

Optimization of temporal sampling for ^{82}Rb rubidium PET myocardial blood flow quantification

Benjamin C. Lee, PhD,^a Jonathan B. Moody, PhD,^a Richard L. Weinberg, MD, PhD,^b James R. Corbett, MD,^{a,c} Edward P. Ficaro, PhD,^{a,c} and Venkatesh L. Murthy, MD, PhD^c

^a INVIA Medical Imaging Solutions, Ann Arbor, MI

^b Division of Cardiovascular Medicine, Department of Internal Medicine, University of Michigan, Ann Arbor, MI

^c Division of Nuclear Medicine, Department of Radiology, University of Michigan, Ann Arbor, MI

Received Nov 15, 2016; accepted Apr 6, 2017

doi:10.1007/s12350-017-0899-7

Background. Suboptimal temporal sampling of left ventricular (LV) blood pool and tissue time-activity curves (TACs) may introduce bias and increased variability in estimates of myocardial blood flow (MBF) and flow reserve (MFR) from dynamic PET myocardial perfusion images. We aimed to optimize temporal sampling for estimation of MBF and MFR.

Methods. Twenty-four normal volunteers and 32 patients underwent dynamic stress/rest rubidium-82 chloride (^{82}Rb) PET imaging. Fine temporal sampling was used to estimate the full width at half maximum (FWHM) of the LV blood pool TAC. Fourier analysis was used to determine the longest sampling interval, T_s , as a function of FWHM, which preserved the information content of the blood phase. Dynamic datasets were reconstructed with frame durations varying from 2 to 20 seconds over the first 2 minutes for the blood phase and 30 to 120 seconds for the tissue phase. The LV blood pool and tissue TACs were sampled using regions of interest (ROI) and fit to a compartment model for quantification of MBF and MFR. The effects of temporal sampling on MBF and MFR were evaluated using clinical data and simulations.

Results. T_s increased linearly with input function FWHM ($R = 0.93$). Increasing the blood phase frame duration from 5 to 15 seconds resulted in MBF and MFR biases of 6-12% and increased variability of 14-24%. Frame durations <5 seconds had biases of less than 5% for both MBF and MFR values. Increasing the tissue phase frame durations from 30 to 120 seconds resulted in <5% biases.

Conclusions. A two-phase framing of dynamic ^{82}Rb PET images with frame durations of 5 seconds (blood phase) and 120 seconds (tissue phase) optimally samples the blood pool TAC for modern 3D PET systems. (J Nucl Cardiol 2017;24:1517-29.)

Electronic supplementary material The online version of this article (doi:10.1007/s12350-017-0899-7) contains supplementary material, which is available to authorized users.

The authors of this article have provided a PowerPoint file, available for download at SpringerLink, which summarizes the contents of the paper and is free for re-use at meetings and presentations. Search for the article DOI on SpringerLink.com.

JNC thanks Erick Alexanderson MD, Carlos Guitar MD, and Diego Vences MD, UNAM, Mexico, for providing the Spanish abstract, and Haipeng Tang MS, Zhixin Jiang MD, and Weihua Zhou PhD, for providing the Chinese abstract.

An audio interview was held January 25th, 2017 between the Associate Editor, Heinrich R. Schelbert, and Benjamin C. Lee, co-author of

this article. An audio file of the interview is available as an .mp3 download at the article webpage on SpringerLink.com, and can be found by searching for the article title or DOI.

Venkatesh L. Murthy and Edward P. Ficaro contributed equally to this work and are co-senior authors.

Reprint requests: Benjamin C. Lee, PhD, INVIA Medical Imaging Solutions, 3025 Boardwalk St., Suite 200, Ann Arbor, MI, 48108; blee@inviasolutions.com

1071-3581/\$34.00

Copyright © 2017 American Society of Nuclear Cardiology.

Spanish Abstract

Antecedentes. Una subóptima adquisición temporal del contenido sanguíneo del ventrículo izquierdo así como de las curvas de actividad tiempo (CAT) tisulares pueden introducir sesgos e incremento de la variabilidad en las estimaciones del flujo sanguíneo miocárdico (FSM) y la reserva de flujo (RFM) a partir de las imágenes dinámicas de perfusión miocárdica con PET. Nuestro objetivo fue optimizar la adquisición temporal para el cálculo de FSM y RFM.

Métodos. 24 voluntarios normales y 32 pacientes se sometieron a un estudio PET con Cloruro de Rubidio-82 (^{82}Rb) y protocolo estrés/reposo. Se utilizó una cuidadosa adquisición temporal para estimar el “full width at half máximo” (FWHM) de la CAT del contenido sanguíneo ventricular izquierdo. Se utilizó el análisis de Fourier para determinar el intervalo de adquisición más largo, IA, en función del FWHM, que preserva el contenido informativo de la fase sanguínea. Los conjuntos de datos dinámicos se reconstruyeron con duraciones de los cuadros que variaron de 2 a 20 segundos durante los primeros dos minutos para la fase sanguínea y de 30 a 120 segundos para la fase tisular. Las CAT tisular y del contenido sanguíneo ventricular se obtuvieron usando regiones de interés (ROI, *por sus siglas en inglés*) y se ajustaron a un modelo compartamental para la cuantificación de FSM y RFM. Los efectos de la adquisición temporal sobre el FSM y RFM se evaluaron utilizando datos clínicos y simulaciones.

Resultados. El IA aumenta linealmente con el aporte de la función FWHM ($R=0.93$). Aumentando la duración de los cuadros de la fase sanguínea de 5 a 15 segundos se incrementó el sesgo del FSM y RFM en la fase de estrés en un 10% así como la variabilidad en el 20-25%. La duración de los cuadros menor a 5 segundos tuvo sesgos de menos del 5% para los valores de FSM y RFM. Aumentando la duración de los cuadros de la fase tisular de 30 a 120 segundos resultó en sesgos menores a 2%.

Conclusiones. Realizar las 2 fases del PET con ^{82}Rb con duraciones de los cuadros de 5 segundos (fase sanguínea) y 120 segundos (fase tisular) es óptimo para adquirir la CAT del contenido sanguíneo ventricular en los nuevos sistemas de PET 3D. (J Nucl Cardiol 2017;24:1517–29.)

Chinese Abstract

背景. 在用动态的 PET 心肌灌注影像评估心肌血流量(MBF)和血流储备(MFR)时,对左心室血池和组织时间活动曲线(TACs)的不恰当时间采样可能会引起结果的偏差和不稳定。本研究拟优化MBF和MFR评估的时间采样方案。

方法. 24名正常志愿者和32名患者采用动态静息/负荷铷-82氯化物(^{82}Rb)PET成像,使用良好的时间采样来评估左心室血池TAC的半峰值处全幅宽度(FWHM)。采用傅里叶变换来确定最长采样间隔(T_s), T_s 作为FWHM的一个函数反映血液相位的信息。随着帧持续时间的改变,从超过前两分钟后的第2到20秒血液相位动态数据集被重建,以及从第30到120秒组织相位数据集被重建。对左心室血池和组织的TACs用兴趣区(ROI)进行采样,然后拟合成隔室模型,实现对MBF和MFR的量化。时间采样对MBF和MFR的影响使用临床数据和计算机模拟进行评估。

结果. T_s 随着输入函数FWHM呈线性增加($R=0.93$)。增加血液相位帧持续时间5到15秒导致负荷MBF和MFR出现10%的偏差并且增加了20%-25%的不稳定性,而小于5秒的帧持续时间导致MBF和MFR值出现小于5%的偏差。增加组织相位帧持续时间30到120秒导致MBF和MFR值小于2%的偏差。

结论. 一个两相位帧的动态(^{82}Rb)PET图像且持续时间为5秒(血液相位)和120秒(组织相位),能够最佳地对现代3DPET系统的血池TAC采样。(J Nucl Cardiol 2017;24:1517–29.)

