

# Free-Breathing Phase Contrast MRI with Near 100% Respiratory Navigator Efficiency Using k-Space-Dependent Respiratory Gating

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**Purpose:** To investigate the efficacy of a novel respiratory motion scheme, where only the center of k-space is gated using respiratory navigators, versus a fully respiratory-gated acquisition for three-dimensional flow imaging.

**Methods:** Three-dimensional flow images were acquired axially using a gradient echo sequence in a volume, covering the ascending and descending aorta, and the pulmonary artery bifurcation in 12 healthy subjects ( $33.2 \pm 15.8$  years; five men). For respiratory motion compensation, two gating and tracking strategies were used with a 7-mm gating window: (1) All of k-space acquired within the gating window (fully gated) and (2) central k-space acquired within the gating window, and the remainder of k-space acquired without any gating (center gated). Each scan was repeated twice. Stroke volume, mean flow, peak velocity, and signal-to-noise-ratio measurements were performed both on the ascending and on the descending aorta for all acquisitions, which were compared using a linear mixed-effects model and Bland–Altman analysis.

**Results:** There were no statistical differences between the fully gated and the center-gated strategies for the quantification of stroke volume, peak velocity, and mean flow, as well as the signal-to-noise-ratio measurements. Furthermore, the proposed center-gated strategy had significantly shorter acquisition time compared to the fully gated strategy ( $13:19 \pm 3:02$  vs.  $19:35 \pm 5:02$ ,  $P < 0.001$ ).

**Conclusions:** The proposed novel center-gated strategy for three-dimensional flow MRI allows for markedly shorter acquisition time without any systematic variation in quantitative flow measurements in this small group of healthy volunteers. **Magn Reson Med 71:2172–2179, 2014. © 2013 Wiley Periodicals, Inc.**

**Key words:** phase contrast MR; blood flow assessment; 3D flow; respiratory navigators; stroke volume; mean flow; intra-scan variability; gating efficiency; cardiac MR

Phase contrast (PC) MRI is widely used to assess blood flow in cardiovascular disease (1–3). Clinically, the velocity component perpendicular to a two-dimensional plane is used for the measurements of blood flow (4,5). Through-plane aortic and pulmonic blood flow are measured and used for the evaluation of cardiac function and output, valvular regurgitation, and shunts. Recent advances have enabled three-dimensional (3D) time-resolved PC-MRI that allows measurement and visualization of all three directional components of blood flow (6). However, 3D PC-MRI requires long scan times, owing to the use of four-point encoding schemes that capture all three directions of velocity encoding, which limits its clinical usage. Cardiac and respiratory variations within the long scan time of 3D PC-MRI also hinder the accuracy and reproducibility of the flow measurements, which are important for quantification of cardiac indices, such as cardiac output or mitral regurgitation.

Various techniques have been proposed for reducing the acquisition time of PC-MRI. Non-Cartesian trajectories, including both radial (7,8) and spiral trajectories (9–12), have been used to reduce scan time 2- to 3-fold compared to Cartesian sampling. Echo planar imaging has also been employed to improve acquisition time (13–15). Parallel imaging has been utilized to reduce the scan time up to 3-fold (16–18). Significant speed-ups have been achieved using methods exploiting spatio-temporal correlations, such as k-t broad-use linear-acquisition speedup technique/sensitivity encoding (19,20), k-t PCA (21,22), and compressed sensing (23–25).

Although image acceleration for PC-MRI has been explored using various techniques, improving respiratory motion compensation during the 3D PC-MRI acquisition has attracted limited attention. Bellows gating (6), respiratory navigators (NAVs) (12,22,26) and self-gating techniques (27) have been utilized for 3D PC-MRI. Bellows gating (28) uses a pressure sensor positioned around the lower chest to measure chest wall expansion with respiration. Self-gating techniques (27,29) sample additional k-space profiles of the imaging volume at certain time intervals to gate the acquisition. Respiratory NAVs are commonly used, and utilize a two-dimensional pencil beam typically positioned on the dome of the right hemi-diaphragm (RHD) to track respiratory motion (30). Due to an approximately linear dependency between the respiratory motion of the heart and that of the RHD, NAV can be used to indirectly monitor the motion of the heart (31). In prospective NAV gating, the k-space lines acquired immediately after the NAV signal are used for

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Grant sponsor: NIH; Grant number: R01EB008743-01A2; Grant sponsor: NIH; Grant number: K99HL111410-01; Grant sponsor: Samsung Electronics, Suwon, South Korea.

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Received 5 March 2013; revised 10 June 2013; accepted 16 June 2013

DOI 10.1002/mrm.24874

Published online 30 July 2013 in Wiley Online Library (wileyonlinelibrary.com).

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FIG. 1. The NAV-gated 3D phase-contrast MRI sequence, and the flow chart for the proposed center-gated strategy with a 7-mm gating window (NAV, respiratory navigator; ACQ, data acquisition).

