MUGA processing: intra and interoperator variability impact using manual and automated methods

Rita Belo¹, Cláudia Alves¹, Cristiana Carvalhal¹, Sérgio Figueiredo²⁻³, Elisabete Carolino^{2,4}, Lina Vieira^{2-3,5}

- Escola Superior de Tecnologia da Saúde de Lisboa, Instituto Politécnico de Lisboa. Lisboa, Portugal. ritabelo96@gmail.com
 H&TRC Centro de Investigação em Saúde e Tecnologia. ESTeSL Escola Superior de Tecnologia da Saúde, Instituto Politécnico de Lisboa. Lisboa, Portugal.
- 3. Unidade de Ensino e Investigação em Fisiologia, Imagem Médica e Terapia, Escola Superior de Tecnologia da Saúde de Lisboa, Instituto Politécnico de Lisboa. Lisboa, Portugal.
- 4. Unidade de Ensino e Investigação em Matemática e Física, Escola Superior de Tecnologia da Saúde de Lisboa, Instituto Politécnico de Lisboa. Lisboa, Portugal.
- 5. CIMOSM, ISEL Centro de Investigação em Modelação e Optimização de Sistemas Multifuncionais. Lisboa, Portugal.

ABSTRACT: Introduction – Multigated acquisition (MUGA) scan is mainly used for the assessment of left ventricular ejection fraction (LVEF) in patients who undergo cardiotoxic chemotherapy drugs. When applying automatic (A) or manual (M) processing methods, some biases in the quantitative metrics can be obtained. The aim of this study is to evaluate the influence of A and M methods, specifically, the inter and intraoperative variability in accordance with the professional experience. Methods - A retrospective study was performed with 14 MUGA exams available in ESTeSL's Xeleris™ Functional Imaging Workstation v. 1.0628 database. Three operators (OP) with no professional experience and two with more than 10 years of experience, processed every study five times for each method, using the EF Analysis™ and the Peak Filling Rate™. To perform the multiple comparisons, the Repeated Measures ANOVA, Friedman, t-test and Wilcoxon tests were used, considering α =0.05. **Results** – Four of the OP presented statistically significant differences between methods in one or more parameters; similar values between experienced OP and between the non-experienced were observed when the A method was applied, and higher discrepancies were present for all parameters obtained by the M mode; higher LVEF, peak filling rate, and peak empying rate values were observed for the M method. Conclusion - Variability was found when comparing M and A processing methods, as well as interoperator variability associated with their level of experience. Despite that, there was a trend of less variability between the two experienced OP and in the A method.

Keywords: Equilibrium radionuclide angiography; Cardiac function; Segmentation; Left ventricular ejection fraction; Diastolic parameters

Processamento de estudos de angiografia de radionuclídeos em equilíbrio: impacto da variabilidade intra e interoperador por métodos manuais e automáticos

RESUMO: Introdução – A angiografia de radionuclídeos em equilíbrio (ARNe) é principalmente realizada para determinar a fração de ejeção do ventrículo esquerdo (FEVE) em doentes submetidos a quimioterápicos cardiotóxicos. Quando aplicados métodos de processamento automáticos (A) ou manuais (M) podem ser obtidas distorções métricas. Este estudo teve como objetivo aferir a influência dos métodos A e M e avaliar a variabilidade inter e intraoperador associada a diferentes experiências profissionais. **Métodos** – Estudo retrospetivo com 14 exames ARNe existentes na base de dados da estação de processamento Xeleris[™] *Functional Imaging Workstation* v. 1.0628 da ESTeSL. Três operadores (OP) sem experiência profissional e dois com mais de dez anos de experiência processaram cada estudo cinco vezes por cada método, recorrendo ao EF Analysis[™] e ao Peak Filling Rate[™]. As múltiplas comparações foram realizadas com os testes

ANOVA de medidas repetidas, *Friedman*, teste-t e *Wilcoxon*, considerando a=0,05. **Resultados** – Quatro dos OP apresentaram diferenças estatisticamente significativas entre métodos para um ou mais parâmetros; foram obtidos valores semelhantes entre os OP experientes e entre os não experientes quando se aplicou o método A e observaram-se maiores discrepâncias para todos os parâmetros obtidos pelo método M; obtiveram-se valores superiores de FEVE, taxas de esvaziamento e preenchimento máximas com o método M. **Conclusão** – Verificou-se variabilidade dos resultados obtidos a partir da comparação dos métodos de processamento M e A, bem como variabilidade do interoperador associada ao seu nível de experiência profissional. Contudo os dois OP experientes apresentaram menor discrepância de valores entre si e para o método A.

