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ORIGINAL RESEARCH MAGGIC Risk Model Predicts Adverse Events and Left Ventricular Remodeling in Non-Ischemic Dilated Cardiomyopathy

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Purpose: We aimed to study the Meta-analysis Global Group in Chronic Heart Failure (MAGGIC) risk model's prognostic value and relationship with left ventricular remodeling in dilated cardiomyopathy.

Patients and Methods: Dilated cardiomyopathy patients were prospectively recruited and underwent clinical assessments. MAGGIC risk score was calculated. Patients were followed up for adverse events and echocardiography. Primary endpoints were all-cause mortality and first rehospitalization due to heart failure. Secondary endpoint was left ventricular remodeling defined as a decline in left ventricular ejection fraction >10% or an increase in left ventricular end-diastolic diameter >10%. Survival status was examined using Cox regression analysis. The model's ability to discriminate adverse events and left ventricular remodeling was calculated using a receiver operating characteristics curve.

Results: In total, 114 patients were included (median follow-up time = 31 months). The risk score was independently related to adverse events (2-year all-cause mortality: hazard ratio [HR] = 1.122; 95% confidence interval [CI], 1.043–1.208; 1-year first rehospitalization due to heart failure: HR = 1.094; 95% CI, 1.032-1.158; 2-year first rehospitalization due to heart failure: HR = 1.088; 95% CI, 1.033-1.147, all P < 0.05). One-year change in left ventricular end-diastolic diameter was correlated with the risk score (r = 0.305, P = 0.002). The model demonstrated modest ability in discriminating adverse events and left ventricular remodeling (all areas under the curve were 0.6-0.7).

Conclusion: The MAGGIC risk score was related to adverse events and left ventricular remodeling in dilated cardiomyopathy.

Keywords: dilated cardiomyopathy, left ventricular remodeling, prognosis, risk model

Introduction

Non-ischemic dilated cardiomyopathy (DCM) is a progressive disease presenting with left ventricle dilation as well as decreased systolic function excluding abnormal loading conditions and coronary artery disease.¹ As an important cause of heart failure (HF), DCM is responsible for a substantial proportion of adverse events such as cardiac morbidity and mortality.^{2,3} Left ventricular remodeling plays an important role in the pathophysiological progression of DCM, which can appear as a change in left ventricular diameter and left ventricular systolic function on echocardiography.^{1,4-6} Drugs currently treating DCM, including angiotensin converting enzyme inhibitors, angiotensin receptor blockers, mineralocorticoid receptor antagonists, and beta blockers are used to halt or reverse left ventricular

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remodeling.^{2,7} Some researchers also found that short-term left ventricular remodeling or left ventricular reverse remodeling was related to long-term prognosis.^{8–11} There is a demand for an easily available tool to predict left ventricular remodeling in DCM patients at baseline to guide treatment.

The Meta-Analysis Global Group in Chronic Heart Failure (MAGGIC) risk score emerged in 2013 and has since been validated by several studies as a good risk stratification tool for predicting the prognosis of HF patients.^{12–15} Moneghetti et al also reported that the MAGGIC risk score was independently related with the outcomes in DCM patients.¹⁶ However, most studies focused on patients with HF of heterogenous etiology and lacked validation of the MAGGIC risk model's prognostic value in DCM patients only. Studies were also lacking on the relationship between MAGGIC risk score and left ventricular remodeling.

Here we aimed to calculate the baseline MAGGIC risk score and study its prognostic value and relationship with left ventricular remodeling in a cohort of DCM patients.

Patients and Methods Study Population

This observational study was approved by Ethics Committee of the First Affiliated Hospital, School of Medicine, Zhejiang University. Written informed consents were obtained from all included patients according to the Helsinki Declaration. DCM patients were prospectively recruited from June 2015 to June 2018. DCM was diagnosed as increasing left ventricular end-diastolic diameter (LVEDD) (>55 mm in men or >50 mm in women) and decreasing left ventricular ejection fraction (LVEF) <45% on echocardiography. Patients with coronary artery disease and abnormal loading conditions, such as hypertension, congenital heart disease, or valvular disease, were excluded.¹⁷ All patients underwent coronary angiography or coronary computed tomography angiography. Patients with a significant stenosis of coronary artery >50% were excluded.¹⁸ The demographic characteristics and clinical data of every patient were collected. All DCM patients received standard drug therapy according to the current guideline.¹⁹

