Visualization of Lenticulostriate Arteries on CT Angiography Using Ultra-High-Resolution CT Compared with Conventional-Detector CT


AJNR Am J Neuroradiol 2020, 41 (2) 219-223
doi: https://doi.org/10.3174/ajnr.A6377
http://www.ajnr.org/content/41/2/219
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ABSTRACT

BACKGROUND AND PURPOSE: The newly developed ultra-high-resolution CT is equipped with a 0.25-mm detector, which has one-half the conventional section thickness, one-half the in-plane detector element width, and one-half the reconstructed pixel width compared with conventional-detector CT. Thus, the ultra-high-resolution CT scanner should provide better image quality for microvasculature than the conventional-detector CT scanners. This study aimed to determine whether ultra-high-resolution CT produces superior-quality images of the lenticulostriate arteries compared with conventional-detector CT.

MATERIALS AND METHODS: From February 2017 to June 2017, thirteen patients with aneurysms (4 men, 9 women; mean age, 61.2 years) who underwent head CTA with both ultra-high-resolution CT and conventional-detector CT were enrolled. Two board-certified radiologists determined the number of all lenticulostriate arteries on the CTA coronal images of the MCA M1 segment reconstructed from 512 matrixes on conventional-detector CT and 1024 matrixes on ultra-high-resolution CT.

RESULTS: There were statistically more lenticulostriate arteries identified on ultra-high-resolution CT (average, 2.85 ± 0.83; 95% CI, 2.50–3.18) than on conventional-detector CT (average, 2.17 ± 0.76; 95% CI, 1.86–2.48) (P < .009) in 16 of the total 26 MCA M1 segments.

CONCLUSIONS: Improvements in lenticulostriate artery visualization were the result of the combined package of the ultra-high-resolution CT scanner plus the ultra-high-resolution scanning protocol, which includes higher radiation doses with lower than the national diagnostic reference levels and stronger adaptive iterative dose-reduction processing. This package for ultra-high-resolution CT is a simple, noninvasive, and easily accessible method to evaluate microvasculature such as the lenticulostriate arteries.

ABBREVIATIONS: AIDR = adaptive iterative dose reduction; C-CT = conventional-detector CT; C-CTA = conventional CTA; CTDIvol = volume CT dose index; LSA = lenticulostriate artery; UHR = ultra-high-resolution

The lenticulostriate arteries (LSAs) are perforating arteries that originate in the MCA and supply the basal ganglia. Accurately assessing the variation and structure of the LSAs reduces the likelihood of postoperative complications when performing resections of insular gliomas and when clipping or coiling MCA M1–2 segment aneurysms.1,2 It is also clinically important to evaluate the features of the LSAs in patients with symptomatic intracranial atherosclerotic stenosis to determine the best therapeutic strategy.3 However, although conventional-detector CT (C-CT), which uses a detector with a range of 0.5 mm, has been used to image the proximal side of the LSA,4 it is not easy to image the entire LSA from the proximal-to-distant side using C-CT.

In 2005, Canon Medical Systems and the National Cancer Center, Japan, jointly developed a prototype ultra-high-resolution CT system (UHR-CT; TSX-304R; Canon Medical Systems, Otawara, Japan). The UHR-CT was equipped with a 0.25-mm detector, which has one-half the conventional section thickness, one-half the in-plane detector element width, and one-half the reconstructed pixel width compared with the C-CT. Kakinuma et al5 reported that the prototype UHR-CT scanner had significantly better image quality for lung nodules than the C-CT scanners. Clinical studies that have tested phantoms using the

Received November 2, 2018; accepted after revision November 12, 2019.
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This work was supported by Grants-in-Aid for Scientific Research (KAKENHI) 18K07694.
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http://dx.doi.org/10.3174/ajnr.A6377

production version of the UHR-CT (Aquilion Precision; Canon Medical Systems) have reported that a higher image quality could be obtained compared with C-CT.\(^6\)\(^7\) Another such study that imaged the artery of Adamkiewicz reported improved images.\(^8\) On the basis of these findings, we predicted that UHR-CTA will have improved imaging quality of the LSAs that exceeds that of conventional CTA (C-CTA). Therefore, the goal of this study was to investigate whether UHR-CTA produced superior images of the LSAs compared with C-CTA.

