Management strategies in heart failure with preserved ejection fraction

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Abstract

The diagnosis and therapy of heart failure with preserved ejection fraction (HFpEF) remain challenging. Currently, there are ongoing discussions on whether the diagnosis of HFpEF should be based solely on left ventricular ejection fraction, which may not account for the heterogeneity of HFpEF syndrome. This aspect has been addressed by the recently proposed HFA-PEFF and the H2FPEF algorithms, which take numerous diagnostic modalities into account to establish the diagnosis of HFpEF. Moreover, this review focuses on the adequate treatment of comorbidities and risk factors in HFpEF that should be an essential part of any HFpEF therapy. Furthermore, the management of fluid level in HFpEF patients is pointed out, as it plays an important role in symptom control. In addition, the value of LCZ696 therapy in HFpEF is discussed. Although LCZ696 had neutral effects in the large PARAGON-HF trial, it had previously been granted an extended indication by the Food and Drug Administration. Since the publication of the EMPEROR-Preserved trial, empagliflozin now represents the first drug to significantly improve the prognosis of HFpEF patients. Therefore, the role of SGLT2 inhibitors in HFpEF management is highlighted. Overall, this review aims to enhance the knowledge on the diagnostic processes and best treatments available for HFpEF patients.

Keywords
Ventricular ejection fraction · Pharmacotherapy · HFpEF · SGLT2 inhibitor · Stroke volume

According to the 2021 European Society of Cardiology (ESC) guidelines [1], heart failure (HF) should be differentiated into three different forms depending on left ventricular ejection fraction (LVEF): heart failure with preserved ejection fraction (HFpEF, LVEF ≥ 50%), heart failure with reduced ejection fraction (HFrEF, LVEF ≤ 40%), and heart failure with mildly reduced ejection fraction (HFmrEF, LVEF 41–49%). Despite the latest breakthroughs in medical treatment, the diagnosis, as well as the therapy of the syndrome HFpEF, remains challenging. This review focuses on patient management strategies in HFpEF to accelerate and to enhance the diagnostic process and to provide the patients with the best treatment possible.

The heterogeneity of heart failure with preserved ejection fraction

As pointed out by the 2021 ESC guidelines [1], the potential causes of HFpEF are numerous. Typically, patients suffering from HFpEF are older and more often female compared with patients who have HFrEF and HFmrEF [2, 3]. Moreover, HFpEF patients commonly exhibit various comorbidities and cardiovascular risk factors such as chronic kidney disease, diabetes, hypertension, obesity, and deconditioning [4]. Furthermore, specific diseases may result in HFpEF. These comprise, for instance, primary cardiomyopathies, storage diseases such as Fabry’s disease and amyloidosis, or pericardial diseases such as constrictive pericarditis. Thus, rather than a single
clinical diagnosis, the term “HFrEF” describes a clinical syndrome with different underlying etiologies that require distinct therapies [5]. As a result, there are ongoing discussions on whether the definition of HFrEF should be based solely on LVEF [6]. To address this issue, two algorithms have been proposed recently—H2FPEF [7] and HFA-PEFF [8]—that consider numerous findings (noninvasive as well as invasive) besides LVEF to establish the diagnosis of HFrEF.

