Cardiac Magnetic Resonance and Computed Tomography in Hypertrophic Cardiomyopathy: an Update

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Abstract

Hypertrophic cardiomyopathy (HCM) is the most common genetic cardiovascular disease and represents the main cause of sudden death in young patients. Cardiac magnetic resonance (CMR) and cardiac computed tomography (CCT) are noninvasive imaging methods with high sensitivity and specificity, useful for the establishment of diagnosis and prognosis of HCM, and for the screening of patients with subclinical phenotypes. The improvement of image analysis by CMR and CCT offers the potential to promote interventions aiming at stopping the natural course of the disease. This study aims to describe the role of CMR and CCT in the diagnosis and prognosis of HCM, and how these methods can be used in the management of these patients.

Introduction

Hypertrophic cardiomyopathy (HCM) is an autosomal dominant disorder that affects 1 in every 500 people.¹⁻³ It is characterized by ventricular hypertrophy with preserved systolic function, in the absence of other conditions that may cause such changes.⁴⁻⁶ The development of HCM is determined by mutations in genes that codify sarcomeric proteins,⁷,²⁻⁷ which cause myocyte disarray and fibrosis that are characteristic of the disease.⁸⁻¹⁰

Clinical manifestations of HCM range from asymptomatic patients to sudden death.¹¹⁻¹⁵ HCM is the main cause of sudden death in adolescents, young adults and athletes.¹²⁻¹⁶ Clinically, the main risk factors for sudden death are nonsustained ventricular tachycardia, syncope, familial history of sudden death or aborted sudden death.¹²⁻¹⁶ Other risk factors include left ventricular wall thickness greater than or equal to 30 mm and left ventricular outflow tract obstruction found in echocardiography.²⁻⁴,¹⁰

Although the diagnosis of HCM may be established by two-dimensional echocardiography, cardiovascular magnetic resonance (CMR) has been the method of choice in the last years, due to its accuracy for determination of morphology, tissue and functional characterization, and detection of myocardial fibrosis by delayed myocardial enhancement (DME).²,⁴

Patients with HCM, in use of implantable cardioverter defibrillators (ICDs), cannot be followed-up by CMR, since the presence of ICD implant may be a contraindication for the exam. In this context, cardiac computed tomography (CCT) may be a useful alternative in the assessment and management of these patients.

This study aimed to provide an updated literature review of current concepts in the use of CMR and CCT for HCM, emphasizing the diagnostic impact of both methods.

Interaction With Echocardiography

Echocardiography is the most available method to assess morphological and functional changes of HCM. The diagnostic criteria for HCM are left ventricular (LV) wall thickness greater than or equal to 15 mm at the end of diastole, and a septal to lateral wall thickness ratio greater than or equal to 1.3 in a non-dilated left ventricle, and in the absence of other conditions that may explain such abnormality.¹⁷⁻¹⁹

Although widely available, the method has some limitations in the evaluation of HCM, such as: patients with poor acoustic window, poor visualization of some regions – basal anterolateral wall of left ventricle, cardiac apex and right ventricle (RV).²⁻³,¹⁰,¹¹ Both CRM and CCT are three-dimensional, multiplanar methods, with excellent spatial resolution, that have been recognized as important tools for the assessment of HCM patients.²,²²

Cardiac Magnetic Resonance

CMR is an excellent method for the evaluation of HCM, since it precisely determines both the localization and extension of hypertrophy and evaluates ventricular function. The method also allows the detection of mechanisms of obstruction of the LV outflow tract, as well as the establishment of the pressure gradient between the LV outflow tract and the aorta. Other advantages include detection of areas of myocardial fibrosis by DME, diagnosis of apical HCM, and the follow-up of patients undergoing septal ablation.²⁻¹⁰,²¹,²²

CMR is more sensitive than echocardiography in the detection of HCM markers, such as myocardial crypts.²⁰,²² It can contribute to the preoperative planning of myectomy and quantification of necrotic tissue by alcohol septal ablation. Due to these characteristics, CMR has been considered a method to be routinely used for the establishment of diagnosis and prognosis of HCM.²⁰

Keywords

Cardiomiopatia Hipertrófica, Espectroscopia de Ressonância Magnética / uso diagnóstico, Tomografia Computadorizada por Raíos X / métodos, Tomografia Computadorizada por Raíos X / tendências.

