Dilated Cardiomyopathy With Mid-Range Ejection Fraction at Diagnosis: Characterization and Natural History

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Background—Limited data are available on mid-range ejection fraction (mrEF) patients with dilated cardiomyopathy. We sought to define the characteristics, evolution, and long-term prognosis of dilated cardiomyopathy patients with mrEF at diagnosis.

Methods and Results—We analyzed all dilated cardiomyopathy patients consecutively evaluated in the Trieste Heart Muscle Disease Registry from 1988 to 2013. mrEF and reduced ejection fraction (rEF) were defined as baseline left ventricular (LV) ejection fraction values between 40% and 49% and <40%, respectively. All-cause mortality or heart transplantation, sudden cardiac death, or major ventricular arrhythmias were considered as outcome measures. Worsening LV ejection fraction (reduction to <40%) during follow-up was also considered to identify possible predictors of adverse remodeling. Among 812 enrolled patients, 175 (22%) presented with mrEF at presentation. At baseline, as compared with the rEF group, mrEF patients had lower rates of moderate–severe mitral regurgitation and restrictive LV filling pattern. During a median follow-up period of 120 (60–204) months, the mrEF group presented a lower rate of death/heart transplantation (9% versus 36%, P<0.001) and sudden cardiac death or major ventricular arrhythmias (4.5% versus 15%, P<0.001) than rEF patients. Moreover, 29 out of 175 mrEF patients (17%) evolved to rEF. Restrictive LV filling pattern emerged as the strongest predictor of rEF development following multivariable analysis.

Conclusions—mrEF identified a consistent subgroup of dilated cardiomyopathy patients diagnosed in an earlier stage with subsequent apparent better long-term evolution. However, 17% of these patients evolved into rEF despite the use of medical therapy. A baseline restrictive LV filling pattern was independently associated with subsequent evolution to rEF. (J Am Heart Assoc. 2019;8:e010705. DOI: 10.1161/JAHA.118.010705.)

Key Words: dilated cardiomyopathy • heart failure • mid-range ejection fraction • prognosis

The European Society of Cardiology has introduced a novel class of heart failure (HF) without a severely reduced ejection fraction (EF), calling it as HF with a mid-range EF. HF with a mid-range EF mainly includes patients with primarily mild systolic dysfunction and features of diastolic dysfunction. This new entity encompasses patients with clinical, laboratory, or instrumental evidence of HF and a left ventricular ejection fraction (LVEF) between 40% and 49%. The definition of this new class of HF could lead to a better understanding of the characteristics, pathophysiology, and treatment options for a group of patients often neglected in randomized clinical trials. Dilated cardiomyopathy (DCM), a heterogeneous myocardial disease with a frequent genetic cause, is characterized by several degrees of reduced LVEF. Notably, DCM patients are typically relatively young, with low comorbidities and long periods of asymptomatic LV systolic dysfunction. They are often diagnosed in an early phase of the disease, through the application of extensive genetic counseling programs and sometimes by way of the results of preparticipation screenings for sport activities. Therefore, at diagnosis, DCM patients could be characterized by the absence of HF symptoms and non-severe systolic dysfunction. However, few data on the characteristics, evolution, and outcomes of DCM patients presenting with mid-range ejection fraction (mrEF) at baseline are available at this time. In the present study, we sought to characterize DCM patients presenting with mrEF at baseline, describing their natural history and prognosis in comparison with reduced EF (rEF) patients.
Clinical Perspective

What Is New?

- This article includes the largest-to-date patient population with dilated cardiomyopathy and mid-range ejection fraction at first presentation.
- Additionally, it provides the first long-term outcome in these patients with a median follow-up period of 120 (60–204) months.

What Are the Clinical Implications?

- Patients with mid-range ejection fraction denotes a consistent (ie, 22%) subgroup of dilated cardiomyopathy deserving careful and reliable risk assessment.
- A non-negligible number of patients (17%) with mid-range ejection fraction at baseline tend to evolve towards worse clinical phenotype, with a subsequent worse prognosis.
- The presence of a restrictive left ventricular filling pattern at baseline might identify those mid-range ejection fraction patients with a greater risk to experience worse outcomes during follow-up.

Methods

The authors declare that all supporting data are available within the article.