Diagnosis of HFrEF

Introduced by the Heart Failure Association (HFA) of the ESC, the HFA-PEFF score suggests a step-wise diagnostic approach to standardize and enhance the diagnosis of HFrEF [8]. The first step (P = pre-test assessment) can be performed in the ambulatory setting and takes into account clinical characteristics (HF symptoms and signs, prevalence of comorbidities) as well as standard diagnostic tests including natriuretic peptides, electrocardiogram, X-ray, and echocardiography. If positive, the authors recommend a risk stratification by natriuretic peptide levels and sophisticated echocardiographic work-up into three different groups: low, intermediate, and high risk (step E). While the diagnosis of HFrEF is confirmed in patients at high risk and is excluded in patients at low risk, patients at intermediate risk should undergo echo stress tests or, if those are inconclusive, invasive hemodynamic measurements in step 3 (F1). Lastly, in all patients confirmed with HFrEF, an etiological work-up should be pursued aimed at identifying specific causes of HFrEF (step F2). However, as acknowledged by the ESC guidelines, the broad application of this score is limited since some of the proposed tests can only be performed in specialized centers [1]. Therefore, the guidelines suggest a simplified algorithm for physicians with no access to this kind of expertise. In accordance with the HFA-PEFF algorithm, the first step of the simplified approach is to determine the pre-test probability according to the following three points: (1) symptoms and signs of HF, (2) LVEF ≥ 50%, and (3) objective evidence indicating the presence of LV diastolic dysfunction or raised LV filling pressures (summarized in Table 1; [1]).

The case of diagnostic uncertainty, a diastolic stress test is recommended. Although invasive hemodynamic exercise testing remains the confirmatory test for the diagnosis of HFrEF, its routine performance is not encouraged by the guidelines, in particular, due to its rather low availability worldwide and the risk of potential complications [1].

Therapy of HFrEF

Treatment of comorbidities and risk factors

Patients with HFrEF exhibit numerous comorbidities and risk factors that have been associated with increased morbidity and mortality [9, 10]. Therefore, the systematic screening and adequate treatment of these comorbidities and risk factors should represent a cornerstone of any HFrEF management strategy [11], as they have an important impact on patient prognosis (Fig. 1); this is also currently studied in the OPTIMIZE-HFrEF trial (NCT02425371). Obesity and deconditioning are frequently present in HFrEF patients. In a post hoc analysis of the I-PRESERVE trial, the majority of the 4109 patients included were overweight or obese according to their body mass index (BMI) of ≥ 26.5 kg/m² [12]. Interestingly, patients with higher BMI were younger, more likely to be female, hypertensive as well as diabetic, and had higher LVEF. Moreover, besides a low BMI of < 23.5 kg/m², the risk for the primary endpoint (death from any cause or hospitalization for HF, myocardial infarction, unstable angina, arrhythmia, or stroke) was significantly increased in obese HFrEF patients with a BMI of ≥ 35 kg/m² [12]. Accordingly, in a pooled analysis with a total of 96,424 HF patients (59,263 with HFrEF and 37,161 with HFrEF), the association between BMI and all-cause mortality was found to be U-shaped for both HFrEF and HFrEF patients, with a similar nadir of risk at a BMI of 32–33 kg/m² [13]. Thus, treating physicians should emphasize the importance of physical activity as well as a hypocaloric diet, which both have been shown to be beneficial in HFrEF. For instance, according to a sub-analysis of the TOPCAT trial, high physical activity led to a significantly decreased risk of the com-

### Table 1: Simplified algorithm for the diagnosis of HFrEF, as recommended by the 2021 ESC guidelines [3]

| Symptoms and signs of HFrEF | Dyspnea (NYHA II–III), orthopnea, reduction in physical capacity, … |
|----------------------------|---------------------------------------------------------------|
| Clinical signs             | Peripheral edema, visible engorgement of the neck veins, ascites, … |
| Comorbidities/risk factors | Hypertension, diabetes, obesity, atrial fibrillation, age, … |
| LVEF ≥ 50%                 | Exclusion of patients with a history of HFrEF who have shown an improvement in LVEF ≥ 50% → patients with recovered LVEF → continuation of treatment for HFrEF |
| LV hypertrophy             | LV mass index ≥ 95 g/m² (female), ≥ 115 g/m² (male) Relative wall thickness < 0.42 |
| LA enlargement             | LA volume index < 34 ml/m² (sinus rhythm) LA volume index < 40 ml/m² (atrial fibrillation) |
| Increased E/e′ ratio       | E/e′ ratio at rest > 9 |
| Increased natriuretic peptides | NT-proBNP > 125 pg/ml or BNP > 35 pg/ml (sinus rhythm) NT-proBNP > 365 pg/ml or BNP > 105 pg/ml (atrial fibrillation) |
| Increased pulmonary pressure | PA systolic pressure > 35 mm Hg TR velocity at rest > 2.8 m/s |

