

exercise-induced myocardial ischaemia and/or for assessing myocardial viability in patients with coronary artery disease. The diagnosis, however, has been limited to qualitative or visual assessment of the physical extent of the defect areas rather than quantitative assessment of physiological functions. Quantitative methods would for example enable longitudinal studies when assessing therapy response and pharmacological interventions. Some groups have already investigated the feasibility of estimating quantitative parameters with dynamic SPECT in the myocardium using ^{201}Tl [1] and $^{99\text{m}}\text{Tc}$ -Teboroxime [1, 2], but these techniques have not yet been applied to clinical practice. This is largely attributed to the fact that quantitative reconstruction programmes are not readily available on commercial SPECT systems.

We have developed a reconstruction programme package for SPECT, which can accurately provide quantitative images of radio-labelled tracer distributions *in vivo*, which is a pre-requisite for absolute physiological parameter estimation. The adequacy and accuracy of these methods have been demonstrated in multiple papers for $^{99\text{m}}\text{Tc}$ and ^{201}Tl in cardiac studies [3–5], and for $^{99\text{m}}\text{Tc}$ and ^{123}I in brain studies [6]. It has also been demonstrated, in brain studies, that physiological parameters such as cerebral perfusion [6] and cerebral flow reactivity [7] obtained using our package were as accurate as those determined by PET. These findings suggest that absolute quantitation of regional myocardial perfusion might also be possible in a clinical setting using commercial SPECT cameras.

^{201}Tl is a potassium analogue, and its kinetics has been extensively investigated in previous studies [8, 9]. Due to the high first-pass extraction fraction (EF) [10] and a large distribution volume, ^{201}Tl has been considered an ideal tracer for quantitation of absolute myocardial blood flow, not only at rest but also at hyperemic conditions. As a clinical implication, quantitative assessment of MBF and coronary flow reserve is important. For instance, coronary micro-vascular dysfunction or impaired endothelial function in patients with coronary risk factors or patients with cardiomyopathy or with heart failure is an un-resolved important issue to answer [11]. Coronary flow reserve can also be reduced in patients with hyper-cholesterolemia without overt coronary stenosis [12]. The low energy and long half-life of ^{201}Tl have, however, seriously limited its use in nuclear cardiology.

The goal of this study was to validate our reconstruction methodology for the estimation of myocardial blood flow using ^{201}Tl and dynamic SPECT using tissue time-activity curves (TTAC) derived from myocardial regions. In addition, we aimed to find the optimal kinetic model configuration and to investigate the factors affecting the estimation of physiological parameters such as the partial volume effect (PVE), appropriate choice of input function, conversion from plasma to blood flow using haematocrit (Hct) and the limited first-pass tracer EF.

Materials and methods

Subjects

A total of 21 dogs were studied in which 8 were in a resting condition, 9 dogs during constant infusion of adenosine for increased MBF, and 4 dogs during constant infusion of beta-blocker. Of the 21 studies, 1 study was unsuccessful and projection data could not be retrieved from the scanner, reducing the number of resting studies to 7 and total dog studies to 20. Adenosine was infused continuously over the study duration at a rate ranging from 140 to 700 mg/kg/h to achieve a range of blood flow increases. An initial dose of beta-blockers ranging from 2 to 6 mg was given, followed by a constant infusion for the duration of the study of 2 or 4 mg/h. The study protocol was approved by the animal ethics committee at the Akita Research Institute of Brain, Akita City, Japan where all experiments were carried out.

SPECT procedures

All dogs were anaesthetised, and the catheters for dose administration and arterial blood sampling were inserted before the study. The SPECT system was a conventional dual-head gamma camera (Toshiba GCA-7200A, Tokyo, Japan) fitted with short focal length fan-beam collimators (LEHR-Fan). The transverse field-of-view (FOV) was 22 cm diameter and axial FOV was 20 cm. The dogs were carefully taped into a cradle to minimise motion during the study, and also to ensure that no truncation occurred. Heart rate and blood pressure were monitored throughout the study and recorded at regular intervals.

Before the injection of any tracer, a 15-min transmission study was carried out in which a rod source filled with approximately 740 MBq of $^{99\text{m}}\text{Tc}$ was placed along the focal line of one of the fan-beam collimators (see Fig. 1). The transmission study was followed by injection of 3 MBq of ^{141}Ce microspheres into the left ventricle via a catheter and blood was withdrawn from the aorta at a constant flow rate of 5 ml/min for 2 min to serve as an input function. For the pharmacological intervention studies, adenosine infusion or beta-blocker injection followed by infusion was commenced before the ^{141}Ce microsphere administration.

Dynamic SPECT was commenced with the start of the 4-min constant infusion of 110 MBq ^{201}Tl . The frame collection rates and 360° rotation times were 10×1 min (rotation time 15 s), 6×2 min (30 s), 3×4 min (60 s) and 5×5 min (60 s) for the first hour for all studies. Resting blood flow studies had an additional 18×10 min (120 s) frames collected for a total study period over 4 h. The shorter total study time for the drug infusion studies was mandated by the difficulties in keeping the dogs stable with prolonged infusions of the drugs used. A 34% energy

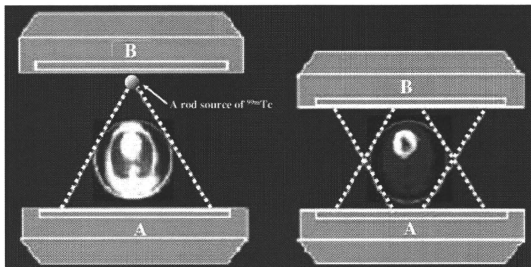


Fig. 1 Schematic diagram of data acquisition using a clinical dual-headed SPECT camera fitted with fan-beam collimators. Transmission scan was performed using a ^{99m}Tc -filled rod source placed at a focal

line of one of the collimators, and only one of the detectors was used (left). Both detectors were used in the emission scan (right)

window centred on 77 keV was used for the ^{201}Tl acquisitions [4, 13].

Arterial blood samples were taken every 20 s for the first 6 min, every 60 s for 6–10 min, 120 s for 10–20 min, 300 s for 20–30 min and 600 s for 30–60 min. For the resting studies, blood samples were also taken every 20 min for 1–2 h and additional samples at 2.5, 3 and 4 h post- ^{201}Tl infusion. In six studies, plasma was separated immediately after sampling by centrifugation, and plasma samples were counted in a well counter cross-calibrated with the SPECT scanner. To minimise the effects of the continued exchange of ^{201}Tl between plasma and red blood cells in the test tubes after sampling, immediate, rapid separation of plasma from whole blood was required. An averaged relationship between plasma and whole blood concentration ratio over time was obtained, and then multiplied with the whole blood curves for all studies to derive a plasma input function.

At the end of the SPECT study, the microsphere blood flow measurement was repeated with ^{51}Cr microspheres. The dogs were then killed by injection of potassium chloride (KCl) and the myocardium was dissected into samples suitable for counting in the well counter. The ^{201}Tl concentration in the tissue samples was derived from the sample weight normalised gamma counter counts. The samples were stored to allow for the decay of ^{201}Tl ($T_{1/2} = 73$ h vs $T_{1/2} = 32.5$ days for ^{141}Ce and 27.8 days for ^{51}Cr) and then counted to measure the ^{141}Ce and ^{51}Cr activities. Separation between ^{141}Ce and ^{51}Cr counts was based on their respective gamma ray energies (145 keV for ^{141}Ce and 323 keV for ^{51}Cr).