Echocardiography

All patients underwent two-dimensional echocardiography using a dedicated ultrasound system (Vivid E9, GE

Vingmed Ultrasound, Horten, Norway). Cardiac structure and function were assessed according to the former scan protocol of the American Society of Echocardiography.²⁰ Left atrial diameter, LVEDD and LVEF were collected to evaluate the status of left heart of DCM patients. Left atrial diameter and LVEDD were obtained using twodimensional ultrasonography in the parasternal long-axis 2-chamber view. Left atrial diameter was obtained at the end of left ventricle systole, while LVEDD was obtained at the end of left ventricle diastole. LVEF was estimated using the biplane Simpson method. All echocardiographic scans and measurements, including scans on baseline and scans during follow-up, were performed by an experienced research sonographer blinded to the patients' other clinical data. Twenty percent of total subjects were randomly selected from the DCM cohort by random numbers to evaluate intra-observer variabilities. The sonographer repeated the measurements by using the identical method at least 6 months later.

MAGGIC Risk Model

The MAGGIC risk score was calculated at baseline when each DCM patient was included. The risk model comprises 13 variables including LVEF (from 0 to 7 points according to 6 different groups of LVEF), LVEF extra for age (from 0 to 15 points according to 7 different groups of age combining with 3 different groups of LVEF), LVEF extra for systolic blood pressure (from 0 to 5 points according to 6 different groups of systolic blood pressure combining with 3 different groups of LVEF), body mass index (from 0 to 6 points according to 5 different groups of body mass index), creatinine (from 0 to 8 points according to 8 different groups of creatinine), New York Heart Association class (from 0 to 8 points according to 4 different groups of New York Heart Association class), male sex (one point), current smoker (one point), diabetes (three points), chronic obstructive pulmonary disease (two points), first diagnosis of HF within the past 18 months (two points), not on beta-blocker use (three points), and not on angiotensin converting enzyme inhibitor/angiotensin receptor blocker use (one point).¹⁵ All patients were divided into two groups, including M1 group and M2 group, by the median of all MAGGIC risk scores.

Follow-Up

Patients were followed up every 6 months to record adverse events. All living patients underwent

echocardiography within 2 weeks of the 1-year follow-up. Primary endpoints were all-cause mortality and first rehospitalization due to HF. First rehospitalization due to HF was defined as worsening symptoms and signs of HF, such as dyspnea or signs of congestion, which require urgent treatment in hospital for the first time after patients' baseline admissions.²¹ The secondary endpoint was left ventricular remodeling during follow-up, which was defined as a decline of LVEF >10% or increase of LVEDD >10%.⁷ The 1-year change of left heart remodeling parameters (left atrial diameter, LVEDD, and LVEF) were calculated using the formula:

$$\frac{1 - year}{change(\%)} = \frac{\left(Value \ at \ 1 - \frac{year}{follow \ up} - \frac{Value \ at}{baseline}\right)}{Value \ at \ baseline}$$

Statistical Analysis

MedCalc version 13.0 (Med-Calc Software; Ostend, Belgium) and GraphPad Prism version 6.0 (GraphPad Software, San Diego, CA, USA) were used to perform statistical analysis. The Kolmogorov-Smirnov test was used to check the distribution of continuous variables. Normally distributed continuous variables were described as mean \pm SD, while non-normally distributed data were presented as median and interguartile range. Intergroup differences of continuous variables were analyzed by Student's t-test or the Mann-Whitney U-test. Differences between two groups of categorical variables were tested by the Chi-square test. Survival curves of 2-year all-cause mortality as well as 1-year and 2-year first rehospitalization due to HF were created using the Kaplan-Meier method. Survival curves between different groups of MAGGIC risk scores were compared using the Log rank test. Univariate and multivariate Cox regression analyses were used to examine relationships between the clinical characteristics and adverse events, and hazard ratio (HR) was calculated. Spearman correlation analysis was performed to evaluate the relationship between left heart remodeling parameter changes and MAGGIC risk scores. The diagnostic ability of the MAGGIC risk model for discriminating adverse events and left ventricular remodeling were calculated by receiver operating characteristics curve, and the areas under the curve were calculated. The intra-observer variability was calculated by mean bias from Bland-Altman analyses, coefficient of variation (COV, %), and intra-class correlation coefficient (ICC). ICC calculation used two-way mixed, absolute

agreement single measure model. Values of P < 0.05 were considered statistically significant.