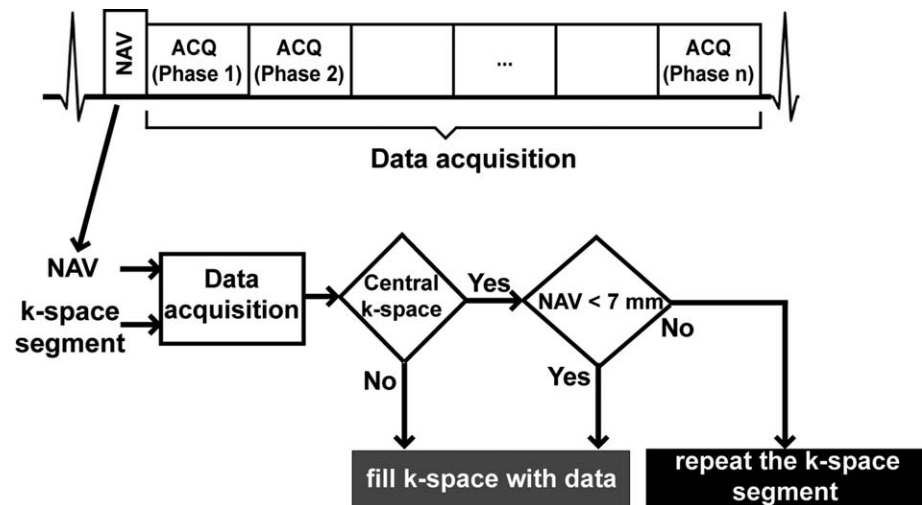


image reconstruction only if the NAV signal is within a predefined gating window. Otherwise, the corresponding k-space lines are rejected and reacquired in the next cardiac cycle. These motion compensation techniques typically lead to 30–60% data acquisition efficiency, depending on the subject's breathing pattern and the gating window that has been specified. Adaptive k-space reordering has also been proposed to further improve gating efficiency (32).

In this study, we sought to reduce the scan time of 3D PC-MRI without compromising reproducibility of flow measurements by improving the efficiency of respiratory motion compensation. We hypothesized that only respiratory gating the center of k-space will yield similar measurements to an acquisition where the full extent of k-space is respiratory gated as the phase information is mainly contained in the central k-space. The proposed technique that only respiratory-gates the central k-space was evaluated in vivo, and compared to fully gated acquisitions in terms of quantitative flow measurements, as well as intrascan variability and reproducibility.

## METHODS

### Center-Gated 3D PC-MRI Strategy

Figure 1 shows the proposed center-gated strategy for data acquisition. Following the preparation (NAV training) phase, where the end-expiration RHD position is determined based on the first 15 RHD positions, the image acquisition is divided into two phases. In the first phase, the central 4% of k-space is acquired with NAV gating and tracking using a predefined 7-mm gating window, similar to the prospective NAV acquisition. Subsequently, the remaining outer k-space data are acquired *without* using any respiratory motion gating, but with end-expiratory tracking. Hence, the central k-space is acquired with motion compensation, whereas the outer k-space contains both motion-corrupted and motion-free k-space segments.

### Free-Breathing PC-MRI

All imaging was performed on a 1.5-T Philips Achieva (Philips Healthcare, Best, The Netherlands) system with a 32-channel cardiac phased-array receiver coil. For this Health Insurance Portability and Accountability Act (HIPPA)-compliant study, the imaging protocol was approved by our institutional review board, and written informed consent was obtained from all participants.

Twelve healthy adult subjects ( $33.2 \pm 15.8$  years; five males) without contraindications to MRI were recruited. Scout images were acquired with a steady-state free precession sequence with in-plane resolution  $3.1 \times 3.1 \text{ mm}^2$  and 10 mm slice thickness, which was used for localization and assignment of the appropriate imaging slab covering the ascending and descending aorta, and the pulmonary bifurcation. A free-breathing electrocardiogram-triggered gradient echo sequence was used for acquisition. The trigger delay was chosen to be 20 ms following the acquisition of the leading NAV signal. Arrhythmia rejection was utilized, allowing the sampling of up to 90% of the cardiac cycle. This resulted in 25–15 phases per cardiac cycle for heart rates varying between 60 and 90 bpm, respectively. The imaging parameters were pulse repetition time/echo time = 5.7/3.4 ms, flip angle =  $10^\circ$ , velocity encoding = 300 cm/s, field of view =  $340 \times 340 \times 40 \text{ mm}^3$ , spatial resolution =  $2.0 \times 2.0 \times 4.0 \text{ mm}^3$  reconstructed to  $1.2 \times 1.2 \times 4.0 \text{ mm}^3$ . All images were acquired axially with right-left phase encoding. Only foot-head velocity encoding with interleaving in the same cycle was used with two lines per segment to provide a temporal resolution of 35 ms, including the default start-up pulses. The nominal scan time for these acquisitions was 13:30 min at a heart rate of 60 bpm (810 heartbeats), assuming no respiratory motion compensation.