SPECT data processing

Projection data were processed according to previously described procedures [5]. Briefly, the transmission data obtained by the fan-beam collimator were first re-binned

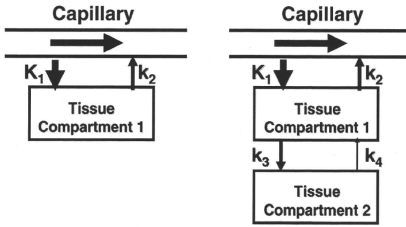
into parallel projections. Transmission projections were normalised by blank projection, re-constructed to generate quantitative maps of the attenuation coefficient for ^{99m}Tc and then linearly scaled to provide attenuation correction maps for ^{201}Tl . Emission data were corrected for detector non-uniformity and also re-binned into parallel projections. The projection data were then corrected for scatter with transmission-dependent convolution subtraction (TDCS) originally proposed by Meikle et al. [14] and further optimised by our group [4, 5]. The emission projection data were re-constructed with the OS-EM reconstruction algorithm [15] using three iterations and ten subsets. The re-constructed images were cross-calibrated with the well counter system.

Data analysis

Re-constructed images were normalised by acquisition time for each frame. Multiple circular regions of interest (ROI) were drawn on the myocardium, and the TTAC of ^{201}Tl were generated for the anterior, apical, lateral, posterior and septal areas of the myocardium. The two-compartment model (one tissue compartment) and three-compartment model (two tissue compartments) shown in Fig. 2 were applied to determine two parameters (K_1 and K_2) for the two-compartment model and four parameters (K_1 – K_4) for the three-compartment model by means of non-linear least squares fitting (NLLSF).

The regional MBF was considered to be related to K_1 obtained from compartment model fits. K_1 is, however, affected by the PVE, Hct and the limited first-pass EF whose effects were corrected according to Eq. 1:

$$\text{MBF} = \frac{\text{PVE}}{\text{EF} \times (1 - \text{Hct})} \times K_1 \quad (1)$$



2-Compartment model

3-Compartment model

Fig. 2 Two- and three-compartment models evaluated in this study. K_1 in units of ml/min/g denotes the regional MBF for both models. Distribution volume (V_d) in units of ml/g is defined as K_1/K_2 for the two-compartment model, and $\frac{K_1}{K_2} \left(1 + \frac{K_3}{K_4}\right)$ for the three-compartment model

The physiological basis for the correction factors in Eq. 1 can be described as follows:

1. TTACs obtained from SPECT images are under-estimated due to the limited spatial resolution relative to the myocardial wall thickness and also due to the myocardial contractile motion. This phenomenon is known as PVE. The PVE correction factor for each TTAC was determined from the ratio of the last SPECT frame counts to the ^{201}Tl myocardial tissue sample counts obtained from the tissue samples taken and measured with the well counter at the end of the SPECT scan.
2. The arterial input function for the compartment model studies was defined from the plasma radioactivity concentration curve, rather than the whole blood radioactivity curve. K_1 is therefore the regional “plasma” flow. Thus, for comparison with the microsphere flow measurements, which estimates the whole blood flow, K_1 was divided by $(1 - \text{Hct})$ to obtain the flow for the total blood.
3. For a tracer with limited first-pass $\text{EF} < 1.0$, flow (MBF) is related to K_1 by $K_1 = \text{EF} \times \text{MBF}$. The first-pass EF is flow-dependent and decreases at high flow. We have applied an empirical formulation for the first-pass EF based on the data by Weich et al. [10] ($\text{EF} = 0.84 - 0.524 \cdot \log_{10}(K_1^*)$) where K_1^* is $K_1 / (1 - \text{Hct})$. The K_1 values obtained with two- and three-compartment models with/without corrections according to Eq. 1 were compared to the average of microsphere blood flow values obtained pre- and post-dynamic SPECT scan.

The distribution volume of ^{201}Tl (V_d) was defined as

$$V_d = \frac{K_1}{K_2} \text{ for the two - compartment model} \tag{2a}$$

$$V_d = \frac{K_1}{K_2} \left(1 + \frac{K_3}{K_4}\right) \text{ for the three - compartment model.} \tag{2b}$$

As mentioned before, the resting studies were collected for 4 h, whilst the adenosine and beta-blocker studies were collected for approximately 1 h. To investigate whether the shorter collection time introduces systematic bias, NLLSF fits restricted to the first 1 h of the resting study data were also performed and compared with the V_d values from the full 4 h resting data set and with the estimates obtained from the beta-blocker and adenosine studies.

Akaike information criterion (AIC) and Schwarz criterion (SC) were calculated for both two-compartment and three-compartment model fits [16] to test the adequacy of the two models. All data are presented as mean \pm 1 SD. Student's t test was employed in the comparison of the V_d values. Pearson's regression analysis was applied to compare K_1 and microsphere flow values. A probability value of < 0.05 was considered statistically significant.

Results

Figure 3 shows the plasma to whole blood concentration ratios in the six dogs with rapid plasma separation and the averaged data. Equilibrium is reached after about 40 min, at which time the mean ratio was found to be 0.76. As expected, relative plasma concentration is highest early on as the tracer is injected into the plasma (and not red blood cells). ^{201}Tl is rapidly cleared from the plasma causing a rapid decline in relative plasma concentration and “undershoot” before equilibrium is established. Samples left for a prolonged period before plasma separation showed the value of approximately 0.78, which was close to the plasma to whole blood concentrations ratio at the equilibrium shown in

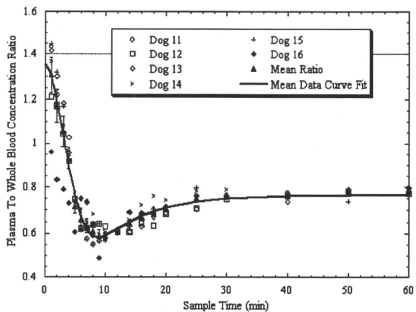


Fig. 3 Individual and mean plasma to whole blood concentration ratios over time for the six dogs with rapid plasma separation. Error bars indicate the standard error of the mean. Solid line is the curve fit to mean ratio data

Fig. 4 A typical example of sequential SPECT images of the myocardium for six representative slices after intravenous injection of ^{201}Tl into a canine at rest

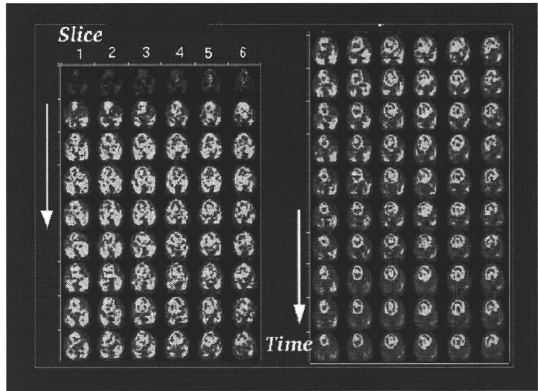


Fig. 3. The plasma to whole blood ratio curves could be approximated by the following equation:

$$R_{pl/wb} = A_0 e^{-\lambda(t+\Delta t)} + A_1 (1 - e^{-\lambda_2(t+\Delta t)}), \quad (3)$$

which resulted in $A_0 = 1.303 \pm 0.045$, $A_1 = 0.7649 \pm 0.0056$, $\lambda_1 = 0.03636 \pm 0.0039 \text{ min}^{-1}$, $\lambda_2 = 0.1263 \pm 0.0077 \text{ min}^{-1}$ and $\Delta t = 0.9516 \pm 0.41 \text{ min}$. The correlation coefficient for the fit was $r = 0.995$.