Palavras-chave: Angiografia de radionuclídeos em equilíbrio; Função cardíaca; Segmentação; Fração de ejeção do ventrículo esquerdo; Parâmetros diastólicos

Introduction

Multigated acquisition (MUGA), also known as equilibrium radionuclide angiography, is a nuclear medicine procedure that uses ^{99m}Technetium labelled erythrocytes to acquire cardiac images in synchronism with an electrocardiogram¹⁻⁸. It is a well-established and noninvasive method used to assess left ventricular (LV) ejection fraction (EF) and is mainly used for serial assessment of LVEF in patients who undergo cardiotoxic chemotherapy drugs since the LV dysfunction is the most common manifestation of cardiotoxicity^{1-3,5-6,8-10}.

The cardiac alterations identified are considered as late injuries and since the deterioration of the diastolic function precedes the systolic one, it is important to have tools available to detect early cardiac damages, which can be achievable by measuring the peak filling rate (PFR) and peak emptying rate (PER) using the same acquisition used to determine LVEF^{1,3,9-11}.

In order to obtain these quantitative metrics, image segmentation is an important processing step for the detection of the LV cavity to determine LVEF and consequently other physiological parameters such as PFR and PER^{11,13}. Specifically, it is essential to apply different regions of interest (ROI), based on automatic (A) and/or manual (M) segmentation¹².

To calculate the EF, three ROI need to be drawn, such as a periventricular one to cover only background (BKG) structures and two around the LV at the end of the diastole (ED) and at the end of systole (ES). Therefore, EF can be determined using the equation^{3-4,13-15}:

%EF = $\frac{(BKG \text{ corrected counts - BKG corrected ES counts})}{BKG \text{ corrected ED counts}} \times 100$

Equation 1. %LVEF calculation.

Generally, a LVEF value greater than 55% is considered normal and a drop greater than 10% is consistent with early cardiotoxicity^{10,16-17}.

In the clinical field, there are different commercial software applications, with both A and M processing approaches available. The EF Analysis[®] program is one of them and it's used for the quantification of the LV function. The A and M processing methods differ in the way that ventricular ROIs are obtained, such that they can be drawn manually by the operator (OP) or automatically defined by the program after the OP centers an elliptical ROI in the LV. In the A mode, an edge detection algorithm is used to determine the ED and ES ROI. ED and ES ROI are drawn in frames with the highest and lowest counts in the LV, respectively. The background ROI is automatically created in the segment with the lowest average count rate, regardless of the processing mode chosen. The LVEF is calculated and a time-activity curve is created^{13,15}.

The Peak Filling Rate[™] application is used as a diagnosis and prognosis tool for early detection of deterioration of LV diastolic function¹⁹, which is an early manifestation of developing coronary disease that if not treated or if the cardiotoxic drugs are not discontinued, it can evolve to systolic disfunction^{1,3,11}. This tool uses the resulting series created by the EF Analysis[™] as input and smooths the LV time-activity curve in order to create a derivative curve^{3,19}. The commonly used diastolic parameters obtained from this curve are the PFR and the PER. The PFR represents the maximum rate of filling and the PER is obtained from the systolic phase of the time-activity curve and determines the maximum rate of LV emptying^{14,20}. These parameters are more sensitive than the EF index, which only decreases when the LV function has adulterated quite significantly, which justifies their importance^{14,16,19,24}.

All these quantitative parameters obtained by MUGA processing applications require an accurate segmentation of the LV. However, when applied for LVEF estimation, some inter and intraoperator biases were obtained as well as between centers and commercial applications systems^{18,24-27}. These facts create certain inconsistencies in some of the quantitative metrics obtained⁴, which need experimental validation. This is especially important regarding the diastolic parameters like PFR and PER, the first ones to change and remit to early deterioration of ventricular function.

The main goal of this study was to evaluate the influence of A and M processing methods on MUGA studies. Specifically, we intend to evaluate the influence of inter and intraoperative variability in the determination of LVEF, PFR and PER parameters obtained by the A and M methods, in accordance with the professional experience.

Methods

A retrospective study was performed considering MUGA studies integrated into the database of the Xeleris[®] Functional

Imaging Workstation v. 1.0628 at ESTeSL. A non-probabilistic sample of 14 patients with a clinical indication for MUGA was used. The patients with a valid left anterior oblique MUGA dataset for processing were included, all the others were excluded.