**MATERIALS AND METHODS**

**Subjects**

Our institutional review board (Fujita Health University) approved this opt-out research design. Opt-out opportunities were provided to all participants included in this retrospective study. Forty-two consecutive patients with intracranial aneurysms who underwent intracranial UHR-CTA between February 2017 and June 2017 and who underwent C-CTA using 320-row detector CT during a follow-up study (14–373 days later; mean interval, 224 days) were selected. Twenty-nine patients were excluded for the following reasons: unavailable datasets, the interval, 224 days) were selected. Twenty-nine patients were provided to all participants included in this retrospective study. Forty-two consecutive patients with intracranial aneurysms who underwent intracranial UHR-CTA between February 2017 and June 2017 and who underwent C-CTA using 320-row detector CT during a follow-up study (14–373 days later; mean interval, 224 days) were selected. Twenty-nine patients were excluded for the following reasons: unavailable datasets, the interval, 224 days) were selected. Twenty-nine patients were included in this retrospective study, Forty-two consecutive patients with intracranial aneurysms who underwent intracranial UHR-CTA between February 2017 and June 2017 and who underwent C-CTA using 320-row detector CT during a follow-up study (14–373 days later; mean interval, 224 days) were selected. Twenty-nine patients were excluded for the following reasons: unavailable datasets, the interval, 224 days) were selected. Twenty-nine patients were included in this retrospective study.

**CTA Protocols for UHR-CT and C-CT**

Contrast-enhanced volume data for UHR-CTA were acquired with a 160-detector row UHR-CT scanner (Aquilion Precision) using helical scanning. The UHR-CT scanner used in this study is a medical device with certification as a CT scanner that is safe for clinical use. The other scan parameters were as follows: section thickness of 0.25 mm, tube voltage of 120 kV, automatic exposure control tube current SD of 5, pitch factor of 0.625, reconstructed FOV size of 190–240 mm, and gantry rotation speed of 1 s/rotation. Contrast-enhanced volume data for C-CTA were acquired with a 320-row detector CT scanner (Aquilion ONE; Canon Medical Systems) using nonhelical scanning. The other scan parameters were as follows: section thickness of 0.5 mm, tube voltage of 120 kV, automatic exposure control tube current SD of 3–5, pitch factor of 0.625, reconstructed FOV size of 180–240 mm, and gantry rotation speed of 1 s/rotation. The contrast medium volume was calculated on the basis of each patient’s body weight, and 250 mg I/kg of iopamidol (Iopamiron 370; Bayer Healthcare, Osaka, Japan) was injected intravenously as a bolus for a fixed infusion duration of 10 seconds, followed by an intravenous bolus injection of 30 mL of physiologic saline solution at the same rate as the contrast medium.

**Image Postprocessing and Data Analysis**

The CTA coronal MIP images of 20-mm thick MCA M1 segments were reconstructed from a pixel matrix of 512 × 512 for C-CTA with a 0.5-mm thickness and a pixel matrix of 1024 × 1024 for UHR-CTA with a 0.25-mm thickness with a strong AIDR 3D. C-CTA was reconstructed with a weak AIDR 3D using the analysis software installed in the CT console. Previous studies evaluating the LSAs on MR imaging and CTA determined the number of LSA branches within 5 mm from the MCA and the number of branches beyond 5 mm.\(^1\)\(^2\)\(^3\)\(^4\)\(^5\)\(^6\)\(^7\)\(^8\)\(^9\)\(^10\) The number of LSAs, defined as those longer than 5 mm, were analyzed.\(^12\) Two board-certified radiologists (K.M. with 14 years of experience; S.S. with 14 years of experience) participated as observers. A standard window level and width were applied during reader observations. The observers were allowed to freely vary window levels and widths.

