Optimal Duration of Acquisition for Dynamic Perfusion CT Assessment of Blood-Brain Barrier Permeability Using the Patlak Model


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Parenchymal hematoma with significant mass effect, the most severe form of hemorrhagic transformation (HT) in acute ischemic stroke, occurs in ≈5.9% of patients treated with tissue plasminogen activator1 and has devastating clinical consequences.2 Because blood-brain barrier (BBB) damage plays a fundamental role in HT,3 early identification of significant BBB damage on admission might predict which patients will develop HT and could constitute a contraindication to thrombolytic therapy.4 In the pursuit of this goal, direct measurements of the BBB permeability surface product (BBBP) have been attempted in the context of acute stroke.5–8

Prior perfusion CT (PCT) studies have applied Patlak modeling to first-pass data to calculate BBBP.7–10 However, a recent study demonstrated that using first-pass data resulted in incorrect BBBP measurements by modeling delayed perfusion rather than permeability.11 This study demonstrated the necessity of using delayed-acquisition data but did not investigate specifically how long the delayed acquisition needed to be. Rather, the authors arbitrarily used an acquisition of ≲4 minutes.

On one hand, the duration of the delayed acquisition should be long enough to meet the assumptions of the Patlak model and to allow an accurate measurement of BBBP; such measurements require distinguishing intravascular and extravascular (extravasated) contrast, which is facilitated by a delayed acquisition not dominated by first-pass intravascular contrast.12 On the other hand, minimizing the duration of the delayed acquisition reduces the likelihood of motion artifacts that may degrade image quality and interfere with BBBP calculation,13 mainly when these motion artifacts are severe and cannot be corrected by image registration.

The goal of our current study was to determine how short the delayed acquisition can be while still meeting the Patlak assumption of linearity.

Materials and Methods

Study Design

Imaging data obtained as part of standard clinical stroke care at our institution were retrospectively reviewed with the approval of the institutional review board. At our institution, patients with suspicion of acute stroke and no history of significant renal insufficiency or con-
Table 1: Study patient characteristics

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Patients with Stroke</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of patients</td>
<td>23</td>
</tr>
<tr>
<td>No. of men (%)</td>
<td>8 (35%)</td>
</tr>
<tr>
<td>Median age (years)</td>
<td>74</td>
</tr>
<tr>
<td>Interquartile range</td>
<td>58.5–82.5</td>
</tr>
<tr>
<td>Range</td>
<td>26–92</td>
</tr>
<tr>
<td>Median time from stroke to PCT (hours)</td>
<td>2.25</td>
</tr>
<tr>
<td>Interquartile range</td>
<td>1.5–5</td>
</tr>
<tr>
<td>Range</td>
<td>1–11.75</td>
</tr>
<tr>
<td>Stroke location</td>
<td></td>
</tr>
<tr>
<td>ACA, MCA territories</td>
<td>3</td>
</tr>
<tr>
<td>MCA territory</td>
<td>20</td>
</tr>
<tr>
<td>ACA territory</td>
<td>0</td>
</tr>
<tr>
<td>PCA territory</td>
<td>0</td>
</tr>
</tbody>
</table>

Note: —ACA indicates anterior cerebral artery; MCA, middle cerebral artery; PCA, posterior cerebral artery; PCT, perfusion CT.

Contrast allergy routinely undergo a stroke CT survey, including noncontrast CT (NCCT) of the brain, PCT at 2 cross-sectional positions, CT angiography (CTA) of the cervical and intracranial vessels, and postcontrast cerebral CT, obtained in this chronologic sequence.

We retrospectively identified a consecutive series of 23 patients admitted to the University of California San Francisco Medical Center from July 2007 to August 2008 who met the following inclusion criteria: 1) admission to the emergency department with signs and symptoms suggesting anterior circulation stroke within 12 hours after symptom onset; 2) documentation of acute ischemic anterior circulation stroke by both admission stroke protocol and clinical examination; 3) no evidence of intracerebral hemorrhage on the admission NCCT; and 4) no significant motion artifacts following application of the registration algorithm. Patients’ charts were reviewed for demographic and clinical data.