Results

Demographic and Clinical Characteristics

A total of 217 patients were recruited according to echocardiography findings. One hundred and three patients were excluded due to coronary artery disease according to coronary angiography or coronary computed tomography angiography findings. Ultimately, 114 DCM patients were included in the study. The median follow-up time was 31 months (interquartile range, 26-34 months). After following up the adverse events of all included DCM patients, 10 patients (<10% of all included patients) died within the first year after inclusion, while 21 patients died within the first 2 years after inclusion. Thirty-one patients experienced the first rehospitalization due to HF in the first year after inclusion, while 44 patients experienced the first rehospitalization due to HF within the first 2 years after inclusion. The patients' demographic and clinical characteristics stratified by endpoint are shown in Table 1. Age, sex, body mass index, systolic blood pressure, N-terminal-pro-brain natriuretic peptide level, ratio of current smokers, diagnosis of chronic obstructive pulmonary disease, angiotensin receptor blocker use, LVEDD, and MAGGIC risk score differed significantly among the endpoint groups.

Relationship Between MAGGIC Risk Score and Adverse Events

Patients in M1 group (5–20 points, n = 58) had significantly better survival status than that in M2 (21–37 points, n = 56) group (Figure 1). Cox regression analysis showed that MAGGIC risk score was independently related to adverse events in DCM patients (Table 2). However, the MAGGIC risk model demonstrated modest ability to discriminate adverse events (Figure 2 and Supplementary Table 1).

Relationship Between MAGGIC Risk Score and Left Ventricular Remodeling Status

Since 10 patients died within the first year of follow-up, 104 patients underwent 1-year follow-up echocardiography. Patients showed significant left heart reverse remodeling after 1 year as demonstrated by the decreased left atrial diameter (baseline vs 1-year follow-up: 45 ± 7 mm vs 42 ± 9 mm, P = 0.017), LVEDD (baseline vs 1-year