Study Population

All DCM patients consecutively enrolled in the Trieste Heart Muscle Disease Registry from January 1988 to December 2013 were considered eligible for inclusion in the present study and were retrospectively analyzed. Enrolled patients were directly hospitalized in our referral center or referred shortly after the onset of symptoms from other centers.

Follow-up ended on December 31, 2015 or at the time of death or urgent heart transplant (D/HTx).

The institutional ethics board approved the study and written informed consent was obtained under the institutional review board policies of the hospital administration.

A diagnosis of DCM was determined according to the currently accepted criteria. Ischemic etiology was excluded with coronary angiography, systematically performed in patients aged >35 years with cardiovascular risk factors. Patients with significant coronary artery disease (>50% stenosis of a major coronary artery), history of severe systemic hypertension (>160/100 mm Hg), biopsy-proven active myocarditis, alcohol intake of >100 g/day, significant organic valve disease, tachycardia-induced cardiomyopathy, peripartum cardiomyopathy, congenital heart disease, or advanced systemic disease affecting their short-term prognosis were excluded.5,10,11

After enrollment, if not contraindicated, all patients received evidence-based pharmacological (at maximum tolerated dosages) and device therapies.1

Per protocol, all patients underwent a scheduled follow-up (clinical and echocardiographic) at 6, 12, and 24 months after enrollment and then every ≥2 years frequently with respect to individual clinical necessity.

We divided our population into 2 groups according to the baseline EF: patients with rEF (ie, EF <40%) and patients with mrEF (ie, EF 40%–49%).

Table 1. Baseline Clinical Characteristics of the Population Divided According to LVEF Categories

| Value | Total 812 pts | rEF DCM 637 pts | mrEF DCM 175 pts | P Value |
|-------|---------------|-----------------|-----------------|---------|
| Age, y | 46 (36;56) | 48 (37;57) | 42 (32;50) | 0.001 |
| Male sex (%) | 565 (70%) | 455 (71%) | 110 (63%) | 0.029 |
| NYHA III–IV, n (%) | 195 (24%) | 186 (29%) | 9 (5) | <0.001 |
| SBP, mm Hg | 124 ± 18 | 123 ± 18 | 126 ± 17 | 0.27 |
| Heart rate, bpm | 78 ± 16 | 80 ± 16 | 70 ± 12 | <0.001 |
| LBBB (%) | 256 (32%) | 221 (35%) | 35 (20%) | <0.001 |

Echocardiography

| Value | Total 812 pts | rEF DCM 637 pts | mrEF DCM 175 pts | P Value |
|-------|---------------|-----------------|-----------------|---------|
| LAESAI, cm²/m² | 14 ± 4 | 14 ± 4 | 11 ± 3 | <0.001 |
| LVEDDI, mm/m² | 36 ± 6 | 37 ± 6 | 32 ± 4 | <0.001 |
| LVEDVI, mL/m² | 99 ± 37 | 106 ± 37 | 71 ± 18 | <0.001 |
| LVEF, % | 31 ± 10 | 27 ± 7 | 44 ± 3 | <0.001 |
| RVD, n (%) | 199 (26%) | 182 (30%) | 17 (10) | <0.001 |
| Moderate–severe MR, n (%) | 285 (36%) | 262 (42%) | 23 (13%) | <0.001 |
| Restrictive LV filling pattern, n (%) | 233 (30%) | 225 (37%) | 8 (5) | <0.001 |
| E/E’ | 14 ± 7 | 15 ± 8 | 10 ± 3 | <0.001 |
| sPAP | 35 ± 14 | 37 ± 15 | 25 ± 5 | <0.001 |

Therapy

| Value | Total 812 pts | rEF DCM 637 pts | mrEF DCM 175 pts | P Value |
|-------|---------------|-----------------|-----------------|---------|
| ACE-inhibitors–ARBs, n (%) | 783 (96%) | 624 (98%) | 159 (91%) | 0.081 |
| Beta-blockers, n (%) | 729 (90%) | 576 (91%) | 153 (88%) | 0.311 |
| Diuretics, n (%) | 607 (75%) | 531 (84%) | 76 (44%) | <0.001 |
| MRAs, n (%) | 239 (30%) | 216 (34%) | 23 (13%) | <0.001 |