Figure 4 shows a typical example of sequential images after the intravenous injection of ^{201}Tl for six representative slices of a dog studied at rest. It can be seen that ^{201}Tl appeared in the ventricular chambers first and then gradually accumulated homogeneously into the left myocardium. The quality of these images is reasonably good, indicating that our approach of estimating the kinetic parameters by NLLSF is feasible without excessive noise

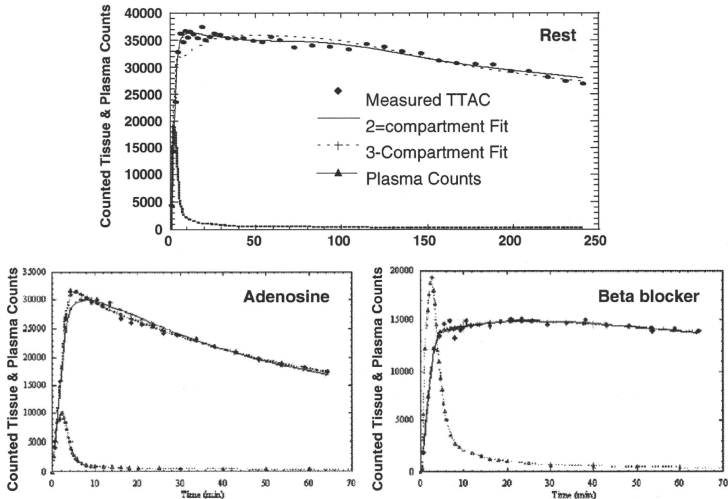


Fig. 5 TTACs and two- and three-compartment model fits for a resting, adenosine (increased MBF) and beta-blocker (reduced MBF) study. Note the different time scales for the resting study because

resting studies were collected for 4 h compared to $\approx 1 \text{ h}$ for the pharmacological intervention studies

amplification. Curve fits to representative TTACs for resting, beta-blocker and adenosine infusion studies are shown in Fig. 5. The height of the TTACs relative to the input function corresponded well with the pharmacological challenges. Compared to the resting studies, peaks of TTACs relative to the arterial input function were higher for adenosine and lower after beta-blocker administration. Results of kinetic fitting by the two- and three-compartment models are also plotted on this figure. Visually, the three-compartment model provided better fits than the two-compartment model to the observed TTACs, which is particularly evident for the initial scan period of the resting and adenosine studies.

Shown in Fig. 6a–e is the comparisons of K_1 obtained by NLLSF (three-compartment model fit) with the microsphere

flow estimates. Values were averaged over the myocardial segments in both axes, thus each point corresponds to a single study. There was good correlation between K_1 and the microsphere flow when no corrections were applied, but K_1 significantly under-estimated the true flow (Fig. 6a). All the corrections improved the K_1 estimates (Fig. 6b–d) and the best agreement between K_1 and microsphere flow was observed when all three factors were corrected as described in Eq. 1 (Fig. 6e). Results of the regression analysis also demonstrated the highest correlation coefficient when all three correction factors were applied. Table 1 summarises the results of the Akaike information criteria (AIC) and Schwartz criteria (SC) obtained from the kinetic fitting analysis for all myocardial segments of all subjects. Both

Fig. 6 Plot of K_1 derived from the three-compartment model fit against the mean of the pre- and post-dynamic SPECT microsphere blood flow measurements. **a** No correction for PVE, limited first-pass EF or conversion from plasma to blood flow has been applied. **b** Correction for PVE has been applied, but not for Hct or limited first-pass EF. **c** Corrections for PVE and Hct have been applied, but not for limited first-pass EF. **d** Corrections for PVE and limited first-pass EF have been applied, but not for Hct. **e** All corrections are applied for PVE, limited first-pass EF and Hct

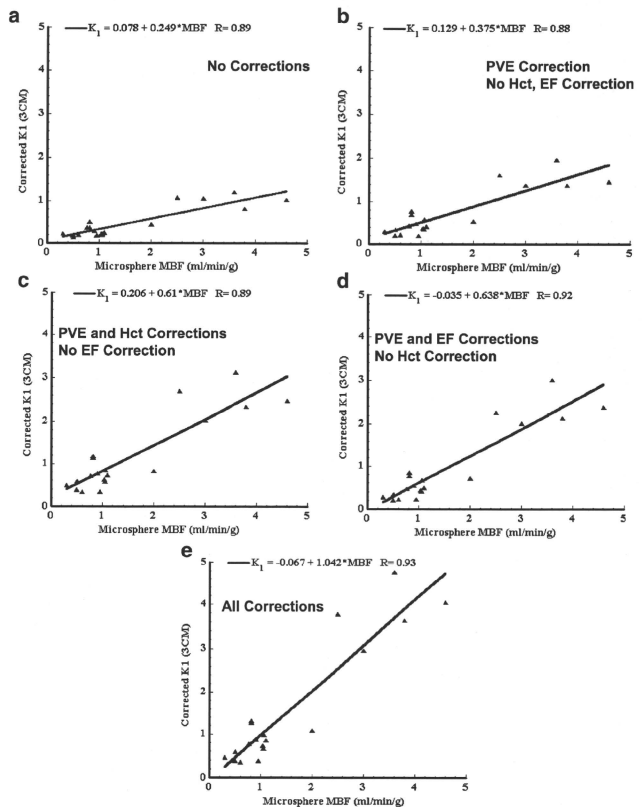


Table 1 Summary of improvement in fit with the three-compartment model over the two-compartment model

Study group	Number of curves	Mean AIC two-compartment	Mean AIC three-compartment	Mean SC two-compartment	Mean SC three-compartment	Number of curves (%) (three-compartment better than two-compartment) ^a
Resting	35	652.4	630.2 ($p < 0.01$)	663.8	638.4 ($p < 0.01$)	24 (69)
Beta-blocker	20	378.4	378.8 ($p = n.s.$)	382.0 ($p < 0.01$)	384.7	3 (15)
Adenosine	45	405.1	393.6 ($p < 0.01$)	408.7	399.5 ($p < 0.01$)	28 (62)

The p value indicates that the value in the cell is significantly lower than the corresponding other value.

AIC: Akaike information criterion, SC: Schwarz criterion

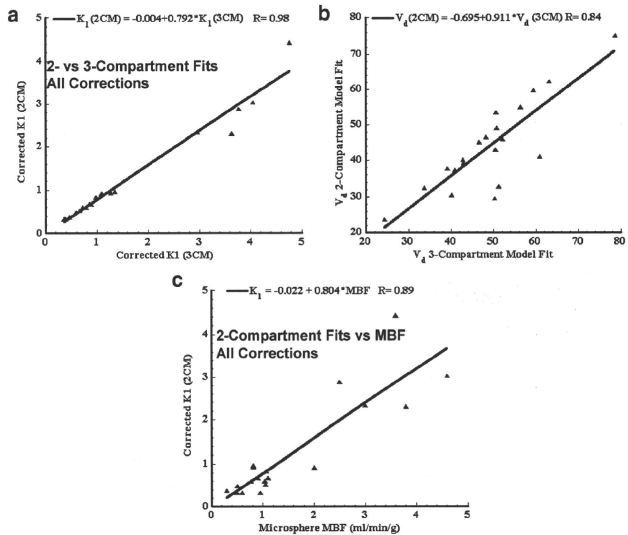
^aThis column gives the number of TTAC fits where the three-compartment model fit provided a significant improvement over the two-compartment fit according to all criteria (AIC, SC).

AIC and SC demonstrated that the three-compartment model fit provided significant improvement over the two-compartment model fit for resting and adenosine studies. For the beta-blocker studies, AIC between the two model fits was not significantly different, whilst SC demonstrated significantly better fit with the two-compartment model. Improved AIC and SC for the three-compartment model fit were observed in 69% of resting TTACs and 62% of adenosine TTACs, but only 15% in beta-blocker TTACs.