Key Words: Myocardial perfusion imaging; PET · myocardial blood flow · myocardial flow reserve · Rubidium-82 · temporal sampling

Abbreviations

FWHM	Full width at half maximum
LV	Left ventricular
MBF	Myocardial blood flow
MFR	Myocardial flow reserve
PET	Positron emission tomography

ROI	Region of interest
T_1	Blood phase frame duration
T_2	Tissue phase frame duration
T_s	Maximum blood phase frame duration
TAC	Time-activity curve

See related editorial, pp. 1530–1534

INTRODUCTION

Measurement of myocardial blood flow (MBF) and myocardial flow reserve (MFR) from dynamic rubidium-82 chloride (^{82}Rb) PET offer incremental prognostic^{1,2} and diagnostic value^{3,4} over conventional relative perfusion imaging. The short 75-second half life of ^{82}Rb mandates careful optimization of tracer dose as well as infusion and imaging protocols in order to balance limitations of PET scanner sensitivity (which favor larger infused doses to preserve image quality of delayed images for visual interpretation) and maximum count rate capabilities due to dead time losses (which favor smaller infused doses for more accurate quantification of the arterial input function). These factors, along with the type of infusion system utilized, affect the measurement of the left ventricle (LV) blood pool and tissue time-activity curves (TAC). Dynamic image series, used to compute TACs, are reconstructed in discrete time bins, typically ranging from ~ 5 seconds to several minutes or more. While shorter time bins will increase temporal resolution, shorter time bins also result in decreased image quality due to lower total counts per image and increased reconstruction time and storage requirements for reconstructed images. Insufficient temporal resolution results in under-sampling of the LV blood pool TAC (input function) and may produce biased estimates of MBF and MFR and increase variability.

Prior studies have evaluated the effects of the shape of the input function and temporal sampling on the accuracy and precision of the K_1 uptake parameter using simulations.^{5,6} These analyses demonstrated that input functions with narrower widths have more high temporal frequency information and therefore require shorter frame durations for accurate measurement.⁴ However, these studies did not evaluate input functions with widths and shapes typically seen with modern ^{82}Rb infusion systems beyond simulations nor did they account for increased temporal sampling achievable with modern scanners using clinical data.

We sought to define an optimal temporal sampling protocol for dynamic ^{82}Rb imaging in modern PET systems, using real-world data from patients and healthy volunteers in addition to simulations alone,⁷ that minimizes the tradeoffs between biases and variability in estimates of MBF and MFR and increased time and storage for image reconstruction.^{5–10} We first found an efficient protocol and then evaluated its accuracy and variability.

MATERIALS AND METHODS

Study Population

We evaluated dynamic ^{82}Rb rest and stress images from 24 prospectively recruited normal volunteers with less than or

equal to 5% pre-test likelihood for coronary artery disease (CAD) (8 men, 16 women, age 51 ± 10 years) and 32 randomly selected patients that were referred for clinically indicated ^{82}Rb rest and stress imaging (22 men, 10 women, age 64 ± 14 years). Pre-screening tests for normal volunteers included detailed cardiovascular history and physical examination, comprehensive metabolic and lipid panels, hemoglobin A1c and/or oral glucose tolerance test, and symptom-limited maximal treadmill exercise EKG test. All subjects provided written informed consent and all exam protocols were approved by the University of Michigan Institutional Review Board.

PET Imaging

All subjects were instructed to avoid caffeine and methylxanthine intake for 24 hours and to fast overnight prior to PET imaging. ^{82}Rb was administered using a weight-adjusted protocol of 12 MBq/kg (0.32 mCi/kg) using the same activity (481–1665 MBq [13–45 mCi]) for both rest and stress. ^{82}Rb was directly eluted from a generator and infused into a brachial vein at 50 mL/min over 5–25 seconds using the Cardiogen-82 infusion system (Bracco Diagnostics, Monroe Township, NJ). Pharmacological stress was achieved using 0.4 mg of intravenous regadenoson. Dynamic PET scans were acquired in 3D list mode on a Biograph mCT whole-body PET/CT scanner (Siemens Healthcare USA, Malvern, PA).

Image Processing

Dynamic attenuation-corrected emission images were reconstructed using the manufacturer recommended protocol (attenuation-weighted iterative 3D ordered-subset expectation-maximization iterative reconstruction or 3D-OSEM with 24 subsets and 3 iterations) without post-filtering during reconstruction to a matrix size of 128×128 and pixel size of 3.18×3.18 mm. Dynamic series were reconstructed from the first 6 minutes of acquired data.¹¹ Temporal sampling protocols were divided into two phases: (1) a fine, uniformly sampled dynamic blood phase that spanned the full width of the input function peak and (2) a coarse, uniformly sampled slow-varying tissue phase. For analysis of optimum temporal sampling during the blood phase, the blood phase frame duration (T_1) was varied from 2 to 20 seconds while the tissue phase frame duration (T_2) remained fixed at 30 seconds. For example, $\{12 \times 10$ seconds, 8×30 seconds $\}$ is a sampling protocol with 10-second blood phase frame durations. For analysis of optimum temporal sampling during the tissue phase, T_1 was fixed at 10 seconds while T_2 was varied from 30 to 120 seconds. Because the peak of the input function may not fall consistently within a single temporal frame,⁶ reconstructions with time shifts of a half frame duration for the blood phase were also analyzed.

LV myocardial surfaces were automatically determined using the 4DM software (INVIA, Ann Arbor, MI) that utilized a myocardial volume summed from the data acquired from 2 to 6 minutes. As needed, manual adjustment of the cardiac axis and the position of the mitral valve plane were performed (30 of the 56 datasets). Minimal smoothing was applied to the image volume using a 1-12-1 weighted averaging filter in three dimensions.

The region-of-interest (ROI) sampling methodology used a 3D box that was centered at the mitral valve plane along the axis of the LV to automatically extract a unique LV blood pool time-activity curve. The size of the box was 2×2 pixels wide (6.4 mm) to minimize spillover from the myocardium and spanned the long axis to include activity in both the LV and left atrium ($6.4 \times 6.4 \times 30$ mm). The myocardial tissue time-activity curves were estimated from the tracer activity midway between the endocardial and epicardial surfaces for each of the three vascular territories of the left anterior descending (LAD), left circumflex (LCX), and right coronary artery (RCA), and averaged for the global tissue TAC.

Input Function Width Estimation

From the dynamic sequence with $T_1 = 2$ seconds, the input function width was estimated from the FWHM of the LV blood pool TAC within the blood phase. The peak height and peak time were approximated with a quadratic fit to the points from the peak frame and its two adjacent neighboring frames. The FWHM was determined using linear interpolation between points in all other frames.