Values are expressed as mean ± SD or median with interquartile range as appropriate, and as percentage. ACE indicates angiotensin-converting enzyme; ARBs, angiotensin receptor blockers; LAESAI, left atrial end systolic area indexed; LBBB, left bundle branch block; LV, left ventricular; LVEDDI, left ventricular end-diastolic diameter indexed; LVEDVI, left ventricular end-diastolic volume indexed; LVEF, left ventricular ejection fraction; MR, mitral regurgitation; MRAs, mineralocorticoid receptor antagonists; NYHA, New York Heart Association; RVD, right ventricular dysfunction; SBP, systolic blood pressure; sPAP, systolic pulmonary artery pressure.
Outcome Measures

The study analyzed the incidence of the following combined outcome measures: (1) all-cause mortality and heart transplantation (D/HTx) and (2) sudden cardiac death (SCD) or major ventricular arrhythmias (MVAs). Specifically, MVAs were defined as sustained (≥30 seconds, hemodynamically symptomatic) ventricular tachycardia, ventricular fibrillation (VF), appropriate treatment of sustained ventricular tachycardia or ventricular fibrillation via implantable cardioverter-defibrillator therapy (shock or antitachycardia pacing for termination of sustained ventricular tachycardia ≥185 bpm), or cardiopulmonary resuscitation after cardiac arrest. Finally, a worsening of LVEF to <40% during follow-up in the group of patients initially presenting with mrEF was also considered to identify possible predictors of adverse remodeling.

Echocardiographic Study

Biventricular dimensions and systolic and diastolic functions were assessed according to international guidelines. Specifically, LV volume and LVEF were calculated by the Simpson biplane method. All volumes were indexed according to body surface area.

Transmitral E- and A-wave velocities were measured using pulsed wave Doppler at the level of the mitral leaflet tips. The LV filling pattern was classified as restrictive in the presence of an E-wave deceleration time of <120 ms or E/A value ≥2 associated with an E-wave deceleration time of ≤150 ms. Right ventricular dysfunction was defined by a right ventricular fractional area change of <35%.

Mitral regurgitation (MR) was assessed using a multiparametric approach applied by measuring the effective regurgitant volume or orifice area as determined by the proximal isovelocity surface area, whenever feasible, or the vena contracta width at color-flow Doppler, and graded on a 3-point scale according to current recommendations. Significant MR was considered if greater than mild. Left atrial end-systolic area was calculated from a 4-chamber view.

All measurements were obtained from the mean of 3 beats (patients in sinus rhythm) or of 5 beats (patients with atrial fibrillation).

Statistical Analysis

Summary statistics of clinical and laboratory variables were expressed as means and standard deviations, medians and interquartile ranges, or counts and percentages, as appropriate. Cross-sectional comparisons between groups were made using the ANOVA test on continuous variables, the Brown–Forsythe statistic when the assumption of equal variances did not hold, or the non-parametric Mann–Whitney test when necessary. The chi-squared or Fisher exact test was used for discrete variables. Repeated measures between baseline and follow-up were evaluated by means of the paired t test for continuous Gaussian-distributed parameters or the Wilcoxon
test as appropriate. For binary variables, the McNemar test was applied. Cumulative incidence curves for SCD/MVAs with respect to the competing risk of death for other causes or heart transplant were estimated and compared between groups. Cause-specific uni- and multivariable Cox regression models were estimated to find factors associated with EF...
worsening (EF <40%), starting from the list of significant and clinically relevant parameters that emerged at univariable analyses and taking into account those with a P value at univariable analysis of <0.1. Cumulative incidence curves for the risk of experiencing EF worsening with respect to the competing risk of death or heart transplant were estimated and compared between groups, stratified by the presence of the only significant risk factor derived from the previous analyses. A time-dependent Cox model was finally estimated to evaluate the impact of experiencing EF worsening on survival during follow-up, and an “extended” version of survival curves was estimated to visually represent results.18

Results
The study population included 812 patients with recently diagnosed DCM (median duration of symptoms of 2.5 [0–9] months). One hundred and seventy-five patients (22%) presented with mrEF (ie, 40%–49%).