In order to calculate the LVEF, data were processed using the EF Analysis[™] software and the Peak Filling Rate[™] was used to obtain the PFR and PER values. Five OP were selected to process the MUGA exams and categorized by their degree of professional experience, respectively: OP1 and OP2 with more than 10 years of experience, and OP3, OP4 and OP5 with basic knowledge of nuclear medicine and without clinical experience. Each OP processed each study five times per method, as exemplified for an experienced OP in Figure 1. The data was processed non-consecutively, and it was guaranteed that the OPs did not know the previously obtained values, as well as the information related to the patient.



Legenda: Dias = Diastolic ROI; Sys = Systolic ROI; Bkgnd = Background ROI; Limit = Limit ROI for boundaries detection.

Figure 1. Automatic ROI (A) and manual ROI (B) drawn by an experienced OP.

The LVEF, PFR and PER values were analyzed using the International Business Machine Statistical Package for the Social Sciences[®] v. 23.0 for macOS. The results were considered significant at the significance level of 5%. The Shapiro-Wilk test was performed to test the normality of the data. When comparing the multiple parameters between OP, the Repeated Measures ANOVA test was used when the normality assumption was verified (p \geq 0.05) and when not verified (p<0.05), the Friedman's test was performed. When comparing the two methods for the same OP, the t-test was used for two paired samples when the normality assumption was verified (p<0.05), the Wilcoxon test was used.

Results

When comparing the obtained LVEF values with the A method, statistically significant differences were detected between OP [Greenhouse-Geisser statistic (2.804)=5.897, p=0.003]. Of the paired multiple comparisons, the differences were between the OP1 and the OP 3 (p=0.013), 4 (p=0.020)

and 5 (p=0.006). Analogously, there were differences between the OP2 and the OP 3 (*p*=0.024), 4 (*p*=0.018) and 5 (*p*=0.004). In this case, not only the experienced OP obtained values were close to each other but they both obtained LVEF values significantly different than the ones obtained by the non-experienced OP. The experienced OP also obtained higher LVEF values as seen in Figure 2. Regarding the LVEF values obtained with the M method, there were statistically significant differences identified between OP [Greenhouse-Geisser statistic (2.460)=12.02, p=0.000]. Of the paired multiple comparisons, the differences found were between the OP1 and the observers OP 2 (p=0.001), 3 (p=0.048) and 4 (p=0.000); between the OP2 and the OP 4 (p=0.018) and 5 (p=0.033); between OP3 and OP4 (p=0.005); and between OP4 and OP5 (p=0.001). It is verified that there is a discrepancy between the entire OP in the values of the LVEF, with higher dispersion in LVEF values for this method (Figure 2).

Comparing both methods, we found statistically significant differences in the LVEF value of OP1 [t(13)=-5.538, p=0.000], OP3 [t(13)=-3.12, p=0.008] and OP5 [t(13)=-6.505, p=0.000], with higher values of LVEF obtained with the M method for all 5 OP.



Figure 2. Comparison of the values of A and M LVEF of each OP.

Concerning the PFR values obtained by the A processing, statistically significant differences were detected between OP $[X_{r}^{2}(4)=16.16]$. From Friedman's multiple comparisons, the differences obtained were between OP1 and the OP 4 (p=0.041) and 5 (p=0.04); and between OP4 and OP5 (p=0.028). It is verified that there are differences between experienced and non-experienced OP, with proximal PFR values between each group. Experienced OP with higher PFR values and with proximal values between non-experienced OP (Figure 3). For the PFR values obtained with the M processing, statistically significant differences were also detected between OP [²(4)=29.943]. From Friedman's multiple comparisons the differences obtained were between OP4 and OP3 (p=0.034); and between OP2 (p=0.023) and OP1 (p=0.001). There are variations in the PFR values between all OP regardless of the processing method used; however, the M one presented higher discrepancies (Figure 3).



Figure 3. Comparison of the values of A and M PFR of each OP.

Concerning the PFR value, differences between methods were found in OP1 (z=3.107, p=0.008), OP3 (z=-2.103, p=0.008) and OP5 (z=-3.171, p=0.013). Values obtained by M processing were also higher than the A ones.