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**Genetic variants (genotype)**

HCM is an autosomal dominant genetic disorder that determines mutations in genes that codify sarcomeric proteins.\(^6,^8\) The most frequent mutations involve beta-myosin heavy chain gene (MYH7), myosin-binding protein C (MYBPC3) and troponin T (TNNT2).\(^6,^7\) Mutation in MYH7 is associated with earlier manifestations of HCM as compared with MYBPC3 gene mutation, whereas TNNT2 gene mutation is more associated with the risk of sudden death.\(^8\)

Until recently, the presence of reverse septal curvature was the only change in CMR imaging that indicated mutations in genes that codify sarcomeric proteins. However, studies conducted in 2012 identified deep basal interseptal crypts as stronger positive predictors of HCM genotype, since they can be found in 81% of patients who are genotype-positive for HCM.\(^21,^24\) Besides, the combination of both changes increases the positive predictive value (98%) for mutation.\(^25\)

Recent studies have demonstrated that other changes are indicative of the presence of a sarcomere gene mutation, including anterior mitral valve leaflet elongation, abnormal trabeculation and smaller LV systolic cavity.\(^26\) The identification of these abnormalities allows the detection of individuals who are carriers of mutations, and hence more susceptible to the development of HCM. This information enables the development and implementation of strategies that can change the natural history of HCM,\(^2,^25\) as well as the development of new imaging sequences that better characterize such changes.\(^25\)

**Morphological variants (phenotype)**

The precise characterization of the HCM phenotype is valuable for the establishment of invasive therapies, including septal myectomy and alcohol septal ablation. The HCM phenotype allows the definition of localization and magnitude of hypertrophy,\(^21\) and characterization of mitral and subvalvular apparatus and papillary muscles.\(^20\) Many morphological variants of HCM have been described using CRM (Figure 1).

a) **Normal variant** – related to relatives of individuals genetically positive for HCM, or those who should be followed up for the risk of developing any other variant throughout life. We should consider the presence of crypts 24, abnormal trabeculation, and anterior mitral valve leaflet elongation.\(^26\)

b) **Asymmetric variant with sigmoid septum** – this is the main presentation of CMH,\(^6,^16\) characterized by a myocardium hypertrophy next to the LV outflow tract and the sigmoid septum ("S" shape). This variant may cause subaortic obstruction and mitral regurgitation.

c) **Asymmetric variant with reverse septal curvature** – characterized by a septum hypertrophy as a reversed “S”, more distant from the LV outflow tract. This presentation does not cause obstruction to the LV outflow. The identification of this variant by CMR is characterized by a septal/free wall thickness ratio greater than 1.3 in the short-axis plane.\(^4\)

d) **Variant with mid-ventricular obstruction with or without a left ventricular apical diverticulum** – characterized by a mid-ventricular hypertrophy that causes a local narrowing and, in severe cases, apical dilatation. In approximately 10% of patients, there may be apical aneurysm formation.\(^5\) Apical aneurysm is better diagnosed by CRM than echocardiography which, in turn, can fail to detect this change in 10% of cases.\(^16\)

e) **Apical variant** – characterized by obliteration of LV cavity at the apex together with an apical wall thickness greater than 15 mm or a ratio between apical and basal LV wall thicknesses greater than or equal to 1.3-1.5 cm\(^4\). This presentation is considered to have a better prognosis than the other variants, although it has been more associated with ischemia and apical myocardial infarction\(^16\) (Figure 2).

f) **Symmetric (concentric) variant** – characterized by hypertrophy of LV wall with reduction of LV cavity. This variant may also be present in other disorders, including amyloidosis, sarcoidosis and Fabry’s disease\(^4,^10\) (Figure 3).

g) **Focal variant** – characterized by hypertrophy located in the myocardium. CMR help distinguishing focal HCM from other cardiac masses, by identifying evidence of myocardial contractility in the former case.\(^4\)

h) **HCM in RV** – occurs in 18% of HCM patients, generally involving the mid-and apical portion of the RV. This may cause right ventricular outflow obstruction in severe situations.\(^4\).

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**Figure 1** – Types of hypertrophic cardiomyopathy. (A) Normal heart. (B) Sigmoid septum. (C) Reverse septal curvature. (D) Mid-ventricular obstruction without apical aneurysm. (E) Apical. (F) Symmetric. (G) Asymmetric hypertrophy in the lateral wall. (H) Hypertrophic cardiomyopathy in right ventricle.
Increased maximum thickness of the right ventricular wall (> 8 mm) has been shown by CMR in approximately 20% of HCM patients. Areas with increased wall thickness are commonly observed in the insertion of the right ventricular wall into anterior and posterior septum, although the entire RV may be involved.

Assessment of ventricular mass
LV hypertrophy is currently one of the clinical requirements for HCM diagnosis, and is associated with poorer prognosis. However, comparative studies of genotypes and phenotypes involved in HCM have reported that varied LV wall thickness may be found in patients with HCM.

