Quantitative perfusion-CMR is significantly influenced by the placement of the arterial input function

Ibnul Mia1,3 · Melanie Le1 · Christophe Arendt1 · Diana Brand1 · Sina Bremekamp1 · Tommaso D’Angelo2 · Valentina O. Puntmann1 · Eike Nagel1,3

Received: 6 July 2020 / Accepted: 26 September 2020 / Published online: 12 October 2020 © The Author(s) 2020

Abstract
The aim of this study is to provide a systematic assessment of the influence of the position on the arterial input function (AIF) for perfusion quantification. In 39 patients with a wide range of left ventricular function the AIF was determined using a diluted contrast bolus of a cardiac magnetic resonance imaging in three left ventricular levels (basal, mid, apex) as well as aortic sinus (AoS). Time to peak signal intensities, baseline corrected peak signal intensity and upslopes were determined and compared to those obtained in the AoS. The error induced by sampling the AIF in a position different to the AoS was determined by Fermi deconvolution. The time to peak signal intensity was strongly correlated ($r^2 > 0.9$) for all positions with a systematic earlier arrival in the basal ($−2153 ± 818$ ms), the mid ($−1429 ± 928$ ms) and the apical slice ($−450 ± 739$ ms) relative to the AoS (all $p < 0.001$). Peak signal intensity as well as upslopes were strongly correlated ($r^2 > 0.9$ for both) for all positions with a systematic overestimation in all positions relative to the AoS (all $p < 0.001$ and all $p < 0.05$). Differences between the positions were more pronounced for patients with reduced ejection fraction. The error of averaged MBF quantification was 8%, 13% and 27% for the base, mid and apex. The location of the AIF significantly influences core parameters for perfusion quantification with a systematic and ejection fraction dependent error. Full quantification should be based on obtaining the AIF as close as possible to the myocardium to minimize these errors.

Keywords Cardiovascular magnetic resonance · Myocardial perfusion · Coronary artery disease · Ejection fraction · Heart failure · Signal intensity · Aortic input function

Abbreviations
3Ch 3-Chamber
AIF Arterial input function
AoS Aortic sinus
CAD Coronary artery disease
CMR Cardiovascular magnetic resonance
LVEF Left ventricular ejection fraction
LV Left ventricle
ROI Region of interest
MBF Myocardial blood flow

Eike Nagel
eike.nagel@cardiac-imaging.org

1 Institute for Experimental and Translational Cardiovascular Imaging, University Hospital Frankfurt, Theodor-Stern Kai 7, 60590 Frankfurt am Main, Germany
2 Department of Biomedical Sciences and Morphological and Functional Imaging, G. Martino University Hospital Messina, Via Consolare Valeria 1, 98100 Messina, Italy
3 Institute for Experimental and Translational Cardiovascular Imaging, University Hospital, Goethe University, Frankfurt am Main, Germany
Introduction

Perfusion cardiovascular magnetic resonance (perfusion-CMR) is an established test for assessing the presence and severity of myocardial ischemia with a class IA indication in the most recent European Guidelines [1]. It has shown to be strongly correlated with fractional flow reserve [2, 3] and has recently been shown to allow for a safe noninvasive guidance of patients with moderate to severe pretest likelihood for coronary artery disease (CAD) which is noninferior to invasive angiography supported by fractional flow reserve [4]. Increasingly fully quantitative methods are employed to determine myocardial blood flow (MBF) and flow reserve [5–8]. Perfusion quantification requires correction for the speed and amount of contrast agent arrival to the coronary arteries with the so called “arterial input function” (AIF). Since the AIF cannot be measured in the coronary arteries with CMR, the use of the same imaging plane as the perfusion measurement itself (e.g. basal short axis view, mid short axis view or apical short axis view) or the use of the basal short axis plane for all other slices has become the accepted standard [9]. As the contrast agent bolus disperses during its passage through the circulation some influence of the location of the AIF on the final results is to be expected.