Imaging Protocol

PCT studies were obtained on 16-section (4 patients) and 64-section (19 patients) CT scanners. Each PCT study involved successive gantry rotations performed in cine mode during intravenous administration of iodinated contrast material. Images were acquired and reconstructed at a temporal sampling rate of 1 image per second for the first 37 seconds and 1 image every 2 seconds for the next 33 seconds. Additional gantry rotations were obtained at 90, 120, 150, 180, 210, and 240 seconds. Acquisition parameters were 80 kilovolt (peak) and 100 mAs. Two successive PCT series at 2 different levels were performed following the NCCT and before the CTA. At each PCT level, two 10-mm-thick sections (16-section CT scanners) or eight 5-mm-thick sections (64-section CT scanners) were assessed. The first PCT series was performed at the level of the third ventricle and the basal ganglia, and the second PCT series, above the lateral ventricles. For each PCT series, a 40-mL bolus of iohexol (300 mg/mL of iodine, Omnipaque; GE Healthcare, Piscataway, NJ) was administered into an antecubital vein by using a power injector at an injection rate of 5 mL per second for all patients. CT scanning was initiated 7 seconds and 1 image every 2 seconds for the next 33 seconds. Additional gantry rotations were obtained at 90, 120, 150, 180, 210, and 240 seconds. Acquisition parameters were 80 kilovolt (peak) and 100 mAs. Two successive PCT series at 2 different levels were performed following the NCCT and before the CTA. At each PCT level, two 10-mm-thick sections (16-section CT scanners) or eight 5-mm-thick sections (64-section CT scanners) were assessed. The first PCT series was performed at the level of the third ventricle and the basal ganglia, and the second PCT series, above the lateral ventricles. For each PCT series, a 40-mL bolus of iohexol (300 mg/mL of iodine, Omnipaque; GE Healthcare, Piscataway, NJ) was administered into an antecubital vein by using a power injector at an injection rate of 5 mL per second for all patients. CT scanning was initiated 7 seconds after start of the injection of the contrast bolus. Data from both boluses were used because prior work has demonstrated that there is no significant parenchymal saturation effect from the first bolus and no underestimation of BBBP values from data from the second bolus.

Image Postprocessing

PCT data were analyzed using commercially available PCT software (Brain Perfusion; Philips Healthcare, Cleveland, Ohio). This software relies on the central volume principle, which is the most accurate for low injection rates of iodinated contrast material.

The software obtains mathematic descriptions of the time-attenuation curves for each voxel, by applying curve fitting by least mean squares, after correcting for motion and noise reduction through an anisotropic edge-preserving spatial filter. A closed-form (noniterative) deconvolution is then applied to calculate the mean transit time (MTT) map. The deconvolution operation requires a reference arterial input function (most often within the anterior cerebral artery), automatically selected by the PCT software within a region of interest drawn by the user. The cerebral blood volume (CBV) map is calculated from the area under the time-attenuation curves, with a correction factor for the leakage of contrast into the interstitial space calculated from the Patlak model applied to the delayed acquisition. The PCT infarct core and salvageable brain tissue are automatically calculated by the software by using CBV thresholds and MTT thresholds reported in the literature as the most accurate (PCT salvageable brain tissue: MTT > 145% of the contralateral side values plus CBV = 2.0 mL × 100 g⁻¹; PCT infarct core: MTT > 145% of the contralateral side values plus CBV < 2.0 mL × 100 g⁻¹).

BBBP measurements were extracted from PCT data by using a second prototype software developed by Philips Healthcare. This software is based on the Patlak model. Applying the Patlak model to PCT involves performing linear regression by using data calculated from the PCT datasets. The slope of these regression lines was interpreted as a blood-to-brain transfer constant and used as an indicator of absolute BBBP values. Relative BBBP values, defined as absolute BBBP values divided by cerebral blood flow values in the corresponding voxels, were also recorded.

