Optimal Plane Selection for Measuring Post-prandial Blood Flow Increase within the Superior Mesenteric Artery: Analysis Using 4D Flow and Computational Fluid Dynamics

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Purpose: 2D cine phase contrast (PC)-MRI is a standard velocimetry for the superior mesenteric artery (SMA); however, the optimal localization of the measurement plane has never been fully discussed previously. The purpose of this Institutional Review Board approved prospective and single arm study is to test whether flow velocimetry of the SMA with combined use of 2D cine PC-MRI and meal challenge is dependent on the localizations of the measurement planes and to seek optimal section for velocimetry.

Methods: Seven healthy volunteers underwent cardiac phase resolved ECG gated 2D cine PC-MRI pre- and 30 min post-meal challenge at three measurement planes: proximal, curved mid section and distal straight section of the SMA at 3T. 4D Flow using 3D cine PC-MRI with vastly undersampled isotropic projection imaging (PC VIPR) was also performed right after 2D cine PC-MRI to delineate the flow dynamics within the SMA using streamline analysis. Two radiologists measured flow velocities, and rated the appearances of the abnormal flow in the SMA on streamlines derived from the 4D Flow and the computational fluid dynamics (CFD).

Results: 2D cine PC-MRI measured increased temporally averaged flow velocity (mm/s) after the meal challenge only in the proximal (129.3 vs. 97.8, \(P = 0.0313\)) and distal section (166.9 vs. 96.2, \(P = 0.0313\)), not in the curved mid section (113.1 vs. 85.5, \(P = 0.0625\)). The average velocities were highest and their standard errors (8.5–26.5) were smallest at the distal straight section both before and after the meal challenge as compared with other sections. The streamline analysis depicted more frequent appearances of vertical or helical flow in the curved mid section both on 4D Flow and CFD (κ: 0.27–0.68).

Conclusion: SMA velocimetry with 2D cine PC-MRI was dependent on the localization of the measurement planes. Distal straight section, not in the curved mid section is recommended for MR velocimetry.

Keywords: 4D Flow, phase contrast image, superior mesenteric artery, flow measurement, post-prandial hyperemia

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**Introduction**

Flow velocity measurement of the superior mesenteric artery (SMA) to assess acute or chronic small bowel ischemia has been one of the classical clinical topics in abdominal radiology.\(^1\)\(^{-4}\) Doppler ultrasonography and 2D cine phase contrast (PC)-MRI are established techniques that have traditionally been utilized with or without meal challenge to measure blood flow velocity.\(^3\)\(^{-4}\) However, there have been few investigations concerning the optimal plane selection for flow velocity measurements. We hypothesized that the measurement of blood flow velocity may be dependent on plane localization, because abnormal blood flow dynamics can be generated by abruptly curved pathways of the SMA.

To visually access the flow dynamics, streamline analysis based on 4D Flow in vivo and computational fluid dynamics (CFD) in silico are two best methods currently available. For in vivo study, we used one of the cutting edge sequences innovated for 4D Flow. Three dimensional cardiac phase resolved phase contrast MRI techniques (4D Flow) utilizing 3D cine phase contrast imaging with vastly undersampled isotropic projection imaging (PC VIPR)\(^5\)\(^{-8}\) uses non-Cartesian 3D radial data with k-space sampling, which has shown its ability to produce high spatial and temporal resolution data of the blood flow velocity, and thereby, visualize 3D streamlines of the blood flow.\(^9\)\(^{-11}\) Recent studies have validated the ability of PC VIPR to visualize and quantify the abdominal aorta as well as visceral arteries\(^5\)\(^{-9}\) including SMA.\(^10\) Typical MRI uses Cartesian data sampling, on the other hand, PC VIPR samples data using a radial trajectory, similar appearances to a “koosh ball”. Since all the data lines includes k-space center or low frequency data components, PC VIPR provides a high level of robustness to motion and undersampling. Resultant short TE allows for avoidance of intravoxel de-phasing, thereby, beneficial for the assessment of the flow analysis. PC VIPR allows for measuring not only time-resolved velocity data during a cardiac cycle, but high definition 3D MRA when maximum intensity projection (MIP) reconstruction of the temporally averaged magnitude images are performed.\(^5\) By using this MR method, temporal and whole spatial velocity as well as the morphology data within the main trunk of SMA or its branches are acquired. For in silico validations, we used CFD as well.

With an aid of 4D Flow and CFD, the purpose of this Institutional Review Board (IRB) approved prospective and single arm study is to test whether flow velocimetry in the SMA with combined use of 2D cine PC and meal challenge is dependent on the localizations of the measurement planes and to seek optimal section for velocimetry.