A NAV placed on the dome of the RHD with 17-ms duration was used for respiratory motion measurement, utilizing prospective real-time correction and 0.6 superior-inferior tracking ratio (31,33). Two gating strategies were utilized with a 7-mm gating window: (1) The proposed center-gated strategy and (2) conventional gating

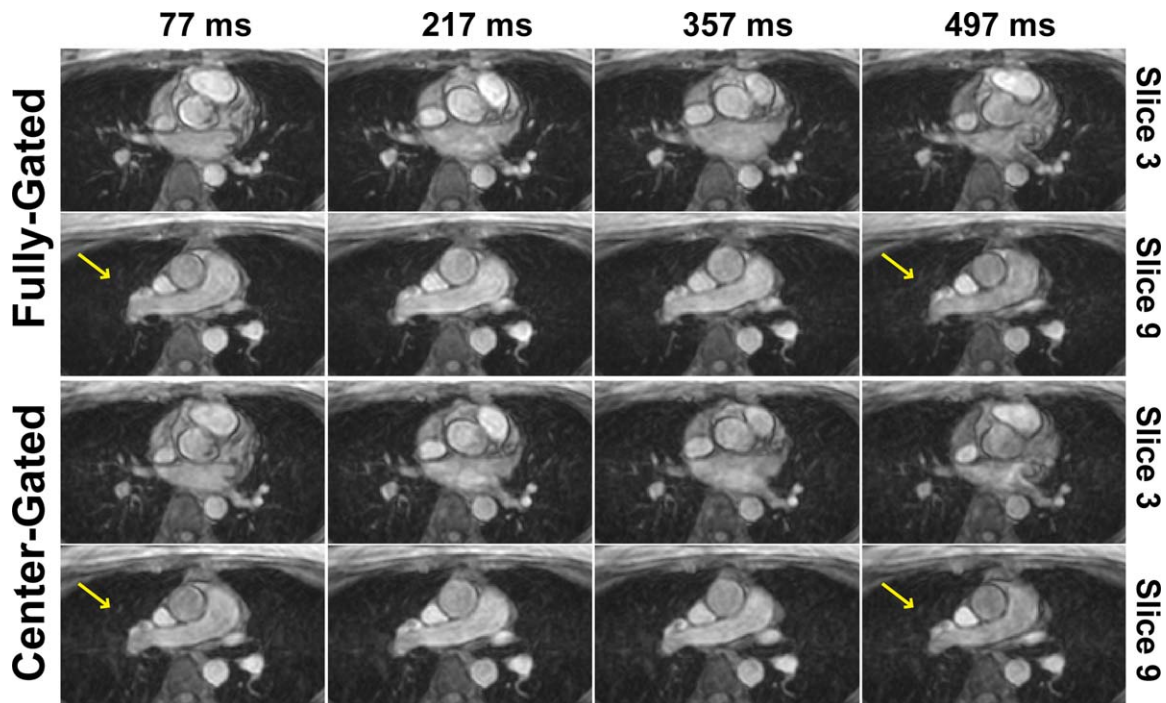


FIG. 2. Example magnitude images for two slices and various cardiac phases (specified by time after the *R*-wave) of 3D PC MRI acquisitions from a subject with a heart rate of 61 bpm, acquired using the center-gated and fully gated NAV strategies, depicting cross-sections across the descending aorta and ascending aorta. The magnitude images are also visualized similarly, with minor ghosting artifacts in the center-gated acquisition owing to motion corruption at the edges of the k-space. However, the center-gated acquisition took 12:13 min, whereas the fully gated one was 25:21 min. In both cases, more motion artifacts are visible in the later phases due to the use of a leading NAV.

and tracking for all k-space data (fully-gated). Central k-space lines were acquired first for both fully gated and center-gated scans. Each type of acquisition was repeated twice to characterize intrascan variability and reproducibility. Hence, a total of four data sets were acquired for each subject. The order of the acquisitions was randomized to cancel out bias owing to changes in the breathing patterns. Acquisition times were recorded for each of the scans.

#### Image and Statistical Analysis

Quantitative image analyses were performed using ViewForum (vR4.2V1L2, Philips Healthcare, Best, NL) to evaluate the two acquisition strategies. For each acquisition, a region-of-interest was manually drawn in the ascending and in the descending aorta on different slices of the 3D volume using the corresponding magnitude images. The region-of-interest was manually corrected throughout the cardiac cycle for cardiac motion. The mean blood flow in each cardiac cycle and the stroke volume were calculated, and the peak velocity was measured for each acquisition. Additionally, signal-to-noise ratio (SNR) was measured at the level of the pulmonary artery bifurcation in both the ascending and the descending aorta. The mean signal intensity was measured in the aortic blood pools. Noise was measured in the nonsignal areas of each image using a larger region-of-interest. The nonsignal areas considered were the anterior to the chest wall and the posterior to the back, as these areas were free

from ghosting artifacts or spurious signals owing to the prescribed scan orientation. SNR was calculated as the ratio of the mean signal to the standard deviation of the noise.