A reliable, accurate, quantitative analysis of LV wall thickness is fundamental, since a measurement greater than 30 mm increases the risk of sudden death. Therefore, this may be a crucial information for the implant of ICD to prevent sudden death in some patients. Measurements of LV wall thickness should be performed in the short-axis plane, at the end of diastole. LV mass, quantified by CMR (indexed for body surface area), greater than 2 standard deviations above the normal range is considered a sensitive predictor of favorable clinical outcomes in HCM. The normal range for LV mass index is 62.5 ± 9.0 g/m² for men and 54.6 ± 12.0 g/m² for women. However, the measurement of LV mass lacks specificity as an indicator of clinical outcomes, probably because in many HCM patients, hypertrophy is limited to a small number of segments of the left ventricle.

CMR is more accurate than echocardiography in diagnosing ventricular hypertrophy, its magnitude and distribution. For this reason, the method is decisive for stratifying the risk of HCM. In addition, CMR provides a better evaluation of hypertrophy distribution, especially when it is localized in the anterolateral region, and in posterior and apical septum of LV, and a more reliable quantification of the myocardial mass in case of asymmetric hypertrophy (Figure 4).

Quantification of myocardial fibrosis
Myocardial fibrosis can be detected by delayed enhancement CMR. This method is based on the property of gadolinium to distribute in the extracellular space between normal and fibrotic tissue, leaving the latter tissue in a slower rate.
There is no standard of reference for DME in HCM, although its presence in the interventricular septum, particularly in the mid- and basal anteroseptal segment, is suggestive of HCM. DME can also occur in LV free wall and hypertrophied segments of the ventricles. These data indicate that analysis of the segments by delayed enhancement CMR is an important parameter for differential diagnosis of HCM.

DME is found in 65% of patients and the distribution of fibrosis typically follows a multifocal, heterogeneous and mid-wall pattern. DME is not commonly found in non-hypertrophic cardiomyopathy, except in advanced stages of the disease, and can be associated with increased myocardial stiffness and reverse remodeling of the left ventricle.

Studies have demonstrated an inverse relationship between DME and LV function. The presence of DME predicts a worse prognosis for HCM patients due to higher risk of sudden death, systolic dysfunction and nonsustained ventricular tachycardia.

Areas of DME may represent the substrate for malignant ventricular tachyarrhythmias (Figure 5). Delayed enhancement CMR is a supporting tool in the decision-making process for primary prevention ICDs in patients in whom high-risk for sudden death remains uncertain after assessment of conventional risk factors.

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DME technique consists in the intravenous administration of approximately 0.2 mmo/kg of gadolinium-based contrast at 1-2 mL/s. Acquisition of images may be started 10 minutes after contrast infusion, at multiple inversion times (TI Scout), which allows the use of a proper TI to null the myocardial signal (typically between 200 and 300 s). DME images (1 cm-sections) are acquired in the short-axis plane, from the base to the apex of the heart, using a T1-weighted inversion recovery gradient-echo sequence. Images in two- and four-chamber plane were also acquired, for better visualization of apical disease (Figure 6).

Recent studies have pointed out that the presence of either extensive late enhancement or late enhancement in the interventricular septum and right ventricular anterior and posterior insertions may be a biomarker of sudden death. Patients in this situation should be classified as high risk. We recommend that quantification of myocardial fibrosis should be performed with visual assessment.

Analysis of diastolic dysfunction

Diastolic dysfunction in HCM results from abnormal dissociation of actin and myosin filaments during diastole, mainly in its early active phase. It can be measured by time-volume curve obtained from cine-magnetic resonance imaging (Figure 7).

Diastolic function can also be evaluated by in-plane phase-contrast CMR imaging, with direct measurement of diastolic relaxation of cardiac muscle. This, in turn, may be obtained by the assessment of LV strain and recovery rates during diastole, yielding the values of fast relaxation (E’) and slow relaxation (A’). In hypertrophied segments, the early diastolic filling velocity is reduced, and there is a decrease in the rate of LV relaxation.

Mitral valve flow velocity and pulmonary artery flow velocity can also be estimated by through-plane phase contrast, allowing the calculation of pressure gradient and flow velocity. Analysis of transmitral flow enables the estimation of early ventricular filling velocity (E-wave) and peak flow velocity at atrial contraction (A-wave). Mild diastolic dysfunction is characterized by a reduction of E/A ratio. As the dysfunction progresses, there is a pseudonormalization and subsequent increase in the E/A ratio, which represents the restrictive physiology of diastolic dysfunction.

Use of myocardial tagging (MT)

Myocardial tagging (MT) is an excellent non-invasive tool for quantifying regional diastolic and systolic myocardial function. Quantification of global myocardial dysfunction is...
not sensitive to detect small reductions in regional ventricular performance, which may occur even in normal LV ejection.\textsuperscript{40,41}

In this context, MT has emerged as a technique that allows the definition of subclinical myocardial dysfunction (Figure 8).