**Materials and Methods**

**Volunteers**

The IRB approved study (#25–40) was performed enrolling seven healthy male volunteers with age ranging from 23 to 53 years. Written informed consent was provided by each volunteers.

**Meal challenge**

After 5 h of fasting, MR images were acquired before and 30 min after the intake of 400 kcal of a dietary supplement (Calorie Mate, Otsuka Pharm Co., Ltd., Japan). The contents of the supplement are as follows; protein: 8 g, fat: 21.9 g, sugars: 41.7 g, dietary fiber: 2 g, sodium: 280 mg, potassium: 110 mg, calcium: 200 mg, iron: 2.5 mg, magnesium: 50 mg, phosphorus: 80 mg, vitamin A: 225 μg, vitamin B1: 0.5 mg, vitamin B2: 0.6 mg, vitamin B6: 0.5 mg, vitamin B12: 1 μg, niacin: 5.5 mg, pantothenic acid: 2.8 mg, folic acid: 100 μg, vitamin C: 40 mg, vitamin D: 2.5 μg, vitamin E: 4 mg. The supplement was provided with 100 mL of mineral water (hardness of 24) in the sitting position on the MR cradle.

**MR scanner**

All studies were conducted on a 3T MR scanner (Discovery MR750, General Electric, Waukesha, WI, USA) with a 12-channel phased array coil.

**2D cine PC MRI**

ECG gated 2D cine PC MRI with velocity encoding (VENC) of 200 cm/s perpendicular to the measurement plane was performed to measure the flow velocities of three sections of the main trunk of the SMA pre- and 30 min post-meal challenge, and also to determine the VENC for 4D Flow. First, the transaxial plane was placed traversing the abdominal aorta above the branch of the celiac axis (CA). The parameters used were as follows: TR (ms)/TE (ms)/FA (°)/Number of excitations: 5.4/2.8/20/1, RBW (kHz): ±31.25, FOV (cm): 30, thickness (mm): 4, matrix: 160 × 160, R–R phase resolution: 30. The same parameters were used for the other measurement planes using 2D cine PC. Volunteers were requested to keep quiet and regular respiratory cycles while the imaging. The flow profile was assessed immediately to rule out aliasing, and to separate systole and diastole, and the maximum flow velocity was measured. Referring to the slice line on the scout view of the 2D cine PC MRI, identical measurement sections have been placed on the SMA before and after the meal challenge. Three series of 2D cine PC MRI were then performed in 1–2 cm intervals by placing the measurement planes perpendicular to the SMA at the proximal segment closest to its orifice, curved mid segment and the distal straight section of the main trunk of SMA (Fig. 1). Same parameters were used for each acquisitions.

**4D Flow MRI**

After 2D cine PC MRI, 4D Flow using PC VIPR was performed using the following parameters; axial slab orientation, FOV (cm): 32 × 32 × 80, slice thickness: 1.25 mm, locs/slab: 8, TR (ms)/TE (ms)/(FA °): 7.0–7.1/3.3–3.4/8, matrix: 256 × 256, RBW (kHz): ±62.5, acquisition time (min): 9:29–9:36, retrospective ECG gating, prospective respiratory gated
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Fig. 1 Three measurement planes were set perpendicular to the superior mesenteric artery (SMA). The scheme shows Pro: proximal section, CurvMid: curved mid section, DistStra: distal straight section on the MR angiography of the upper abdomen.

(adaptive acceptance window: 50%), projection number: 10000, and VENC (cm/s): 100–200 (The optimum VENC for 4D Flow was determined individually based on the +20% over the maximum velocities that was measured with 2D cine PC MRI at the abdominal aorta above CA.). 4D Flow was also performed pre- and post-meal challenge. Twenty cardiac phase resolved images were obtained for all cases. Temporally averaged magnitude signals were post-processed with maximum intensity projection algorithm and the MRA was used to verify the morphologies of the visceral branches.

Flow velocity measurements with 2D cine PC
Using dedicated flow analysis software (FLOVA II, R'Tech, Hamamatsu, Japan), velocity measurements were performed by two diagnostic radiologists (MS and WY). They measured cardiac phase resolved flow velocities by placing the largest possible region of interest (ROI) (26–40 pixels) on each cross-section of the SMA at (1) proximal section of the SMA, (2) curved mid section of the SMA and (3) distal straight section before branching the first jejunal branch, and then, time velocity curves were automatically drawn. The ROIs were copied from the systolic cardiac phase of the image, and pasted on the other cardiac phase resolved sections on the SMA, and then misregistrations were manually corrected. The acquired values were then averaged. Same procedures were repeated before and after the meal.

