Tracer Delay–Insensitive Algorithm Can Improve Reliability of CT Perfusion Imaging for Cerebrovascular Steno-Occlusive Disease: Comparison with Quantitative Single-Photon Emission CT

BACKGROUND AND PURPOSE: Reliability of CT perfusion (CTP) algorithms has not been fully validated. We investigated whether the cerebral blood flow (CBF) values obtained by using a dynamic CTP technique with a tracer delay–insensitive deconvolution algorithm are more accurate than those obtained by using CTP with delay-sensitive algorithms in unilateral cerebrovascular steno-occlusive disease, when compared with those generated by quantitative single-photon emission CT (SPECT).

MATERIALS AND METHODS: Using CTP and iodine-123-N-isopropyl-p-iodoamphetamine SPECT with an autoradiographic quantification technique, we examined 20 patients with suggested hemodynamic ischemia due to stenosis or occlusion of the unilateral internal carotid or middle cerebral artery. The algorithms used for CTP included delay-insensitive block-circulant singular value decomposition (SVD) (bSVD) and delay-sensitive standard SVD (sSVD) and box-modulation transfer function (bMTF).

RESULTS: Absolute CBF values obtained by using CTP with bSVD were significantly lower than those obtained with SPECT, but the ratios to the nonaffected side were significantly correlated to the quantitative SPECT values with significant agreements, particularly when the arterial input function was obtained from the unaffected side. Contrastingly, CBF ratios with sSVD and bMTF were significantly underestimated, and no significant agreement was determined between CTP with sSVD or bMTF and SPECT, though there were substantial correlations between them in some parameters.

CONCLUSIONS: With the CTP technique, the insensitivity of the deconvolution algorithm to the tracer-delay effect appears to be essential for estimating semiquantitative CBF values in patients with unilateral steno-occlusive lesions.
a power injector at a flow rate of 4 mL/s. Five seconds after the injection, dynamic scanning was performed at the basal ganglia level by using the following parameters: 4 contiguous 8-mm-thick sections, 80 kVp, 40 mA, 1.5 s/rotation (60 mAs), and 30 rotations (total scanning time, 45 seconds). Radiation doses of the scanning protocol were as follows: volume CT dose index, 150 mGy; dose-length product, 480 mGy cm; and effective dose, 1.34 mSv.

Postprocessing of the acquired data was performed by using the CTP analysis software obtained from a commercial workstation (M900 Quadra; Ziosoft, Tokyo, Japan). In this software, 3 types of deconvolution algorithms were implemented. A region of interest of the arterial input function (AIF) was manually applied at single branch of the insular segment of the MCA at either the contralateral or the ipsilateral side of the affected hemisphere. A region of interest of the venous output function was also established on the superior sagittal sinus. The mean transit time (MTT) was obtained as a first moment of the residue function by using 3 types of deconvolution techniques: block-circulant singular value decomposition (bSVD), which is theoretically delay insensitive; standard SVD (sSVD), which is widely used and known as delay sensitive; and box-modulation transfer function (bMTF), which is delay-sensitive and used in many institutions in Japan. The threshold of the bSVD and sSVD was set as 20%.

Cerebral blood volume (CBV) values were obtained by dividing the area under the curve of the brain tissue by the area under the curve of the venous output function after automatic pixel-by-pixel determination of the start point and discard of the second bolus. Subsequently, CBF values were calculated by dividing the CBV by the MTT, according to the central volume theorem. Vascular pixels were defined as those with CBV values >8 mL/100 mg and were automatically eliminated. CBF values of the bilateral cortical areas of MCA territories in the single section at the level of the basal ganglia and thalamus were measured by using a pair of line-symmetric polygon-shaped regions of interest obtained with cursor-driven manual tracing (Fig 1). Color maps were generated after determining the range of the color scale as follows: The maximum value of the range was set to twice the average pixel value of the entire unaffected hemisphere and the minimum value was set to zero.