*BNP brain natriuretic peptide, HFrEF heart failure with preserved ejection fraction, LA left atrium, LV left ventricle, LVEF left ventricular ejection fraction, NP natriuretic peptides, NT-proBNP N-terminal prohormone of brain natriuretic peptide, PA pulmonary artery, TR tricuspid regurgitation

*Patients diagnosed with HFrEF have to exhibit (1) symptoms and signs of HFrEF, (2) LVEF ≥ 50%, and (3) cardiac structural and/or functional abnormalities.

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postive primary endpoint (HF hospitalization, cardiovascular mortality, or aborted cardiac arrest) in HFpEF patients [14]. Furthermore, physical activity in HFpEF has been assessed by the prospective, randomized controlled Ex-DHF pilot trial [15]. Compared to usual care, supervised exercise training reduced diastolic dysfunction in terms of E/e′ and improved exercise capacity, regardless of chronotropic competence. However, as recently shown in the OptimEx-Clin trial, advising HFpEF patients to be physically active is more important than creating mandatory training programs and permanent supervision of physical activity, which may be difficult to realize in clinical practice [16]. Accordingly, one-time advice on physical activity as recommended by the guidelines resulted in a similar increase in peak oxygen consumption (peak VO2) as opposed to high-intensity interval training (3 x 38 min/week) and moderate continuous training (5 x 40 min/week) among 180 HFpEF patients. To date, the data on caloric restriction in HFpEF patients are rather sparse and are mainly derived from a single study. In this randomized, prospective trial with 100 obese HFpEF patients, a 20-week hypocaloric diet led to a 7% decrease in body weight and an improvement in peak VO2 of 1.3 ml/kg/min [17]. In patients that additionally were assigned to perform supervised exercise training, body weight was decreased by even 10% and peak VO2 increased by 2.5 ml/kg/min.

Moreover, hypertension as well as diabetes are common comorbidities and risk factors in patients with HFpEF [5, 11]. Particularly since the publication of the EMPEROR-Preserved trial [18], the SGLT2-inhibitor empagliflozin should be the primary treatment option for diabetes in HFpEF patients. In March 2022, the European Commission granted an indication of empagliflozin for all adults with symptomatic chronic HF, which includes patients across the full spectrum of LVEF [19]. The EMPEROR-Preserved trial [18] and the value of SGLT2-inhibitors in HFpEF patients are discussed in greater detail in Sect. “Sodium glucose cotransporter 2 inhibitors”. As hypertension may lead to cardiac decompensation [20] and mainly contributes to the development of HFpEF, an adequate hypertension treatment represents an essential part of any HFpEF therapy. Although there is uncertainty about the optimal hypertensive therapy in HFpEF patients, the 2021 ESC guidelines recommend considering the same treatment strategy as in patients with HFrEF [1]. In a meta-analysis including 75 prospective, randomized comparative studies, beta-blockers or diuretics caused less LV mass regression than angiotensin receptor blockers (ARBs), angiotensin-converting enzyme inhibitors (ACEi), and calcium channel blockers (CCBs; [21]).

Furthermore, iron deficiency is commonly present in patients with HFpEF. According to a small study with 190 symptomatic HFpEF patients, severe diastolic dysfunction was more likely to be found in iron-deficient patients [22]. Also, the presence of iron deficiency correlated with reduced exercise capacity and quality of life (QOL). The effects of intravenous iron administration in HFpEF are currently unknown, while it has been shown to have beneficial effects on symptoms and QOL in HFrEF [23]. However, therapy with intravenously administered iron in HFpEF is being evaluated in the two ongoing randomized controlled trials: FAIR-HFpEF (NCT03074591) and PREFER-HF (NCT03833336).