MT evaluates myocardial deformation during the cardiac cycle. The most used method is a coordinate system,\textsuperscript{42} which takes into account three axes of myocardial contraction: radial, circumferential, and longitudinal.\textsuperscript{39,42}

- **Radial strain** – describes myocardial stiffness, oriented toward the central long axis of the left ventricle.
- **Circumferential strain** – describes circumferential shortening of myocardium, in the short-axis plane, tangential to the epicardial wall.
- **Longitudinal strain** – represents LV shortening from the base to the apex, in the long-axis plane.

- **Myocardial rotation** – represents the analysis of mid-myocardial deformation, in degrees, in the short-axis plane during systole or diastole. In a normal cardiac cycle, myocardial rotation is generated by a clockwise rotation of the base and a counterclockwise rotation of the apex during systole. During diastole, the opposite rotation to the normal position promotes the release of potential energy stored, and a suction force in the left ventricle during isovolumic relaxation. A change in the rotation pattern may be used to detect subtle systolic and diastolic dysfunctions in many cardiac diseases.\textsuperscript{39} Myocardial twist (in degrees) can be calculated as the difference between apical and basal rotation during all cardiac cycle. We can calculate torsion (in degrees per cm) as a type of twist normalization by ventricular longitudinal length. Also, the peak of systolic or diastolic rotation, twist and torsion, can be also calculated, but they are only used in scientific studies.

Figure 5 – Asymmetric hypertrophic cardiomyopathy. Asymmetric parietal hypertrophy with septal predominance. Presence of important myocardial fibrosis in the mid anteroseptal segment (white) in the delayed enhancement technique. (A) Cine-magnetic resonance and (B) Delayed myocardial enhancement.

Figure 6 – End-stage hypertrophic cardiomyopathy. Diffuse parietal hypertrophy, and hypertrophy in both ventricles, associated with dilation and important dysfunction. Presence of well-defined myocardial fibrosis (white) in delayed enhancement technique. (A) Cine-magnetic resonance and (B) Delayed myocardial enhancement.
Assessment of the LV outflow track

Obstruction of LV outflow tract (induced or at rest) is found in approximately 70% of patients with HCM.\(^4\) During systole, LV outflow is hampered by basal septal hypertrophy and displacement of the papillary muscles and mitral leaflets.\(^16\)

Obstruction of LV outflow with a maximum gradient at rest greater than 30 mmHg is a strong predictor of sudden death. In this case, interventions to reduce such gradient, including septal miectomy and alcohol septal ablation by catheterization, may be justified.\(^10,43\)

Two-D transthoracic echocardiography is currently the method of choice for anatomic assessment and flow measurement in case of LV outflow obstruction.\(^8,10,22\) However, CMR enables a more accurate evaluation of the mitral valve structure, including changes in the papillary muscle anatomy.

Although the quantification of the LV outflow obstruction gradient may be performed by CMR, it still represents a challenge, since it requires an accurate image plane alignment to prevent the loss of high-velocity areas. Besides, the presence of turbulent outflow causes signal loss and phase error.\(^10\) New sequences in ultrashort echo time CMR imaging have been currently used and may make the evaluation of LV outflow obstruction easier and more reliable.\(^10\)
Assessment of myocardial ischemia

Myocardial ischemia caused by microvascular disease is an etiopathogenic hypothesis for the development of HCM. The early detection of ischemia before cardiac remodeling may represent a promising therapeutic target that can change the natural history of HCM.

Contraindications and limitations

CMR imaging in HCM patients depends on the technical quality of image acquisition and requires interpretation of the images by experienced physicians. The method has some relative and absolute contraindications, which have been reassessed every year. This includes the performance of CMR in patients with pacemaker, ICD, brain clips, cochlear implants, and metallic fragments in the eyes. Today, the test can be performed in some pacemakers and patients.

Limitations of CMR include nephrogenic systemic fibrosis (which causes systemic tissue fibrosis and is associated with the use of gadolinium in patients with stage 4 and 5 chronic renal failure) and patients with hepatorenal syndrome. In addition, it is worth mentioning that, despite the increase in accessibility to the CRM, the test is available in a small number of centers.

Cardiac Computed Tomography

CCT provides a clear delineation of the myocardium and accurate measurement of cardiac wall thickness, ventricular volumes, and LV mass ejection fraction, which are well correlated with the CMR findings. Additionally, CCT allows the assessment of coronary arteries and cardiac valves. The European Guidelines on HCM recommends that CCT should be considered in patients with poor acoustic window for echocardiography or contraindications for CMR (class IIa).

CCT has a wide range of clinical applications due to anatomic and functional properties. However, they are indicated only in case of diagnostic doubt, poor acoustic window to perform echocardiography or contraindications to perform CMR. Therefore, CCT is rarely used as the first method of choice for patients with HCM. Table 1 describes the main clinical applications of CCT in HCM.