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	lotal	All-Cause Mortality	rtality					First Rehospi	First Rehospitalization Due to HF	to HF			
		l Year			2 Year			l Year			2 Year		
		+ (N=I0)	-(N=I04)	٩	+ (N=2I)	-(N=93)	٩	+ (N=31)	-(N=83)	٩	+ (N=44)	-(N=70)	4
Age (vears)	61+13	63+17	60+13	0.547	63+15	60+13	0.310	66+12	58+13	0.005	64+14	58+13	0.035
(22)	65 (74/114)	50 (5/10)	66 (69/104)	105.0	(12/21) (9	(26/19/99	0.749	52 (16/31)	70 (58/83)	0.069	50 (22/44)	74 (52/70)	0008
BMI (kg/m ²)	23+4	20+3	24+4	900.0	22+4	24+4	0.075	22+4	24+3	0.078	22+4	24+3	0.005
SBP (mmHø)	118+16	8 +	119+15	0.161	110+16	120+15	0.005	117+17	119+15	0.554	14+16	120+15	0.049
DBP (mmHg)	70±10	66±7	71±11	0.129	68±10	71±11	0.289	68±8	71±11	0.122	69±10	71±11	0.347
HR (min ⁻¹)	76±14	70±14	76±14	0.186	74±15	76±14	0.580	71±10	78±15	0.020	75±12	77±17	0.475
Cr (µmol/l)	81 (66–95)	97 (62–118)	80 (68–95)	0.386	85 (69–110)	80 (66–95)	0.250	80 (65–102)	81 (66–95)	0.688	81 (66–100)	81 (66–95)	0.930
NT-Pro-BNP (pg/dl)	704	2190	620	0.026	1762	653	0.069	1205	572	0.119	1312	564	0.026
	(198–2170)	(585–9000)	(171–1876)		(368–5914)	(154–1803)		(360–3068)	(167–1584)		(391–3213)	(148–1540)	
NYHA classification (%)													
_	0 (1/114)	0 (0/10)	I (I/104)	0.757	0 (0/21)	1 (1/93)	0.633	0 (0/31)	1 (1/83)	0.539	0 (0/44)	1 (1/70)	0.434
=	47 (53/114)	(01/1) 1	50 (52/104)	0.015	29 (6/21)	51 (47/93)	0.068	35 (11/31)	51 (42/83)	0.150	41 (18/44)	50 (35/70)	0.343
=	44 (50/114)	(01/6) 06	39 (41/104)	0.002	57 (12/21)	41 (38/93)	0.174	55 (17/31)	40 (33/83)	0.149	48 (21/44)	41 (29/70)	0.509
١٧	9 (10/114)	0 (0/10)	10 (10/104)	0.305	14 (3/21)	8 (7/93)	0.323	10 (3/31)	8 (7/83)	0.835	II (5/44)	7 (5/70)	0.438
First diagnosis of HF in the past 18	61 (70/114)	40 (4/10)	63 (66/104)	0.145	57 (12/21)	62 (58/93)	0.657	71 (22/31)	58 (48/83)	0.200	61 (27/44)	61 (43/70)	0.994
months (%)													
Current smoker (%)	69 (79/114)	20 (2/10)	74 (77/104)	<0.00	29 (6/21)	78 (73/93)	<0.001	35 (11/31)	82 (68/83)	<0.00	30 (13/44)	94 (66/70)	<0.00
Diabetic (%)	13 (15/114)	0 (01/0) 0	14 (15/104)	0.197	14 (3/21)	13 (12/93)	0.866	16 (5/31)	12 (10/83)	0.566	18 (8/44)	10 (7/70)	0.208
Diagnosis of COPD (%)	4 (5/114)	10 (1/10)	4 (4/104)	0.364	14 (3/21)	2 (2/93)	0.014	10 (3/31)	2 (2/83)	0.092	7 (3/44)	3 (2/70)	0.315
Medication (%)													
BB	85 (97/114)	(01/6) 06	85 (88/104)	0.648	90 (19/21)	84 (78/93)	0.443	87 (27/31)	84 (70/83)	0.713	88 (39/44)	83 (58/70)	0.399
ACEI	50 (57/114)	40 (4/10)	51 (53/104)	0.508	38 (8/21)	53 (49/93)	0.227	35 (11/31)	55 (46/83)	0.058	45 (20/44)	53 (37/70)	0.442
ARB	23 (26/114)	0 (0/10)	25 (26/104)	0.072	5 (1/21)	27 (25/93)	0.029	16 (5/31)	25 (21/83)	0.299	14 (6/44)	29 (20/70)	0.064
MRA	72 (82/114)	60 (6/10)	73 (76/104)	0.258	62 (13/21)	74 (69/93)	0.379	71 (22/31)	72 (60/83)	0.889	73 (32/44)	71 (50/70)	0.881
Device	34 (39/114)	30 (3/10)	35 (36/104)	0.769	24 (5/21)	37 (34/93)	0.266	29 (9/31)	36 (30/83)	0.476	27 (12/44)	38 (27/70)	0.216
implantation*													
LAD (mm)	45±8	45±4	45±8	0.930	46±6	45±8	0.624	46±5	45±9	0.791	46±6	45±9	0.538

	64+8	6+69	64+8	0.038	67+8	64+8	0.040	65+8	64+8	0412	65+8	64+8	0.438
LVEF (%)	32±11	33±8	32±11	0.805	31±9	32±11	0.566	31±11	32±11	0.757	31±10	32±11	0.536
MAGGIC risk score 21±6	21±6	24±8	21±6	0.110	24±7	20±6	0.006	24±7	20±6	0.001	23±6	20±6	0.005
Note: *Device implantation includes pace maker, implantable cardioverter-defibrillator (ICD), cardiac resynchronization therapy (CRT) or cardiac resynchronization therapy defibrillator (CRT-D).	ion includes pace	maker, implantable	e cardioverter-defi.	brillator (IC	CD), cardiac resync	chronization ther	apy (CRT)	or cardiac resynchr	onization therapy-	defibrillator	· (CRT-D).		
Abbreviations: +, presenting adverse events; -, no adverse events; ACEI, angiotensin converting enzyme inhibitor; ARB, angiotensin receptor blocker; BB, beta blocker; BM, body mass index; COPD, chronic obstructive pulmonary disconting to the mass index; COPD, chronic obstructive pulmonary disconting to the mass index; COPD, chronic obstructive pulmonary disconting to the mass index; COPD, chronic obstructive pulmonary disconting to the mass index; COPD, chronic obstructive pulmonary disconting to the mass index; COPD, chronic obstructive pulmonary disconting to the mass index; COPD, chronic obstructive pulmonary disconting to the mass index; COPD, chronic obstructive pulmonary disconting to the mass index; COPD, chronic obstructive pulmonary disconting to the mass index; COPD, chronic obstructive pulmonary disconting to the mass index; COPD, chronic obstructive pulmonary disconting to the mass index; COPD, chronic obstructive pulmonary disconting to the mass index; COPD, chronic obstructive pulmonary disconting to the mass index; COPD, chronic obstructive pulmonary disconting to the mass index; COPD, chronic obstructive pulmonary disconting to the mass index; COPD, chronic obstructive pulmonary disconting to the mass index; CoPD, chronic obstructive pulmonary disconting to the mass index; CPP, chronic obstructive pulmonary disconting to the mass index; CPP, chronic obstructive pulmonary disconting to the mass index; CPP, chronic obstructive pulmonary disconting to the mass index; CPP, chronic obstructive pulmonary disconting to the mass index; CPP, chronic obstructive pulmonary disconting to the mass index; CPP, chronic obstructive pulmonary disconting to the mass index; CPP, chronic obstructive pulmonary disconting to the mass index; CPP, chronic obstructive pulmonary disconting to the mass index; CPP, chronic obstructive pulmonary disconting to the mass index; CPP, chronic obstructive pulmonary disconting to the mass index; CPP, chronic obstructive pulmonary disconting to the mass index; CPP, chronic obstruct	enting adverse eve	ents; –, no adverse d procenso: HE box	events; ACEI, ang	iotensin co.	No loft strict diam	hibitor; ARB, ang	giotensin re	ceptor blocker; BB	, beta blocker; BM	II, body mas	is index; COPD, o	hronic obstructive	e pulmonary