**SPECT Study and Data Processing**

Iodine-123-N-isopropyl-p-iodoamphetamine (123I-IMP) SPECT examination was conducted by using a ring-type scanner (Headtome-SET080, Shimadzu, Kyoto, Japan). During 20 minutes after the initiation of intravenous infusion of 123I-IMP (5 mL, 222 MBq) and subsequent saline infusion (5 mL) at a rate of 5 mL/min, data were acquired with a midscanning time of 30 minutes. At 10 minutes after the initiation of infusion, 2-mL arterial blood was drawn from the brachial artery. For quantitative analysis, radioactivity was measured by using an autoradiography model, as described previously. The CBF of the MCA cortical areas in the section corresponding to that of CTP was measured by using a polygon-shaped region of interest that was carefully drawn to imitate the corresponding region of interest on CTP (Fig 1). Obtained CBF values were calibrated to those equivalent to CBF values in positron-emission tomography (PET) by using the following equation: 

\[ \text{CBF}_{\text{PET}} = 2.04 \times \text{CBF}_{\text{SPECT}} - 12.26 \]

Color maps were generated by using the same method as that used in CTP.

**Statistical Analysis**

The difference in quantitative CBF and semiquantitative CBF (a ratio of affected side to the healthy contralateral side) values was examined among bSVD, sSVD, bMTF, and SPECT by using a repeated measures analysis of variance (ANOVA) followed by the Tukey test. Correla-
tions of these values between CTP and SPECT were examined by using linear regression analysis. Sensitivity and specificity of the semi-quantitative CBF on CTP with respect to SPECT were calculated on a region-by-region basis when the cutoff line for the misery perfusion was set as <80% of the contralateral hemisphere, as applied for SPECT in the Japanese External Carotid-Internal Carotid Bypass Trial.\(^\text{19}\) Intermodality agreements were also examined by intraclass correlation coefficients (ICCs). The \(\alpha\) level for all the analyses was 0.05.

**Results**

The absolute CBF values varied significantly among bSVD, sSVD, and bMTF, particularly when the AIF was obtained from the affected side (Figs 1 and 2). Differences in the CBF values were statistically significant among these algorithms (\(P < .01\), repeated measures ANOVA with the Tukey test) except for those between bSVD and sSVD in the affected hemisphere (Fig 2). Among them, the CBF values obtained by using sSVD with the AIF from the affected side exhibited a good correspondence with the values generated by SPECT. However, the values obtained by using sSVD with the AIF from the unaffected side and those obtained by using bSVD in any condition were significantly lower than the values generated by SPECT (\(P < .01\)), whereas those obtained by using bMTF except in the affected hemisphere with the unaffected AIF were significantly greater than those obtained by using SPECT (\(P < .01\)) (Fig 2).

With regard to the correlation of the absolute CBF values between CTP and SPECT, the CBF values obtained by using the bSVD with the unaffected AIF (unaffected and affected hemispheres: \(r^2 = 0.58\) [\(P = .008\)] and 0.46 [\(P = .04\)], respectively; linear regression analysis) as well as those obtained by using the sSVD of the unaffected hemisphere (unaffected and affected AIFs: 0.52 [\(P = .02\)] and 0.46 [\(P = .04\)], respectively) correlated significantly to the SPECT values. Besides these combinations, no combination exhibited a statistically significant correlation (Table). Intertechnique agreements of the absolute CBF values between CTP and SPECT were found only when using sSVD (unaffected hemisphere with the affected AIF, 0.43 [\(P < .05\)], respectively; interclass correlation ICC) (Table).

Semi-quantitative values (CBF ratios of the affected hemisphere to the corresponding unaffected hemisphere) obtained by using bSVD were significantly different from those obtained with sSVD or bMTF (\(P < .01\), repeated measures ANOVA with the Tukey test) and corresponded to those generated by SPECT (Fig 3). The values obtained by using bSVD correlated significantly to those generated by SPECT (unaffected and affected AIF; \(r^2 = 0.70\) [\(P < .001\)] and 0.60 [\(P = .005\)], respectively), and the agreements between them were also substantially high (0.64 [\(P < .001\)] and 0.58 [\(P = .004\)], respectively; ICC). Correlations between sSVD and bMTF and SPECT were statistically significant only when the AIFs were
obtained from the unaffected side ($r^2 = 0.50$ [$P = .02$] and $0.57$ [$P = .008$], respectively), but their ICCs were remarkably low (Table).