However, the influence of the location on the AIF has not been systematically assessed. We determined the presence and extent of error induced by positioning the AIF in the basal, mid and apical level. These positions are routinely used for the position of the short axis views for perfusion imaging in our Institute. Care was taken to avoid any papillary muscles. A fourth ROI was placed in the aortic sinus (AoS) at the level of the coronary arteries (Fig. 1). After baseline correction, signal intensity time curves were constructed for each ROI generating four AIFs. Time to peak was determined as the time between arrival of the contrast agent (signal increase above 2 SD of the contrast free images) and the time of the peak signal intensity. The change of signal from baseline to peak signal intensity was determined and noted as peak signal intensity. Upslope was defined as the highest upslope of three consecutive heartbeats of the AIF. For quantitative analysis Fermi deconvolution was performed with R studio 1.2.5001 using a least square fit based on the Levenberg-Marquardt algorithm.

Differences versus the AIF in the AoS were then calculated in [ms] for time to peak and in [%] for peak signal intensity and the upslope.

To calculate the error induced by the different locations, four average AIFs of all patients were calculated after baseline correction and normalization for time to peak and peak signal intensity using the data obtained from the four sampling positions. A single myocardial response function was created from a patient with an ejection fraction closest to the mean of all patients. The difference in MBF based on the AIF obtained in the three ventricular positions versus the AIF obtained in the AoS was calculated and provided in [%].

Ejection fraction was measured using automatically generated contours with final user correction and approval (SuiteHeart, NeoSoft, Pewaukee, Wi, USA). Patients were categorized into preserved ejection fraction (EF > 50%) or reduced ejection fraction (EF ≤ 50%).

Methods

In 39 patients with suspected nonischemic cardiomyopathy and a wide range of ejection fractions a weight adapted contrast agent bolus (0.00375 mmol/kg body weight in 95% saline, Gadovist R, Bayer AG, Leverkusen, Germany) was injected with a flow rate of 3 ml/sec followed by a 20 ml saline chaser (Accutron MR, Medtron, Saarbrücken, Germany) (adapted from [10]). During the first pass of the contrast agent a 3-chamber view was obtained in mid-diastole of every heartbeat using the identical sequence usually used for stress-perfusion imaging in our Institute (Echo-Time: 1.11-2.0 ms, Repetition-Time: 3.5 ms, Flip Angle: 35–50°, typical acquired spatial resolution 2 × 2.5 × 8 mm, 3 T, 90° saturation prepulse, with 100 ms prepulse delay) with the patients holding their breath in expiration. Two proton density images per slice were acquired at the beginning of each acquisition for estimating the coil sensitivity profile [11].

All patients were scanned at 3 T (Skyra, Siemens AG, Erlangen, Germany) using 32-channel surface coils. All patients gave written informed consent and the study was approved by the local ethics committee (T1 Mapping Registry CMR Studie, Business Nr.: 1/16 and Decipher HFpEF, Business Nr.: 273/17).

Image analysis

Using MEDIS Suite 3.2 (Medis, Leiden, The Netherlands), three regions of interest (ROI) were placed in each dynamic of the first pass perfusion 3-chamber images at a basal, mid and apical level. These levels are defined as 25%, 50% and 75% of the endsystolic distance from the mid of the mitral valve to the tip of the apical blood pool of the LV blood [12]. These positions are routinely used for the position of the short axis views for perfusion imaging in our Institute. Care was taken to avoid any papillary muscles. A fourth ROI was placed in the aortic sinus (AoS) at the level of the coronary arteries (Fig. 1). After baseline correction, signal intensity time curves were constructed for each ROI generating four AIFs. Time to peak was determined as the time between arrival of the contrast agent (signal increase above 2 SD of the contrast free images) and the time of the peak signal intensity. The change of signal from baseline to peak signal intensity was determined and noted as peak signal intensity. Upslope was defined as the highest upslope of three consecutive heartbeats of the AIF. For quantitative analysis Fermi deconvolution was performed with R studio 1.2.5001 using a least square fit based on the Levenberg-Marquardt algorithm.

Differences versus the AIF in the AoS were then calculated in [ms] for time to peak and in [%] for peak signal intensity and the upslope.