As shown in Fig. 7a and b, the K_1 and V_d values derived from the two-compartment model fit showed significant differences compared with those by the three-compartment model. Both K_1 and V_d were under-estimated with the two-compartment model fit compared with the three-compartment

model fit. It should, however, be noted that there was a good correlation between the two- and three-compartment models for K_1 , thus the bias introduced by the two-compartment model fit can potentially be corrected. K_1 values by the three-compartment model fit with all three corrections were 0.86 ± 0.36 , 2.71 ± 1.64 and 0.55 ± 0.24 ml/min/g corresponding to rest, adenosine infusion (with constant infusion at 140–700 mg/kg/h) and beta-blocker (with 2–6 mg administration), respectively. Difference in V_d was less than 10% and again this bias can potentially be corrected by the regression equation. The K_1 obtained with the two-compartment model also demonstrated a good correlation with the microsphere flow (Fig. 7c), though there was again a systematic under-estimation in K_1 .

Fig. 7 a Plot of K_1 estimates derived from the two-compartment model fit against those from the three-compartment model fit. b Plot of V_d estimates derived from the two-compartment model fit against those from the three-compartment model fit. c Plot of K_1 values derived from the two-compartment model fit against mean of the pre- and post-dynamic SPECT microsphere blood flow measurements



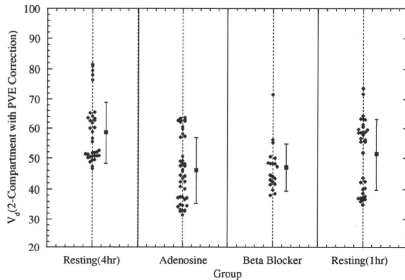


Fig. 8 V_d values obtained from the two-compartment model fit to the full 4 h resting data, adenosine and beta-blocker infusion 1 h curves and fit to first 1 h only of the resting study curves. Data from the multiple individual myocardial regions are shown

Figure 8 plots the V_d values for all evaluated myocardial segments for the fit to 4 h resting data, adenosine and beta-blocker infusion 1 h data and fit to only the first 1 h of resting data. The 4-h resting V_d values are significantly higher ($p < 0.01$) compared with the adenosine, beta-blocker values and compared with the fit to the first 1 h of resting data. However, the 1-h resting values are not significantly different from the beta-blocker V_d values nor the adenosine values.

Discussion

This study demonstrates that the kinetic analysis of quantitatively assessed myocardial ^{201}Tl accumulation (build-up and washout in healthy canines) provided quantitative MBF values, which agreed well with flows obtained using microspheres for a wide physiological range of flows. The size of the TTACs relative to the arterial plasma concentration corresponded well to the pharmacological stresses induced by adenosine and beta-blocker challenges. The compartmental model approach could reproduce these TTACs to make the determination of kinetic parameters, such as K_1 and V_d , possible. The three-compartment model gave results which were generally higher than the two-compartment model and which were statistically significantly better in terms of AIC, SC for the resting and adenosine studies, and this was in line with the visual inspection of the TTAC model fit curves. It should, however, be noted that the differences were only small between the two- and three-compartment model approaches, approximately 20% for K_1 and 10% for V_d . The bias associated with the two-compartment model could be corrected by a linear regression as shown in Fig. 7a–c. This opens the possibility of using the more reliable two-compartment model fit due to its reduced number of parameters for routine clinical studies. The improved reliability of the two-compartment model fit in

the clinical setting is particularly important if one intends to shorten the study time or generate parametric images.

The three corrections for PVE, Hct and first-pass EF proved to be important. The PVE correction method used in this work cannot, however, be applied to clinical studies, and the PVE correction in the beating heart still remains a considerable challenge in clinical studies. PVE may be reduced by gating the data, which may not, however, be feasible for the already noisy and large dynamic SPECT data sets. PVE may also be reduced by including resolution recovery as part of the reconstruction process [17–20]. Alternatively, it may also be possible to include PVE as part of the kinetic model fitting [21–25]. However, this adds extra fitting parameters and requires some parameters to be assumed fixed.

The input function is an important component in compartment model fitting. In this study, rapid arterial blood sampling was performed, and the plasma was separated by centrifugation. A number of important insights were gained by performing rapid separation of plasma in a subset of samples and dogs. It was found that ^{201}Tl enters the red blood cells as observed from the rapid separation of plasma in a subset of samples and dogs, which is not un-expected as potassium is also known [22] to be taken up by the red blood cells. The exchange of ^{201}Tl between red blood cells and plasma is relatively slow compared to the passage of blood through the capillary bed and hence direct uptake of activity from the red blood cells into tissue is believed to be negligible. Hence, tissue uptake will be dominated by the activity in the plasma during passage through the capillary bed and plasma in the substrate being measured. As a consequence, the flow measurement obtained with ^{201}Tl is plasma flow, which is in contrast to the microsphere studies, which measure whole blood flow. Conversion of plasma to blood flow was achieved by dividing the plasma flow by $(1 - \text{Hct})$, as shown in Eq. 1, which then allowed the direct comparison with the microsphere measurements.

Rigorous estimation of the input function requires frequent arterial blood sampling. This is not only considered invasive, but also labor intensive. In addition, it has been shown in this study that rapid separation of the plasma for at least the first 30–40 min post- ^{201}Tl administration is required to obtain accurate plasma concentration. If the separation of plasma is delayed, then the true plasma concentration at the time of sampling cannot be measured, which results in biased K_1 estimates. An empirical relationship of plasma to whole blood ratio as a function of time was developed and was found to be sufficiently consistent between dogs (Fig. 3) to allow the mean curve to be applied with minimal bias. Thus, in clinical practice, whole blood samples may be counted and converted to plasma concentration using the empirical relationship. This also potentially allows the input function to be obtained

non-invasively from the SPECT data using, for example, a curve derived from a left ventricular region. However, it should be noted that the relationship between plasma and whole blood counts in this study was derived for a 4-min infusion protocol and may be different for other injection protocols, such as bolus injection. Previously, it has been shown that population-based input functions calibrated with one or two blood samples could avoid the need for frequent arterial blood samples [26–28]. There is also a potential for applying this approach to ^{201}Tl studies. This is beyond the scope of this study and a systematic study should be designed to confirm this in clinical settings.

^{201}Tl has a high trans-capillary EF and thus the initial regional uptake of this tracer predominantly reflects the regional blood flow [10]. Use of a tracer that has a high first-pass EF is essential when one intends to quantitatively assess MBF at a high flow range or the coronary flow reserve. The EF of ^{201}Tl is reported as >0.8 [10] for a wide flow range and is known to be higher than $^{99\text{m}}\text{Tc}$ -labelled tracers such as tetrofosmine and sestamibi [29]. The physical characteristics of ^{201}Tl are unfortunately not ideal as low energy emission increases the attenuation factor and the scatter in the image. In addition, the relatively long half-life limits the administered activity to about a tenth of that with $^{99\text{m}}\text{Tc}$ tracers. Despite these shortcomings, the physiological characteristics of having high first-pass EF make ^{201}Tl an interesting tracer particularly for the absolute quantitation of MBF and the coronary flow reserve. This study demonstrates that quantitative physiological parameters can be derived from dynamic ^{201}Tl SPECT studies, despite its less than ideal imaging characteristics.

Whilst the quantitative physiological parameter estimation removed the systematic bias between MBF estimated by ^{201}Tl dynamic SPECT and by microspheres, the spread of data points around the regression line was rather large (Figs. 6e and 7c). This is not only due to possible errors in the estimation of MBF from the ^{201}Tl , but there was also considerable variation in flow estimated by the microspheres at the beginning and end of the study. Thus, at least part of the variability is attributable to errors in microsphere flow measurement, and particularly for the pharmaceutical intervention studies, flow may not have remained constant throughout the entire study duration, which may also account for some of the differences seen between the various flow measurements.