**Statistical Analysis**

We compared the mean visible number of LSAs recorded by 2 blinded and independent observers between the C-CTA and UHR-CTA images. A Wilcoxon matched-pairs signed rank test was used to calculate P values for comparisons. For all statistical analyses, a P value of <.05 indicated a statistically significant between-group difference. The statistical analyses were performed using commercially available statistical software (GraphPad Prism software, Version 6, GraphPad Software, San Diego, California; BellCurve for Excel, Version 2.11, Social Survey Research Information, Tokyo, Japan).

**Radiation Dose**

The estimated volume CT dose index (CTDIvol) for CTA displayed on the CT scanner console was recorded for each patient. The estimated dose-length product was calculated as CTDIvol × Scan Length. The radiation dose was also compared against the diagnostic reference level of the Japan Network for Research and Information on Medical Exposure (J-RIME) (http://www.radher.jp/J-RIME/report/DRLhoukokusyoEng.pdf). The J-RIME is the network used to collect data related to medical exposure and to construct a framework within Japan for appropriate medical radiation doses. The J-RIME suggested that national diagnostic reference levels established in 2015 for routine brain CT examination were 85 mGy (CTDIvol) and 1350 mGy × cm (dose-length product).

**RESULTS**

The number of LSAs determined by the 2 observers is summarized in Fig 1. There were statistically more LSAs identified on UHR-CT (average, 2.85 ± 0.83; 95% CI, 2.509–3.183) than on C-CT (average, 2.17 ± 0.76; 95% CI, 1.866–2.480) (P = .009). There were also more LSAs identified on UHR-CT than on C-CT in 16 of the total 26 MCA M1 segments. Figures 2 and 3 show representative images that clearly show bilateral LSAs from the proximal-to-distal tip in UHR-CTA compared with LSAs imaged by C-CTA, respectively.

The CTDIvol obtained with C-CT was 9.7–37.7 mGy (mean, 27.2 mGy), whereas it was 44.7–63.1 mGy (mean, 53.8 mGy) with UHR-CT. The estimated dose-length product range was calculated as CTDIvol × Scan Length, which was determined as 124.9–488.1 mGy × cm (mean, 266.5 mGy × cm) with C-CT and 685.8–942.5 mGy × cm (mean, 740.8 mGy × cm) with UHR-CT. The radiation doses for both C-CT and U-HRCT
Our results suggest that UHR-CT provides higher quality images of intracranial CTA than C-CT because of improved spatial resolution and partial volume effect. UHR-CTA is a simple, noninvasive, and easily accessible method to evaluate microvasculature such as the LSAs.

LSA imaging has various important clinical roles. Surgically, it is important to accurately assess the anatomic structures of the LSAs before insular glioma surgery and coiling and clipping of MCA aneurysms.13 Additionally, LSA imaging is an important step toward reducing postoperative complications.9 Patients with hypertension have significantly fewer LSAs compared with healthy volunteers.14 It is also important to evaluate LSA features in patients with symptomatic intracranial atherosclerotic stenosis.3 Investigations using flow-sensitive black-blood MRA with MR imaging have confirmed that it is superior for visualizing LSA branches in patients with stroke.11,15 Other studies using 7T MR imaging have reported advanced noninvasive methods for visualization of the cerebral microvasculature, including the LSAs.16,17 However, it is not easy to image the LSA when using TOF-MRA with conventional 1.5T MR imaging systems, which is the most prevalent technique currently available. Digital subtraction angiography is considered the criterion standard of cerebral vascular imaging, and 3D rotational angiography on an x-ray angiographic system is useful for identifying an important anatomic relationship between the MCA trunk and LSAs;18 however, it is an invasive procedure and has some risk of complications. Therefore, noninvasive CTA or MR imaging is necessary for imaging microvasculature, such as LSAs.