Group in Chronic Heart Failure; MRA, mineralocorticoid receptor anagonist; NT-Pro-BNP, N-terminal Pro brain natriuretic peptide; NYHA, New York Heart Association; SBP, systolic blood pressure.

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follow-up: $64 \pm 8 \text{ mm vs } 59 \pm 9 \text{ mm}, P < 0.001$) and increased LVEF (baseline vs 1-year follow-up: $32 \pm 11\%$ vs 41 \pm 14%, P < 0.001) (Supplementary Figure 1). Comparison of the 1-year changes in left heart remodeling parameters between M1 and M2 groups revealed that change in LVEDD was significantly less over 1 year in the M1 group than in the M2 group (M1 (n = 54) vs M2 (n = 50): -9 [-16 to -2] % vs -3 [-10 to 4] %, P = 0.006)(Supplementary Figure 2). One-year change of left atrial diameter and LVEF did not differ significantly between the two groups. Spearman correlation revealed that the 1-year change in LVEDD was significantly correlated with MAGGIC risk score (Supplementary Table 2). The MAGGIC risk model demonstrated modest ability to discriminate left ventricular remodeling (Figure 3 and Supplementary Table 3). Echocardiography parameters measurement demonstrated good consistency in 23 randomly selected patients (Supplementary Table 4).

Discussion

In this study, we found that MAGGIC risk score was independently related to adverse events in a cohort of DCM patients and appeared to have modest ability to discriminate adverse events in DCM patients. Meanwhile, MAGGIC risk score was also independently related to left ventricular remodeling parameters of DCM patients and had modest ability to discriminate left ventricular remodeling in DCM patients.

MAGGIC Risk Score and Adverse Events

Since its inception in 2013, the MAGGIC risk score has been validated several times for its ability to discriminate adverse events in HF patients with the ejection fraction reducing, ejection fraction preserving, and mixed types.^{14,15,22–25} Here we validated the prognostic value of MAGGIC risk score for adverse events of DCM patients. Our results were consistent with those of DCM patients reported by Moneghetti et al demonstrating that MAGGIC risk score was independently related to adverse events.¹⁶ However, the MAGGIC risk model demonstrated modest ability to discriminate adverse events in our study. Sawano et al reported that the MAGGIC risk score showed modest ability to discriminate 1-year mortality (area under the curve = 0.71), while Sartipy et al reported that the MAGGIC risk score was able to discriminate 1-year mortality (area under the curve = 0.743) in HF patients.^{13,25} The MAGGIC risk model could not predict adverse events in DCM patients in this cohort as strongly as in HF