V_d estimated in this study could serve as an index of viability, as viable myocytes are required to maintain the large concentration gradient between plasma and myocardium at equilibrium. There was no significant difference in V_d values between rest, beta-blocker and adenosine studies when fitted for 1 h (Fig. 8). The significant difference between the 1- and 4-h fit for resting data could be explained by the limitation of the two-compartment model.

Considerable spread in the V_d values observed over all dog studies on the other hand was partially attributed to the short (insufficient) scan time for reliable estimates of V_d . With the exception of the large, outlying V_d values in all 5 regions of 1 dog, the resting V_d values fell within a relatively narrow range of 47 to 65 (mean \pm SD=55 \pm 6). Given the sufficiently long scan time, significant reduction in V_d in infarcted areas may be detected. However, this would need to be tested with a suitable study design.

The scan time of 4 h required to achieve reliable V_d estimates is not practical in the routine clinical setting. As has been shown by Lau et al. [30], the scan period may be split into two sessions, an early dynamic scan for 30 min followed by a single static scan at approximately 3 h. This scheme is not more onerous than current rest/re-distribution protocols and hence could be practical. In addition, it may be possible to simplify the scanning protocol further to two static scans by using the table look-up method for the two-compartment model, which has been successfully employed for other SPECT tracers with relatively slow kinetics similar to ^{201}Tl [27, 31, 32]. This warrants further investigation.

This study relies on established, rigorous attenuation and scatter correction in SPECT [5] and availability of multi-detector SPECT systems capable of performing dynamic acquisition. To our knowledge, this is the first report that has demonstrated that it is possible to obtain quantitative physiological parameter estimates of K_1 and V_d in the myocardium using a clinical SPECT scanner and ^{201}Tl . This work suggests that it is feasible to apply our technique to clinical studies. Further studies are, however, needed to validate the proposed approach in the clinical setting. Incomplete motion correction is one possible error source, particularly in patients. Dynamic SPECT is probably more sensitive to the possible movement of patients during the study. Shortened clinical protocol is preferred, but this requires additional development to improve the reliability of parameter estimates. In addition, two scanning sessions are needed to assess the coronary flow reserve. We have recently demonstrated a technique to assess two cerebral blood flow images, one at rest and another after a vasodilating drug, from a single session of a SPECT scan in conjunction with split dose administration of ^{123}I -iodoamphetamine and dynamic SPECT [7]. As a clinical implication, the quantitative assessment of MBF and coronary flow reserve is important. For instance, coronary micro-vascular dysfunction or impaired endothelial function in patients with coronary risk factors or patients with cardiomyopathy or with heart failure is an un-resolved important issue to answer [11]. Coronary flow reserve can be reduced in patients with hypercholesterolemia without overt coronary stenosis [12]. A systematic study should be carried out to validate this approach for assessing MBF at rest and after adenosine from a single session of a scan.

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NOTE

Acceleration of Monte Carlo-based scatter compensation for cardiac SPECT

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Abstract

Single proton emission computed tomography (SPECT) images are degraded by photon scatter making scatter compensation essential for accurate reconstruction. Reconstruction-based scatter compensation with Monte Carlo (MC) modelling of scatter shows promise for accurate scatter correction, but it is normally hampered by long computation times. The aim of this work was to accelerate the MC-based scatter compensation using coarse grid and intermittent scatter modelling. The acceleration methods were compared to un-accelerated implementation using MC-simulated projection data of the mathematical cardiac torso (MCAT) phantom modelling ^{99m}Tc uptake and clinical myocardial perfusion studies. The results showed that when combined the acceleration methods reduced the reconstruction time for 10 ordered subset expectation maximization (OS-EM) iterations from 56 to 11 min without a significant reduction in image quality indicating that the coarse grid and intermittent scatter modelling are suitable for MC-based scatter compensation in cardiac SPECT.

1. Introduction

The quality of single proton emission computed tomography (SPECT) is degraded by attenuation, collimator blurring and scatter. Whereas attenuation and collimator blurring can nowadays be corrected in clinically acceptable times, accurate and efficient scatter correction has been proven to be a more difficult problem. This is mainly due to the fact that the shape of the scatter point-spread function depends on the location inside the object and can be very difficult to parametrize.

One very promising recently presented scatter compensation method is the Monte Carlo (MC)-based scatter correction by Beekman *et al* (2002). In this method, a MC simulator is used as a forward-projector for scatter in the ordered subset expectation maximization

(OS-EM) algorithm (Hudson and Larkin 1994). MC-based scatter modelling is expected to be especially advantageous in areas where the attenuating media is highly non-uniform such as the thorax, because it can faithfully produce the complex shape of the scatter response function. The MC-based scatter compensation has been shown to outperform the common triple-energy window scatter compensation method in terms of contrast and myocardial lesion detectability (Xiao *et al* 2006). Unfortunately, despite the considerable advances made in MC-based scatter compensation its widespread utilization in clinical practice can still be limited by long computation times.

Kadmas *et al* (1998) have presented two simple and effective acceleration methods for reconstruction-based scatter compensation. The first of these approaches is the coarse grid scatter modelling method, which calculates the scatter contribution using sparser grid than is used in the actual reconstruction. The second approach is the intermittent scatter modelling method, where scatter is modelled only during the first couple of OS-EM iterations and then held as a constant additive factor for the later iterations. Both of these approaches are based on the assumption that the scatter response contains mainly low-frequency components and can thus be modelled with a relatively large voxel size and not during the last OS-EM iterations. Kadmas *et al* presented their acceleration methods for the effective scatter source estimation method (Frey and Tsui 1996) and showed that the performance of the accelerated scatter compensation was very similar to the performance of the standard (un-accelerated) scatter correction.

The goal of this work was to implement the coarse grid and intermittent scatter modelling methods for MC-based scatter compensation and to evaluate their performance in case of ^{99m}Tc cardiac SPECT.

2. Materials and methods

2.1. Implementation of the MC simulator

MC simulations are based on sampling radioactive decays within a source volume and following their interactions inside the attenuating media. Our MC simulator uses four different maps in the simulations: emission map for setting photons to be tracked, density map for sampling interactions and primary and scatter maps for storing photon weights. The history of each photon (primary or scatter) in our simple MC simulator is tracked shortly as follows:

- (1) If the simulated photon is a primary the primary map is updated. Otherwise direction cosines are sampled and random walk is started for the scatter photon.
- (2) Photon interaction points are calculated using the delta scattering algorithm (Woodcock *et al* 1965), which does not require ray tracing through the attenuating media. The basic principle of delta scattering is that the path-length P between two 'fictitious' interaction points is sampled as $P = -\ln(R)/\mu_{\max}$, where R is a uniform random number and μ_{\max} is the largest attenuation coefficient in the attenuating media (Ljungberg *et al* 2005). At the end of the sampled path a possible interaction is sampled according to established methods or tracking is continued by sampling a new distance P according to the above-mentioned equation.
- (3) At each real interaction point photon weight is multiplied by the probability that no photoelectric effect occurred and a copy of the original photon is forced to Compton scatter towards the detector and scatter map is updated according to the convolution-based forced detection approach (de Jong *et al* 2001).

- (4) If predetermined number of scattering events has not been reached, the history of the original photon is continued by Compton scatter. New direction cosines are sampled from the Klein–Nishina function and tracking is continued from (2).

After all the photons have been simulated the primary and scatter maps are multiplied by probability that photon emitted from a certain voxel reaches the detector without interaction. Then these maps are convolved with depth-dependent detector response function, which is assumed to be Gaussian. The blurred primary and scatter maps are finally forward-projected to primary and scatter projections.