At baseline, in comparison with the rEF group, mrEF patients presented features of a less-advanced stage of disease, including smaller LV end-diastolic volume, lower rates of New York Heart Association functional classes III to IV, left bundle branch block, right ventricular dysfunction, moderate–severe MR, and restrictive LV filling pattern. No differences in the administration of angiotensin-converting enzyme inhibitors and beta-blockers were found; on the contrary, mrEF patients less frequently took diuretics and mineralocorticoid receptor antagonists (Table 1).

During a median follow-up time of 120 (60–204) months, 247 patients (30%) died (n=168) or underwent HTx (n=79). One hundred and five patients (13%) experienced an episode of SCD (n=59) or had MVAs (n=46). At a median follow-up of 120 months, as compared with the rEF patients, the mrEF group presented lower rates of all-cause D/HTx (10% versus 30%, P<0.001) and of SCD/MVAs incidence (4% versus 13%, P<0.001) (Figure 1). A lower incidence of events in terms of all-cause D/HTx and SCD/MVAs was reported in mrEF patients than in rEF patients; this was true also upon stratifying the analysis by enrollment period (ie, before 2006 and from 2007, see Figure 2).

Evolution in rEF
Twenty-nine of 175 mrEF patients (17%) evolved to the rEF group at a median follow-up of 69 (24–91) months. Patients with worsened systolic function more frequently showed restrictive LV filling pattern at baseline in comparison with patients with persistent mrEF. Consequently, they were more frequently treated with diuretics and mineralocorticoid receptor antagonists (Table 2). In the time-dependent Cox model, DCM patients with persistent mrEF showed a significantly lower risk of long-term D/HTx than did those with baseline rEF (considering rEF as the reference category, hazard ratio: 0.08; 95% CI, 0.03–0.19; Figure 3). Conversely, DCM patients with mrEF at baseline who, during follow-up, experienced a shift to rEF status presented a significantly higher risk of long-term D/HTx than did DCM patients with baseline rEF (hazard ratio: 2.27; 95% CI, 1.23–4.18; Figure 3). The presence of a restrictive LV filling pattern (hazard ratio: 4.0; 95% CI, 1.39–11.57, P=0.01) emerged as the strongest baseline predictor independently associated with EF worsening after the stepwise selection starting from

Table 2. Baseline Clinical Characteristics of the mrEF DCM Group Divided According to Evolution/No Evolution to rEF (LVEF <40%)

|                                    | No Evolution to rEF Group (n=146) | Evolution to rEF Group (n=29) | P Value |
|------------------------------------|----------------------------------|------------------------------|---------|
| Age, y                             | 42±6                             | 42±15                        | 0.84    |
| Male sex (%)                       | 91 (62%)                         | 19 (65%)                     | 0.74    |
| NYHA III–IV, n (%)                 | 6 (4)                            | 3 (10)                       | 0.15    |
| SBP, mm Hg                         | 125±16                           | 129±18                       | 0.17    |
| Heart rate, bpm                    | 69±12                            | 73±12                        | 0.14    |
| LBBB (%)                           | 29 (19%)                         | 6 (20%)                      | 0.94    |

**Echocardiography**

|                               | No Evolution to rEF Group | Evolution to rEF Group | P Value |
|-------------------------------|---------------------------|------------------------|---------|
| LAESAI, cm²/m²                | 9±3                       | 9±3                    | 0.82    |
| LVEDDI, mm/m²                 | 31±3                      | 32±4                   | 0.72    |
| LVEDVI, mL/m²                 | 70±18                     | 70±15                  | 0.67    |
| LVEF, %                       | 44.1±2.6                  | 44.3±3.0               | 0.70    |
| RVD, n (%)                    | 12 (8)                    | 5 (17)                 | 0.15    |
| Moderate–severe MR, n (%)     | 16 (11)                   | 7 (24)                 | 0.057   |
| Restrictive LV filling pattern, n (%) | 4 (2) | 4 (14) | 0.02 |
| E/E                            | 10±3                      | 11±4                   | 0.11    |
| sPAP                           | 24±5                      | 29±5                   | 0.017   |