Maximum Frame Duration Definition

Fourier analysis was used to determine the minimum temporal sampling to maintain 95% of underlying information contained in the input function. The minimum temporal sampling defines the maximum limit for the frame duration during the blood phase (defined as T_S), which is also related to the Nyquist sampling rate which preserves 95% of the signal information. Linear mixed-effects modeling, with robust estimation and with per subject error terms to account for correlation between stress and rest data from a given subject, was used to assess the relationship between T_S and FWHM. Similar analyses were also conducted for the tissue phase frame duration using the LV tissue TAC. These analyses defined an efficient temporal sampling protocol. The minimum temporal sampling definition was also used to analyze the number of phases and their phase intervals that result in the fewest number of frames across all datasets.

Blood Flow Estimation

Both LV blood pool input function and LV tissue TAC were fit to a 1-tissue compartment to obtain estimates for uptake rate K_1 , washout rate k_2 , and LV blood pool to myocardium spillover f_V . Myocardial blood flow was computed from the estimated K_1 using a previously validated K_1 -MBF relationship for ^{82}Rb .¹² All temporal frames were frame duration weighted in the kinetic fitting.

Verification by Simulation

Using previously validated methods,⁵ noiseless and noisy simulated TACs were generated to confirm the effects of

varying input function widths and blood phase frame durations on MBF and MFR. A model for the baseline time-activity curves from the infusion system was selected from the available ^{82}Rb normal volunteer data with a blood pool frame duration of 2 seconds and a narrow FWHM of 9.8 seconds for stress. An additional input function with a greater width was simulated by a convolution with rectangular function of width of 15 seconds as a dispersion function⁵ with a resulting FWHM of 16.3 seconds. The baseline TACs were interpolated to 1-second frames.

The myocardial tissue TACs were generated using a 1-tissue compartment model including spillover from the blood pool TAC to the myocardial tissue TAC. The kinetic parameters used in the 1-tissue compartment model were chosen from the averaged kinetic fits of the ^{82}Rb normal volunteer data during stress: $K_1 = 1.11$ (MBF = 2.95) $\text{mL}\cdot\text{min}^{-1}\cdot\text{g}^{-1}$, $k_2 = 0.12 \text{ min}^{-1}$, and $f_V = 0.43$.

To generate noisy simulated TACs, time-varying Gaussian noise (100 noise realizations per frame) with zero mean and with standard deviation varying with the tissue activity and globally scaled to 5% of the mean tissue activity post 2 minutes was added to the baseline tissue TACs. Both noiseless and noisy 1-second frame duration TACs were rebinned to TACs with the blood phase frame duration T_1 varying from 1 to 20 seconds with a fixed tissue phase frame duration of 30 seconds. During the rebinning process, the blood phase frames were also time-shifted from 1 to 20 seconds to allow varying proportions of the peak of the input function to fall within one or more temporal frames. MBF estimates were averaged over all noise realizations and all frame shifts per sampling protocol.

Time Sampling Effects on MBF and MFR

Relative differences in global and regional MBF or MFR by varying frame durations were computed for simulated and clinical data. The MBF and MFR values from the 2-second blood phase frame duration and the 30-second tissue phase frame duration sampling protocols were the reference values for computing relative differences. Changes in MBF and MFR were used to select the optimal temporal sampling protocol that improved accuracy and reduced variability.

Statistical Analysis

Statistical significance was assessed with Wilcoxon tests and Fisher exact tests for continuous and dichotomous variables, respectively. Two-sided values of $P < 0.05$ were considered significant. All statistical analyses were performed with R 3.3.1 (The R Foundation for Statistical Computing).

RESULTS

Study Population

Characteristics of patients and normal volunteers evaluated are given in Table 1.

Table 1. Subject characteristics

Characteristic	Normals (n = 24)	Patients (n = 32)	All (n = 56)	P
Age (y)	50.5 [44.5–58.8]	64.2 [57.4–73.0]	58.3 [46.3–69.0]	0.0002
Weight (kg)	74.5 [61.7–83.7]	88.2 [72.3–99.5]	82.3 [65.4–90.3]	0.014
Height (m)	1.70 [1.63–1.75]	1.64 [1.63–1.75]	1.67 [1.63–1.75]	0.72
Male sex	8 (33)	22 (69)	30 (54)	0.014
Hypertension	0 (0)	26 (81)	26 (48)	<0.0001
Dyslipidemia	0 (0)	29 (91)	29 (52)	<0.0001
Diabetes	0 (0)	17 (53)	17 (30)	<0.0001

Continuous variables are presented as median [interquartile range]
 Dichotomous variables are presented as number (%)
 P-values compare normals and patients

Input Functions

TACs from a representative normal volunteer are presented in Figure 1 who received a 1000 MBq (27 mCi) injected dose from a 4-day-old generator. The relatively narrow input function FWHM of 8.3 seconds requires sampling of less than 8.2 seconds to accurately capture the peak (Figure 1A). With a 5-second frame duration, a modest 1.0 second (12%) increase in the FWHM is observed. With a 10-second frame duration, the measured FWHM is markedly increased at 13.9 seconds due to insufficient temporal

sampling. Figure 1B shows for the same representative volunteer, the tissue phase of the tissue TAC with 30-, 60-, and 120-second frame durations maintain their similar shape with relative root mean squared differences of 0.6% for 60 seconds and 1.7% for 120 seconds both with respect to the 30-second frame duration tissue TAC.

Increasing protocol complexity to include one or more additional transition phases with intermediate length frame durations between the blood pool and tissue phases (Figure 2A) did not result in significantly

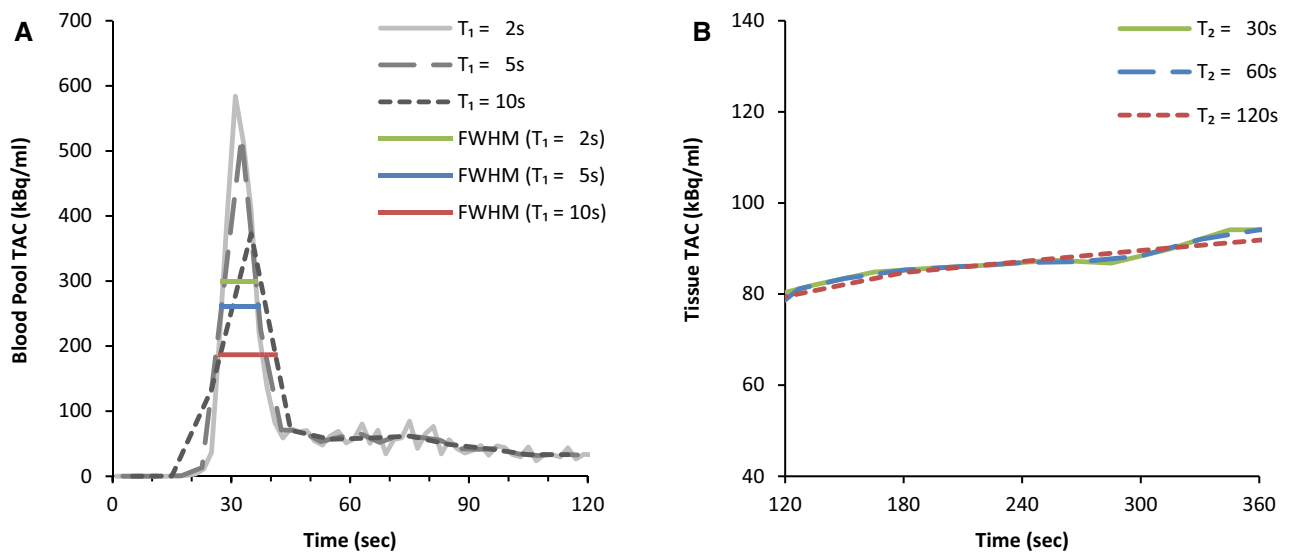


Figure 1. Time-activity curves from a representative normal volunteer with time points from initiation of the scan to 120 seconds (blood phase), panel A and from 120 to 360 seconds (tissue phase), panel B. In panel A, TACs from reconstructions with frame durations of 10 seconds demonstrate a dramatically different shape from those with shorter durations: the 10-second reconstructions resulted in a substantially lower and wider TAC. With increasing frame durations of 2, 5, and 10 seconds, there were increasing FWHM values of 8.3, 9.3, and 13.9 seconds, respectively. In panel B, the tissue TAC shows successive longer tissue phase frame durations of 30, 60, and 120 seconds with only minor differences in shape.