Using the prototype software, we calculated BBBP values from the gold standard delayed acquisition dataset, which incorporated all time points from 90 to 240 seconds. Additionally, we calculated BBBP values by using truncated datasets based on analysis windows from 90 to 210 seconds (excluding the data from the last gantry rotation), 90 to 180 seconds (excluding the data from the last 2 gantry rotations), 90 to 150 seconds (excluding the data from the last 3 gantry rotations), and 90 to 120 seconds (excluding the data from the last 4 gantry rotations). BBBP values were recorded in the infarct and penumbra, as automatically delineated by the software, and in mirrored regions of interest in the contralateral nonischemic tissue (also automatically delineated by the software).

Statistical Analysis

We calculated the correlation between permeability values from each of the truncated analysis windows and permeability values from the gold standard 90- to 240-second delayed acquisition.

We measured the quality of the linear fit to quantify how well the assumptions of the Patlak model were met by data extracted from the datasets of 90–240, 90–210, 90–180, 90–150, and 90–120 seconds. To do this, we used the root-mean-square error (\(\sqrt{\text{MSE}}\)), which is a measure of variability of data points around a fitted straight line: A value close to zero indicates a smaller spread of data points around the line, corresponding to a better fit.

Absolute and relative BBBP values and corresponding \(\sqrt{\text{MSE}}\) values extracted from different durations of delayed PCT analysis windows in ischemic and nonischemic regions of interest were compared by using generalized estimating equation models with robust variance estimation, with fixed effects for patients, type of CT scanner, and type of regions of interest. Because the distribution of the parameters was not normal but skewed, we report estimated mean values (ob-
Radiation Dose and Motion Artifacts
We calculated the effective radiation dose associated with the gold standard 90- to 240-second acquisition and the percentage decrease in radiation associated with truncated analysis windows (datasets of 90–210, 90–180, 90–150, and 90–120 seconds).

The number of cases with motion artifacts interfering with the processing was recorded for the gold standard 90- to 240-second acquisition and for the truncated analysis windows (datasets of 90–210, 90–180, 90–150, and 90–120 seconds).

Results
Patients and Imaging Studies
Twenty-three patients who matched our inclusion criteria were retrospectively identified. Patient characteristics are summarized in Table 1. The median time from symptom onset to PCT was 2.25 hours (range, 1–11.75 hours). The Patlak analyses were performed in a total number of 230 ischemic regions of interest and 230 nonischemic regions of interest.

BBBP Measurements for Different Durations of the Delayed PCT Acquisition
BBBP values for truncated analysis windows of 90–210 seconds and 90–180 seconds were highly correlated with the gold standard 90- to 240-second delayed acquisition (Table 2). Shorter analysis windows (90–150 seconds and 90–120 seconds) were less correlated, especially in the infarct core (Table 2).

Absolute (Table 3) and relative (Table 4) BBBP values were overlapping for the gold standard 90- to 240-second delayed acquisition and for the 90- to 210-second analysis window. For shorter analysis windows (90–180 seconds and, especially, 90–150 and 90–120 seconds), absolute and relative BBBP values were overestimated in the infarct, penumbra, and non-ischemic regions of interest.
ischemic tissue (Fig 1). In terms of mean absolute BBBP values, the differences between the gold standard 90- to 240-second delayed acquisition and the 90- to 210-second analysis window were 12% and 15% in the infarct and penumbra, respectively. The differences in terms of mean absolute BBBP values in the infarct and penumbra jumped sharply to 36% and 42%, 76%, and 89%, and 146% and 166% for the 90- to 180-, 90- to 150-, and 90- to 120-second analysis windows, respectively. Very similar differences were observed for the relative BBBP results.

**Radiation Dose and Motion Artifacts**

The effective radiation dose associated with the gold standard 90- to 240-second delayed acquisition and for the 90- to 210-second analysis window. For shorter analysis windows (including 90–180 seconds and, especially, 90–150 and 90–120 seconds), MSE measurements were higher, indicating a poorer linear fitting.