All statistical analyses were performed using SAS (v9.3, SAS Institute, Cary, NC). To assess the similarity of the measurements, including interscan variability, a linear model analysis was used for all the measurement parameters, including stroke volume, mean flow, and SNR in the ascending and the descending aorta. To capture the interscan (within-subject) variability, the data from the two fully gated acquisitions were structured into one single vector with an indicator variable for time within-subject. Center-gated data were structured the same way. The overall slope between these two vectors and the 95% confidence intervals was computed taking into account the correlation of the two within-subject measurements. The acquisitions were considered to be equivalent if the confidence interval covered 1.0. The correlation (variance-covariance) structure was assumed to be compound symmetry. This yielded the within- and between-subject variance components that were used in the estimation of the slope and confidence interval via a linear mixed-effects model. Additionally, Bland-Altman analysis was performed in comparing the individual fully gated and center-gated scans using the mean flow and stroke volume measurements through the ascending and descending aorta. The peak velocity measurements through the ascending and descending aorta were compared by first averaging over the two fully gated and



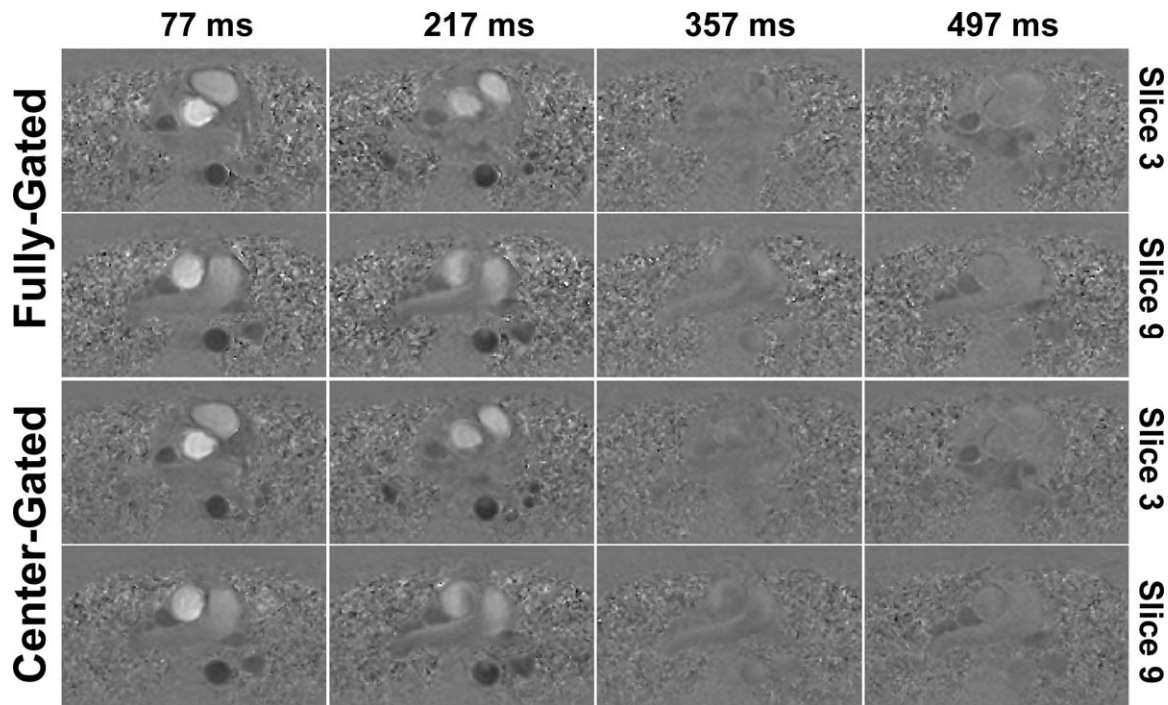


FIG. 3. Example of phase images for two slices and various cardiac phases (specified by time after the *R*-wave) of 3D PC MRI acquisitions from the same subject of Figure 2. The phase images show similar flow patterns through the ascending and the descending aorta, and any motion corruption at the edges of the k-space does not hinder the visualization of these patterns.

center-gated acquisitions, respectively, and then using a paired Student's *t*-test. A *P*-value of  $<0.05$  was considered to be significant.