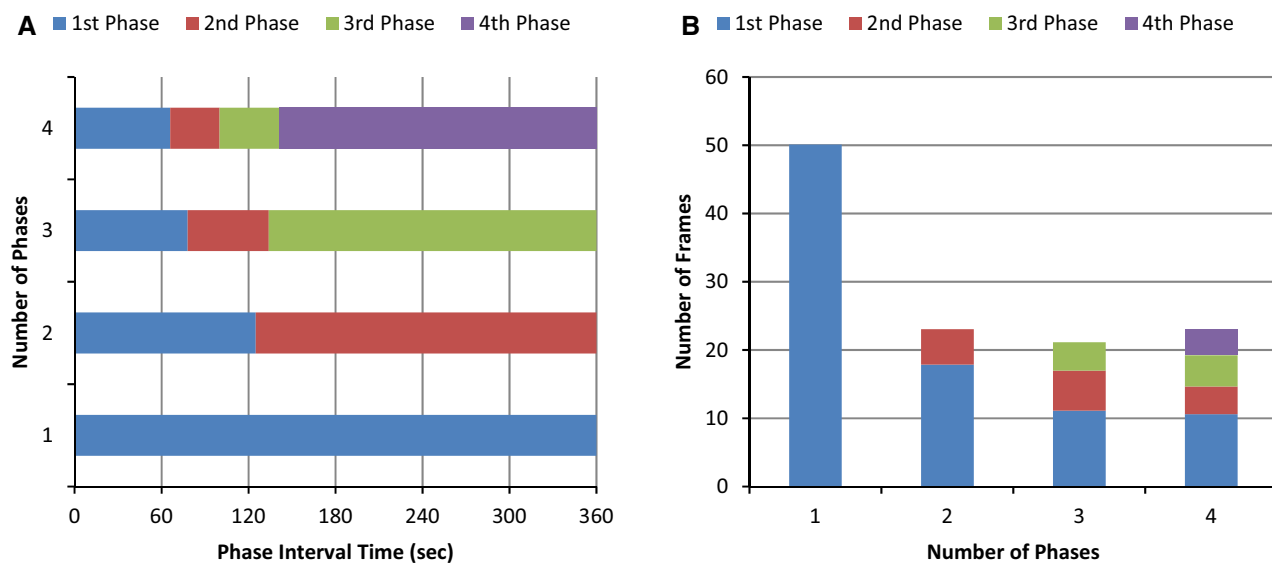


Figure 2. **A** Interval times for 1, 2, 3, and 4 uniformly sampled phases with the fewest number of frames across all datasets. **B** Fewest number of frames for protocols with different numbers of phases.

Table 2. Imaging parameters

Parameter	Normals (n = 24)	Patients (n = 32)	All (n = 56)	P
Stress cardiac output (mL/min)	6.7 [5.8-7.8]	5.9 [4.9-7.2]	6.3 [5.0-7.3]	0.045
Rest cardiac output (mL/min)	4.3 [3.5-5.3]	4.7 [3.6-5.5]	4.5 [3.6-5.4]	0.32
Stress ejection fraction (%)	69.0 [65.3-74.0]	50.0 [37.3-59.0]	58.1 [47.5-69.0]	0.0008
Rest ejection fraction (%)	61.5 [57.3-65.0]	50.5 [38.0-63.0]	55.2 [45.8-65.0]	0.016
Stress ejection fraction <45%	0 (0)	12 (38)	12 (21)	0.0006
Rest ejection fraction <45%	0 (0)	14 (44)	14 (25)	0.0001
Generator age, range (d)	16.8 [7.3-23.8]	16.3 [9.3-24.0]	16.5 [8.3-24.0]	0.90
Stress time to peak, range (s)	32.3 [29.9-34.0]	40.6 [34.3-43.3]	37.0 [32.3-39.0]	<0.0001
Rest time to peak, range (s)	37.5 [34.3-40.4]	41.5 [36.3-45.1]	39.8 [35.6-42.5]	0.018
Stress FWHM, range (s)	10.4 [8.4-12.7]	12.9 [8.2-15.3]	11.8 [8.3-14.0]	0.16
Rest FWHM, range (s)	12.8 [10.7-14.4]	13.9 [10.6-16.9]	13.4 [10.6-14.9]	0.74
Stress max. frame duration T_S (s)	11.0 [8.9-12.7]	12.1 [8.4-14.9]	11.7 [8.8-13.7]	0.55
Rest max. frame durations T_S (s)	12.5 [10.5-14.0]	12.5 [9.8-15.4]	12.5 [10.2-14.4]	0.84

Continuous variables are presented as median [interquartile range]
Dichotomous variables are presented as number (%)
P-values compare normals and patients

improved protocol efficiency (Figure 2B). Similarly, a four-phase protocol did not result in greater efficiency than achieved by a two-phase protocol.

The mean time of peak blood pool activity was 37 ± 9.2 seconds for all stress studies and 39 ± 6.4 seconds for all rest studies, with a maximum of 85 seconds (Table 2). The mean FWHM was 11.8 ± 5.6 seconds for all stress studies and 13.4 ± 5.2 seconds for all rest studies, with a maximum of 38 seconds. Consequently, an empiric transition point between the blood pool and tissue phases at the maximum time to peak activity plus the maximum

FWHM ($85 + 38 = 123 \approx 120$ seconds) was selected which confirms the results in Figure 2A. This duration was sufficient to fully capture the blood phase in 100% of normal volunteers and 98% of patients (Figure 3A). The frame duration during the blood phase which retained $\geq 95\%$ of the blood pool TAC signal information for all volunteers and patients was 5 seconds (Figure 3C), including at rest and stress (Figure 3D). Analogously, the phase frame duration during the tissue phase which retained $\geq 95\%$ of the tissue TAC signal information for all volunteers and patients was 125 seconds (Figure 3B).