Depictions of the streamline patterns with 4D Flow datasets
Using FLOVA II, the boundary of the vessel was segmented using the reconstruction planes of temporary averaged magnitude signals derived from the PC VIPR. Somewhat larger boundaries were selected by setting lower thresholds and/or region growing method in order not to omit any velocity vectors inside the SMA. The 3D flow velocity vectors were interpolated using a spatial resolution of 1.25 × 1.25 × 1.25 mm³ from the three-dimensional data sets. Streamlines were then generated using the Runge–Kutta method.¹¹

Depictions of the streamline patterns using computational fluid dynamics
Streamlines were also depicted with computer simulations. CFD simulations were performed using a general purpose solver that is capable of performing thermal fluid simulations (Ansys CFX ver. 14.5, Ansys, PA, USA). The simulation was performed by solving the Navier–Stokes equations, which is a set of equations describing the motion of fluids in a finite volume. The simulation was based on several assumptions as follows: rigid vessel walls, an incompressible Newtonian fluid, and zero flow velocity at the vessel wall. Viscosity and density values were set to the standard blood values (3.8e⁻³ kg/m * s, 1054.0 kg/m³). 3D MR angiography was reconstructed with maximum intensity projection algorithm using temporally averaged magnitude images, which was exported by FLOVA II in STereo Lithography (STL) format. The STL data was then verified and moderated by 3D modeling software (MIMICS 3-Matic, Materialize, Leuven, Belgium). Time resolved inflow volume data measured at the abdominal aorta above the CA was used for the boundary conditions that was determined from the 4D Flow measurement. For the outflow condition, the pressure at the main trunk of the SMA as well as each measurable branches of the SMA, bilateral renal arteries, the main trunk of the CA, and the aortic section below the visceral arteries were set to null. The rationale for the delineations of the streamline using CFX is elaborated elsewhere.¹² The seeding points was 25.

3D flow dynamics assessment
Two diagnostic radiologists (MS and WY) with more than 5 years of experience each in flow analysis assessed the flow patterns over the cardiac cycle within the SMA based on the streamline post-processed by PC VIPR as well as the simulation data derived from the CFD. Viewing the cine loops of the streamlines from arbitrary directions on a monitor display of the workstation, abnormal flow patterns were visually rated on three point scale as 0: mostly laminar (parabolic or Wormersley) flow, 1: mild vortex and/or helical flow and 2: severe vortex and/or helical flow. Systole and diastole were rated separately (Fig. 2a–2d).

Statistics
The increase of blood flow velocity at each plane measured with 2D cine PC MRI was statistically analyzed. Temporally averaged flow velocity was calculated for each section before and after meal challenge. The increase in temporally averaged flow velocity at each plane was statistically analyzed by Wilcoxon test of paired samples for its significance. Cardiac phase resolved flow velocities pre- and post-meal were also compared and the results were analyzed by Wilcoxon test.
Inter-observer and inter-modality agreements in flow pattern analysis was assessed with linear weighted kappa statistics. κ-values below 0.2 were interpreted as poor, 0.21–0.40 as fair, 0.41–0.60 as moderate, 0.61–0.8 as good, and 0.81–1.0 as very good agreements. Statistical application (MedCalc Statistical Software version 14.10.2, MedCalc Software bvba, Ostend, Belgium) was used for the analysis. P-value <0.05 was considered significant.

Results

All the MR examinations were successfully performed. No obstruction or deformity were found within the SMA in six out of seven volunteers by MR angiography. One volunteer had replaced right hepatic artery (rRHA) as a normal vascular variant. Cardiac rate reflected by R–R interval did not change significantly between pre- and post-meal challenge, which were pre: 1.176 (CI: 1.051–1.182) vs. post: 1.111 (CI: 1.004–1.116), P = 0.125.

The velocity data measured by 2D cine PC MRI showed an increase in blood flow velocity after the meal challenge (Fig. 3); however, significant increase of temporally averaged velocities was observed only at the proximal (97.8 ± 23.5 mm/s vs. 129.3 ± 35.6 mm/s; P = 0.0313) and the distal section (96.2 ± 8.5 mm/s vs. 166.9 ± 20.5 mm/s; P = 0.0313). Significant increase of the velocity during diastole was observed only at distal straight section (20.3 ± 5.0 mm/s vs. 64.7 ± 14.1; P = 0.0313) (Table 1). The velocity value after meal was highest at the distal straight section irrespective of cardiac phase.