**Therapy**

|                               | No Evolution to rEF Group | Evolution to rEF Group | P Value |
|-------------------------------|---------------------------|------------------------|---------|
| ACE-inhibitors-ARBs, n (%)    | 146 (100)                 | 29 (100)               | 0.34    |
| Beta-blockers, n (%)          | 127 (88)                  | 26 (88)                | 0.82    |
| Diuretics, n (%)              | 58 (40)                   | 18 (62)                | 0.02    |
| MRAs, n (%)                   | 15 (10)                   | 8 (27)                 | 0.02    |

Values are expressed as mean±SD or median with interquartile range as appropriate, and as percentage. ACE indicates angiotensin-converting enzyme; ARBs, angiotensin receptor blockers; LAESAI, left atrial end systolic area indexed; LBBB, left bundle branch block; LV, left ventricular; LVEDDI, left ventricular end-diastolic diameter indexed; LVEDVI, left ventricular end-diastolic volume indexed; LVEF, left ventricular ejection fraction; MR, mitral regurgitation; MRAs, mineralocorticoid receptor antagonists; NYHA, New York Heart Association; RVD, right ventricular dysfunction; SBP, systolic blood pressure; sPAP, systolic pulmonary artery pressure.
the list of significant parameters at univariable analyses (Table 3 and Figure 4).

Discussion

The existence of patients with mrEF is well-known in clinical practice. However, these patients have been included in clinical trials on preserved LVEF by using different EF cutoff values (eg, >40%, >45%, or >50%), regardless of ischemic or non-ischemic HF etiology. Following the introduction of the HFmrEF class in the last European Society of Cardiology guidelines, the need for a deeper understanding of this specific subgroup has been suggested. Nevertheless, recent data reported on HFmrEF often refer to evaluations in different moments of underlying disease course. Therefore, the real impact of mrEF at the time of diagnosis of HF is still lacking. Here, we described the natural history of mrEF from the time of disease presentation in a recently diagnosed DCM population. This is a strength of the present study, because limited information is available at this time in the literature on this subject. DCM patients represent a particular subgroup of the HF population. In fact, they are frequently asymptomatic or mildly symptomatic for HF at the time of diagnosis, often despite severe LV remodeling. Asymptomatic DCM patients deserve a careful and reliable determination of risk, because they could suffer rapid disease progression or die suddenly early after disease identification. This study provides a deep insight into the characterization and natural history of a large and well-selected cohort of DCM patients presenting with mrEF.

Incidence and Long-Term Outcome of DCM Patients With mrEF

Our findings show that almost one-quarter of DCM patients (22%) had mrEF at baseline, with most of them (95%) presenting with New York Heart Association functional classes I or II. Similarly, in the CHARM (Candesartan in Heart Failure: Assessment of Reduction in Mortality and Morbidity) trial, a consistent quota was found: 1295 of 7955 (17%) HF patients had an LVEF between 43% and 52%. Thus, about one-fifth of newly diagnosed HF patients could present with mrEF, and they should not be neglected.

The main clinical and echocardiographic characteristics at baseline found in this study indicate that the largest portion of mrEF DCM patients are diagnosed in a less-advanced stage of disease. An early diagnosis is probably supported by family screening programs or results from diagnostic tests performed in subjects being screened for sports participation. The resulting early optimal medical treatment probably has...
led to a significantly better prognosis in mrEF patients in comparison with rEF ones. Nevertheless, the benign prognosis of mrEF DCM patients is not generalizable to all cases. Importantly, a consistent percentage (ie, 17%) of mrEF DCM patients evolved towards rEF at a median time of 5 years. There are a subgroup of DCM patients in which mrEF at presentation represented a transition zone in the way towards the rEF group. These data highlight the importance of an adequate early comprehensive characterization and of individualized long-term follow-up with recommended treatments performed in all DCM patients, including those apparently belonging to low-risk classes. In fact, a malignant genotype could be expressed without severe LV remodeling, such as mutations in LMNA A/C.22 Moreover, a “restrictive” phenotype has been included in the clinical spectrum of desmin-related cardiomyopathies.23