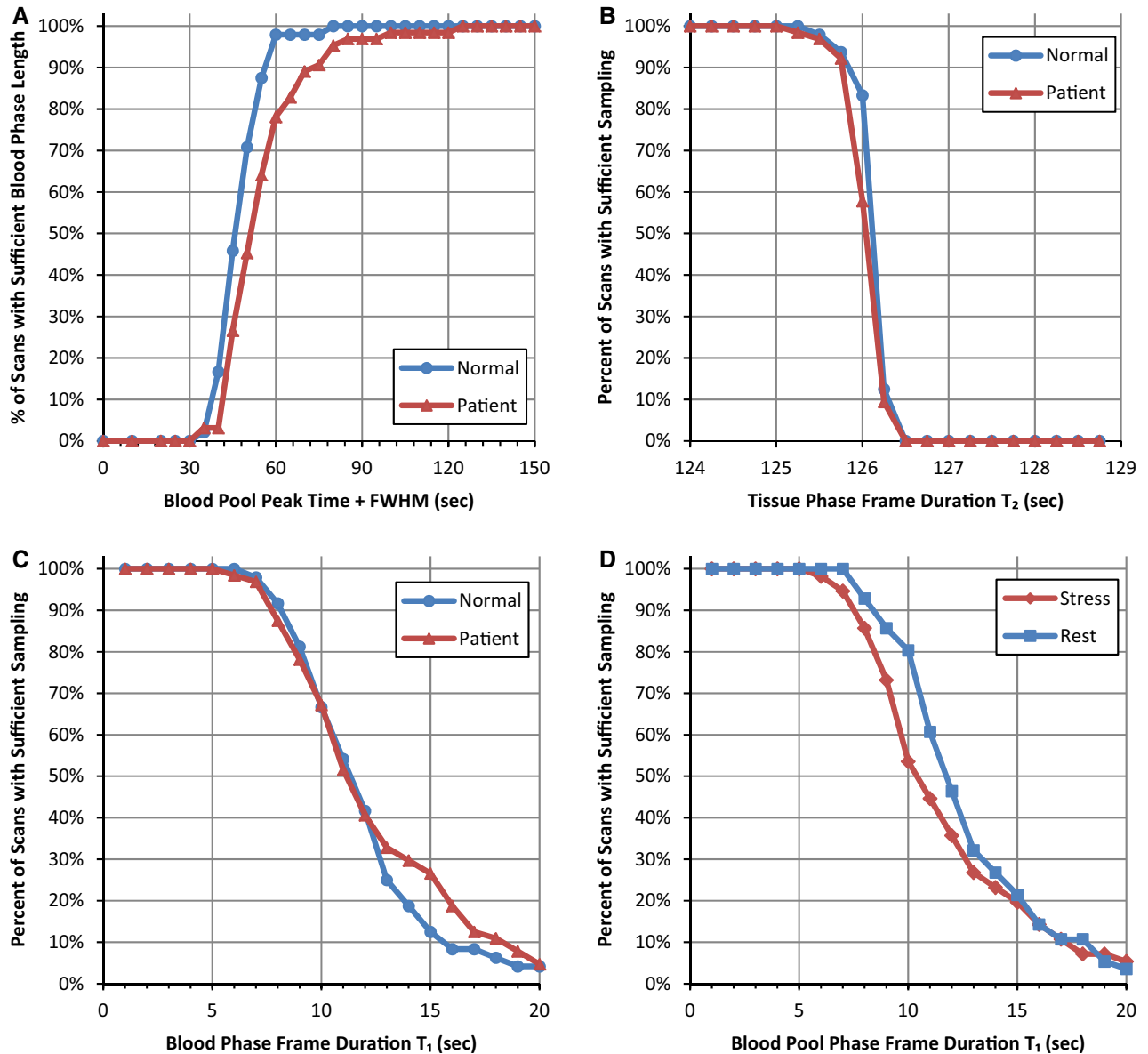


Figure 3. **A** Percentage of PET datasets with sufficient blood phase length that includes the blood pool peak time from start of injection plus FWHM. **B** Percentage of sufficiently sampled tissue TAC in the tissue phase. **C, D** Percentage of sufficiently sampled blood pool TAC where $T_1 < T_S$ as a function of initial frame duration. All data are shown separated into **(A, B, C)** normal and patient datasets and **(D)** stress and rest series.

Fitting for Infusion System and Scanner

We found that the minimum sampling rate (frame duration) was proportionally correlated to the FWHM ($R = 0.926$). The minimum sampling rate could be reliably predicted from the linear mixed-effects model fit as seen in the scatter plot in Figure 4. Generally, input functions with width and frame duration that fall above this regression line will be under-sampled, while those

below will be over-sampled. Those on or near the line will be near optimally sampled.

Simulated Time Sampling Effects on MBF and MFR

In simulations, the mean relative MBF error slowly increases as a function of frame duration for different

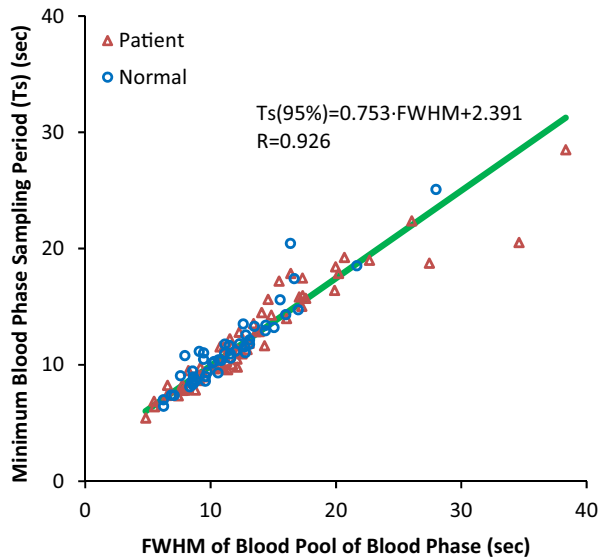


Figure 4. Linear relationship between the frame duration of the blood phase and the peak FWHM of the input curve that preserves 95% of the blood pool signal information.

input function widths of 9.8 (Figure 5A) and 16.3 seconds (Figure 5C). The variability in measurement dramatically increases when the blood phase frame duration T_1 is greater than the maximum frame duration T_S , which preserves $\geq 95\%$ of the information content of the input function. The 95% confidence intervals substantially increase beyond 8.9 seconds for FWHM = 9.8 and 15.0 seconds for FWHM = 16.3 seconds. Furthermore, where T_1 is substantially greater than T_S , the average measured MBF is meaningfully different from the true value. When noise is incorporated into simulations (Figure 5B, D), the mean MBF error is nearly identical to the noise-free simulation model. With the 95% confidence intervals beginning at approximately 5%, the MBF error variability begins to increase slowly with frame durations after the maximum frame duration T_S and increases more sharply 5 seconds after T_S .

Clinical Time Sampling Effects on MBF and MFR

In Figure 6A, C, E, the mean relative differences in global MBF and MFR of the 56 studies increased as the blood phase frame duration T_1 increased. The standard deviations of relative MBF and MFR values also increased with increasing frame duration T_1 beyond the recommended frame duration T_S for each dataset.

Sudden increases in the variability are more clearly seen beyond T_S . The global MBF and MFR mean relative differences from varying T_2 were within 5% with little change in variability compared to the reference sampling protocol with T_2 as 30 seconds seen in Figure 6B, D, F.

Mean magnitude differences in global MBF and MFR as a function of frame durations T_1 are re-categorized by dataset type in Table 3. For sufficiently sampled protocols ($T_1 \leq 5$ seconds), the average relative magnitude differences of stress MBF, rest MBF, and MFR are less than 5%. For insufficiently sampled protocols ($T_1 > 5$ seconds), the average relative magnitude differences for stress MBF for all datasets exceed 5% at approximately 6-second frame durations at an increasing rate of 12 percentage points per 10 seconds of frame duration. The average relative magnitude difference for rest MBF and MFR for all datasets also exceeds 5% at approximately 8-second frame durations but at an increasing rate of 6 percentage points per 10 seconds of frame duration.