For the A PER values, statistically significant differences were detected between OP [$_{\rm F}^{2}(4)$ =13.200]. From Friedman's multiple comparisons the differences obtained were between OP5 and the two experienced OP (p=0.008, p=0.005); and between the OP4 and the two experienced ones (p=0.028, p=0.019). As we can verify, there is a difference between experienced and non-experienced OP, besides, experienced OP obtained closer values and the non-experienced obtained values closer to each other. For the M PER values, statistically significant differences were detected between OP [$_{\rm F}^{2}(4)$ =29.943]. From the Friedman multiple comparisons, the differences obtained were between OP4 and OP5 (p=0.028) and OP1 (p=0.000); and between OP3 and OP1 (p=0.008). There is a discrepancy between the entire OP in the value of the PER, with more impact in OP4 (Figure 4).



Figure 4. Comparison of the values of A and M PER of each OP.

Concerning the PER values, differences between methods were found in OP1 (z=-2.668, p=0.002), OP2 (z=-1.977, p=0.056), OP3 (z=-2.668, p=0.035) and OP5 (z=-2.480, p=0.002), with the PER M values being higher than the A.

Discussion

Concerning the LVEF parameter, we observed similar values between experienced OP and between the non-experienced,

when the A method was applied. Regarding the M method, higher discrepancies were found between all OP. These results tend to correlate with a study conducted by Bresser *et al.*²⁶ since they also obtained M LVEF values with larger variability when compared to the A ones and higher discrepancies in the values by the less experienced OP, and <u>in that aspect</u>, our results were in agreement. On the other hand, Hains *et al.*²⁵ obtained significantly higher A LVEF values than the M ones. Nonetheless, we observed higher LVEF values when the M processing was performed by all OP. This can be related to the different ROI dimensions and geometry used, which tend to overestimate de LV edges, resulting in higher LVEF values. This should be taken into consideration since the sum of the counts in a ROI is assumed to be a proportional measure of a clinically relevant factor²⁸.

Additionally, the difficulty to keep the BKG ROI in exactly the same position may be a limitation, regardless of the method applied and the OP involved. This is important because it can be correlated with the variations obtained between OP for the M method, higher values of LVEF for the M method and consequent changes in the diastolic parameters. We realized, particularly for the operators with basic knowledge of nuclear medicine, in both processing methods, that occasionally the BKG ROI overlaid the diastolic ROI, which is not supposed to happen since it may result in overestimated values. Therefore, it is important to evaluate the final results of the processed data and check if BKG ROI is in an overlay position. Nevertheless, it is important to analyze its size as well, since the smaller the BKG area, the higher the %LVEF value is⁸, and consequently change in the diastolic parameters can introduced and possibly generate biases results.

The determination of the diastolic function parameters is more vulnerable and sensitive^{11,19,25}, which may justify the fact that most of the OP presented values significantly different between both methods for these parameters. Considering they are determined by the geometric slope of the final LVEF curve²⁸, if the LVEF value is affected, there is a chance that the PFR and PER values are artefactual, which leads us to reinforce the idea that the segmentation process is reflected in the determination of important physiological parameters.

Between methods, the OP1, OP3, and OP5 showed significant differences for all ventricular function parameters, and most discrepancies were produced by the less experienced OP. However, we did not expect to see these differences in one experienced OP. While analyzing the results from the exams processed by OP1, we saw that polygonal ROI were used to draw the limits manually so these variations can be correlated with the ROIs' shape, dimensions, and area²⁸.

More, with the exception of the OP4, which did not present any discrepancies, all other OP presented statistically significant differences between methods in one or more parameters. However, the application of different processing methods should result in different physiological measurements since the M method should only be used when the values obtained by the other are not consensual^{3,14-15}. The analyzed data results from exams processed by OP4 showed an agreement between A and M ROI, which justifies similar results between the two processing methods. Regarding the processing methods available for MUGA exams and as documented by Boudraa *et al.*¹², the A method should be used preferably if available since it is more reproducible, in comparison with the M method that presents greater interoperator variability^{15,27}, which is consistent with our results. However, in clinical practice, there are some studies where the bounding boxes created by the A mode do not correspond to the real LV edges¹⁵ and in those cases reprocessing by M mode should be performed. For example, in patients with heterogeneous ventricular contraction, it is difficult to accurately segment the ventricular cavity from other cardiac structures²⁷, thus sometimes there is the need to delineate manually the structure of interest.

In cases of cardiotoxicity studies, patients usually perform multiple MUGA scans and there is the chance the LV function values differ significantly from the previous values. So, in those cases, there is also the need to reprocess using the M mode to analyze their reliability^{3,14-15}.