Table 3. Univariable Cause-Specific Cox Regression Analysis for EF Worsening

| Age, y | HR 0.981 to 1.037 0.525 | SPB 0.988 to 1.032 0.368 | NYHA III–IV 0.716 to 7.873 0.158 | LBBB 0.477 to 2.880 0.730 | LAESAI 0.913 to 1.186 0.551 | LV 0.976 to 1.013 0.563 | LVEF 0.899 to 1.182 0.660 | Moderate–severe MR 1.001 to 5.498 0.050 | Restrictive LV filling pattern 4.814* 1.662 to 13.947* 0.004* | sPAP 1.047 to 1.354 0.008 | RVD 0.848 to 5.887 0.104 | LAESAI 0.900 to 1.184 0.650 |

| Unadjusted HRs | HR | 95% CI | P Value |
|----------------|-----------------|-----------------|----------|
| Age, y         | 1.009           | 0.981 to 1.037  | 0.525    |
| SPB            | 1.010           | 0.988 to 1.032  | 0.368    |
| NYHA III–IV    | 2.374           | 0.716 to 7.873  | 0.158    |
| LBBB           | 1.172           | 0.477 to 2.880  | 0.730    |
| LAESAI         | 1.041           | 0.913 to 1.186  | 0.551    |
| LV              | 0.995           | 0.976 to 1.013  | 0.563    |
| LVEF           | 1.031           | 0.899 to 1.182  | 0.660    |
| Moderate–severe MR | 2.346         | 1.001 to 5.498  | 0.050    |
| Restrictive LV filling pattern* | 4.814* | 1.662 to 13.947* | 0.004* |
| sPAP           | 1.191           | 1.047 to 1.354  | 0.008    |
| RVD            | 2.234           | 0.848 to 5.887  | 0.104    |
| LAESAI         | 1.032           | 0.900 to 1.184  | 0.650    |

HR indicates hazard ratio; LAESAI, left atrial end-systolic area indexed; LBBB, left bundle branch block; LV, left ventricular; LVEF, left ventricular ejection fraction; MR, mitral regurgitation; NYHA, New York Heart Association; RVD, right ventricular dysfunction; SBP, systolic blood pressure; sPAP, systolic pulmonary artery pressure.

*Stepwise Cox regression: starting from a list composed by moderate-severe MR, sPAP, and a restrictive LV filling pattern; this latter remained the only significant parameter, with adjusted HR: 2.896; 95% CI, 1.726–15.216; P=0.003 (see Methods section, Statistical analysis subsection).

The Prognostic Role of Diastolic Dysfunction

Our analysis underlines the prognostic role of diastolic dysfunction in apparently low-risk DCM patients. At first evaluation, the presence of severe diastolic dysfunction (ie, restrictive LV filling pattern), associated with non-severe systolic dysfunction, emerged in our cohort as the strongest predictor of subsequent EF worsening (Figure 4). Importantly, patients with mrEF evolving to rEF presented a significantly higher risk of long-term major events as compared with baseline rEF patients. As previously reported in patients with non-dilated hypokinetic cardiomyopathy, the presence of significant diastolic dysfunction brought to light a subgroup with a high risk of adverse outcomes.8,10 This similar behavior could suggest a possible overlap between mrEF and non-dilated hypokinetic cardiomyopathy. A comprehensive and systematic genetic characterization, in association with a stricter follow-up regimen, may be necessary to better characterize these patients. In fact, a malignant genotype could be expressed without severe LV remodeling, such as mutations in LMNA A/C.22 Moreover, a “restrictive” phenotype has been included in the clinical spectrum of desmin-related cardiomyopathies.23

Limitations

Our study population, partially referred to a tertiary center, along with the retrospective nature of the analysis, may have

![Figure 4. Cumulative incidence curves for ejection fraction worsening of mid-range ejection fraction dilated cardiomyopathy patients with or without a restrictive left ventricular filling pattern. The presence of a restrictive left ventricular filling pattern emerged as the strongest baseline predictor independently associated with ejection fraction worsening. LV indicates left ventricular; mrEF, mid-range ejection fraction; rEF, reduced ejection fraction.](image-url)
created a degree of selection bias. Our results are applicable only to DCM patients and should not be generalized to the whole population of patients with HF. Although the study population represents, to the best of our knowledge, one of the largest existing mrEF DCM populations evaluated at baseline and during follow-up with a complete assessment, the low number of patients evolving to rEF underpowers the statistical analysis with only three potential predictors in the multivariable model. Dosages of medical HF therapy were not systematically available. However, according to the internal protocol, all patients titrated the therapy to the maximum tolerated dosage.