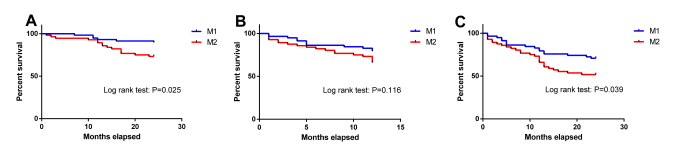


Figure I Kaplan-Meier curves of adverse events in the cohort of DCM patients (N = 114) during follow-up. (A) All-cause mortality within 2-year follow-up. (B) First rehospitalization due to heart failure within the I-year follow-up. (C) First rehospitalization due to heart failure within the 2-year follow-up. MI-M2: MAGGIC risk score of all DCM patients divided into two groups by median. Abbreviations: DCM, dilated cardiomyopathy; MAGGIC, Meta-Analysis Global Group in Chronic Heart Failure.

patients. Due to the small sample size of our study, future studies might examine this difference in predictive ability of adverse events in DCM versus HF patients.

MAGGIC Risk Score and Left Ventricular Remodeling

There has been a lack of studies examining the relationship between MAGGIC risk score and left ventricular remodeling in DCM patients. Some researchers reported that some biomarkers such as tenascin-c and galectin-3 were predictors of left ventricular remodeling or reverse remodeling in DCM patients.^{26,27} Other researchers reported that genotypes were predictors of left ventricular reverse remodeling in DCM patients.7,28,29 However, these biomarkers require extra examination and increase patient financial burdens, while all MAGGIC model risk factors are obtained from routine evaluations of DCM patients. We found that the MAGGIC risk score was related to left ventricular enlargement and demonstrated modest discrimination ability on left ventricular remodeling in DCM patients. This implies that the MAGGIC risk score could become a reference for the short-term evaluation of left ventricular remodeling status which is an indicator of the long-term prognosis of DCM patients.⁸⁻¹¹

MAGGIC Risk Model and Risk Factors Included

Risk factors included in the MAGGIC risk model were proven associated with adverse events in HF patients as well as DCM patients.^{16,30–33} And we could also infer that these risk factors were associated with left ventricular remodeling in DCM patients as well, for some of these risk factors were previously proven associated with left ventricular remodeling.^{6,34,35} Thus, the MAGGIC risk model generates a good combination of these risk factors for predicting adverse events and left ventricular remodeling. The easy availability of the model makes it a choice for evaluating DCM patients. Although there is limited translational benefit from the model's use in clinical practice, this study has inspirational meaning for future research on the model's usage. Notably, some researchers adjusted the MAGGIC risk model by including N-terminal-pro-brain natriuretic peptide or brain natriuretic peptide in the MAGGIC model, and the revised models demonstrated better prognostic value than original MAGGIC model in HF patients.^{12,13,36} Exploration of a unified model combining natriuretic peptide with the original MAGGIC risk model and validation of the adjusted model's prediction ability in DCM patients are also works that need to be done, especially in a DCM cohort with large sample size, in the future.

Limitations

This study has some limitations. First, it was a singlecenter study with a small sample size. And since the discriminative capacity of the model was simply weak, there is limited translational benefit from the model's use in clinical practice for now. More studies with large sample size and multi-center studies are needed to uncover the accurate discrimination ability of the model in adverse events and left ventricular remodeling. Second, DCM patients included in the study did not use a unified treatment plan, which could have created bias in our study. However, since personalized medicine becoming increasingly more popular, it would be difficult to ensure that all DCM patients use the same treatment approach. Our study could reveal a meaningful regular pattern in the real world. Third, the follow-up time in this study was short. The life expectancy of DCM patients would be prolonged with treatment advances. The MAGGIC risk model's ability to predict long-term survival could be tested in the future.