When the cutoff line of the semiquantitative CBF ratio for the misery perfusion was set as $<80\%$ of the contralateral hemisphere, the sensitivity and specificity of the CBF ratio in bSVD with respect to SPECT were relatively high, 75\% and 100\%, respectively, with the unaffected AIF, but 25\% and 94\%, respectively, with the affected AIF. In sSVD and bMTF, however, the specificities of this ratio were remarkably low, though the sensitivities were high: The sensitivity and specificity of the unaffected AIF were 100\% and 31\%, respectively, in sSVD and 100\% and 13\%, respectively, in bMTF. Those with the affected AIF were 100\% and 38\%, respectively, in sSVD and 75\% and 19\%, respectively, in bMTF (Fig 3).

**Discussion**

In this study, we successfully demonstrated that the semiquantitative CBF values obtained by using CTP with the bSVD algorithm, which is known as tracer delay–insensitive, correlated well with those generated by $^{129}$I-IMP SPECT with autoradiography quantification, particularly when the AIF was obtained from the second part of the MCA of the unaffected side. However, the CBF values obtained by using CTP with sSVD and bMTF varied significantly and were not consistent with the values generated by SPECT. Besides this study, to our knowledge, there were few investigations performing a comparison among the different deconvolution algorithms for CTP and elucidating the advantages of tracer delay–insensitive algorithms over other common algorithms. Nonetheless, several reports on DSC-MR imaging have stated that tracer-arrival delay can result in the overestimation of MTT prolongation and CBF reduction and that its compensation can correct this error.$^{11-14}$

In the field of DSC-MR imaging, deconvolution algorithms for the intravascular tracer model have been thoroughly investigated. The sVD technique is widely used for the quantitative evaluation of MTT and CBF; however, it is known to be susceptible to the delay in the arrival of contrast materials to the brain tissue, leading to a miscalculation of MTT and CBF.$^{12,13}$ Algorithms that compensate for the delayed arrival time of the tracer$^{12}$ and those that are inherently insensitive to tracer delay have already been proposed, such as bSVD$^6$ and fast Fourier transform$^{13}$ to overcome this issue.

On the other hand, few studies concerning CTP have focused on deconvolution algorithms.$^{15,16}$ Although different algorithms are implemented in commercial software packages provided by CT and workstation vendors, to our knowledge, a direct comparison between algorithms or software packages has hardly been performed. Among these algorithms, sSVD is commonly used and bSVD, bMTF, and fast Fourier transform are used in a limited number of software packages. One of the vendors uses a maximum slope model$^{16}$ instead of a deconvolution technique. The previously mentioned variety of analysis algorithms can result in considerable differences in the generated perfusion maps; however, this problem has not been investigated enough. Thus, we performed a direct comparison of 3 different algorithms by using otherwise identical parameters and by using a software program that is the commercial package available with both tracer delay–sensitive and tracer delay–insensitive algorithms.

It is generally believed that CTP has an advantage over DSC-MR imaging with respect to its 1) quantitativeness, due to the linear relationship between the concentration of the contrast agent and the CT value; and 2) the easy acquisition of AIF.$^{15,16}$ However, we revealed that the absolute CBF values on CTP varied remarkably among algorithms and were significantly different from those generated by SPECT in patients with hemodynamic ischemia with unilateral steno-occlusive lesions. Previously, several studies have demonstrated a good correlation in CBF values between CTP and PET or Xenon CT. However, these studies included no subjects with hemody-
namic ischemia in whom substantial tracer-arrival delay occurred. On the other hand, the ratios of the CBF in the affected hemisphere to the contralateral hemisphere correlated well with those generated by SPECT only when using bSVD, though its absolute values were apparently smaller than those generated by using sSVD, bMTF, and SPECT, presumably due to oscillations of the residue function caused by large image noises. Our results suggest that the accuracy of the semiquantitative CTP values in unilateral steno-occlusive conditions strongly depend on the deconvolution algorithms, particularly on the insensitivity to tracer-arrival delay.