2.2. Implementation of the reconstruction algorithm with MC-based scatter compensation

The developed MC simulator was included into an OS-EM reconstruction algorithm as forward-projector for the scattered counts as originally proposed by Beekman *et al* (2002). The OS-EM is given by

$$f_j^{\text{new}} = \frac{f_j^{\text{old}}}{\sum_{i \in S_n} a_{ij}} \sum_{i \in S_n} a_{ij} \frac{p_i}{\sum_k a_{ik} f_k^{\text{old}} + s_i}, \quad (1)$$

where f is the reconstructed image, p is the measured projections, j (or k) is the reconstruction voxel index, i is the projection pixel index, a_{ij} is the probability that emission from voxel j is detected in pixel i , s is the MC-based scatter projections and S_n is the n th subset. The image update in OS-EM consists of sequential forward- and back-projection operations. The estimated projections are obtained by forward-projecting the current image estimate ($\sum_k a_{ik} f_k^{\text{old}}$), and correction terms that are used to update the old image are formed by back-projecting the ratio of the measured and estimated + scatter projections ($\sum_{i \in S_n} a_{ij} \frac{p_i}{\sum_k a_{ik} f_k^{\text{old}} + s_i}$).

In this work the forward- and back-projectors were implemented as rotation based (Di Bella *et al* 1996). The back-projector included attenuation and detector response compensation and the forward-projector attenuation, detector response and MC-based scatter compensation. Attenuation correction factors for each voxel were calculated simply by summing the rotated attenuation map along columns. Detector response, on the other hand, was modelled by convolving each plane of the reconstruction matrix parallel to the projection plane with collimator response kernel, which was assumed to be Gaussian. The scatter projection for each projection angle was obtained by MC-based forward-projection of the current image estimate.

2.3. Acceleration of MC-based scatter compensation

The coarse grid scatter modelling method was implemented by simply down-sampling the current image estimate (and the attenuation map) into a sparser matrix before the MC-based forward-projection and linearly interpolating the scatter projections back to the original size after forward-projection was finished. In the intermittent acceleration approach the MC-based scatter modelling is performed only in the few early OS-EM iterations and after that the scatter projections are kept fixed for remaining iterations.

2.4. Performance tests

Performance tests were performed using a female version of the mathematical cardiac torso (MCAT) phantom (Tsui *et al* 1994), and the camera parameters are shown in table 1. The MCAT phantom modelled normal human tissue densities in the thorax region and the source distribution presented the $^{99\text{m}}\text{Tc}$ uptake with the following relative activities per

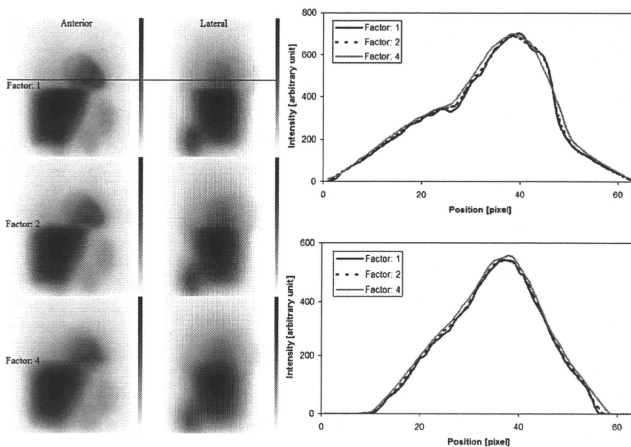


Figure 1. Comparison of scatter projections and profiles obtained with different down-sampling factors (factor 1 is scatter projection without down-sampling). Upper profile is obtained from the anterior projection and lower from the lateral projection. Locations of the profiles are shown as horizontal lines.

Table 1. Camera parameters for Monte Carlo simulations.

Low-energy high-resolution collimator (4.0 cm hole length, 0.178 cm hole diameter)
0.4 cm intrinsic detector resolution
9.0% energy resolution
15.0% energy window centred on 140 keV
23.0 cm radius of rotation

voxel: myocardium 100, liver 50, kidney 100, spleen 80, lung 5 and rest of the body 2.5. Two lesions (anterior and inferior) with the relative uptake of 2.5 were included in the left myocardium.

The effects of different down-sampling factors in the coarse grid scatter modelling scheme were studied by performing MC simulations using the MCAT phantom. The pixel size was set to 0.625 cm (64×64 projection and $64 \times 64 \times 64$ image matrix size). Down-sampling factors of 2 ($64 \times 64 \times 64$ image matrix down-sampled to $32 \times 32 \times 32$ matrix) and 4 ($64 \times 64 \times 64$ image matrix down-sampled to $16 \times 16 \times 16$ matrix size) were investigated. Figure 1 presents anterior and lateral scatter projection images of the MCAT phantom. According to this figure factor 4 produces slightly distorted scatter projections, and thus the down-sampling factor was set to 2 for all the reconstructions accelerated with the coarse grid scatter modelling.

The effect of the number of scatter update iterations was studied by using MCAT projection data simulated using the SIMIND MC simulator (Ljungberg and Strand 1989). The camera parameters for this simulation are again shown in table 1. The number of simulated noise-free

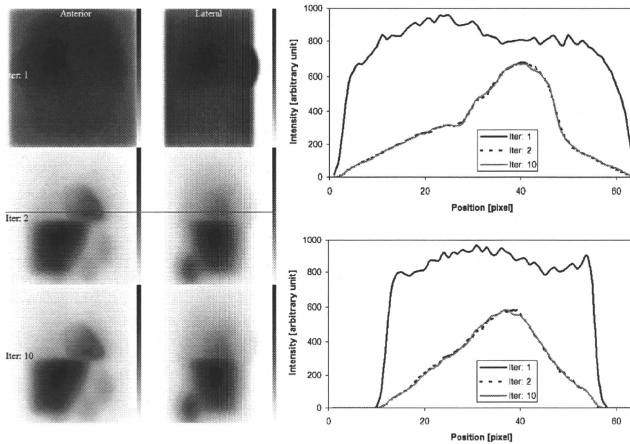


Figure 2. Comparison of different number of scatter update iterations. Upper profile is obtained from the anterior projection and lower from the lateral projection. Locations of the profiles are shown as horizontal lines.

projections was 64 on a 360° orbit and pixel size was 0.3125 cm (128×128 projection and $128 \times 128 \times 128$ image matrix size). The projections were collapsed to a 64×64 matrix size before the reconstruction. Reconstruction using the noise-free projection data was performed with 16 subsets and 1–10 iterations, and the MC forward-projected scatter projections were saved after every iteration. Figure 2 presents scatter projection images and profiles for different iteration numbers, and these images show that the scatter projections do not change markedly after two iterations. Therefore, in the following performance tests only two scatter iterations were used when intermittent scatter modelling was applied.

The acceleration methods were tested using the same MCAT projection data that were used to find the scatter iterations stopping point. Poisson noise was added to the noise-free projections by setting number of total counts to 5 million. The projections were reconstructed without scatter compensation, with the MC-based scatter compensation but without acceleration, with coarse grid scatter modelling (down-sampling factor of 2), with intermittent scatter modelling (scatter updated during the first two iterations) and with both accelerations applied at the same time. Number of MC simulated photons per projection angle was set to 1.0 million and 16 subsets with 10 iterations were used. After reconstruction images were filtered with a 3D Butterworth filter (order: 5, cutoff: 1.0 cm^{-1}), transverse slices were zoomed and reoriented into short-axis slices. Regions of interest (ROI) were drawn on the normal myocardium, ventricle and on the anterior/inferior defect areas. Normal myocardium to ventricle and normal myocardium to lesion contrasts were calculated.