The mean magnitude differences in global and regional MBF and MFR as a function of the tissue phase frame durations T_2 are within 3% for T_2 of 60 seconds and within 5% for T_2 of 120 seconds in Table 4. Nearly all of the MBF and MFR relative differences in each of the vascular territories of LAD, LCX, and RCA were not significantly different from the global relative differences in spite of their lower tissue count statistics.

We minimized MBF differences to within 5% with a reduction of 53% of the number of frames when using 5-second frame durations in the blood phase compared to the least efficient frame duration of 2 seconds. Alternatively, using frame durations less than 5 seconds resulted in steeply increasing requirements for storage space and image reconstruction time (Figure 7). Ensuring sufficient sampling, the tradeoff that remains in selecting the appropriate sampling protocol is between quantification accuracy and computational efficiency.

DISCUSSION

This study optimizes dynamic time frame binning during image reconstruction for quantification of MBF and MFR using ^{82}Rb PET. Frame durations were defined to minimize bias, variability, and demands on computation and storage. Compared to the other protocols used in this study, these protocols result in improvements of 2-8%, 3-15%, and 16-53% in bias, variability, and computational efficiency, respectively. These optimal

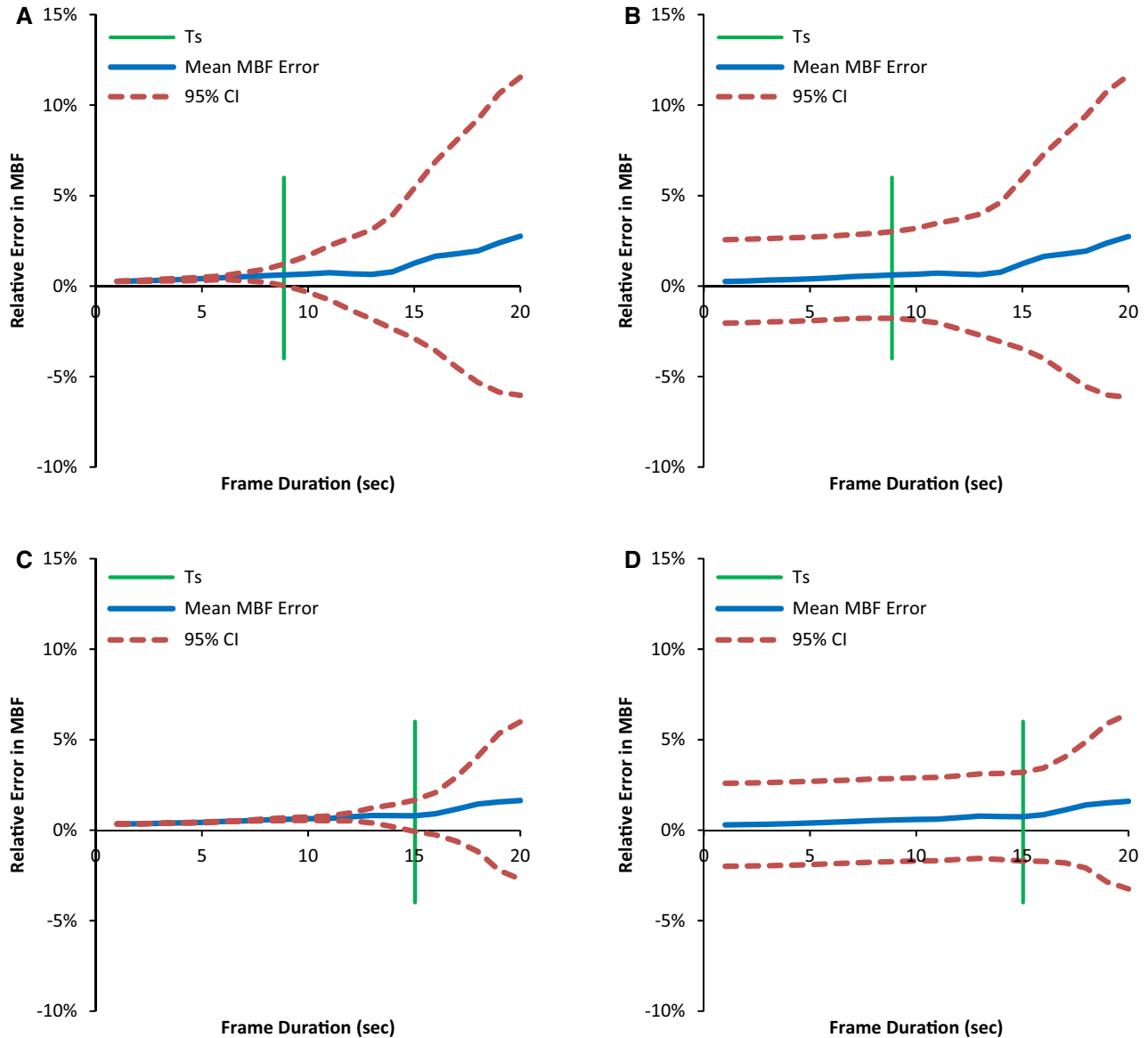


Figure 5. Relative MBF error vs blood phase frame duration in simulated data with FWHM of 9.8 seconds and T_S of 8.9 seconds, indicated by *green line*, (A) noiseless and (B) with simulated noise, and FWHM of 16.3 seconds and T_S of 15.0 seconds, (C) noiseless and (D) with simulated noise. *Blue line* indicates mean and *red lines* indicate range of errors (CI confidence intervals).

protocols can readily be implemented with existing software packages on clinical PET instruments.

Current System Recommendations

For injections performed by an automated constant flow ^{82}Rb infusion system such as the Bracco Cardigen-82, the input function width is not directly controllable by the user and is determined by multiple factors including requested activity, generator age, and

the physiology of the individual patient. Given substantial potential variability in input function duration, frame durations should be determined based on the lower end of the range of input function durations to be expected in clinical practice. For example, when using a Bracco infusion system with a Siemens Biograph mCT, a 5-second blood phase frame duration would be sufficient as seen in Figure 3C.

With a current-generation 3D scanner with high count rate capabilities, we showed that data with blood