Concerning this intra and inter-variability and compared with other imaging techniques, echocardiography became a routine to assess LVEF decrease. Although the equipment required being smaller, cheaper, more mobile and more available, MUGA was shown to be more reproducible than echocardiography^{4,6,29}. Additionally, with the emerge of myocardial perfusion imaging, SPECT cardiac imaging became the state of the art in nuclear cardiology²⁹. But will MUGA have a comeback? Chen et al.³⁰ compared LVEF, ED and ES volumes derived from conventional 24-frame gated planar MUGA with the same LV function parameters derived from 24-frame gated CZT SPECT MUGA and 24-frame gated reprojected at 45° CZT planar MUGA. Good overall correlations between each data were found but planar MUGA LVEF and CZT reprojected LVEF values were lower than the CZT SPECT LVEF, suggesting that reprojection of 3D CZT gated data indeed may substitute planar MUGA.

In addition, the results of our study may be influences by some factors. The small and random sample may be a limitation, despite the fact that we tried to overcome this aspect by processing each exam five times per mode. Besides, there are some influencing variables that weren't taken into consideration such as the gender and weight of the patients that can induce some attenuation in female and obese patients³. Also, the best septal angle and detector tilt can influence the final results, since these factors when optimized, may enhance the accuracy of this technique^{9,14}.

Conclusion

MUGA scans play an important role in the assessment of cardiotoxicity. On the other hand, PFR and PER values predict early cardiac damages. The evaluation of measurement errors is irrefutable in the search for a better diagnostic quality in clinical practice.

Varying levels of OP professional experience in clinical practice along with the application of different processing methods can lead to discrepancies in the values obtained by the MUGA technique. Variability was found when comparing M and A processing methods, as well as variability interoperator associated with their level of experience. Despite the overall interoperator oscillation, there was a trend of less variability between the two experienced OP for each processing method. However, the observations of any inequalities existing between OP or within operators may need further studies in the clinical field, in order to obtain a reliable impact on diagnosis and patient management, especially concerning the diastolic parameters.

References

- Cruz M, Duarte-Rodrigues J, Campelo M. Cardiotoxicidade na terapêutica com antraciclinas: estratégias de prevenção [Cardiotoxicity in anthracycline therapy: prevention strategies]. Rev Port Cardiol. 2016;35(6):359-71. Portuguese
- Adão R, de Keulenaer G, Leite-Moreira A, Brás-Silva C. Cardiotoxicidade associada à terapêutica oncológica: mecanismos fisiopatológicos e estratégias de prevenção [Cardiotoxicity associated with cancer therapy: pathophysiology and prevention strategies]. Rev Port Cardiol. 2013;32(5):395-409. Portuguese
- Hesse B, Lindhardt TB, Acampa W, Anagnostopoulos C, Ballinger J, Bax JJ, et al. EANM/ESC guidelines for radionuclide imaging of cardiac function. Eur J Nucl Med Mol Imaging. 2008;35(4):851-85.
- Foley TA, Mankad SV, Anavekar NS, Bonnichsen CR, Morris MF, Miller TD, et al. Measuring left ventricular ejection fraction: techniques and potential pitfalls. Eur Cardiol. 2012;8(2):108-14.
- Scheiner J, Sinusas A, Wittry MD, Royal HD, Machac J, Balon HR, et al. Society of Nuclear Medicine procedure guideline for gated equilibrium radionuclide ventriculography [Internet]. Society of Nuclear Medicine Procedure Guidelines Manual; 2002 [cited 2018 Jun]. Available from: <u>http://citeseerx.ist.psu.edu/viewdoc/ summary?doi=10.1.1.172.514</u>
- 6. Mitra D, Basu S. Equilibrium radionuclide angiocardiography: its usefulness in current practice and potential future applications. World J Radiol. 2012;4(10):421-30.
- Zaret B, Beller G. Clinical nuclear cardiology: state of the art and future directions. 4th ed. Mosby; 2010. ISBN 9780323085724
- 8. Sharp PF, Gemmell HG, Murray AD. Practical nuclear medicine. 3rd ed. London: Springer; 2005. ISBN 9781846280184
- 9. Wagner RH, Halama JR, Henkin RE, Dillehay GL, Sobotka PA. Errors in the determination of left ventricular functional parameters. J Nucl Med. 1989;30(11):1870-4.
- D'Amore C, Gargiulo P, Paolillo S, Pellegrino AM, Formisano T, Mariniello A, et al. Nuclear imaging in detection and monitoring of cardiotoxicity. World J Radiol. 2014;6(7):486-92.
- Reuvekamp EJ, Bulten BF, Nieuwenhuis AA, Meekes MR, de Haan AF, Tol J, et al. Does diastolic dysfunction precede systolic dysfunction in trastuzumab-induced cardiotoxicity? Assessment with multigated radionuclide angiography (MUGA). J Nucl Cardiol. 2016;23(4):824-32.
- 12. Boudraa AO, Zaidi H. Image segmentation techniques in nuclear medicine imaging. In: Zaidi H, editor. Quantitative

analysis in nuclear medicine imaging. Boston: Springer; 2006. p. 308-57.