In addition to Monte Carlo simulations the acceleration methods were also tested using a clinical $^{99\text{m}}\text{Tc}$ myocardial stress/rest perfusion study acquired with the Siemens Symbia SPECT/CT scanner. Low-energy high-resolution parallel hole collimators were used, and

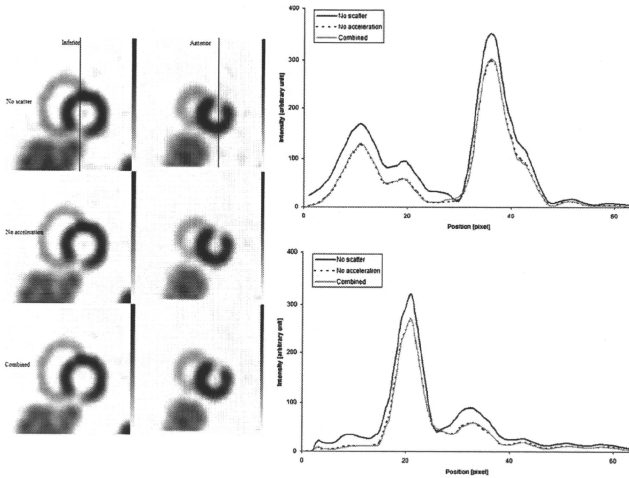


Figure 3. Short-axis slices and profiles through the heart of the MCAT phantom for reconstruction without scatter compensation (no scatter) and with MC-based scatter compensation without acceleration (no acceleration) and with coarse grid + intermittent scatter modelling (combined). Results for other acceleration methods are not shown, because they overlap the coarse grid + intermittent scatter compensation profiles. Upper profile is obtained from the image with inferior lesion and lower with anterior lesion. Locations of the profiles are shown as vertical lines.

Table 2. Lesion-to-normal myocardium and ventricle-to-normal myocardium contrasts for different MC-based scatter compensation acceleration schemes.

Scatter compensation method	Acceleration method	Contrast		
		Anterior	Inferior	Ventricle
No compensation		0.57	0.51	0.80
MC-based	No acceleration	0.64	0.58	0.91
	Coarse grid	0.65	0.59	0.93
	Intermittent	0.62	0.60	0.93
	Coarse grid+intermittent	0.64	0.59	0.93

60 projection angles over a 180° arc were acquired using a symmetric 15% energy window centred at 140 keV.

3. Results

Figure 3 shows short-axis slices and profiles through the left ventricle of the MCAT phantom for reconstruction without scatter compensation and with the MC-based scatter compensation without acceleration and with coarse grid + intermittent scatter modelling. Profiles for the

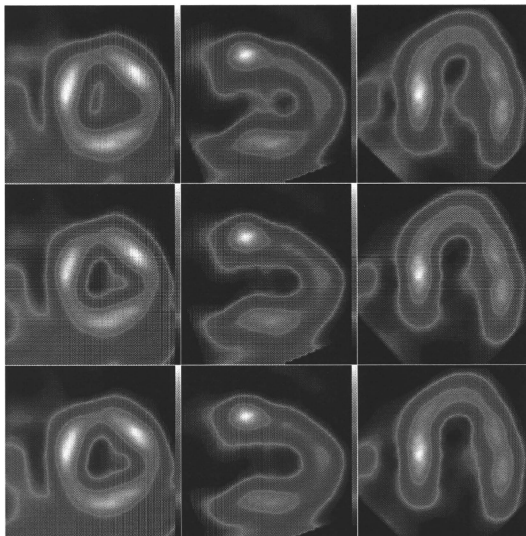


Figure 4. Example short-axis (first column), vertical long-axis (second column) and horizontal long-axis (third column) slices from a clinical rest myocardial perfusion study for reconstruction without scatter compensation (first row), with MC-based scatter compensation without acceleration (second row) and with coarse grid +intermittent scatter modelling (third row).

un-accelerated and accelerated scatter compensation practically overlap indicating similar performance. This same conclusion can also be drawn from table 2, which presents the lesion and ventricle contrasts. All the scatter compensation methods offer very similar contrast values and clearly improve contrast when compared to reconstruction without scatter compensation.

Figure 4 presents results for the resting clinical study (for the stress study the findings in image quality were identical to the resting study). It can be seen that the images reconstructed with and without MC-based scatter compensation acceleration are nearly identical. Table 3 presents the approximate reconstruction times for different scatter compensation methods.

4. Discussion

The aim of this study was to accelerate the MC-based scatter compensation using the coarse grid and intermittent scatter modelling methods. Both of these acceleration methods proved to provide clear improvement in execution times without any marked degradation in image quality when compared to un-accelerated version of MC-based scatter-compensation (tables 2 and 3, figures 3 and 4). The greatest improvement in terms of speed was obtained by combining the two acceleration methods.

Table 3. Approximate reconstruction times (10 iterations, 16 subsets and 1 million photons per projection) for different MC-based scatter compensation acceleration schemes. Reconstructions were performed using 2.33 GHz Pentium processor with 8 GB RAM.

Scatter compensation method	Acceleration method	Time (min)
No compensation		0.8
MC-based	No acceleration	55.6
	Coarse grid	53.4
	Intermittent	11.6
	Coarse grid + intermittent	11.2

The comparison studies for the scatter compensation methods were performed with $64 \times 64 \times 64$ matrix sizes and coarse-grid down-sampling factor of 2 was used, because larger down-sampling factors could not fully preserve the details of the scatter projections (figure 1). With larger matrix sizes than $64 \times 64 \times 64$ larger down-sampling factors could probably be used for higher acceleration. This was not studied in this work, because in the case of cardiac SPECT $64 \times 64 \times 64$ matrix size is usually considered the standard.

The acceleration achieved with the coarse grid scatter modelling depends also on the implementation of the MC simulator that is used in the forward-projection. As mentioned, our MC simulator is based on the delta scattering technique (Woodcock *et al* 1965), which allows calculation of the photon interaction points without time-consuming ray tracing. Therefore, the coarse-grid scatter modelling provided acceleration mainly because the collimator response and attenuation modelling were performed with the sparser matrix. In the case of 1 million simulated photons per projection the time needed to model the collimator response and attenuation is, however, only a small fraction of the time that is spent in sampling the various probability distributions in the MC calculations. Thus the speed-up provided by the coarse-grid scatter modelling is quite small, but can be much greater if a smaller number of simulated photons are used. In addition, if a ray-tracing-based MC algorithm is used instead of delta scattering, coarse-grid scatter modelling will probably increase the speed much more, because the efficiency of ray tracing depends heavily on the matrix size.

In this study the scatter projections were not noticed to change markedly after two iterations, and thus in the intermittent scatter modelling scheme scatter projections were updated only during the first two OS-EM iterations. The number of scatter iterations needed may, however, depend on the imaging situation as mentioned by Kadrmaz *et al* (1998) and should therefore be checked before using intermittent scatter modelling as an acceleration method.

One important thing that was not considered in detail in this work is the number of simulated photons per projection. We chose the 1 million photons per projection according to our preliminary studies, where we compared the quality of images reconstructed using different number of simulated photons and noticed that using over 1 million photons per projection does not lead to a significant increase in image quality. The performance of MC-based scatter compensation is, however, quite complicated because it is not only influenced by the number of simulated photons but also by the interplay of noise in attenuation map/projection data and the number of simulated photons. Therefore, a detailed study of noise effects, e.g. similar to the one presented by de Wit *et al* (2005), is probably needed in the future. If different number of photons per projection are to be used the speed-up factors might not be the same as in table 2.