The source of AIF is another important factor that influences CBF values obtained by using CTP. Although an AIF closest to the brain tissue is ideal, it cannot be distinctly obtained. Therefore, a surrogate AIF is usually measured at the intracranial major arteries such as the MCA. Its location (affected or unaffected side) can thus result in significant variation in CBF quantification, mainly with regard to tracer-arrival delay, bolus dispersion, and partial volume effects. In this study, the quantitative and semiquantitative CBF values generated by using the MCA ipsilateral to the affected hemisphere as the source of AIF were substantially different from those generated by using the contralateral MCA; this difference was observed particularly when tracer delay–sensitive sSVD and bMTF were used. The bSVD algorithm appears to afford relatively robust results regardless of the difference in AIF locations, but quantitative CBF values with the AIF from the affected side could not show a significant correlation with those generated by SPECT. We speculate that dispersion of the first bolus curve, partial volume effect of AIF, and/or image noise can deteriorate the accuracy of CBF values. A sophisticated technique for automatically obtaining an appropriate surrogate AIF is required to improve further reproducibility of the CTP results even when using bSVD.

One of the limitations of this study was that only a single software program was used for the evaluations and intervendoor differences were not examined. The quantitative and semiquantitative CBF values may be different among different software packages, even when the same algorithm is implemented. Currently, we are investigating this issue by analyzing identical source data by using a number of commercial software packages. Another limitation of our study was that a few patients with severe ischemia were included. Only 4 of the 20 patients were reported to have <80% CBF ratio by SPECT. The calculated specificity and sensitivity of 3 algorithms against SPECT with the cutoff line of 80% included substantial statistical errors. The third limitation was that no acetazolamide challenge test was performed in CTP because of ethical issues. It has not been elucidated whether deconvolution algorithms affect the results of the acetazolamide challenge test, which estimates regional cerebrovascular reserve capacity in cases of hemodynamic ischemia. We speculated that the tracer delay–insensitive algorithm would be preferable because acetazolamide may affect the AIF and the time taken by the contrast medium to reach the brain tissue. Another limitation was that we only evaluated CBF values in the cortical areas of the MCA territory but did not assess differences in the volume showing CBF abnormalities among deconvolution techniques, which appear to be important clinically, because there were a few patients showing remarkable decrease in the CBF. Further investigation with a large number of patients with severe hemodynamic ischemia is needed to elucidate how the volume with CBF decrease varies among the algorithms.

In this study, CBF quantified not by PET but by the 123I-IMP autoradiography method by using SPECT was defined as the control, because the value obtained by this method significantly correlates with that obtained by the H215O PET method. In particular, the fit to the regression line of the 2 values is high in the hypoperfusion (lower than 35 mL·100 mg⁻¹·min⁻¹ on SPECT) areas. Although PET is widely used as a gold standard for quantitative CBF, we believe that the recent technical development in SPECT with regard to quantification enables its use as a gold standard.

Correlation coefficients of the CBF values between CTP and SPECT in this study were substantially low even when using the bSVD method. The low radiation dose that we applied might deteriorate accuracies during deconvolution processing, though deconvolution algorithms are said to be relatively robust to the image noise. Further optimization of the deconvolution algorithms can also improve the accuracy. The manual tracing method we used for quantitative measurement would cause some additional biases, so some sophisticated methods using coregistration techniques may improve the results.

CTP is widely applied for assessing not only chronic ischemia but also acute ischemic stroke. Previous studies concerning DSC-MR imaging have already revealed that tracer delay–sensitive algorithms tend to overestimate MTT prolongation and CBF reduction in patients with embolic occlusion of the major arteries. We believe that tracer delay–insensitive algorithms such as bSVD are much more appropriate for generating CTP maps that are used for assessing patients with acute stroke, including candidates for thrombolytic therapy. Evaluation of perfusion abnormalities may be crucial for expanding the time window of the thrombolytic therapy beyond 3 hours. The use of the deconvolution algorithm that is insensitive to the tracer-delay effect will facilitate the decrease in patient selection bias and will improve patient outcome.

Conclusions

We demonstrated that delay-sensitive deconvolution algorithms are inaccurate for evaluating CBF changes by using CTP and that the delay-insensitive algorithms are much more suitable for the purpose when semiquantitative ratios, not absolute values, are applied. The insensitivity to the delay of tracer arrival in CTP algorithms appears to be essential for reliable assessment of perfusion abnormalities in patients with unilateral steno-occlusive lesions.