## RESULTS

3D PC-MRI using both strategies was completed in all subjects, without complication. The average scan time for the center-gated acquisitions was  $13:19 \pm 3:02$  min (range, 8:35–20:04 min) and  $19:35 \pm 5:02$  min (range, 12:48–31:00 min) for the fully gated ones ( $P < 0.001$ ). The average scan times in heartbeats were  $1278 \pm 313$  (range, 890–1950) for the fully gated scans and  $859 \pm 70$  for the center-gated ones (range, 795–1002). The respiratory gating efficiencies were  $65.0 \pm 16.1$  and  $59.8 \pm 15.8$  (ranges, 40–90 and 41–82%, respectively) for the first and second fully gated scans, and  $94.4 \pm 2.5$  and  $93.8 \pm 2.2$  (ranges, 90–97% for both) for the first and second center-gated scans. The respiratory amplitudes, measured as the difference of the maximum and minimum NAV positions, are  $24.5 \pm 13.9$  and  $33.2 \pm 15.0$  mm for the first and second fully gated scans, respectively, and  $25.6 \pm 15.4$  and  $27.8 \pm 14.1$  mm for the first and second center-gated scans. The difference between the nominal scan time and the actual average scan time is due to the differences in breathing patterns, the prescribed FOV, as well as the heart rates of the subjects.

Figures 2 and 3 show examples of magnitude and phase images, respectively, using the two different NAV gating strategies. In the center-gated magnitude images, some residual ghosting artifacts are visible; however, these do not hinder visualization as the center of k-space was acquired within a 7-mm gating window. Furthermore, in both cases, more motion artifacts are visible in

the later phases due to the use of a leading NAV. The flow in the foot-head direction is visualized similarly both for the ascending and for the descending aorta with both gating strategies, and the effects of any motion on the edges of k-space are not easily identifiable. The center-gated scan was half as long as the fully gated scan for this subject (12:13 vs. 25:21 min). Figure 4 shows the velocity curves from these acquisitions. Although the mean length of the *R*–*R* interval has changed between the two acquisitions, the curves show the same trends, including the peak velocity values in systole.

The absolute stroke volume measurements and mean flow measurements for the ascending and descending aorta for both fully gated and center gated acquisitions are listed in Table 1. The linear mixed model analysis lead to overall slopes and corresponding confidence intervals of  $1.04 \pm 0.10$  and  $1.02 \pm 0.12$  for the stroke volumes through ascending and descending aorta, respectively. Similarly, for the mean flow, the slopes and the corresponding confidence intervals from the analysis were  $1.08 \pm 0.08$  and  $0.98 \pm 0.07$  for the ascending and descending aorta, respectively. Thus, the two acquisitions were statistically equivalent in terms of the quantification of these cardiac indices. Figures 5 and 6 show the reproducibility of the stroke volume and mean flow measurements, respectively, through the ascending and descending aorta, for all four acquisitions. The peak velocity measurements yield a peak descending aorta velocity of  $-70.3 \pm 13.9$  cm/s for the fully gated acquisition and  $-71.1 \pm 13.6$  cm/s for the center-gated acquisitions ( $P = 0.38$ ). The corresponding peak ascending aorta velocity measurements were  $61.6 \pm 11.2$  cm/s for the fully gated acquisitions and  $62.2 \pm 12.5$  for the center-