Systematic data on natriuretic peptides, 3-dimensional echocardiography, strain and strain rate, cardiac magnetic resonance, and genetic characterization findings were not available. They should be the focus of future larger studies.

Conclusions

mrEF denotes a consistent (ie, 22%) subgroup of DCM patients who are generally diagnosed in an earlier stage and who present an apparently better clinical evolution. However, some mrEF DCM patients evolve into rEF despite medical therapy, with a subsequently worse prognosis. The presence of a restrictive filling pattern might be an important clue for identifying high-risk DCM patients presenting with mrEF.

Disclosures

None.

References

1. Ponikowski P, Voors AA, Anker SD, Bueno H, Cleland JGF, Coats AJS, Falk V, Gonzalez-Juanatey JR, Harjola V-P, Jankowska EA, Jessup M, Linde C, Nihoyannopoulos P, Parissis JT, Pieske B, Riley JP, Rosano GMC, Rulope LM, Ruschitzka F, Rutten FH, van der Meer P. 2016 ESC guidelines for the diagnosis and treatment of acute and chronic heart failure. Eur J Heart Fail. 2016;18:2129–2206.
2. Merlo M, Cannatà A, Gobbo M, Stocco D, Elliott PM, Sinagra G. Evolving concepts in dilated cardiomyopathy. Eur J Heart Fail. 2018;20:228–239.
3. McNally EM, Mestroni L. Dilated cardiomyopathy: genetic determinants and mechanisms. Circ Res. 2017;121:731–748.
4. Seidman JG, Seidman CE. The genetic basis for cardiomyopathy: from mutation identification to mechanistic paradigms. Cell. 2001;104:557–567.
5. Merlo M, Pivetta A, Pinamonti B, Stocco D, Zecchin M, Barbati G, Di Lenarda A, Sinagra G. Long-term prognostic impact of therapeutic strategies in patients with idiopathic dilated cardiomyopathy: changing mortality over the last 30 years. Eur J Heart Fail. 2014;16:317–324.
6. Moretti M, Merlo M, Barbati G, Di Lenarda A, Brun F, Pinamonti B, Gregori D, Mestroni L, Sinagra G. Prognostic impact of familial screening in dilated cardiomyopathy. Eur J Heart Fail. 2010;12:922–927.
7. Tsuji K, Sakata Y, Nochioka K, Morita M, Tsuchida T, Minamiya M, Chiba Y, Yamasaki T, Kuroda K, Kitai K, Yamashita S, Shimokawa H. Characterization of heart failure patients with mid-range left ventricular ejection fraction-a report from the CHART-2 Study. Eur J Heart Fail. 2017;19:1258–1269.
8. Pinto YM, Elliott PM, Arbustini E, Adler Y, Anastasakis A, Bohm M, Duboc D, Gimeno J, de Groot P, Imazio M, Heymans S, Klingel K, Komajda M, Limongelli G, Linhart A, Mogensen J, Moon J, Pieper PG, Seferovic PM, Schueler S, Zamorano JL, Catrof ALP, Charron P. Proposal for a revised definition of dilated cardiomyopathy, prokoine-dependent cardiomyopathy, and its implications for clinical practice: a position statement of the ESC working group on myocardial and pericardial diseases. Eur Heart J. 2016;37:1850–1858.
9. Elliott P, Andersson B, Arbusinini E, Bliliszka Z, Cecchi F, Charron P, Dubourg O, Kuhl U, Maisch B, McKenna WJ, Monserrat L, Pankuweit S, Rapezzi C, Seferovic P, Tavazzi L, Keren A. Classification of the cardiomyopathies: a position statement from the European Society of Cardiology Working Group on Myocardial and Pericardial Diseases. Eur Heart J. 2008;29:270–276.
10. Gigli M, Stocco D, Merlo M, Barbati G, Ramani F, Brun F, Pinamonti B, Sinagra G. Insights into mildly dilated cardiomyopathy: temporal evolution and long-term prognosis. Eur J Heart Fail. 2017;19:531–539.
11. Merlo M, Ammirati E, Gentile P, Artico J, Cannata A, Finocchiaro G, Barbati G, Sormani P, Verrenti A, Fabris E, Aleksova A, Bussani R, Petrella D, Ciceri M, Raineri C, Fragasso G. Persistent left ventricular dysfunction after acute lymphocytic myocarditis: frequency and predictors. PLoS One. 2019;14:e0214616.