Delayed myocardial enhancement

Since the demonstration that CCT allows the visualization of fibrotic areas in patients with acute myocardial infarction similarly with CMR, a similar DME technique has been developed to identify fibrosis in HCM patients. Studies have shown a good correlation between DME by CCT and DME by MCR.

DME technique should be performed in multidetector computed tomography scans, capable to connect cardiac acquisition to electrocardiography. This is the case of some of the 16-channel multidetector scanner and most of the 64-channel devices. Iodinated contrast (150 mL) is appropriate for this application.

Table 1 – Clinical applicability of computed tomography in the assessment of hypertrophic cardiomyopathy

| Parameters                      | Clinical applicability                                              |
|---------------------------------|---------------------------------------------------------------------|
| Diameters / volumes             | Indicated in case of diagnostic doubt                               |
| Wall thickness                  |                                                                     |
| Ventrices                       |                                                                     |
| Global systolic function        | Indicated in case of diagnostic doubt                               |
| Regional systolic function      |                                                                     |
| Diastolic function              | Not indicated                                                       |
| Volumes                         |                                                                     |
| Function                        |                                                                     |
| Systolic anterior motion        | It may be indicated                                                 |
| Valves                          |                                                                     |
| Mitral regurgitation            | Not indicated. It may be used only for static evaluations, such as valve planimetry and morphological characterization. |
| Pharmacological stress          |                                                                     |
| Ischemia / perfusion defects    | It may be indicated especially in combination with anatomiac features of coronary arteries |
| Coronary                        |                                                                     |
| Luminal reduction               | Deve ser utilizada                                                   |
| CAD                             |                                                                     |
| Pressures / Velocities          |                                                                     |
| Gradients                       | Not indicated                                                       |
| Fibrosis                        |                                                                     |
| Delayed myocardial enhancement | It may be used in case cardiac magnetic resonance is contraindicated |

Adapted from Nagueh et al. CAD: coronary artery disease.
intravenously administered by automatic injector at 3 mL/s. Seven minutes later, the images are acquired in retrospective ECG-synchronized helical mode, during a 10-second breath-holding, and rebuilt during the same diastolic cycle.

**Contraindications and limitations**

In comparison with CMR, CCT has inferior temporal resolution and soft tissue characterization. Radiation exposure from a 64-channel computed tomography scanner (mean of 6.7 ± 2.07 mSv) is a drawback of the method as compared with CMR. Nevertheless, the development of new techniques and CCT scanners involving lower radiation doses may provide safer conditions for myocardial fibrosis investigation.

**Perspectives**

**T1 mapping by CMR**

Conventional techniques to assess DME that evaluates the presence of focal myocardial fibrosis may underestimate the distribution and extension of fibrosis. The technique of T1 mapping is more sensitive to detect fibrotic areas, since it amplifies regional variations in gadolinium distribution, making it easier to detect diffuse interstitial myocardial fibrosis.

Recent studies have demonstrated that HCM genotype- and phenotype- positive patients, and genotype-positive / phenotype-negative patients have increased levels of biomarkers of collagen deposition. This indicates that increased levels of profibrotic markers are also found in patients without ventricular hypertrophy, suggesting that hypertrophy is preceded by fibrosis process. In this case, T1 mapping may be used at the initial phase of HCM for the early detection of diffuse myocardial fibrosis in these patients.

Today, the modified Look-Locker inversion recovery (MOLLI) has been the most studied sequence for the assessment of T1 mapping. MOLLI uses electrocardiogram-gated image acquisition at end-diastole and merges 11 images in a 17-heartbeat breathhold. Many protocols have been published and may be used in the assessment of interstitial myocardial fibrosis by CMR.

Recently, Nacif et al. have developed a method to quantify interstitial fibrosis by CCT that may be useful in HCM.

**Spectroscopy**

Spectroscopy by 31 P-CMR may be used to assess the energy status in the myocardial tissue. Patients with HCM have decreased myocardial energy status that directly correlates with extension of hypertrophy and severity of diastolic function. Limitations of spectroscopy include prolonged scanning time, low spatial resolution and requirement of dedicated surface coils.

**Aortic stiffness**

The assessment of aortic stiffness may be an important parameter for risk stratification in HCM. Studies have demonstrated that HCM patients have increased aortic stiffness, especially those with myocardial fibrosis detected by DME technique.

**Conclusion**

HCM is the most common genetic cardiovascular disease with potential for high mortality, since it is the most common cause of sudden death in young patients. The development and improvement of imaging analysis by CMR and CCT has enabled the early establishment of diagnosis and prognosis of HCM. In the future, interventions aimed at stopping the natural history of the disease can be made. Both CMR and CCT are validated as methods with high sensitivity and specificity, with few contraindications and minimal risks of adverse effects, and should be used in the management of patients with HCM.