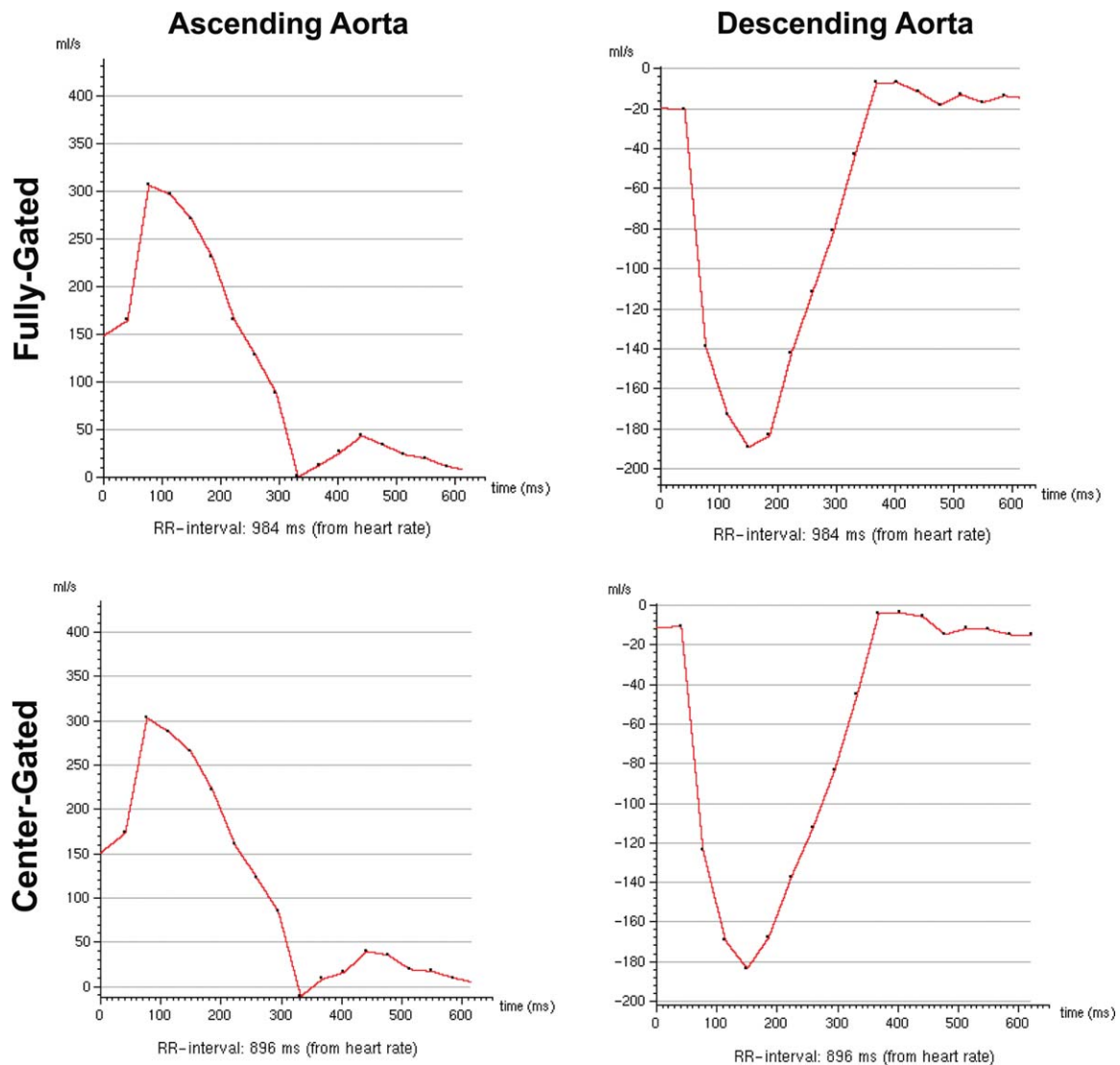


FIG. 4. Example of velocity curves through the ascending and descending aorta of the 3D PC MRI acquisitions from the same subject of Figure 2. The curves show the same trends, including the peak velocity values, although the mean value of the  $R$ - $R$  interval length has changed between the two acquisitions. Due to prospective electrocardiogram triggering, early part of the  $R$ - $R$  interval is not sampled densely.

gated acquisitions ( $P=0.57$ ), showing no significant difference between the temporal flow patterns in terms of the peak velocity.

The fully gated acquisitions had SNR of  $30.7 \pm 6.8$  and  $30.5 \pm 4.7$  in the ascending and the descending aorta, respectively. The respective SNR measurements for the

center-gated acquisitions yield an SNR of  $31.0 \pm 7.0$  and  $30.7 \pm 4.6$ . The linear mixed model analysis resulted in overall slopes and corresponding confidence intervals of  $1.00 \pm 0.06$  and  $0.96 \pm 0.09$  in the ascending and the descending aorta, respectively, indicating no significant difference between the two acquisitions in terms of SNR.

Table 1

Average Stroke Volume and Mean Flow Measurements through the Ascending Aorta and Descending Aorta for Fully Gated and Center-Gated Scans, and the Corresponding 95% Confidence Intervals for the Overall Slopes from the Linear Mixed Model Analysis<sup>a</sup>

	Stroke volume (mL)		Mean flow (mL/s)	
	Ascending aorta	Descending aorta	Ascending aorta	Descending aorta
Fully gated	$96.8 \pm 23.5$	$61.5 \pm 13.3$	$106.6 \pm 26.4$	$69.4 \pm 18.6$
Center gated	$95.5 \pm 25.0$	$61.6 \pm 16.5$	$104.6 \pm 28.5$	$69.0 \pm 17.8$
CI for slopes	$1.04 \pm 0.10$	$1.02 \pm 0.12$	$1.08 \pm 0.08$	$0.98 \pm 0.07$

<sup>a</sup>In all cases, the confidence intervals include 1.0, indicating statistical equivalence.