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	All-Cause Mortality				First Rehospitalization Due to HF	tion Due	to HF					
	2 Year (N = 21)				I Year (N=31)				2 Year (N=44)			
	Univariate		Multivariate		Univariate		Multivariate		Univariate		Multivariate	
	HR (95% CI)	Ь	HR (95% CI)	Ь	HR (95% CI)	Ч	HR (95% CI)	d	HR (95% CI)	A	HR (95% CI)	d
Age (years)	1.020 (0.984–1.059)	0.281			1.049 (1.015–1.085)	0.005*			1.031 1.005–1.059)	0.022*		
Male (%)	1.180 (0.489–2.846)	0.713			1.949 (0.963–3.994)	0.064			2.209 (1.222–3.993)	0.009*	2.418 (1.327-4.405)	0.004
BMI (kg/m ²)	0.888 (0.783–1.006)	0.063			0.909 (0.819–1.009)	0.073			0.878 (0.801–0.963)	0.006*		
SBP (mmHg)	0.955 (0.923-0.988)	0.007*	0.952 (0.921–0.983)	0.003	0.994 (0.971–1.018)	0.646			0.982 (0.962–1.003)	0.093		
HR (min ⁻¹)	0.991 (0.959–1.023)	0.576			0.968 (0.941–0.996)	0.025*	0.969 (0.942–0.997)	0.029	0.989 (0.969–1.010)	0.312		
Lg[NT-pro-BNP	1.838 (0.956–3.536)	0.068			1.567 (1.004–2.445)	0.048			1.626 (1.038–2.548)	0.034		
[(Ip/gd)												
Current	1.625 (0.656–4.027)	0.294			1.438 (0.689–3.001)	0.333			1.978 (1.034–3.781)	0.039		
smoker (%)												
Diagnosis of	4.128 (1.213–14.052)	0.023			0.395 (0.120-1.302)	0.127			0.513 (0.159–1.622)	0.266		
COPD (%)												
ARB (%)	6.564 (0.881–48.917)	0.066			1.644 (0.631–4.282)	0.309			2.079 (0.879–4.920)	0.096		
LVEDD (mm)	1.059 (1.004–1.118)	0.037			1.016 (0.972–1.062)	0.485			1.014 (0.977–1.053)	0.460		
LVEF (%)	0.990 (0.949–1.032)	0.633			0.997 (0.964–1.031)	0.860			0.993 (0.964–1.022)	0.611		
MAGGIC risk	1.104 (1.029–1.184)	0.006*	1.122 (1.043–1.208)	0.002	1.096(1.034–1.161)	0.002*	1.094 (1.032–1.158)	0.002	1.077 (1.026–1.131)	0.003*	1.088 (1.033–1.147)	0.002
score												
Note: *Variables in	Note: *Variables included into multivariate Cox regression model.	ox regression	on model.		Portonic of	structivo a	ulmonary discosso: HE hoo				NEDD Joff vontrij	- Puo

Abbreviations: ARB, angiotensin receptor blocker; BMI, body mass index; CJ, confidence interval; COPD, chronic obstructive pulmonary disease; HF, heart failure; HR, hazard ratio; HR (min⁻¹), heart rate; LYED, left ventricular end-diastolic diameter; LVEF, left ventricular ejection fraction; MaGGIC, Meta-Malysis Global Group in Chronic Heart Failure; NT-Pro-BNP, N-terminal Pro brain natriuretic peptide; SBP, systolic blood pressure.

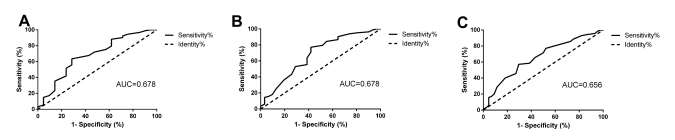


Figure 2 Ability of MAGGIC risk model to discriminate adverse events in dilated cardiomyopathy patients (N = 114). (A) All-cause mortality within the 2-year follow-up. (B) First rehospitalization due to heart failure within the 1-year follow-up. (C) First rehospitalization due to heart failure within the 2-year follow-up. Abbreviations: AUC, area under the curve; MAGGIC, Meta-Analysis Global Group in Chronic Heart Failure.

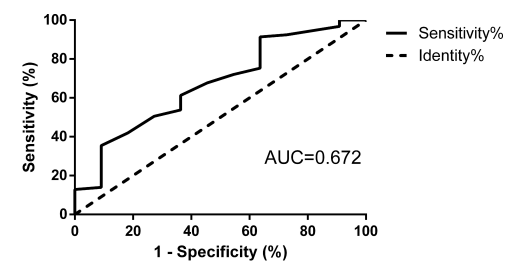


Figure 3 Ability of MAGGIC risk model to discriminate left ventricular remodeling within 1 year in dilated cardiomyopathy patients (N = 104). Abbreviation: MAGGIC, Meta-Analysis Global Group in Chronic Heart Failure.

Conclusions

In conclusion, the MAGGIC risk score was related to adverse events and left ventricular remodeling but demonstrated modest ability to discriminate adverse events and left ventricular remodeling in DCM patients. It has the potential to be a choice for evaluating the prognosis of DCM patients.

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Disclosure

The authors report no conflicts of interest in this work.

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