**Author contributions**

Conception and design of the research: Santos AASMD, Nacif MS. Acquisition of data: Oliveira DCL, Assunção VB, Nacif MS. Analysis and interpretation of the data: Oliveira DCL, Assunção VB, Nacif MS. Writing of the manuscript: Oliveira DCL, Assunção VB, Nacif MS. Critical revision of the manuscript for intellectual content: Santos AASMD, Nacif MS.

**Potential Conflict of Interest**

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References

1. Maron BJ. Hypertrophic cardiomyopathy: a systematic review. JAMA. 2002;287(10):1308-20.

2. Sara L, Szarf G, Tachibana A, Shiozaki AA, Villa AV, Oliveira AC, et al; Sociedade Brasileira de Cardiologia, Colegio Brasileiro de Radiologia. [II Guidelines on Cardiovascular Magnetic Resonance and Computed Tomography of the Brazilian Society of Cardiology and the Brazilian College of Radiology. Arq Bras Cardiol. 2014;103(6 Suppl 3):1-86.

3. Shiozaki AA, Kim RJ, Parga JR, Tassi EM, Arteaga E, Rochitte CE. Cardiovascular magnetic resonance in hypertrophic cardiomyopathy. Arq Bras Cardiol. 2007;88(2):243-8.

4. Nourredin AR, Liu S, Nacl M, Judge DP, Halushka MK, Abraham TP, et al. The diagnosis of hypertrophic cardiomyopathy by cardiovascular magnetic resonance. J Cardiovasc Magn Reson. 2012;14:17.

5. Lima JA, Desai MY. Cardiovascular magnetic resonance imaging: current and emerging applications. J Am Coll Cardiol. 2004;44(6):1164-71.

6. Bittencourt MJ, Rocha RM, Filho FM. Cardiomiopatia hipertrófica. Rev Bras Cardiol. 2010;20(3):17-24.

7. Ho CY, Lopez BA, Coelho-Filho OR, Lakdawala NK, Cirillo AL, Jarolim P, et al. Myocardial fibrosis as an early manifestation of hypertrophic cardiomyopathy. N Engl J Med. 2010;363(6):552-63.

8. Yingchoncharoen T, Tang WW. Recent advances in hypertrophic cardiomyopathy. P1000 Prime Rep. 2014;6:12.

9. Anan R, Greve G, Thierfelder L, Watkins H, McKenna WJ, Solomon S, et al. Prognostic implications of novel beta cardiac myosin heavy chain gene mutations that cause familial hypertrophic cardiomyopathy. J Clin Invest. 1994;93(1):280-5.

10. Hoey ET, Teoh JK, Das I, Ganeshan A, Watkin RW, Simpson H, et al. The role of cardiovascular MRI for risk stratification in hypertrophic cardiomyopathy. Clin Radiol. 2014;69(3):221-30.

11. Anan R, Greve G, Thierfelder L, Watkins H, McKenna WJ, Solomon S, et al. Prognostic implications of novel beta cardiac myosin heavy chain gene mutations that cause familial hypertrophic cardiomyopathy. J Clin Invest. 1994;93(1):280-5.

12. Hoey ET, Teoh JK, Das I, Ganeshan A, Watkin RW, Simpson H. The role of cardiovascular MRI for risk stratification in hypertrophic cardiomyopathy. J Cardiovasc Magn Reson. 2012;14:17.

13. Maron BJ, McKenna WJ, Danielson GK, Kappenberger LJ, Kuhn HJ, et al. Sudden death in young competitive athletes: clinical, demographic, and pathologic findings. JAMA. 1996;276(3):199-204.

14. Maron BJ. Sudden death in young athletes. N Engl J Med. 2003;349(11):1064-75.

15. Maron BJ, McKenna WJ, Danielson GK, Kappenberger LJ, Kuhn HJ, Seidman CE, et al; American College of Cardiology/European Society of Cardiology clinical expert consensus document on hypertrophic cardiomyopathy. A report of the American College of Cardiology Foundation Task Force on Clinical Expert Consensus Documents and the European Society of Cardiology Committee for Practice Guidelines. J Am Coll Cardiol. 2003;42(9):2167-72.

16. Elliott PM, Poloniecki J, Dickie L, Mathenre R, Roberts WC, Mueller FO, et al. Sudden death in young competitive athletes: clinical, demographic, and pathologic findings. JAMA. 1996;276(3):199-204.

17. Elliott PM, Poloniecki J, Dickie L, Mathenre R, Roberts WC, Mueller FO, et al. Sudden death in young competitive athletes: clinical, demographic, and pathologic findings. JAMA. 1996;276(3):199-204.

18. Elliott PM, Poloniecki J, Dickie L, Mathenre R, Roberts WC, Mueller FO, et al. Sudden death in young competitive athletes: clinical, demographic, and pathologic findings. JAMA. 1996;276(3):199-204.

