver agreement of the relative IA-CBCTA diagnostic value scores (absolute difference between overall IA-CBCTA and DSA diagnostic value scores), and relative IA-CBCTA treatment-planning value scores were also evaluated using the Kendall τ coefficient. Good or excellent agreement defined the efficacy of IA-CBCTA relative to DSA in the diagnosis and surgical planning of mAVMs, respectively. The τ coefficient varied between 0 and 1, with 0 representing no agreement and 1 representing complete agreement. τ values of >0.8, >0.5–0.8, >0.2–0.5, and ≤0.2 were considered to indicate excellent, good, fair, and poor agreement, respectively. A P value <.05 indicated statis-
IA-CBCTA Diagnostic and Treatment-Planning Observer Studies

Blinded observer analysis was performed on DSA and multiplanar IA-CBCTA images from 10 patients with suspected intracranial mAVMs (Table 1). Initial DSA evaluation revealed a micronidus with excellent visibility in only 2 patients (20%) by both observers (patients 8 and 9). In contrast, IA-CBCTA clearly identified the micronidus in all 10/10 (100%) patients, with both observers scoring the nidus as grade 2 (excellent visualization) with perfect interobserver agreement. Interobserver agreement in grading the rest of the individual anatomic parameters with IA-CBCTA or DSA ranged from good to excellent (τ coefficient between 0.55 and 1.0). Interobserver agreement for overall diagnostic value scores of IA-CBCTA and DSA was good (τ coefficient = 0.57 and 0.7, respectively).

Both neuroradiologists assigned significantly higher scores to IA-CBCTA for overall diagnostic value (both observers: P < .05; observer 1: P = .007; observer 2: P = .016), confirming the relative diagnostic value of IA-CBCTA. Although no significant differences were seen between DSA and IA-CBCTA scores when evaluating venous drainage (observer 1: P = .157; observer 2: P = .564), both observers assigned significantly higher scores to IA-CBCTA when evaluating arterial feeders (observer 1: P = .02; observer 2: P = .03) and the micronidus (observer 1: P = .009; observer 2: P = .009), resulting in relative IA-CBCTA diagnostic value scores with good interobserver agreement (τ coefficient = 0.66, P = .018).

Both neurosurgeons recorded more confidence in the treatment plan (grades 1–2) for nearly all patients after review of the IA-CBCTA and agreed that it provided better delineation and precise anatomic localization of the micronidus (Table 2). For example, IA-CBCTA led to both neurosurgeons altering their treatment plan (grade 2) in patient 1 (Fig 1, where the nidus could only be identified on IA-CBCTA) and recommending superselective catheterization and possible embolization before proceeding to an operation (grade 1) in patient 4. Overall, they reported that IA-CBCTA would increase confidence in their microsurgical or radiosurgery treatment by importing these data directly into a
neuronavigation system. Interobserver agreement between the 2 vascular neurosurgeons for the relative IA-CBCTA treatment-planning value was good ($\tau$ coefficient = 0.73, $P = .025$).

### Illustrative Cases

**Patient 1.** A 30-year-old woman presented with a 3-year history of epilepsy, periventricular heterotopia was diagnosed with a presumed right cerebellar developmental venous anomaly after emergent outside imaging evaluation including DSA. Due to acute presentation with severe headache, visual impairment, quadriplegia, and rapid deterioration to unconsciousness, she required emergent craniectomy and decompression of a cerebellar parenchymal hemorrhage. Although initial MR imaging/MRA failed to detect a vascular malformation, a 3-month delayed DSA demonstrated findings suspicious for a pial AVF (Fig 1A, B). At 6 months, repeat DSA with 3DRA re-demonstrated arteriovenous shunting and early venous drainage but also identified an occult <5-mm micronidus under the lateral right cerebellar surface, adjacent to the craniectomy site, and only visualized with IA-CBCTA reconstructions (Fig 1C–D).

**Patient 4.** A 21-year-old woman presented to our institution with sudden headache and dizziness for 1 hour. Initial CT/CTA and MR imaging/MRA brain studies revealed an acute paramedian right frontoparietal intracerebral hemorrhage and a distal pericallosal aneurysm with suspicion of an underlying occult vascular malformation, but no abnormal vascular flow voids or nidus was seen (Fig 2A). Subsequent DSA with the 3DRA/IA-CBCTA technique was performed 2 days later confirming a 2-mm pericallosal-splenial artery aneurysm/pseudoaneurysm with suggestion of an early draining vein (Fig 2B, C), but a definitive arterial feeder and <5-mm nidus were only visualized on IA-CBCTA multiplanar reconstructions to confirm the diagnosis of a micro-AVM (Fig 2D–G).