To calculate the error induced by the different locations, four average AIFs of all patients were calculated after baseline correction and normalization for time to peak and peak signal intensity using the data obtained from the four sampling positions. A single myocardial response function was created from a patient with an ejection fraction closest to the mean of all patients. The difference in MBF based on the AIF obtained in the three ventricular positions versus the AIF obtained in the AoS was calculated and provided in [%].

Ejection fraction was measured using automatically generated contours with final user correction and approval (SuiteHeart, NeoSoft, Pewaukee, Wi, USA). Patients were categorized into preserved ejection fraction (EF > 50%) or reduced ejection fraction (EF ≤ 50%).

Statistical analysis

The following statistical tests were used as appropriate: r ANCOVA for repeated measurements with F-test for
The International Journal of Cardiovascular Imaging (2021) 37:1023–1031

significance, unpaired Welch two sample t-test for differences between group categories, linear regression with correlation coefficient for comparing two continuous variables (Pearson’s method for normally distributed data and Spearman’s method for not normally distributed data as determined by Shapiro-Wilk normality test). All tests were 2-tailed. Statistical significance was reached at p < 0.05. Statistical analysis was performed with R studio (Version 1.2.5001).

Results

Patient characteristics are shown in Table 1.

Strong correlations were observed between the respective parameters (peak signal intensity, time to peak, upslope) in the AoS, and the three left ventricular positions ($r^2$ between 0.91 and 0.99, $p<0.01$ for all, Fig. 2a–c).

Highly significant differences between the AoS and the three other locations were observed for each parameter (all $p<0.001$, except for the upslope of the midslice versus AoS with $p<0.05$; Table 2).

The largest differences to the AoS were found for the time to peak at the base, for the peak signal intensity at the apex and for the upslopes at the base (Table 2).

All differences were significantly more pronounced in patients with reduced LVEF in comparison to patients with preserved LVEF (Fig. 3a–c).

The differences for the shape of the input function resulted in errors of 8%, 13% and 27% for the base, the mid slice and the apex in comparison to the AoS.

Discussion

Our data shows, that the position of the region of interest used to determine the arterial input function (AIF) for perfusion quantification has significant effects on its shape. Some of these differences are systematic, however, there is also
large random error. Most importantly, the magnitude of the error correlates negatively with the ejection fraction resulting in larger errors in patients with reduced ejection fraction. This is of specific importance, as one of the core reasons to use the AIF is to control for differences of contrast agent delivery in patients with reduced ejection fraction.

For two decades myocardial perfusion-CMR studies were mainly reported visually by assessing the speed and amount of signal increase within the myocardium during the first arrival of the contrast agent bolus during maximal vasodilation. Until recently, most attempts to quantify MBF based on perfusion-CMR were restricted to academic research sites and the effort for full quantification, in combination with a large amount of assumptions, was tedious. If quantification was performed, frequently a semiquantitative approach based on differences of specific parameters (e.g. upslope)
between stress and rest was used. This has the advantage to cancel out some of the systematic effects. In addition, local normal ranges or a comparison between segments of a given patient were applied. These approaches, however, have the disadvantage to not allow provision of absolute ranges of normal with objective and transferable cut-off values, require stress and rest imaging and are limited in their ability to address changes, e.g. with therapy. More recently fully quantitative methods have become readily available and started to be used in clinical practice. However, when providing full quantification of MBF in ml/g/min myocardium any error will be misleading, as the claim for an absolute number inherently infers, that all assumptions in the processing chain of the parameter are sufficiently controlled.

The role of the position of the input function has been analyzed previously, mainly for cardiac positron emission tomography where a similar dependency of MBF on the position of the input function was found [13, 14]. Its importance for perfusion CMR has been discussed [15], but not systematically analysed.

The obtained data in the current report provide important knowledge:

First, the position of the AIF creates a systematic error. This error could be addressed by either using normal values for the specific pathway used to obtain the results, by applying correction factors or potentially, by calculating a myocardial perfusion reserve based on the quantitative data, rather than providing MBF itself. The excellent
correlation between the various locations for each parameter demonstrate, that the systematic effect is strong and can be accounted for. Importantly, the largest systematic difference was found for time to peak, which is least important for full quantification.