12. Spezzacatene A, Sinagra G, Merlo M, Barbati G, Graw SL, Brun F, Slavov D, Di Lenarda A, Salcedo EE, Towbin JA, Safitzte J, Marcus F, Zareba W, Taylor MR, Mestroni L; Familial Cardiomyopathy Registry. Arhythmogenic phenotype in dilated cardiomyopathy: natural history and predictors of life-threatening arrhythmias. J Am Heart Assoc. 2015;4:e002149. DOI: 10.1161/JAHA.115.002149.
13. Lang RM, Badano LP, Mor-Avi V, Alfialo J, Armstrong A, Emilande L, Flachskampf FA, Lancellotti P, Goldstein SA, Kuznetsova T, Lancellotti P, Murat D, Picard MH, Rietzschel ER, Rudski L, Spencer KT, Tsang W, Voigt JU. Recommendations for cardiac chamber quantification by echocardiography in adults: an update from the American Society of Echocardiography and the European Association of Cardiovascular Imaging. J Am Soc Echocardiogr. 2015;28:31–39.e14.
14. Pinamonti B, Zecchin M, Di Lenarda A, Gregori D, Sinagra G, Camerini F. Persistence of restrictive left ventricular filling pattern in dilated cardiomyopathy: an ominous prognostic sign. J Am Coll Cardiol. 1997;29:604–612.
15. Rudski LG, Lai WW, Alfialo J, Hua L, Handschumacher MD, Chandrasekaran K, Solomon SD, Louie EK, Schiller NB. Guidelines for the echocardiographic assessment of the right heart in adults: a report from the American Society of Echocardiography endorsed by the European Association of Echocardiography, a registered branch of the European Society of Cardiology, and the Canadian Society of Echocardiography. J Am Soc Echocardiogr. 2010;23:685–688.
16. Zoghbi WA, Adams D, Bonow RO, Enriquez-Sarano M, Foster E, Grayburn PA, Hahn RT, Han Y, Hung J, Lang RM, Little SH, Shah DJ, Sherman S, Thavendiranathan P, Thomas JD, Weissman NJ. Recommendations for noninvasive evaluation of native valvular regurgitation: a report from the American Society of Echocardiography developed in collaboration with the Society for Cardiovascular Magnetic Resonance. J Am Soc Echocardiogr. 2017;30:303–371.
17. Stocco D, Merlo M, Pinamonti B, Poli S, Gigli M, Barbati G, Fabris E, Di Lenarda A, Sinagra G. Early improvement of functional mitral regurgitation in patients with idiopathic dilated cardiomyopathy. Am J Cardiol. 2015;115:1137–1143.
18. Snapinn SM, Jiang Q, Iglewicz B. Illustrating the impact of a time-varying covariate with an extended Kaplan–Meier estimator. J Clin Oncol. 2016;34:5196–5201.
19. Yusuf S, Pfeffer MA, Swedberg K, Granger CB, Held P, McMurray JJV, Missionel EL, Olofsson B, Ostergren J. Effects of candesartan in patients with chronic heart failure and preserved ejection fraction. Am Heart J. 2006;152:1307–1316.
20. Pitt B, Pfeffer MA, Assmann SF, Boineau R, Anand IS, Braunwald E, Chaae K, Clark K, DiMarco JP, Giugliano D, Gomis R, Hertzer NR, Kostis WB, Lanasino U, Linhart A, Mogensen J, Moon J, Piepoli MF, Fonseca C, Mebazaa A, Lund L, Ambrosio GA, Coats AJ, Ferranti R, Ruschitzka F, Maggioni AP, Filippatos G. Epidemiology and one-year outcomes in patients with chronic heart failure and preserved, mid-range and reduced heart failure ejection fraction: an analysis of the ESC Heart Failure Long-Term Registry. Eur J Heart Fail. 2017;19:1574–1585.
21. Chl L, Benson L, Dahlstrom U, Edner M. Association between use of renin-angiotensin system antagonists and mortality in patients with heart failure and preserved ejection fraction. JAMA. 2012;308:2108–2117.

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