**Patient 9.** A 50-year-old man presented to an outside institution with confusion followed by right hemiparesis. Initial CT/MR imaging discovered a left parietal cortical hemorrhage, which was

### Table 2: Treatment-planning scoring of IA-CBCTA relative to DSA<sup>a</sup>

| Patient | Neurosurgeon 1 | Relative CBCTA Treatment Value | Neurosurgeon 2 | Relative CBCTA Treatment Value |
|---------|----------------|-------------------------------|----------------|-------------------------------|
| 1       | Not sufficient | Operation                     | 2              | Not sufficient | Operation                     |
| 2       | Radiosurgery   | Radiosurgery                  | 1              | Radiosurgery | Radiosurgery                  |
| 3       | Radiosurgery   | Radiosurgery                  | 1              | Radiosurgery | Radiosurgery                  |
| 4       | Operation      | Operation                     | 1              | Operation | Operation                     |
| 5       | Operation      | Operation                     | 1              | Operation | Operation                     |
| 6       | Radiosurgery   | Radiosurgery                  | 1              | Radiosurgery | Radiosurgery                  |
| 7       | Operation      | Operation                     | 1              | Operation | Operation                     |
| 8       | Radiosurgery   | Radiosurgery                  | 1              | Radiosurgery | Radiosurgery                  |
| 9       | Operation      | Operation                     | 1              | Operation | Operation                     |
| 10      | Operation      | Operation                     | 1              | Operation | Operation                     |

<sup>a</sup> $\tau$ coefficient = 0.73, $P = .025$.

**FIG 1.** Anteroposterior DSA images demonstrate left vertebral artery injections in the early (A) and late (B) arterial phases, identifying arteriovenous shunting and early venous drainage into a hypertrophied superior cerebellar hemispheric vein (arrow) with tentorial venous outflow, but both observers were unable to appreciate an occult micronidus. Although arterial feeders are suggested to project into this region on DSA from the right anterior cerebellar artery and SCA (A, arrowheads), the SCA feeder is better appreciated on IA-CBCTA multiplanar axial reconstructions (C, arrowhead). Moreover, both observers identified a <5-mm micronidus under the lateral cerebellar surface and adjacent to the craniectomy site (C, arrow) with adjunctive IA-CBCTA reconstructions, consistent with an mAVM. IA-CBCTA coronal reconstruction (D) shows the nidal outflow to the early draining superior cerebellar hemispheric vein (arrow).
conservatively managed. Delayed 2-month MR imaging findings were negative, but subsequent DSA reported a left parietal pial AVF, and embolization was planned. At our institution, a repeat DSA with the 3DRA/IA-CBCTA technique revealed a superficial left parietal mAVM with a nidus measuring \( \frac{1}{2} \) cm (Fig 3A, B). Left frontoparietal paramedian craniotomy and microsurgical resection were performed with IA-CBCTA imaging integrated into the neuronavigation system (Fig 3C, D).

**DISCUSSION**

Prior literature emphasized the importance of identifying an mAVM and characterizing its anatomy for effective treatment. However, the diagnosis of mAVMs can be challenging. Alén et al\(^3\) reported the role of high-resolution MR imaging fast spin-echo T2-weighted sequences in detecting subtle flow voids corresponding to small nidus volumes. In addition, they noted the supplementary value of contrast-enhanced MR imaging/MRA over...
Some have reported initially described a modified CBCTA imaging technique in 3 patients performed using an intra-arterial diluted iohedrol contrast agent (35%) in the ascending aorta that lasted 22 seconds at a rate of 8 mL/s, which allows imaging of an entire AVM nidus receiving blood supply from >1 cervical artery in a single CBCTA acquisition. In this study, our intra-arterial technique was modified with shorter acquisition times (5–8 seconds) to avoid venous contamination and single cervical artery infusion of nondiluted contrast to optimize visualization of the micronidus and early draining vein. Rahal and Malek reported a higher sensitivity of IA-CBCTA over DSA in identifying the occult nidal anatomy and angioarchitecture of brain AVMs in 3 patients presenting with intracranial hemorrhage as well providing anatomic guidance for either surgical or radiosurgery treatment. However, none of these studies included or discussed the value of CBCTA in cerebral AVMs with a micronidus of <1 cm.

Additionally, IA-CBCTA has been shown to improve the anatomic delineation and localization of both intracranial and spinal dural arteriovenous fistulas for preoperative or endovascular planning. Radvany et al described a modified CBCTA imaging technique in 3 patients performed using an intra-arterial diluted iohedrol contrast agent (35%) in the ascending aorta that lasted 22 seconds at a rate of 8 mL/s, which allows imaging of an entire AVM nidus receiving blood supply from >1 cervical artery in a single CBCTA acquisition. In this study, our intra-arterial technique was modified with shorter acquisition times (5–8 seconds) to avoid venous contamination and single cervical artery infusion of nondiluted contrast to optimize visualization of the micronidus and early draining vein. Rahal and Malek reported a higher sensitivity of IA-CBCTA over DSA in identifying the occult nidal anatomy and angioarchitecture of brain AVMs in 3 patients presenting with intracranial hemorrhage as well providing anatomic guidance for either surgical or radiosurgery treatment. However, none of these studies included or discussed the value of CBCTA in cerebral AVMs with a micronidus of <1 cm.