- Faber TL, Folks RD. Computer processing methods for nuclear medicine images. J Nucl Med Technol. 1994;22(3):145-63.
- 14. Corbett JR, Akinboboye OO, Bacharach SL, Borer JS, Botvinick EH, DePuey EG, et al. ASNC imaging guidelines for nuclear cardiology procedures: equilibrium radionuclide angiocardiography. J Nucl Cardiol. 2006;13:e56-e79.
- 15. GE Medical Systems. Ejection fraction analysis operator guide. Direction 2364209-100 rev. 1. 2003. chapter 1: 1-8.
- Fair JR, Heintz PH, Telepak RJ. Evaluation of new data processing algorithms for planar gated ventriculography (MUGA). J Appl Clin Med Phys. 2009;10(3):173-9.
- Puchal-Añé R, Guirao-Marín S, Domènech-Vilardell A, Rodríguez-Gasén A, Bajén-Lázaro M, Ricart-Brulles Y, et al. Calculation of the left ventricular ejection fraction: comparison between 4 different instruments. Rev Esp Med Nuclear. 2008;27(6):418-23.
- Yang SN, Sun SS, Zhang G, Chou KT, Lo SW, Chiou YR, et al. Left ventricular ejection fraction estimation using mutual information on technetium-99m multiple-gated SPECT scans. Biomed Eng Online. 2015;14:119.
- 19. GE Medical Systems. Peak filling rate operator guide. Direction 2364209-100 rev. 1. 2003. chapter 2: 1-34.
- Carboni GP. Depressed exercise peak ejection rate detected on ambulatory radionuclide monitoring reflects end-stage cardiac inotropic reserve and predicts mortality in ischaemic cardiomyopathy. <u>Cardiol Res</u>. 2012;3(4):164-71.
- 21. Hamilton DI, Diagnostic nuclear medicine: a physics perspective. Springer; 2004. ISBN 9783662065884
- 22. Movahed A, Gnanasegaran G, Buscombe J, Hall M. Integrating cardiology for nuclear medicine physicians. Springer; 2009. ISBN 9783540786740

- Akincioglu C, Berman DS, Nishina H, Kavanagh PB, Slomka PJ, Abidov A, et al. Assessment of diastolic function using 16-frame ^{99m}Tc-sestamibi gated myocardial perfusion SPECT: normal values. J Nucl Med. 2005;46(7):1102-8.
- 24. Steyn R, Boniaszczuk J, Geldenhuys T. Comparison of estimates of left ventricular ejection fraction obtained from gated blood pool imaging, different software packages and cameras. Cardiovasc J Afr. 2014;25(2):44-9.
- 25. Hains AD, Al-Khawaja I, Hinge DA, Lahiri A, Raftery EB. Radionuclide left ventricular ejection fraction: a comparison of three methods. Br Heart J. 1987;57(3):242-6.
- 26. Bresser P, de Beer J, de Wet Y. A study investigating variability of left ventricular ejection fraction using manual and automatic processing modes in a single setting. Radiography. 2015;21(1):e41-4.
- 27. Boudraa AE, Arzi M, Sau J, Champier J, Hadj-Moussa S, Besson JE, et al. Automated detection of the left ventricular region in gated nuclear cardiac imaging. IEEE Trans Biomed Eng. 1996;43(4):430-7.
- 28. International Atomic Energy Agency. Quantitative nuclear medicine imaging: concepts, requirements and methods. Vienna: IAEA; 2014. ISBN 9789201415103
- 29. Wackers FJ. Equilibrium gated radionuclide angiocardiography: its invention, rise, and decline and... comeback? J Nucl Cardiol. 2016;23(3):362-5.
- Chen YC, Ko CL, Yen RF, Lo MF, Huang YH, Hsu PY, et al. Comparison of biventricular ejection fractions using cadmium-zinc-telluride SPECT and planar equilibrium radionuclide angiography. J Nucl Cardiol. 2016;23(3):348-61.

Conflito de interesses

Os autores declaram não ter quaisquer conflitos de interesse. Artigo submetido em 05.06.2019 e aprovado em 07.02.2020