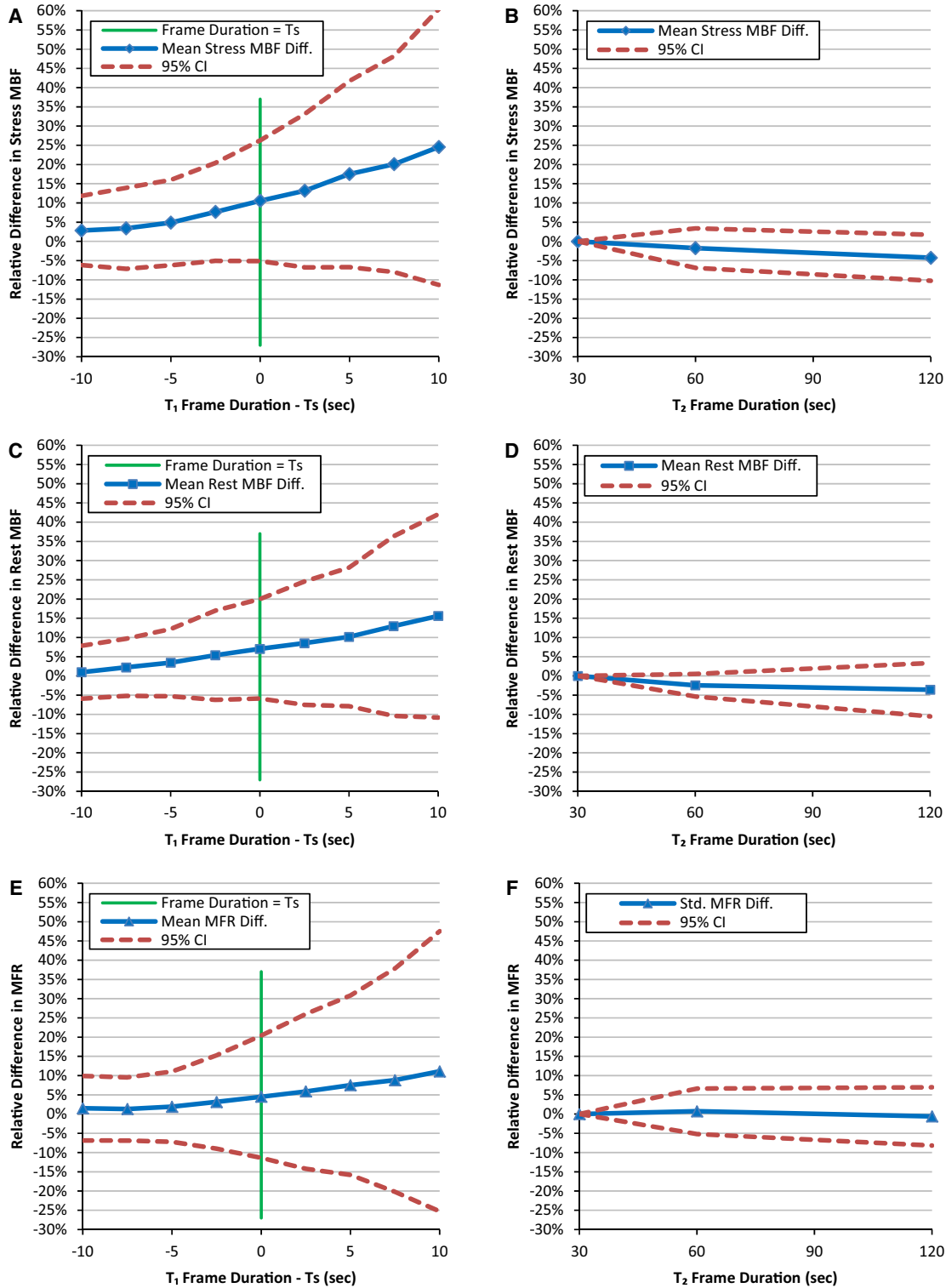


Figure 6. Mean relative differences in global (A) stress MBF, (C) rest MBF, and (E) MFR of all datasets vs blood phase frame duration minus T_S of each dataset. The 95% confidence interval lines denote increasing variability after T_S . Reference values are global MBF and MFR values using the $T_1 = 2$ second frame duration. Mean relative differences in global (B) stress MBF, (D) rest MBF, and (F) MFR of all datasets vs tissue phase frame duration. Reference values are global MBF and MFR values using the $T_2 = 30$ second frame duration.

Table 3. Effects of blood phase sampling on global MBF and MFR means

Dataset	Quantity T_1	Reference ^a 2 s	Magnitude difference mean at T_1 (%)						
			4 s	5 s	8 s	10 s	12 s	15 s	20 s
Normal	Stress MBF	2.6 ± 0.6	3.9	4.6	9.3	12.1	15.6	20.0	28.0
	Rest MBF	0.8 ± 0.2	2.2	2.5	4.0	4.6	5.7	6.9	9.6
	MFR	3.4 ± 0.8	2.6	2.9	5.9	7.9	10.4	13.2	18.0
Patient	Stress MBF	1.7 ± 0.8	3.8	4.4	7.6	9.3	11.1	13.7	18.6
	Rest MBF	1.1 ± 0.4	3.1	4.3	6.3	8.1	9.6	11.4	15.8
	MFR	1.7 ± 0.8	2.8	3.3	3.6	4.7	5.1	6.2	7.6
All	Stress MBF	2.1 ± 0.8	3.8	4.5	8.3	10.5	13.0	16.4	22.7
	Rest MBF	1.0 ± 0.4	2.7	3.5	5.3	6.6	7.9	9.4	13.2
	MFR	2.4 ± 1.2	2.7	3.1	4.6	6.1	7.3	9.2	12.1

^a Reference values come from 2-second frame duration result averages and in mL·min⁻¹·g⁻¹ for MBF and unitless for MFR

Table 4. Effects of tissue phase sampling on global and regional MBF and MFR means for all datasets

Quantity T_2	Region	Reference ^a 30 s	Magnitude difference mean at T_2 (%)			<i>P</i>	
			60 s	120 s	60 s	120 s	
Stress MBF	Global	2.5 ± 1.2	2.3	4.4			
	LAD	2.5 ± 1.1	2.4	4.8	0.77	0.061	
	LCX	2.5 ± 1.3	2.4	4.2	0.82	0.0044	
	RCA	2.8 ± 2.8	2.4	4.4	0.64	0.33	
Rest MBF	Global	1.0 ± 0.4	2.6	4.4			
	LAD	1.1 ± 0.4	2.9	4.5	0.54	0.88	
	LCX	1.0 ± 0.4	2.5	4.2	0.082	0.080	
	RCA	1.0 ± 0.4	3.0	5.2	0.40	0.25	
MFR	Global	2.7 ± 1.6	2.1	2.8			
	LAD	2.6 ± 1.4	2.5	3.0	0.20	0.32	
	LCX	2.6 ± 1.6	2.5	3.6	0.26	0.87	
	RCA	3.1 ± 2.1	2.7	4.0	0.25	0.083	

^a Reference values come from 30-second tissue phase frame duration result averages and in mL·min⁻¹·g⁻¹ for MBF and unitless for MFR

P-values compare regional (LAD, LCX, RCA) with global relative magnitude differences

pool FWHM of 5 seconds or greater were adequately sampled at 5-second frame durations for accurate MBF and MFR quantification to within 5% that of blood phase frame durations of less than 5 seconds. Variability in generator age or patient physiology (increasing LV dysfunction) typically results in variability in the FWHM of the blood pool TAC. Frame durations lower than 5 seconds increase storage space and reconstruction needed up to twofold (Figure 7). Consequently, 5-second sampling remains the recommended protocol, provided adequate counting statistics are obtained.

With respect to the tissue phase, we found that 125-second duration frames were sufficient to retain 95% of signal information for all tissue TACs and that T_2 frame durations of 30, 60, or 120 seconds can be accommodated with less than 2% bias in MBF and MFR values. Longer tissue phase frame durations might also be sufficient but were not evaluated in this study. Therefore, {24 × 5 seconds, 2 × 120 seconds} is one sampling protocol that sufficiently samples the blood phase and the tissue phase. A summary of sampling requirements is listed in Table 5.

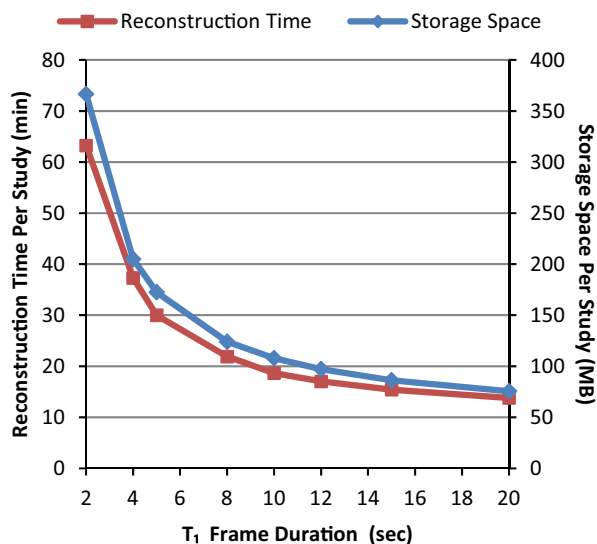


Figure 7. Reconstruction time and storage space per study vs blood phase frame duration.