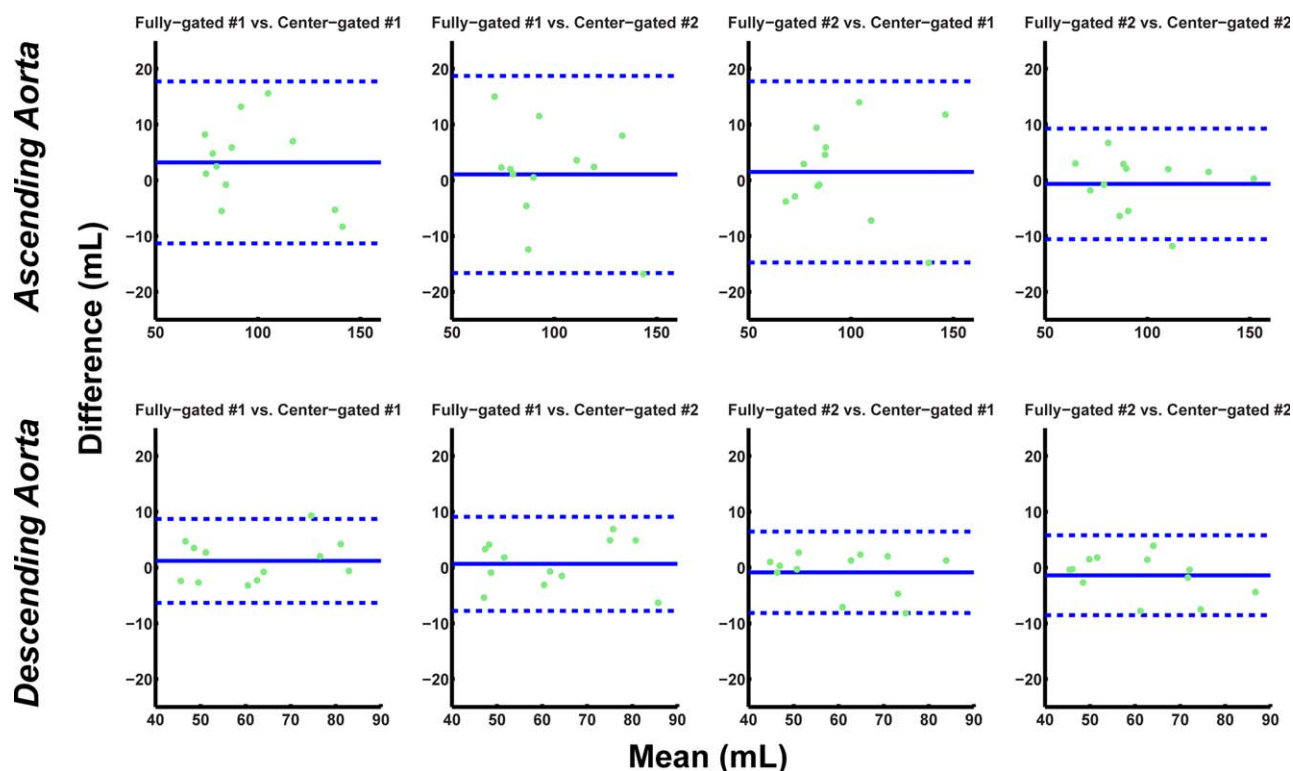


FIG. 5. Bland-Altman analysis of the stroke volume through the ascending and descending aorta for scan variability of fully gated scans versus the center-gated scans (dotted lines indicate  $\pm 1.96$  times the standard deviation of the mean difference). The variation observed between the four different scans are within the acceptable range in all cases. [Color figure can be viewed in the online issue, which is available at [wileyonlinelibrary.com](http://wileyonlinelibrary.com).]