In contrast, the precise cross-sectional localization of mAVMs, especially for surgical treatment planning, can be readily facilitated by IA-CBCTA.

Our results are in keeping with previous studies that have shown the feasibility and superiority of using IA-CBCTA in both preoperative planning and as an intraoperative reference for the microsurgical resection or stereotactic radiosurgery of larger brain AVMs.

In a case series of 16 patients with cerebral AVMs, Srinivasan et al used CBCTA in a neuronavigation system, allowing complete and safe surgical resection of all lesions. In another case series by Safain et al of 22 patients, CBCTA was used to target cerebral AVMs with radiosurgery, with only 4/22 lesions that had a micronidus of 1 cm and resulted in improved visualization of AVM components and treatment planning. Radvany et al described a modified CBCTA imaging technique in 3 patients performed using an intra-arterial diluted iohedrol contrast agent (35%) in the ascending aorta that lasted 22 seconds at a rate of 8 mL/s, which allows imaging of an entire AVM nidus receiving blood supply from >1 cervical artery in a single CBCTA acquisition. In this study, our intra-arterial technique was modified with shorter acquisition times (5–8 seconds) to avoid venous contamination and single cervical artery infusion of nondiluted contrast to optimize visualization of the micronidus and early draining vein. Rahal and Malek reported a higher sensitivity of IA-CBCTA over DSA in identifying the occult nidal anatomy and angioarchitecture of brain AVMs in 3 patients presenting with intracranial hemorrhage as well providing anatomic guidance for either surgical or radiosurgery treatment. However, none of these studies included or discussed the value of CBCTA in cerebral AVMs with a micronidus of <1 cm.

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In our series, even 2D and 3DRA techniques without IA-CBCTA reconstructions were relatively inferior in delineating the AVM arterial feeders and nidus, consistent with mAVMs possibly being very small occult lesions (<5 mm). We have shown that

FIG 3. Lateral DSA (A) and coronal IA-CBCTA reconstruction (B) images both demonstrate a small <1 cm micronidus (arrows), supplied by tortuous parietal branches of the pericallosal anterior cerebral artery and inferior division of the MCA (arrowheads), with early venous drainage into bifurcating cortical veins (double asterisks). Although both observers did not report the improved diagnostic value of IA-CBCTA in this case, both neurosurgeons reported increased confidence in treatment planning, and IA-CBCTA was incorporated into the neuronavigation system for microsurgical resection. 3D IA-CBCTA and MR imaging datasets were merged with sagittal overlay (C) delineating the micronidus (arrow) and draining cortical vein (double asterisks) complex in relation to the adjacent hemorrhage. Both datasets were imported into the intraoperative neuronavigation system (D, BrainLAB), allowing anatomic localization of the micronidus within a specific sulcus guiding the surgical approach as well as the presumed deep nidal rupture site abutting the hematoma (black arrow).
Flat panel detectors improve the signal intensity-to-noise ratio and suppress geometric distortion, enabling 3DRA and IA-CBCTA to produce high spatial and contrast-resolution imaging with radiation and contrast-dose reduction. Isotropic CBCT volumetric datasets allow multiplanar and 3D reconstructions analogous to multidetector CT scanners. Advantages over traditional intravenous CT angiography include submillimeter-spatial-resolution isotropic data with versatile postprocessing software analysis, selective intra-arterial contrast injections for territorial vascular analysis, rapid and variable acquisitions (5–8 seconds) to optimize early venous opacification versus venous contamination, subtraction and hemodynamic flow analysis (3DRA), and unsubtracted cross-sectional localization relative to the osseous structures and soft tissues (IA-CBCTA).

Our study is subject to the inherent limitations of a retrospective series and qualitative grading of DSA images in comparison with IA-CBCTA as an adjunctive imaging technique. We attempted to partly mitigate this bias with blinded observer analysis and a protocol for grading DSA and IA-CBCTA imaging in series on a de-identified workstation that retained the ability to analyze IA-CBCTA multiplanar reconstructions with or without tissue subtraction. Furthermore, memory bias was limited, with an elapsed period of 6 years to accumulate our cases and only 2 observers in each arm of the study from a total of 6 neurointerventionalists involved in the initial diagnostic evaluation and 5 vascular neurosurgeons involved in the surgical/radiation treatment. Due to the rare incidence of mAVMs, sample sizes in all studies of this pathology remained small, and future studies with a larger population would require a multi-institutional cohort or registry.

CONCLUSIONS

We compared DSA and adjunctive IA-CBCTA imaging to assess the relative efficacy of IA-CBCTA in the diagnosis and treatment planning of mAVMs. IA-CBCTA improves the diagnostic accuracy of mAVMs, particularly through enhanced anatomic identification and localization of subtle arterial feeders and/or a micronidus, and enables more confident surgical/radiosurgery treatment planning.

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