The streamline delineated mostly laminar, undisturbed antegrade flow at the distal straight segment of the SMA; however, disturbed (i.e., vortex and/or helical) flow was dominant at the curved mid section. At the proximal section
Table 1 2D cine PC MRI measured increased temporally averaged flow velocity after the meal challenge only in the proximal and distal section, not in the curved mid section. The average velocities were highest and their standard errors were smallest at the distal straight section both before and after the meal challenge as compared to other sections irrespective of cardiac phase.

|                  | AUC (mm/s × 1 cardiac cycle) (median ± SE) | P     |
|------------------|------------------------------------------|-------|
|                  | Before meal | CI | After meal | CI |       |
| Whole cardiac cycle | Proximal | 97.8 ± 23.5 | 27.7–171.8 | 129.3 ± 35.6 | 70.4–270.8 | 0.0313* |
|                  | Mid | 85.5 ± 38.6 | 56.1–272.1 | 113.1 ± 36.9 | 92.8–296.2 | 0.0625 |
|                  | Distal | 96.2 ± 8.5 | 68.1–115.9 | 166.9 ± 26.5 | 76.2–234.0 | 0.0313* |
| Systole | Proximal | 76.6 ± 11.7 | 48.0–119.1 | 92.0 ± 18.9 | 70.9–177.1 | 0.0625 |
|                  | Mid | 66.9 ± 23.9 | 51.9–183.3 | 78.9 ± 25.1 | 66.7–202.2 | 0.0625 |
|                  | Distal | 73.3 ± 4.2 | 62.5–87.9 | 101.6 ± 14.3 | 62.4–148.7 | 0.0625 |
| Diastole | Proximal | 21.2 ± 12.0 | 20.3–52.7 | 37.3 ± 17.0 | -0.5–93.9 | 0.0625 |
|                  | Mid | 18.6 ± 14.8 | 4.2–88.8 | 34.2 ± 12.1 | 26.1–94.0 | 0.0938 |
|                  | Distal | 20.3 ± 5.0 | 5.6–32.2 | 64.7 ± 14.1 | 13.8–101.8 | 0.0313* |

*Significant increase of flow velocity (P < 0.05). AUC, area under the curve.
of the SMA, partially turbulent or helical flow patterns that were not parallel to the vessel were also delineated (Fig. 4a and 4b). The CFD also delineated disturbed vortex and/or helical flow patterns within the curved mid section, and helical flow at the proximal section (Fig. 5a and 5b). The streamline analysis based on CFD simulation *in silico* validated the result based on PC VIPR measurement *in vivo* by showing the same trend in blood flow patterns (Fig. 6).

The inter-observer agreement concerning the assessments of the SMA flow patterns indicated that the visual evaluation of the streamlines was moderate to good agreements for both 4D Flow (systole; \( k = 0.43 \), diastole; \( k = 0.56 \)) and CFD simulation (systole; \( k = 0.42 \), diastole; \( k = 0.68 \)).

The inter-modarity agreement concerning the assessments of the SMA flow patterns indicated that the visual evaluation of the streamlines was also fair to good agreements; systole: \( k = 0.44–0.67 \) (moderate to good), diastole: \( k = 0.27–0.29 \) (fair).

**Discussion**

Phase contrast velocimetry is an established technique; however, many concerns are advocated when it comes to the flowmetry in non-laminar flow. O’Brien et al. found that phase contrast MR flowmetry is robust in non-stenosed flow path; however, it suffered from more velocity errors in the
stenotic flow path. Whereas, the errors reduced with shorter TEs. They speculated that this type of errors are based on the intravoxel phase dispersions, which is reflected by “Salt and pepper artifacts” on the phase images.

Postprandial blood flow increase in the SMA seen on phase contrast MRI is a widely recognized physiological reaction. According to a meta-analysis, significant increase of blood flow ranging from 58% to 250% has been reported. Wide range of the values is probably due to the amount of meal and measured timing. In the current study, MR velocimetry measured increased phase averaged post-prandial flow velocity at the proximal section and the distal straight section; however, failed to show statistical increase at the curved mid section of the main trunk of the SMA. The standard errors of the measured velocity values at the curved mid section were consistently larger than those in proximal or distal straight section (Table 1). We presume this was a failure of velocimetry at the section where rotational flow was dominant due to the sharp turn of the SMA. Those disturbed flow patterns seen at the proximal and curved mid section were considered to have prevented accurate measurements of the flow velocity. Visceral arteries such as SMA typically form abrupt or sharp curve and/or off-branching variant vessels in their pathway. In such vessels, the hemodynamics are not as simple as parabolic or Wormersley flow that is usually seen in the straight flow path. The blood flow within the curved vessels may generate disturbed flow such as vortex and/or helical flow. As a result, fluctuated flow velocities create larger standard errors in the measured data, and thereby, obscure significant post-prandial hyperemia as temporally averaged flow velocities.