Faster reconstruction times with MC-based scatter compensation than those shown in table 3 have been presented in the literature (e.g. de Wit *et al* (2005) and Xiao *et al* (2006)). This

speed difference is probably mainly related to algorithm implementation. Our reconstruction algorithm is not yet fully optimized, and we believe that we can greatly reduce the execution time of our reconstruction. In addition to direct code optimization, we are also planning to parallelize our code for multi-core processors. The acceleration methods presented in this work should be very suitable for different parallelization schemes, because they do not affect the general structure of the OS-EM reconstruction algorithm.

5. Conclusions

We conclude that both the coarse grid and the intermittent scatter modelling methods are suitable for accelerating MC-based scatter compensation, and with these methods MC-based scatter compensation is a promising alternative for clinical cardiac SPECT.

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Comparison of Gd-DTPA-Induced Signal Enhancements in Rat Brain C6 Glioma among Different Pulse Sequences in 3-Tesla Magnetic Resonance Imaging

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Sato H, Enmi J, Teramoto N, Hayashi T, Yamamoto A, Tsuji T, Naito H, Iida H. Comparison of gd-dtpa-induced signal enhancements in rat brain c6 glioma among different pulse sequences in 3-tesla magnetic resonance imaging. *Acta Radiol* 2007;000:1–8.

Background: T1-shortening contrast media are routinely used in magnetic resonance (MR) examinations for the diagnosis of brain tumors. Although some studies show a benefit of 3 Tesla (T) compared to 1.5T in delineation of brain tumors using contrast media, it is unclear which pulse sequences are optimal.

Purpose: To compare gadopentetate dimeglumine (Gd-DTPA)-induced signal enhancements in rat brain C6 glioma in the thalamus region among different pulse sequences in 3T MR imaging.

Material and Methods: Five rats with a surgically implanted C6 glioma in their thalamus were examined. T1-weighted brain images of the five rats were acquired before and after Gd-DTPA administration (0.1 mmol/kg) using three clinically available pulse sequences (spin echo [SE], fast SE [FSE], fast spoiled gradient echo [FSPGR]) at 3T. Signal enhancement in the glioma (E_T) was calculated as the signal intensity after Gd-DTPA administration scaled by that before administration. Pulse sequences were compared using the Tukey-Kramer test.

Results: E_T was 1.12 ± 0.05 for FSE, 1.26 ± 0.11 for FSPGR, and 1.20 ± 0.11 for SE. FSPGR showed significantly higher signal enhancement than FSE and comparable enhancement to SE.

Conclusion: FSPGR is superior to FSE and comparable to SE in its ability to delineate rat brain C6 glioma in the thalamus region.

Key words: Brain; contrast agents; MR imaging

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T1-shortening contrast media are routinely used in magnetic resonance (MR) examinations for the diagnosis of brain tumors. Some studies show a benefit of 3 Tesla (T) compared to 1.5T in delineation of brain tumors using contrast media (1–5). However, it is unclear which pulse sequences are optimal. The conventional spin-echo (SE) technique has been most frequently used for T1-weighted (T1W) imaging of tumors after contrast media administration. The gradient-echo (GRE) technique, which is faster than SE, was introduced initially at 1.5T or lower field strength (6–11). Some

studies have reported that GRE techniques compare favorably with the SE technique for delineation of brain tumors (8–10), while other studies have reported that GRE techniques do not show contrast enhancement as well as SE (6, 7, 11). At 3T, as at 1.5T or lower field strength, the issue of whether GRE techniques are effective compared to SE has not been determined. In 16 patients, NÖBAUER-HUMANN et al. reported that 3D GRE with magnetization preparation (MPRAGE) was comparable to T1W SE in tumor-to-brain contrast at 3T, although the parameters of T1W SE were not

0 optimized for 3T (1). In 12 patients, FISCHBACH et al. compared four T1W sequences: SE, inversion
recovery fast SE (IR-FSE), 2D GRE, and
5 MPRAGE at 3T. They observed that SE and IR-FSE provided higher contrast enhancement of brain
tumors than 2D GRE and MPRAGE. Furthermore, their impressions showed that the
visual quality of SE was superior to that of the other
10 three sequences (12).

In order to compare pulse sequences, it would be preferable if the pathological and physiological conditions of subjects were constant across scans. One possible model system is the widely used rat brain glioma model (4, 5, 13–15). In most studies with small animals, MR imaging systems with small magnets are widely used. The pulse sequences available on the scanner designed for small animals, however, are different from those on a clinical scanner. By using a scanner designed for humans, we can compare diagnostic values of practical clinical pulse sequences. To our knowledge, no studies have been reported comparing pulse sequences on a 3T human scanner using a rat brain glioma model.

The purpose of the current study was to elucidate the optimal pulse sequence that provides the highest obtainable signal enhancement using gadopentetate dimeglumine (Gd-DTPA) in a rat brain C6 glioma model on a 3T human whole-body scanner.

Material and Methods

0 Protocols of all animal procedures were approved by the ethics committee for animal research at the National Cardiovascular Center. Male Sprague-Dawley rats (Japan SLC, Inc., Shizuoka, Japan) were used. Rats had free access to food and water, and were kept in uncrowded conditions (two/cage) in a light-, temperature-, and humidity-regulated room (light on 07.00–19.00, $23 \pm 3^\circ\text{C}$, and $50 \pm 20\%$).

Study design

4 T1 measurements in the brains of three normal rats and phantom studies were performed to identify pulse sequences, among which Gd-DTPA-induced signal enhancements in rat C6 brain gliomas were compared, and to determine pulse sequence parameters. Using the determined pulse sequences and parameters, we examined five rats with developed gliomas out of 20 rats that received C6 glioma implantation in their thalamus region.

MR imaging system

5 All scanning was performed on a 3T whole-body scanner (Signa LX VH3M4; GE Healthcare, Milwaukee, Wisc., USA) equipped with the manufacturer's gradient system (maximum gradient strength 40 mT/m; slew rate 150 mT/m/s).

For imaging rat brains, we built a three-turn solenoid coil with a diameter of 42 mm and a length along the cylindrical axis of 18 mm. The diameter and length of this coil were adjusted to rat head size. The helical pitch of the coil was wide enough to pass the ear bars used to secure the rat's head. The coil was capable of transmission and reception, and was tuned to an impedance of 50Ω at a resonant frequency of 127.76 MHz. Capacitance was divided into six elements in series, which were put at each half turn. The coil was mounted on a fixing apparatus (Narishige Co., Ltd., Tokyo, Japan) using an acrylic jig specially designed for the coil (Fig. 1). Rats were placed prone on the fixing apparatus. Rat heads were secured using an incisor hook and ear bars. All components of the fixing apparatus consisted of non-magnetic materials. During imaging, the fixing apparatus, on which the rat and the coil were mounted, was placed in the gantry so that the cylindrical axis of the coil and the cranial-to-caudal direction of the rat were perpendicular to a static magnetic field, and the center of the rat brain was positioned at the magnet isocenter.

Measurement of T1 in normal rat brain

This measurement was performed to establish the normal T1 value in the transplantation site (thalamus) of the C6 glioma cells. T1 values in the brain of three normal rats (9–13 weeks old, 380 ± 50 g) were

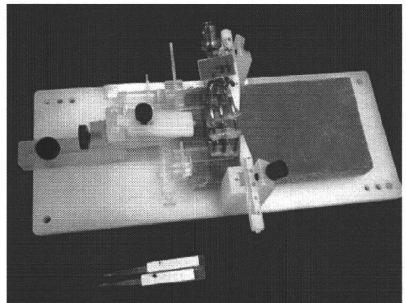


Fig. 1. The three-turn solenoid coil and the fixing apparatus used for the imaging of rats in the present study. The coil was mounted on the fixing apparatus using the specially designed acrylic jig.