Angiotensin receptor neprilysin inhibitor (LCZ696)

Besides the inhibition of the angiotensin II receptor type 1, LCZ696 inhibits the metalloprotease neprilysin, which results in increased levels of natriuretic peptides, and thus causes vasodilatation and increased diuresis [11]. Moreover, inhibition of neprilysin promotes the formation of cGMP, therefore interfering with the NO-
cGMP-PKG ("nitrergic oxide—cyclic guanosine monophosphate–protein kinase") axis, which is thought to play an essential part in the pathophysiology of HFpEF [24, 25].

However, in the randomized controlled PARAGON-HF trial [26] with 4822 HFpEF patients included (LVEF of at least 45%), treatment with LCZ696 failed to significantly reduce the risk of the primary endpoint of total HF hospitalizations or CV death.

Interestingly, data from a retrospective analysis question the pathophysiological rationale for neprilysin inhibition in HFpEF. Accordingly, in 144 HFpEF patients, increased neprilysin levels were not associated with a greater rate of HF hospitalizations or death [27]. By contrast, in patients with HFrEF, elevated neprilysin levels correlated with an unfavorable prognosis. In 1069 HFrEF patients, the risk of the combined endpoint of HF hospitalization and CV death was increased by 20% if the neprilysin serum level was above the median [28].

Nevertheless, a nonsignificant but nominal risk reduction of 13% was observed in the treatment arm of the PARAGON-HF study [26]. This was initially attributed to a more efficient blood pressure reduction with LCZ696, which is approximately twice as high as with valsartan [29]. However, a post hoc analysis of the PARAGON-HF trial showed that the aforementioned nominal risk reduction in the primary endpoint did not result from the greater decrease in systolic blood pressure with LCZ696 therapy [30].

Furthermore, in the PARAGON-HF study [26], the definition of HFpEF and thus the criteria for study inclusion were primarily based on the recorded LVEF, which had to be at least 45%. In addition, patients did not have to receive a mandatory etiological work-up prior to enrolment to exclude specific causes of HFpEF such as amyloidosis and Fabry’s disease, which could be resistant to treatment with LCZ696. Therefore, the inclusion criteria of the PARAGON-HF study did not allow for the heterogeneity of the HFpEF syndrome, representing a potential limitation.

This aspect appears to be further underlined by the results of the subgroup analyses of the PARAGON-HF study. Whereas women appeared to benefit significantly from the LCZ696 therapy, particularly through a reduction in HF hospitalizations, this was not the case in men [31]. Moreover, in patients with an LVEF between 45% and 57%, the risk of the primary endpoint was significantly decreased by 22% through LCZ696 compared with valsartan only. By contrast, in patients with an LVEF above the median of 57%, there were no significant differences regarding the primary endpoint between the intervention and the control arm [26].

Recently, data from PARAGON-HF and PARADIGM-HF were pooled in order to examine LCZ696 treatment effects across the spectrum of LVEF. In this analysis, the administration of LCZ696 led to a significant decrease of the primary endpoint up to an LVEF of 55% ([32]; Fig. 1). Similar results were obtained from post hoc analyses of other large HFpEF studies, suggesting that candesartan (CHARM-Preserved, [33]) and spironolactone (TOPCAT, [34]) have a significant treatment benefit for patients with an LVEF below 55% [35]. Furthermore, positive therapeutic effects have been demonstrated for beta-blocker therapy, up to an LVEF of <50% [36]. However, it should be noted that the LVEF range of 45–55% according to the ESC definition includes not only patients with HFpEF, but also those with HFrEF ([1]; Fig. 2).