Second, the position of the AIF creates a random error demonstrated by the large confidence intervals in the Bland–Altman analyses. While it may be possible to reduce these effects by better fitting algorithms and

Table 2 Differences relative to the Aortic sinus

|                     | Base         | p value | Mid          | p value | Apex          | p value |
|---------------------|--------------|---------|--------------|---------|---------------|---------|
| Delta time to peak  | –2153 ± 818  | <0.001  | –1429 ± 928  | <0.001  | –477 ± 741    | <0.001  |
| (dTTP) (ms)         |              |         |              |         |               |         |
| Delta peak signal   | 21.8 ± 14.0  | <0.001  | 40.4 ± 35.3  | <0.001  | 52.1 ± 27.0   | <0.001  |
| intensities (dpSI)  |              |         |              |         |               |         |
| (%)                 | 11.6 ± 16.1  | <0.001  | 3.0 ± 10.75  | <0.05   | 6.91 ± 9.17   | <0.001  |
| Delta upslope (%)   |              |         |              |         |               |         |
|                     | 8            | 13      | 27           |         |               |         |
| Delta flow (%)      |              |         |              |         |               |         |

Fig. 2 (continued)
modern machine learning optimization, they highlight the need to minimize this error during data acquisition.

Third, the error created by different positions of the AIF varies with important core parameters, such as ejection fraction. Again, it may be possible to partially account for this by taking the ejection fraction or other parameters of cardiac function into account, this observation is especially disturbing, as one of the core functions of the AIF was to correct for physiological differences, such as cardiac function. Given that normal values for myocardial perfusion are mainly based on healthy controls which also have normal ejection fraction, this error may cause
a misunderstanding of underlying pathophysiologies, or systematic misclassification of the presence and extent of myocardial ischemia in specific subgroups of patients.

In order to determine the best AIF, it is necessary to consider and analyze the systematic errors mentioned above and the limitations of the current study mentioned below. Diseases that affect the ejection fraction or cardiac blood flow may have a strong impact on the AIF.

A first opportunity to generate further data in the future is to examine stress images (taking the aspects mentioned above into account) and compare them with the data obtained here. After that a first concept for the standardization of CMR perfusion studies can be proposed.

Limitations

Several limitations apply. The study was meant to highlight the need to standardize the location of the AIF and obtain it in the aortic sinus whenever possible. As such, we did not address other potential confounders, such as valvular heart disease, arrhythmia, or aortic pathologies. It can be speculated, that even larger errors may be observed in these patient groups. We did not perform stress, calculate diagnostic accuracy, or perform full quantification of MBF. Extending our observations to these topics would add data, but not broaden the core results underlined by strong significant differences. Furthermore, gender-specific differences, e.g. regarding the ejection fraction, were not investigated. Our study did not provide any clear signs for this. Investigating this would be relevant from our point of view, but would go beyond the scope of this scientific work. In addition, due to the small number of patients, an adequate statement to that topic could not be made here.

Conclusions

The location of the AIF significantly influences core parameters for perfusion quantification such as peak signal intensity, time to peak SI and upslope. The placement of the AIF creates systematic errors, random errors and ejection fraction dependent errors. Full quantification should therefore be based on obtaining the AIF as close as possible to the myocardium to minimize these errors.

Acknowledgements We acknowledge the support of cardiac radiographers as well as the supporting staff at Goethe CVI Frankfurt for their contribution to clinical data collection.

Author contributions All authors have made relevant contributions to this manuscript as outlined by the Committee of Medical Journal Editors and have read and approved the final manuscript. Substantial contributions to the conception of the work or acquisition, analysis or interpretation of data for the work: IM, TD, VP, EN. EN was a major contributor in writing the manuscript.

Funding Open Access funding enabled and organized by Projekt DEAL. EN and VP receive funding from the German Centre for Cardiovascular Research (DZHK) Rhein/Main as well as from the CardioPulmonary Institute (CPI, EXC 2026, Project ID: 390649896).

Compliance with ethical standards

Conflict of interest The authors declare that they have no competing interests.