The world’s first UHR-CT scanner is equipped with a 0.25-mm detector of the physical size scaled back to the isocenter that has pixels one-quarter the size of those of the C-CT scanner.5 Kakinuma et al5 reported that the prototype UHR-CT scanner had a significantly better image quality for lung nodules than conventional high-resolution CT scanners. As has been observed in previous phantom tests and clinical studies of other organs,5–8 the results of our investigation showed that UHR-CTA improved the image quality of the LSAs compared with C-CTA (Figs 2 and 3). This difference is likely

DISCUSSION

were still lower than the diagnostic reference levels according to J-RIME.
attributable to the spatial resolution of the UHR-CT and its associated reduction in the partial volume effects for the following reasons: First, UHR-CT provides significantly higher quality images of intracranial CTA using detectors with half the thickness in the z-axis direction (0.25 mm × 160 detectors) and twice as many channels in the x- and y-axis directions (1792 channels) on the super-high-resolution data-acquisition mode compared with C-CT. Second, the focal spot size of the UHR x-ray tubes (0.4 × 0.5 mm) is smaller than that of C-CT x-ray tubes (0.9 × 0.8 mm). Finally, UHR-CT is reconstructed by a higher resolution matrix (1024 × 1024) compared with C-CTA (matrix size, 512 × 512). On the basis of these factors, UHR-CT appears to provide significantly higher quality images of intracranial CTA than C-CT. Thus, UHR-CTA allows us to accurately assess the variation and structure of LSAs.

Differences in the imaging quality of C-CTA and UHR-CTA include the difference between helical and nonhelical scanning in addition to the spatial resolution as mentioned above; these differences influence the image quality of contrast-enhanced images. Because UHR-CT must use helical scanning, UHR-CTA image homogeneity with helical scans can be inferior to that of C-CTA with nonhelical scanning. Furthermore, it may be difficult to obtain images with optimal contrast-enhanced timing because UHR-CTA requires a longer scanning time. UHR-CTA is significantly superior to C-CTA for imaging the LSAs despite these disadvantages; this difference implies the greater advantages of UHR-CTA compared with C-CTA. In some cases, the number of LSAs on UHR-CT was the same or fewer than on C-CT for the following reasons: CTA scanning on UHR-CT must use the super-high-resolution mode, which has the smallest size of scan focus. The super-high-resolution mode has the technical upper limit of radiation output power; therefore, all cases showed nearly the same values for CTDI and dose-length product. As a result, we predicted that this occurred when the radiation doses were not sufficient for UHR-CT imaging to visualize LSAs in some cases.

This study has several limitations. First, a small number of subjects were included in this retrospective analysis; future prospective studies should include more subjects. Second, the scan protocols such as the FOV, reconstruction, and AIDR 3D were not the same between UHR-CTA and C-CTA because these scan protocols include the difference between helical and nonhelical scanning. It was difficult to visualize LSAs in some cases, the number of LSAs on UHR-CT was the same or fewer than on C-CT for the following reasons: CTA scanning on UHR-CT must use the super-high-resolution mode, which has the smallest size of scan focus. The super-high-resolution mode has the technical upper limit of radiation output power; therefore, all cases showed nearly the same values for CTDI and dose-length product. As a result, we predicted that this occurred when the radiation doses were not sufficient for UHR-CT imaging to visualize LSAs in some cases.

CONCLUSIONS

Improvement in LSA visualization was the result of the combined package of the UHR-CT scanner plus the UHR scanning protocol, which includes a higher radiation dose with lower than the national diagnostic reference levels and stronger AIDR processing. This package for UHR-CT is a simple, noninvasive, and easily accessible method to evaluate microvasculature such as the LSAs.

ACKNOWLEDGMENT

The authors would like to thank Maruzen-Yushodo Co Ltd (https://kw.maruzen.co.jp/kousei-honyaku/) for the English language editing.

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