19. Elliott PM, Poloniecki J, Dickie L, Mathenre R, Roberts WC, Mueller FO, et al. Sudden death in young competitive athletes: clinical, demographic, and pathologic findings. JAMA. 1996;276(3):199-204.

20. Elliot PM, Anastasakis A, Borger MA, Borggreve M, Cecchi F, Charron P, et al. 2014 ESC Guidelines on diagnosis and management of hypertrophic cardiomyopathy: The Task Force for the Diagnosis and Management of Hypertrophic Cardiomyopathy of the European Society of Cardiology. Eur Heart J. 2014;35(39):2733-79.

21. Rickers C, Wilke NM, Jerosch-Herold M, Casey SA, Panse P, Panse N, et al. Utility of cardiac magnetic resonance imaging in the diagnosis of hypertrophic cardiomyopathy. Circulation. 2005;112(6):855-61.

22. Bogaert J, Olivoto I. MR imaging in hypertrophic cardiomyopathy: from magnet to bedside. Radiology. 2014;273(2):329-48.

23. Maron MS, Rowin EJ, Lin D, Appelbaum E, Chan RH, Gibson CM, et al. Prevalence and clinical profile of myocardial crypts in hypertrophic cardiomyopathy. Circ Cardiovasc Imaging. 2012;5(4):441-7.

24. Brouwer WP, Gernars T, Head MC, Velden JV, Heymans MW, Chiariens I, et al. Multiple myocardial crypts on modified long-axis view are a specific finding in prehypertrophic HCM mutation carriers. Eur Heart J Cardiovasc Imaging. 2012;13(4):292-7.

25. Deva DP, Williams LK, Care M, Siminovich KA, Moshonov H, Wintersperger B, et al. Deep basal interstitial crypts occur more commonly in patients with hypertrophic cardiomyopathy due to disease-causing myofilament mutations. Radiology. 2013;269(1):68-76.

26. Captur G, Lopes LR, Mohun TJ, Patel V, Li C, Basset P, et al. Prediction of sarcomere mutations in subclinical hypertrophic cardiomyopathy. Circ Cardiovasc Imaging. 2014;7(6):863-71.

27. Maron MS, Maron BJ, Harrigan C, Burnos J, Gibson CM, Olivotto I, et al. Hypertrophic cardiomyopathy phenotype revisited after 50 years with cardiovascular magnetic resonance. J Am Coll Cardiol. 2009;54(3):220-8.

28. Chun EJ, Choi SJ, Jin KN, Kwang HJ, Kim YJ, Choi BW, et al. Hypertrophic Cardiomyopathy: assessment with MR imaging and multidetector CT. Radiographics. 2010;30(5):1309-28.

29. Zhang L, Mmagu O, Liu L, Li D, Fan Y, Baranchuk A, et al. Hypertrophic cardiomyopathy: can the noninvasive diagnostic testing identify high risk patients? World J Cardiol. 2014;6(8):764-70.

30. Hoey ET, Elassaly M, Ganesan A, Watkin RW, Simpson H. The role of magnetic resonance imaging in hypertrophic cardiomyopathy. Quant Imaging Med Surg. 2014;4(5):397-406.

31. Kalil Filho R. A resonância nuclear magnética na análise da hipertrofia miocárdica. Rev Soc Cardiol Estado de São Paulo. 1994;4(4):369-75.

32. O’Hanlon R, Grasso A, Roughton M, Moon JC, Clark S, Wage R, et al. Prognostic significance of myocardial fibrosis in hypertrophic cardiomyopathy: J Am Coll Cardiol. 2010;56(11):867-74.

33. Bruder O, Wagner A, Jensen CJ, Schneider S, Ong P, Kispert EM, et al. Myocardial scar visualized by cardiovascular magnetic resonance imaging predicts major adverse events in patients with hypertrophic cardiomyopathy. European Heart J. 2014;35(39):2733-79.

34. Arriagada C, Peters DC, Gibson CM, Maron BJ, Manning WJ, Maron MS, et al. Hypertrophic cardiomyopathy: quantification of late gadolinium enhancement with contrast-enhanced cardiovascular MR imaging. Radiology. 2011;258(1):128-33.

35. Gersh BJ, Maron BJ, Bonow RO, Dearani JA, Fifer MA, Link MS, et al. ACCF/AHA Guideline for the Diagnosis and Treatment of Hypertrophic Cardiomyopathy: a Report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines. Circulation. 2011;124(24):2733-831.

36. Kłopotowski M, Kukula K, Malek LA, Świątek M, Polanka-Skoczylak M, et al. The value of cardiac magnetic resonance and distribution of late gadolinium enhancement for risk stratification of sudden cardiac death in patients with hypertrophic cardiomyopathy. J Cardiol. 2015 Sep 9. [Epub ahead of print].
37. Chan RH, Maron BJ, Olivetto I, Pencina MJ, Assenza CE, Haas T, et al. Prognostic value of quantitative contrast-enhanced cardiovascular magnetic resonance for the evaluation of sudden death risk in patients with hypertrophic cardiomyopathy. Circulation. 2014;130(6):484-95.