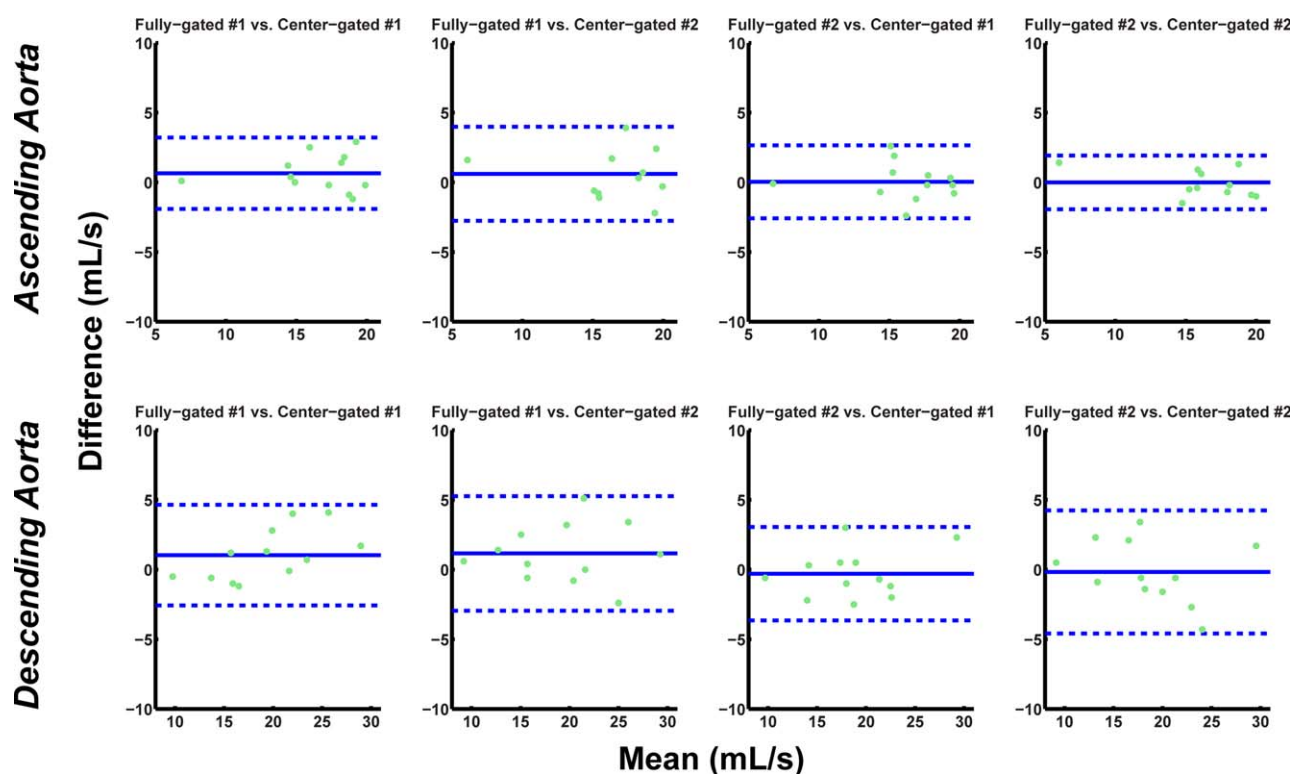


FIG. 6. Bland-Altman analysis of the mean flow through the ascending and descending aorta for scan variability of fully gated scans versus the center-gated scans (dotted lines indicate  $\pm 1.96$  times the standard deviation of the mean difference). The variation observed between the four different scans are within the acceptable range in all cases. [Color figure can be viewed in the online issue, which is available at [wileyonlinelibrary.com](http://wileyonlinelibrary.com).]

## DISCUSSION

In this study, we introduced and evaluated a novel respiratory motion compensation technique for PC MRI, which utilizes NAV gating and tracking for the central part of k-space only. We showed that the proposed technique allows for increased scan efficiency compared to fully gated acquisitions, significantly reducing the acquisition time, while resulting in statistically equivalent quantification of the cardiac indices, including stroke volume and mean flow.

Several approaches have been proposed to improve the NAV gating efficiency by using additional information, such as two-dimensional or 3D NAVs (34–37), binning strategies based on NAV data (38), or self-navigating approaches (29,39,40) in coronary MRI, as well as in 4D flow imaging (27). Although close to 100% gating efficiency may be achieved, these methods mostly rely on non-Cartesian acquisitions and motion correction based on parametric models. The concept of using different gating strategies for central and outer k-space has also been previously explored in coronary MRI (41,42). However, these studies either utilize a smaller gating window for central k-space for increased accuracy, or use motion-correction for the outer k-space lines that are acquired with a larger gating window. In our study, we focus on phase images, which are largely dependent on the central k-space, and we use a lower spatial-resolution than coronary MRI. These differences enable the utilization of the motion-corrupted outer k-space lines without any retrospective processing or respiratory gating for the quantification of cardiac output indices.

There are significant variations among the breathing patterns of subjects, which result in unpredictable scan durations, especially for 3D flow imaging. The main advantage of the proposed technique is that respiratory gating is only applied to 4% of the central k-space. Thus, even with variable breathing patterns, the data acquisition is completed within a near fixed scan time, removing the uncertainty of acquisition duration associated with NAV gating. We emphasize that the range of scan times reported for the proposed center-gated scheme was mainly due to the varying heart rates.

As no further processing is required in our technique, the technique can be combined with the existing accelerated imaging techniques without any modifications. Even a relatively modest 3- or 4-fold acceleration would enable acquisition of all three velocity-encoding directions with a field-of-view covering the whole-heart in approximately 6 min with a  $\sim 45$  ms temporal resolution. In our study, this was not explored to ensure that any artifact introduced between the different acquisitions was only due to respiratory motion, as well as to avoid potential ghosting, noise enhancement, or blurring artifacts that may be caused by image acceleration. Further studies are needed to investigate which existing acceleration techniques are well suited for the proposed center-gated 3D PC-MRI acquisition.

Only a small cohort of healthy subjects with regular breathing patterns and high respiratory efficiencies were studied in this study. We note that the average NAV gating efficiency of most patient populations are around

30–50%, which is lower than that of the fully gated scans acquired in the healthy cohort in this study. Furthermore, the respiratory amplitudes of the healthy cohort were around 3 cm, which may be smaller than patient populations. The lower gating efficiency and larger respiratory amplitudes of patient populations with irregular breathing patterns may lead to a degradation of image quality for the center-gated acquisitions. Further studies are needed to study the clinical evaluation of this approach in a larger cohort of patients.

In our study, we have used diaphragmatic NAV for characterizing respiratory motion. The primary goal is to acquire the central k-space with minimal motion and to acquire outer k-space with no motion tracking, which can also be applied when using respiratory bellows or fat NAV. The size of the central k-space to be gated was empirically chosen in our study based on the results of (42) on the effect of the central k-space size in compressed sensing-based motion correction. Further studies are needed to systematically study the minimum size of central k-space that is center gated.

## CONCLUSIONS

We have proposed and evaluated a novel respiratory motion compensation technique for PC-MRI that utilizes NAV gating for the central k-space and no gating for the outer k-space. This approach increases scan efficiency while providing quantification of cardiac indices that are clinically identical to fully gated scans.

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