References

1. Lev MH, Segal AZ, Farkas J, et al. Utility of perfusion-weighted CT imaging in acute middle cerebral artery stroke treated with intra-arterial thrombolysis: prediction of final infarct volume and clinical outcome. Stroke 2001;32:2021–28
2. Koenig M, Kraus M, Theek C, et al. Quantitative assessment of the ischemic brain by means of perfusion-related parameters derived from perfusion CT. Stroke 2001;32:431–37
3. Chen A, Shyr MH, Chen YJ, et al. Dynamic CT perfusion imaging with acetazolamide challenge for evaluation of patients with unilateral cerebrovascular steno-occlusive disease. AJNR Am J Neuroradiol 2006;27:1876–81
4. Ostergaard L, Weisskoff RM, Chester DA, et al. High resolution measurement of cerebral blood flow using intravascular tracer bolus passages. Part I. Mathematical approach and statistical analysis. Magn Reson Med 1996;36:715–25
5. Ostergaard L, Sorensen AG, Kwong KK, et al. High resolution measurement of cerebral blood flow using intravascular tracer bolus passages. Part II. Experimental comparison and preliminary results. Magn Reson Med 1996;36:726–36
6. Wu O, Ostergaard L, Weisskoff RM, et al. Tracer arrival timing-insensitive technique for estimating flow in MR perfusion-weighted imaging using singular value decomposition with a block-circulant deconvolution matrix. Magn Reson Med 2003;50:164–74
7. Nambu K, Takehara T, Kodama M, et al. A method of regional cerebral blood perfusion measurement using dynamic CT with an iodinated contrast medium. Acta Neurol Scand Suppl 1996;166:28–31
8. Meier P, Zierler KL. On the theory of the indicator-dilution method for measurement of blood flow and volume. J Appl Physiol 1954;6:731–44
9. Ogasawara K, Ito H, Sasoh M, et al. Quantitative measurement of regional cerebrovascular reactivity to acetazolamide using 123I-N-isopropyl-p-iodoamphetamine autoradiography with SPECT: validation study using H215O with PET. J Nucl Med 2003;44:520–25
10. Minoura S, Nakagawa I, Takahashi M, et al. Three-dimensional display in staging hemodynamic brain ischemia for JET study: objective evaluation using SEE analysis and 3D-SSP display. Ann Nucl Med 2004;18:13–21
11. Calamante F, Gadian DG, Connelly A. Delay and dispersion effects in dynamic susceptibility contrast MRI: simulations using singular value decomposition. Magn Reson Med 2000;44:666–73
12. Ibaraki M, Shimosegawa E, Toyoshima H, et al. Tracer delay correction of cerebral blood flow with dynamic susceptibility contrast-enhanced MRI. J Cereb Blood Flow Metab 2005;25:378–90
13. Smith MR, Lu H, Trochet S, et al. Removing the effect of SVD algorithmic artifacts present in quantitative MR perfusion studies. Magn Reson Med 2004;51:631–34
14. Willats L, Connelly A, Calamante F. Improved deconvolution of perfusion MRI data in the presence of bolus delay and dispersion. Magn Reson Med 2006;56:146–56
15. Wintermark M, Maeder P, Thiran JP, et al. Quantitative assessment of regional cerebral blood flows by perfusion CT studies at low injection rates: a critical review of the underlying theoretical models. Eur Radiol 2003;11:1220–30
16. Tomandl BF, Klotz E, Handschu R, et al. Comprehensive imaging of ischemic stroke with multi-section CT. Radiographics 2003;23:565–92
17. Wintermark M, Thiran J, Maeder P, et al. Simultaneous measurement of regional cerebral blood flow by perfusion CT and stable Xenon CT: a validation study. AJNR Am J Neuroradiol 2001;22:905–14
18. Kudo K, Terae S, Katoh C, et al. Quantitative cerebral blood flow measurement with dynamic perfusion CT using the vascular-pixel elimination method: comparison with H215O positron emission tomography. AJNR Am J Neuroradiol 2003;24:419–26
19. Soustiel JF, Mor N, Zaaroor M, et al. Cerebral perfusion computerized tomography: influence of reference vessels, regions of interest and interobserver variability. Neuroradiology 2006;48:870–77. Epub 2006 May 23
20. Latchaw RE, Yonas H, Hunter GI, et al. Guidelines and recommendations for perfusion imaging in cerebral ischemia: a scientific statement for healthcare professionals by the writing group on perfusion imaging, from the Council on Cardiovascular Radiology of the American Heart Association. Stroke 2003;34:1084–104