This type of velocimetry failure due to complex flow is even more prevalent in the pathological status. When the orifice of visceral arteries are narrowed, the time velocity curves measured are sometimes oscillatory. This phenomenon is sometimes seen at the post-stenotic portions of the SMA of the patients suffering from chronic intestinal ischemia, or at the narrowed renal artery of the patients with renovascular hypertension. The reasons for these oscillations or fluctuations have never been explained reasonably. We presume this was created by the jet flow through the stenotic channel and subsequent vortex and/or helical flow in the dilated post-stenotic portion.

4D Flow datasets with an aid of streamline analysis visually indicated that the vortex and/or helical flow were dominant at the curved mid section during systole, which were not at the distal straight section where the stream lines were mostly laminar with partial helical flow. During diastole, helical or disturbed flow were observed in all three sections; however, they were more prominent at the curved mid section. Besides the complex non-laminar disturbed flow in the curved mid section, slightly turbulent flow pattern that was not parallel to the vessel was also delineated at the proximal section of the SMA. This was probably due to the sharp rise of the branching angle of the SMA from the abdominal aorta that might have generated in and out flow caused by contiguous high velocity flow of the abdominal aorta striking the branching orifice of the SMA. Abnormal flow patterns at the proximal section was also depicted in the SMA of the volunteer with rRHA. In the SMA with rRHA, moderately vortex and/or helical flow was observed at the proximal and the curved mid section during systole. In the previously published textbook for quantification of blood flow, SMA orifice has been recommended for the measurement plane, because the plane was supposed to be less affected by respiratory motion of the SMA. However, the results of our hemodynamic analysis implicate that distal straight section is more appropriate than proximal section for correct velocimetry.

Extending the results of our study, not limited to SMA, as a general rule, avoidance of disturbed flow is required for the accurate measurement of the blood flow velocity of the visceral arteries. Recruiting 43 children with aortic
regurgitation, Iwamoto et al. found that the large variations in PC measurement for the systolic forward, diastolic retrograde and net forward flow volumes at different levels of the aortic valve and ascending aorta. They therefore concluded that the PC measurement for aortic regurgitation volume and fraction is inaccurate and should be abandoned. 16 We presume that streamline analysis using 4D Flow data before measurement plane selection may be a beneficial solution for more accurate flowmetry in the complexed flow.

It might sound weird to separately see streamline on 4D Flow and measure flow velocities with 2D cine PC; however, the latter is currently more precise in velocimetry, because of its robustness in measuring accelerated flow velocities, and in better cardiac phase resolutions. We deem the difference is based on the lengthy three directional velocity encoding in 4D Flow vs. brief one velocity encoding perpendicular to the plane for 2D cine PC MRI. In near future; however, visual flow delineations and velocimetry can be checked together only with 4D Flow when the precision of its velocimetry is further improved.

In this study, CFD was used for a validation of the flow patterns measured within the SMA. CFD reproduced disturbed flow most frequently in the curved mid section. Therefore, our hypothesis that there are optimal measurement planes in the SMA has also been supported in silico. Obviously, curved mid section should be avoided for the measurement.

We have dealt with flow velocity instead of flow volume, because we wished to eliminate the uncertainty of the variations between areas of the measurement planes. However, as long as average velocities were measured, the average flow velocities and flow volumes are equally affected by the variations of the areas of the measurement sections. To get rid of as much uncertainty as possible of the varying areas mainly due to the pulsation of the SMA, we have adjusted the ROIs of each cardiac phase manually.

The limitations of this study are (1) single center study using a relatively small sample size and (2) CFD was performed based on the assumption that the vessel wall was rigid; however, in reality, the vessel wall is elastic with a certain level of compliance.

**Conclusion**

In conclusion, for accurate measurement of blood flow velocity within the SMA, distal straight section should be selected. Curved mid section should be avoided for non-laminar flow. Flow depictions using 4D Flow is helpful in selecting optimal measurement planes for the blood flow velocimetry by depicting streamlines within the SMA.

**Conflicts of Interest**

The corresponding author Yasuo Takehara is an endowed chair of Nagoya University supported by a private company; however, the status is irrelevant to the contents of the paper.

Coauthors Tetsuya Wakayama, Atsushi Nozaki and Hiroyuki Kabasawa are employees of GE Healthcare.

Other coauthors have nothing to disclose related to the current study.

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