Considerations for Other Systems

For count-limited scanners, slower protocols relative to the sampling can bias MBF estimates due to poor count statistics per frame.⁶ Conversely, fast bolus injections may result in count losses,¹³ leading to underestimation of the input function. For older PET systems with lower sensitivity and increased risk of count losses, careful optimization may require lengthening the injection duration using a controllable pump¹³ to increase the FWHM of the blood pool TAC. Then using the T_S and FWHM relationship in Figure 4, an optimum sampling protocol for the blood phase may be determined.

For scanners with a minimum frame duration, such as 5-second frame durations, for example, the injection should be performed >3 seconds again using the

relationship in Figure 4 in order to avoid under-sampling of the resulting input function.

For cases where the both the infusion system and scanner are constrained, bias and variability in MBF and MFR estimates may be unavoidable (Figures 5 and 6).

Limitations

Due to this retrospective analysis of the experimental data in the study, a reference arterial blood curve was not available and the 2-second blood phase frame duration and 30-second tissue phase frame duration sampling protocols were used as the reference standards. Another limitation was acquiring the data from one PET scanner and one infusion system. Differences in scanner types such as 2D or 3D acquisition and count rate performances could affect the image quality of short duration frames and the resulting image-derived arterial blood curves. Similarly, utilizing a different infusion system with a constant activity rate allowing for slower infusions¹³ instead of one with a constant flow rate may yield more relaxed recommendations for the blood phase frame duration. Variations in other methodological factors such as scatter correction, prompt gamma correction, image reconstruction and post-filtering, patient motion, and tracer kinetic modeling could also affect the results.¹⁴

CONCLUSIONS

A simple two-phase framing of dynamic ^{82}Rb PET images where the blood phase has frame durations of 5 seconds and a phase length of 120 seconds optimally samples the blood pool TAC for modern 3D PET systems. Frame durations for the tissue phase can be as large as 120 seconds with minimal effect on flow estimates. For imaging systems incapable of the required temporal sampling, the injection duration should be increased accordingly. If required temporal sampling is not possible

Table 5. Summary of sampling requirements

Parameter	Optimal	Alternate	Effect of alternate parameter values
Number of phases	2	3 or 4	Less than 8% space and time savings
Blood phase interval	0–120 s		
Tissue phase interval	120–360 s		
Blood phase frame duration (T_1)	5 s	10 s	Up to 12% MBF and MFR change
Tissue phase frame duration (T_2)	125 s	120, 60, or 30 s	Less than 5% MBF and MFR change

and injections cannot be lengthened, increased variability and bias in MBF and MFR estimates are expected.

NEW KNOWLEDGE GAINED

Under-sampling of the blood phase of TACs from dynamic ^{82}Rb PET images with frame durations longer than 5 seconds significantly increases MBF and MFR estimates and their variability. Tissue phase frame durations can be as large as 120 seconds.

Disclosures

B. C. Lee and J. B. Moody are employees of INVIA Medical Imaging Solutions. R. L. Weinberg has no disclosures. J. R. Corbett and E. P. Ficaro are owners of INVIA Medical Imaging Solutions. V. L. Murthy has received consulting fees from Ionetix, Inc. and Bracco Diagnostics and owns stock in General Electric, Mallinckrodt, and Cardinal Health. V. L. Murthy is supported by 1R01HL136685 from the National Heart, Lung, Blood Institute.

References

1. Murthy VL, Naya M, Foster CR, Hainer J, Gaber M, Di Carli G, et al Improved cardiac risk assessment with noninvasive measures of coronary flow reserve. *Circulation* 2011;124:2215-24. doi:10.1161/CIRCULATIONAHA.111.050427.
2. Ziadi MC, deKemp RA, Williams KA, Guo A, Chow BJW, Renaud JM, et al Impaired myocardial flow reserve on ^{82}Rb positron emission tomography imaging predicts adverse outcomes in patients assessed for myocardial ischemia. *J Am Coll Cardiol* 2011;58:740-8. doi:10.1016/j.jacc.2011.01.065.
3. Naya M, Murthy VL, Taqueti VR, Foster CR, Klein J, Garber M, et al Preserved coronary flow reserve effectively excludes high-risk coronary artery disease on angiography. *J Nucl Med* 2014;55:248-55. doi:10.2967/jnumed.113.121442.
4. Ziadi MC, deKemp RA, Williams K, Guo A, Renaud JM, Chow BJW, et al Does quantification of myocardial flow reserve using ^{82}Rb positron emission tomography facilitate detection of multi-vessel coronary artery disease? *J Nucl Cardiol* 2012;19:670-80. doi:10.1007/s12350-011-9506-5.
5. Raylman RR, Caraher JM, Hutchins GD. Sampling requirements for dynamic cardiac PET studies using image-derived input functions *J Nucl Med*. 1993;34:440-7.
6. Ross SG, Welch A, Gullberg GT, Huesman RH. An investigation into the effect of input function shape and image acquisition interval on estimates of washin for dynamic cardiac SPECT. *Phys Med Biol* 1997;42:2193-213. doi:10.1088/0031-9155/42/11/014.
7. Kolthammer JA, Muzic RF. Optimized dynamic framing for PET-based myocardial blood flow estimation. *Phys Med Biol* 2013;58:5783. doi:10.1088/0031-9155/58/16/5783.
8. Klein R, Beanlands R, deKemp R. Quantification of myocardial blood flow and flow reserve: Technical aspects. *J Nucl Cardiol* 2010;17(4):555-70.
9. Mazoyer BM, Huesman RH, Budinger TF, Knittel BL. Dynamic PET data analysis. *J Comput Assist Tomogr* 1986;10:645-53.
10. Herrero P, Markham J, Bergmann SR. Quantitation of myocardial blood flow with H_2^{15}O and positron emission tomography: Assessment and error analysis of a mathematical approach. *J Comput Assist Tomogr* 1989;13:862-73.
11. Efsaef M, Klein R, Ziadi MC, Beanlands RS, deKemp RA. Short-term repeatability of resting myocardial blood flow measurements using ^{82}Rb PET imaging *J Nucl Cardiol*. 2012;19:997-1006.
12. Lortie M, Beanlands RSB, Yoshinaga K, Klein R, Dasilva JN, DeKemp RA. Quantification of myocardial blood flow with ^{82}Rb dynamic PET imaging. *Eur J Nucl Med Mol Imaging* 2007;34:1765-74.
13. Klein R, Adler A, Beanlands RS, deKemp RA. Precision-controlled elution of a $^{82}\text{Sr}/^{82}\text{Rb}$ generator for cardiac perfusion imaging with positron emission tomography. *Phys Med Biol* 2007;52:659-73. doi:10.1088/0031-9155/52/3/009.
14. Moody J, Lee B, Corbett J, Ficaro E, Murthy V. Precision and accuracy of clinical quantification of myocardial blood flow by dynamic PET: A technical perspective. *J Nucl Med* 2015;22:935-51. doi:10.1007/s12350-015-0100-0.