38. Schwarz F, Schwab E, Beckmann BM, Schuessler R, Zinsser D, Golz T, et al. Magnetic resonance imaging of hypertrophic cardiomyopathy: evaluation of diastolic function. Radiology. 2013;331(1):15-23.

39. Shehata ML, Cheng S, Chuman NF, Bluemken DA, Lima JA. Myocardial tissue tagging with cardiovascular magnetic resonance. J Cardiovasc Magn Reson. 2009;11:55.

40. Jeung MY, Germain P, Croisille P, Elghannudi S, Roy C, Gagne A. Myocardial tagging with MR imaging: overview of normal and pathologic findings. Radiographics. 2012;32(5):1381-98.

41. Kim YJ, Choi BW, Hur J, Lee HJ, Seo JS, Kim TH, et al. Delayed enhancement in hypertrophic cardiomyopathy: comparison with myocardial tagging MRI. J Magn Reson Imaging. 2008;27(5):1054-60.

42. O'Rourke MF, Weber T, Adji A. Aortic stiffness in hypertrophic cardiomyopathy. J Am Coll Cardiol. 2010;55(5):504-5.

43. Naghavi SF, Bierig SM, Budoff MJ, Desai M, Dilsizian V, Eidem B, et al; American Society of Echocardiography; American Society of Nuclear Cardiology; Society for Cardiovascular Magnetic Resonance; Society of Cardiovascular Computed Tomography. American Society of Echocardiography Clinical Recommendations for Multimodality Cardiovascular Imaging of Patients with Hypertrophic Cardiomyopathy: Endorsed by the American Society of Nuclear Cardiology, Society for Cardiovascular Magnetic Resonance, and Society of Cardiovascular Computed Tomography. J Am Soc Echocardiogr. 2011;24(5):473-98.

44. Nordbeck P, Erl G, Ritter O. Magnetic resonance imaging safety in pacemaker and implantable cardioverter defibrillator patients: how far have we come? Eur Heart J. 2015;36(24):1505-11.

45. Leite CC. Gadolinio e fibrose nefrogênica sistêmica: o que todo médico deve saber. Radiol Bras. 2007;40(4):iv-v.

46. Gerber BL, Belge B, Legros GJ, Lim P, Poncelet A, Pasquet A, et al. Characterization of acute and chronic myocardial infaracts by multidetector computed tomography: comparison with contrast-enhanced magnetic resonance. Circulation. 2006;113(6):823-33.

47. Lardo AC, Cordeiro MA, Silva C, Amado LC, George RT, Saliaris AP, et al. Contrast-enhanced multidetector computed tomography viability imaging after myocardial infarction: characterization of myocyte death, microvascular obstruction, and chronic scar. Circulation. 2006;113(3):394-404.

48. Shiozaki AA, Sensa T, Arteaga E, Pita CG, Martinelli Filho M, Ávila LE, et al. Myocardial fibrosis in patients with hypertrophic cardiomyopathy and high risk for sudden death. Arq Bras Cardiol. 2010;94(4):535-40.

49. Shiozaki AA, Sensa T, Arteaga E, Martinelli Filho M, Pita CG, Ávila LE, et al. Myocardial fibrosis detected by cardiac CT predicts ventricular fibrillation/ventricular tachycardia events in patients with hypertrophic cardiomyopathy. J Cardiovasc Comput Tomogr. 2013;7(3):173-81.

50. Esposito A, Colantoni C, De Cobelli F, De Cobelli F, Del Vecchio A, Palmisano A, et al. Multidetector computed tomography for coronary stents imaging: high-voltage (140-kVp) prospective ECG-triggered versus standard-voltage (120-kVp) retrospective ECG-gated helical scanning. J Comput Assist Tomogr. 2013;37(3):395-401.

51. Gottlieb I, Camargo GC, Derenne ME. Ressonância magnética em cardiomiopatia hipertrófica. Arq Bras Cardiol: Imagem cardiovasc. 2014;27(3):202-7.

52. Nacif MS, Kawel N, Lee JH, Chen X, Yao J, Zavodni A, et al. Interstitial myocardial fibrosis assessed as extracellular volume fraction with low-radiation-dose cardiac CT. Radiology. 2012;264(3):876-83.

53. Mazaev VV, Stukalova OV, Ternovoi SK, Chazova IE. [Evaluation of myocardial energy metabolism by 31P magnetic resonance spectroscopy in patients with hypertrophic cardiomyopathy versus healthy individuals]. Vestn Rentgenol Radiol. 2012;6:8-12.