In February 2021, the US Food and Drug Administration (FDA) granted an expanded indication to LCZ696 for patients with LVEF below normal (<50%) [37]. However, the FDA also pointed out that, “LVEF is a variable measure and the use of clinical judgment is essential in deciding whom to treat.” At the present time, the authors believe that therapy with LCZ696 may be beneficial in selected individuals. The decision, however, should not only include the reported LVEF, but should be based on as many diagnostic modalities as possible, such as those included in the HFA-PEFF [8] or H2PFEF [7] score.

### Sodium glucose cotransporter 2 inhibitors

SGLT2-inhibitors, such as empagliflozin, canagliflozin, or dapagliflozin, inhibit the sodium-glucose cotransporter 2 (SGLT2) of the proximal renal tubules, thereby reducing glucose reabsorption from the primary urine, and ultimately causing renal glucosuria. Thus, a decrease in blood glucose levels can be achieved. Consequently, these SGLT2-inhibitors (SGLT2i) have been initially studied as anti-diabetic drugs. As recently shown in two large randomized trials, treatment with either empagliflozin (EMPEROR-Reduced, [38]) or dapagliflozin...
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(DAPA-HF, [39]) is associated with a significant decrease in morbidity and mortality in patients suffering from HFrEF, irrespective of diabetes status. The underlying mechanisms remain the subject of ongoing discussions within the scientific community. These include an improvement in cardiac energetics through increased ketone body production and a decrease in preload and afterload [40, 41]. In 2021, the EMPEROR-Preserved trial investigated the effects of empagliflozin in 5988 symptomatic (NYHA stage III–IV) HFrEF patients (LVEF ≥ 40%), regardless of the presence or absence of diabetes [18]. Compared to placebo, therapy with empagliflozin resulted in a significant reduction of the primary endpoint (composite of CV death or HF hospitalization). According to the sub-analyses, the treatment effects were similar in HFrEF patients with and without diabetes. Furthermore, health-related QOL measured by the KCCQ score was significantly improved in the intervention arm. Of note, this effect was documented early on and was sustained for 1 year or longer [42]. In March 2022, the European Commission approved treatment with empagliflozin for all patients with symptomatic chronic HF, including patients across the full spectrum of LVEF [19]. Currently, there is another large randomized controlled trial (DELIVER; NCT03619213) studying the effects of dapagliflozin on a composite primary endpoint (CV death, HF hospitalization, urgent HF visit) in HFrEF patients (LVEF ≥ 40%).

To date, empagliflozin represents the first drug to significantly improve outcomes in HFrEF. It will be exciting to see whether these effects apply equally to other SGLT2i, in particular dapagliflozin, which will be clarified in the DELIVER trial [58]. Overall, the authors are convinced that empagliflozin should be the cornerstone of any HFrEF medical treatment, especially due to the lack of other evidence-based options.