**Quality of the Linear Regression according to the Patlak Model**

√MSE measurements (Table 5) were low and overlapping for the gold standard 90- to 240-second delayed acquisition and for the 90- to 210-second analysis window. For shorter analysis windows (including 90–180 seconds and, especially, 90–150 and 90–120 seconds), √MSE measurements were higher, indicating a poorer linear fitting.

**Discussion**

The Patlak model is a relatively simple and frequently applied way of calculating BBBP from PCT studies. It involves using arterial and parenchymal contrast-enhancement curves to calculate the rate of contrast extravasation from an intravascular-to-extravascular compartment. A previous study demonstrated that a delayed PCT acquisition is required to verify the assumptions of the Patlak model because applying the Patlak model to first-pass PCT data leads to overestimated and inaccurate BBBP values.

In our study, we found that the duration for the delayed PCT acquisition should be at least 210 seconds because acquisitions shorter than 210 seconds lead to significantly overestimated BBBP values. The question of whether 210 seconds or 240 seconds is preferable is debatable. On one hand, several factors favor the 240-second acquisition. The 240-second acquisition is 12%–15% more accurate, and the additional 30 seconds of scanning time and the 1.7% increase in radiation dose is very likely clinically insignificant. On the other hand, the 210-second acquisition might be associated with a slightly lower risk of motion artifact (some patients might not move up to 210 seconds but may move during the last 240-second gantry rotation of the PCT series). However, we did not observe any such patients in our small series, and registration algorithms can often correct motion artifacts as long as they are not too severe.

We acknowledge several limitations to our work. First, because the goal of our study was to evaluate the minimum duration of the delayed acquisition to achieve valid Patlak-derived BBBP results, we did not assess other non-Patlak models that can be applied to PCT data to calculate BBBP, such as the distributed parameter model. Future studies are needed to determine whether our conclusions regarding an acceptable duration for the delayed acquisition also hold true for alternative models.

Second, we did not address the relationship between permeability and variables, such as admission infarct volume, time from symptom onset to scanning, or collateral flow. Such questions were beyond the scope of this study and will be addressed in a future work.

In conclusion, the duration for the delayed PCT acquisition should be at least 210 seconds. Shorter acquisitions lead to significantly overestimated BBBP values. Further research is required to investigate the relevance of delayed-phase PCT BBBP values in terms of the predictive value for hemorrhagic transformation in patients with acute stroke.

**References**


**Table 5:** √MSE calculated for the fitting of the Patlak line for the gold standard 90- to 240-second delayed acquisition and the truncated delayed acquisitions

<table>
<thead>
<tr>
<th></th>
<th>90–120 Seconds</th>
<th>90–150 Seconds</th>
<th>90–180 Seconds</th>
<th>90–210 Seconds</th>
<th>90–240 Seconds</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nonischemic</td>
<td>4.84 (3.95–5.95)</td>
<td>3.13 (2.65–3.71)</td>
<td>2.16 (1.84–2.54)</td>
<td>1.66 (1.41–1.95)</td>
<td>1.35 (1.14–1.59)</td>
</tr>
<tr>
<td>Infarct</td>
<td>5.17 (4.10–6.52)</td>
<td>3.38 (2.73–4.19)</td>
<td>2.31 (1.87–2.84)</td>
<td>1.75 (1.43–2.15)</td>
<td>1.41 (1.15–1.73)</td>
</tr>
<tr>
<td>Penumbra</td>
<td>5.31 (4.31–5.55)</td>
<td>3.42 (2.86–4.09)</td>
<td>2.35 (1.99–2.78)</td>
<td>1.79 (1.51–2.12)</td>
<td>1.44 (1.21–1.72)</td>
</tr>
</tbody>
</table>

Note: √MSE indicates root-mean-square error.
13. Miles KA. Perfusion CT for the assessment of tumour vascularity: which protocol? Br J Radiol 2003;76(spec no 1):s36–42