Ethical approval and consent to participate The study protocol was reviewed and approved by the respective institutional ethics committees and written informed consent was obtained from all participants. All procedures were carried out in accordance with the Declaration of Helsinki (2013). Ethics Committee Goethe University Frankfurt: Ethics number 1/16 and 273/17.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References

1. Knuuti J, Wijns W, Saraste A, Capodanno D, Barbato E, Funck-Brentano C et al (2019) ESC guidelines for the diagnosis and management of chronic coronary syndromes. Eur Heart J 100(k504):106
2. Watkins S, McGeoch R, Lyne J, Steedman T, Good R, McLaughlin M-J et al (2009) Validation of magnetic resonance myocardial perfusion imaging with fractional flow reserve for the detection of significant coronary heart disease. Circulation 120(22):2207–2213
3. Lockie T, Ishida M, Perera D, Chiribiri A, De Silva K, Kozerke S et al (2011) High-resolution magnetic resonance myocardial perfusion imaging at 3.0-Tesla to detect hemodynamically significant coronary stenoses as determined by fractional flow reserve. J Am Coll Cardiol 57(1):70–75
4. Nagel E, Greenwood JP, McCann GP, Bettencourt N, Shah AM, Hussain ST et al (2019) Magnetic resonance perfusion or fractional flow reserve in coronary disease. N Engl J Med 380(25):2418–2428
5. Morton G, Chiribiri A, Ishida M, Hussain ST, Schuster A, Indermuehle A et al (2012) Quantification of absolute myocardial perfusion in patients with coronary artery disease: comparison between cardiovascular magnetic resonance and positron emission tomography. J Am Coll Cardiol 60(16):1546–1555
6. Engblom H, Xue H, Akil S, Carlsson M, Hindorf C, Oddstig J et al (2017) Fully quantitative cardiovascular magnetic resonance myocardial perfusion ready for clinical use: a comparison between
cardiovascular magnetic resonance imaging and positron emission tomography. J Cardiovasc Magn Reson 19(1):78

7. Hsu L-Y, Groves DW, Aletras AH, Kellman P, Arai AE (2012) A quantitative pixel-wise measurement of myocardial blood flow by contrast-enhanced first-pass CMR perfusion imaging: microsphere validation in dogs and feasibility study in humans. JACC 5(2):154–166

8. Xue H, Brown LAE, Nielles-Vallespin S, Plein S, Kellman P (2020) Automatic in-line quantitative myocardial perfusion mapping: processing algorithm and implementation. Magn Reson Med 83(2):712–730

9. Schulz-Menger J, Bluemke DA, Bremerich J, Flamm SD, Fogel MA, Friedrich MG et al (2013) Standardized image interpretation and post processing in cardiovascular magnetic resonance: society for cardiovascular magnetic resonance (SCMR) board of trustees task force on standardized post processing. J Cardiovasc Magn Reson 15:1–19

10. Ishida M, Schuster A, Morton G, Chiribiri A, Hussain S, Paul M et al (2011) Development of a universal dual-bolus injection scheme for the quantitative assessment of myocardial perfusion cardiovascular magnetic resonance. J Cardiovasc Magn Reson 13(1):2812

11. Miller CA, Hsu L-Y, Ta A, Conn H, Winkler S, Arai AE (2015) Quantitative pixel-wise measurement of myocardial blood flow: the impact of surface coil-related field inhomogeneity and a comparison of methods for its correction. J Cardiovasc Magn Reson 17(1):11–19

12. Selvadurai BSN, Puntmann VO, Bluemke DA, Ferrari VA, Friedrich MG, Kramer CM et al (2017) Definition of left ventricular segments for cardiac magnetic resonance imaging. JACC 11(6):926–928

13. Otaki Y, Lassen ML, Manabe O, Eisenberg E, Gransar H, Wang F et al (2019) Short-term repeatability of myocardial blood flow using 82Rb PET/CT: the effect of arterial input function position and motion correction. J Nucl Cardiol 59:273–278

14. Vasquez AF, Johnson NP, Gould KL (2013) Variation in quantitative myocardial perfusion due to arterial input selection. JACC 6(5):559–568

15. Jerosch-Herold M (2010) Quantification of myocardial perfusion by cardiovascular magnetic resonance. J Cardiovasc Magn Reson 12(1):57

Publisher's Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.