Management of fluid level in HFrEF patients

Fluid overload may rapidly lead to signs and symptoms of congestion in HFrEF patients. Thus, diuretics remain an essential part of any HFrEF management. However, the monitoring of patients’ fluid status can be challenging and is often limited to traditional assessment tools such as daily weight controls [43, 44, 45]. Some telemonitoring devices on the other hand allow for the detection of impending cardiac decompensation at an early stage, and can therefore help to avoid HF hospitalizations. For instance, telemonitoring of the pulmonary arterial pressure using the CardioMEMS device has proven to be an effective therapeutic measure. After the convincing results of the U.S. CHAMPION study [46], the method was evaluated in Europe in the MEMS-HF trial [47]. This was a prospective, but not placebo-controlled, study in which 234 symptomatic HF patients were included. Of note, the study inclusion was independent of LVEF. Consequently, patients with HFrEF (LVEF < 40%) as well as HFrmEF and HFrEF (LVEF ≥ 40%) were included. Hospitalization rates due to HF were reduced by 62% after implantation of the CardioMEMS device [47]. In addition, an improvement in QOL was demonstrated after the initiation of CardioMEMS-guided therapy, which was objectively measured using the KCCQ score. Complications from CardioMEMS implantation and sensor failure were documented in only four patients (1.7%) and did not result in death. For further evaluation of hemodynamic-guided HF management, the randomized, placebo-controlled GUIDE-HF trial was initiated [48]. In this trial, 1022 symptomatic HF patients (NYHA II–IV) with either a recent HF hospitalization or elevated natriuretic peptides were enrolled. Surprisingly, CardioMEMS-guided HF management did not decrease the risk of the primary endpoint, which was a composite of all-cause mortality and total HF events (HF hospitalizations and urgent HF hospital visits) at 12 months [48]. However, these results may have been influenced considerably by the COVID-19 pandemic, thus representing an important limitation of the GUIDE-HF trial. Accordingly, in a pre-COVID-19 impact analysis, the risk of the primary endpoint was significantly lower in the intervention arm compared with usual care. This was primarily driven by the reduction of HF hospitalizations. During the COVID-19 pandemic, the primary event rate in the control arm decreased by 21%, while it did not change significantly in the intervention arm. As a result, the pre-COVID-19 treatment effects diminished [48]. Of note, these differences cannot be explained by changes in provider- or disease-dependent factors [49]. Thus, the outcomes in GUIDE-HF were most likely affected by changes in patient-dependent factors due to COVID-19, such as changes in behavioral patterns. Therefore, the impact of hemodynamic-guided HF management needs to be investigated in further trials.

Assumably, telemonitoring of fluid status to guide diuretic treatment in HF patients (HFrEF, HFrmEF, and HFrEF) will become increasingly important in the future. The authors encourage the implementation of home monitoring in current HFrEF management strategies, if available.

Interventional therapy of HFrEF

There are various interventional approaches to improve HFrEF therapy, which comprise, for instance, the implementation of atrial shunt devices (ASD), cardiac contractility modulation (CCM), cardiac resynchronization therapy (CRT), and the catheter-based denervation of renal sympathetic nerves [5]. Of these, the implementation of ASDs to decrease left atrial pressure by generating artificial left–right shunts has been the most promising. In the REDUCE LAP-HF I trial, this method has been shown to significantly reduce left atrial pressure during exercise [50] and to be safe [51]. However, just recently, the results of the REDUCE LAP-HF II trial [52] were published. In this randomized, multicenter, sham-controlled trial, a total of 626 symptomatic HFrEF patients (LVEF ≥ 40%) with increased pulmonary capillary wedge pressure (PCWP) during exercise (≥ 25 mmHg) were enrolled. The primary endpoint was a hierarchical composite of CV death or non-fatal ischemic stroke at 12 months, rate of total HF events up to 24 months, and change in KCCQ overall summary score at 12 months. Compared with sham-control, the intervention did not have any significant therapeutic effects, neither on the primary composite endpoint nor on its individual components [52].
Management of atrial fibrillation in HFrEF

Atrial fibrillation (AF) is a common and prognostically unfavorable concomitant disease in patients with HFrEF [1]. In a post hoc analysis of the TOPCAT trial, the occurrence of AF was independently associated with an increased risk of CV events (CV mortality, aborted cardiac arrest, or HF hospitalization; [53]).

Catheter-based pulmonary vein isolation (PVI) in patients with HFrEF has emerged as a cornerstone in AF therapy and improves clinical outcomes compared to medical treatment [54]. However, it is currently unclear whether patients with HFrEF can benefit equally from catheter ablation. Interestingly, data from the German ablation registry suggest that PVI for rhythm control is frequently performed in HFrEF patients [55]. Thus, randomized trials are desperately needed to clarify the role of PVI in patients suffering from HFrEF.

According to the ESC guidelines [56], if no further therapeutic attempts are made to preserve sinus rhythm and both the treating physician and the patient accept the presence of AF by consensus, so-called permanent AF is present. In this case, the primary objective of AF therapy is the control of ventricular heart rate, which should not exceed 110 bpm. In patients with HF, beta-blockers and digitalis glycosides are available for this purpose. The RATE-AF study [57] randomly compared the effects of bisoprolol and digoxin in patients with permanent AF and HF (defined as NYHA stage II–IV). The study included mostly patients with HFrEF (LVEF ≥ 50%), whereas only about 19% of patients had LVEF < 50%. With regard to the primary endpoint (increase in QOL objectively by the 36-item Short Form Health Survey physical component summary score), no significant differences were demonstrated in either treatment group. However, a decrease in NT-pro-BNP levels as well as serious clinical events (death, unplanned hospitalizations, treatment-related adverse events, primary care visits) was observed in patients treated with digoxin [57].

Conclusion

Heart failure with preserved ejection fraction (HFrEF) is a heterogeneous syndrome and a diagnosis based solely on LVEF may be insufficient. The diagnostic process should ideally use the recently introduced HFrEF and HFA-PEFF algorithms. HFrEF therapy must include the adequate treatment of comorbidities and risk factors, as they influence prognosis. Avoiding fluid overload by diuretic treatment to increase quality of life is an essential part of HFrEF therapy. If available, telemonitoring should be incorporated into HFrEF management to detect fluid overload before signs and symptoms of congestion. LCZ696 has been granted an expanded indication for patients with LVEF < 50% by the FDA. Empagliflozin is the first drug to significantly reduce morbidity and mortality in HFrEF patients and should be the cornerstone of any HFrEF treatment. The DELIVER trial will investigate whether this also applies to dapagliflozin. Further research is needed to enhance our understanding of the complex syndrome of HFrEF and help improve HFrEF management.

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Behandlungsstrategien bei Herzinsuffizienz mit erhaltener Ejektionsfraktion

Bis heute stellen Diagnose und Therapie der Herzinsuffizienz mit erhaltener linksventrikulärer Ejektionsfraktion (HFpEF) eine Herausforderung dar. Aufgrund der Heterogenität des HFpEF-Syndroms wird aktuell zunehmend kritisch darüber diskutiert, ob die Diagnose ausschließlich auf der linksventrikulären Ejektionsfraktion beruhen sollte. Dieser Aspekt wurde durch die kürzlich vorgeschlagenen HFA-PEEF- und H2FPEF-Diagnosealgorithmen aufgegriffen, die verschiedene Untersuchungsverfahren bei der HFpEF-Diagnose berücksichtigen. Darüber hinaus konzentriert sich diese Übersicht auf die adäquate Behandlung von Komorbiditäten und Risikofaktoren bei Patienten mit HFpEF, die eine Grundlage jeder HFpEF-Therapie darstellen sollten. Außerdem wird auf das Management des Volumenhaushalts bei HFpEF-Patienten eingegangen, durch das Symptome reduziert werden können. Zudem wird der Wert der LCZ696-Therapie bei HFpEF diskutiert. Obwohl LCZ696 in der großen PARAGON-HF-Studie nur einen neutralen Effekt aufwies, wurde die Zulassung in den USA erst kürzlich von der Food and Drug Administration (FDA) angepaßt und erweitert. Seit der Veröffentlichung der EMPEROR-Preserved-Studie ist Empagliflozin nun das erste Medikament, das die Prognose bei HFpEF-Patienten nachweislich signifikant verbessert. Daher wird die Rolle der SGLT2-Hemmer bei der Behandlung von Patienten mit HFpEF genauer beschrieben. Das Ziel dieser Übersicht besteht darin, Kenntnisse über eine optimale Diagnostik sowie eine bestmögliche Therapie für HFpEF-Patienten zu vermitteln.

Schlüsselwörter
Ventrikuläre Ejektionsfraktion · Pharmakotherapie · HFpEF · SGLT2-Inhibitor · Schlagvolumen
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