Heart failure (HF) is an important cardiovascular disease because of its increasing prevalence, significant morbidity, high mortality, and rapidly expanding health care cost. The number of HF patients is increasing worldwide, and Korea is no exception. There have been marked advances in definition, diagnostic modalities, and treatment of HF over the past four decades. There is continuing effort to improve risk stratification of HF using biomarkers, imaging and genetic testing. Newly developed medications and devices for HF have been widely adopted in clinical practice. Furthermore, definitive treatment for end-stage heart failure including left ventricular assist device and heart transplantation are rapidly evolving as well. This review summarizes the current state-of-the-art management for HF and the emerging diagnostic and therapeutic modalities to improve the outcome of HF patients.

Keywords: Heart failure; Diagnosis; Management; Prognosis

INTRODUCTION

Heart failure (HF) is an important cardiovascular disease due to its increasing prevalence and high mortality rate. HF is associated with a diverse range of complications, such as hospitalization, lethal arrhythmia, and death during the disease progression. In addition, HF can be the terminal condition of many cardiovascular diseases, including myocardial infarction (MI), valvular heart disease, and various cardiomyopathies. Due to these unique characteristics, various pharmacological and non-pharmacological treatments have been developed, not only to improve underlying cardiac disease but also to prevent hospitalization and death. In this review, we will address the state-of-the-art management of HF and ongoing studies.
treatment, and prognosis of HFmrEF remain unclear [1].

The increasing amount of research in this area has gradually led to elucidation of the characteristics of HFmrEF, although there is still a great deal of debate. HFmrEF is a heterogeneous group comprised of at least three subsets: HFmrEF improved group (prior LVEF < 40%), HFmrEF unchanged group (prior LVEF 40% to 49%), and HFmrEF deteriorated group (prior LVEF ≥ 50%). The improved and deteriorated groups account for 90% of the total, with the unchanged group accounting for only 10% of cases. Therefore, most patients classified as HFmrEF have various features of HFrEF and HFpEF, rather than belonging to groups with unique characteristics. These results indicate the limitations of the current HF classification system based on LVEF and, therefore, a novel taxonomy is required [4].

Several terms are often used to describe the status of HF patients. Patients with low LVEF and no symptoms or signs of HF can be described as having asymptomatic left ventricular (LV) systolic dysfunction. Patients experiencing HF symptoms and signs are described as having chronic HF, and those whose disease status does not change for at least 1 month are diagnosed as having stable HF. Patients with chronic stable HF showing unexpected deterioration are described as having decompensated HF. The New York Heart Association (NYHA) functional classification has been used to describe the severity of HF symptoms, and the Killip classification is used to describe the disease severity in patients after acute MI.

Epidemiology
The prevalence of HF differs according to definition and region, but has been estimated to be approximately 1% to 2% in developed countries (Fig. 1). The prevalence rate tends to increase with age, and it is > 10% among people > 70 years old [5]. The epidemiological and etiological profiles of HFrEF and HFpEF are different. In comparison with HFrEF, patients with HFpEF are older, show female predominance, and often show hypertension and atrial fibrillation (AF) with a lower rate of MI [6].

In Korea, the prevalence of HF in 2013 was 1.53% according to the National Health Insurance Service-National Sample Cohort [7]. The increasing prevalence of HF with age in Korea is consistent with the worldwide trend, and the rate was reported to be 1.0% for individ-
There is considerable evidence on how to prevent the development of overt HF. Hypertension is one of the major risk factors of HF. Many studies have indicated that proper control of blood pressure can prevent HF. In the Systolic Blood Pressure Intervention Trial (SPRINT), intensive blood pressure control (systolic blood pressure < 120 mmHg) was more beneficial for the prevention of cardiovascular diseases than standard treatment, but controversy over optimal blood pressure persists because the study population had a high risk of cardiovascular disease [9]. With regard to diabetes mellitus, recent studies have shown that sodium-glucose cotransporter-2 (SGLT-2) inhibitors reduced mortality and HF hospitalization in type 2 diabetic patients. In patients with ST-segment elevation MI, rapid primary percutaneous coronary intervention and administration of angiotensin-converting enzyme inhibitor (ACEI), β-blocker, mineralocorticoid receptor antagonist (MRA), and statin can reduce HF hospitalization and mortality. In addition, ACEI and β-blocker can decrease mortality in patients with coronary artery disease (CAD) without LV systolic dysfunction and reduce HF hospitalization in patients with asymptomatic LV systolic dysfunction, regardless of the etiology [1].

EMERGING CARDIAC IMAGING AND OTHER DIAGNOSTIC TESTS

Cardiac imaging enables us to visualize and quantify the structure and function of the heart. LVEF is not only the most widely used indicator of systolic function but is also one of the most powerful prognostic markers in patients with HF. As LVEF is crucial for diagnosis, classification, and establishment of an appropriate management plan, transthoracic echocardiography is the most important tool in the initial evaluation of HF patients. Recently, strain measured by echocardiography and other modalities such as cardiovascular magnetic resonance (CMR) and computed tomography (CT) have attracted attention. In this section, we review emerging imaging modalities in the field of HF.
Strain imaging

Left ventricular strain
LVEF can divide the HF population into two groups according to prognosis and drug responsiveness: HFrEF vs. HFpEF. In HF patients with LVEF ≥ 45%, mortality does not show a negative correlation with LVEF [10,11]. Moreover, most drugs used for HF show a mortality benefit only in patients with HFrEF, and not in those with HfmrEF and HFpEF [12]. This is because LVEF cannot discriminate subclinical LV dysfunction from normal systolic function. In addition, substantial numbers of patients with cardiomyopathy other than ischemic cardiomyopathy have preserved systolic function. Therefore, there is a need for a more accurate prognostic marker to classify HF patients, and LV global longitudinal strain (GLS) is expected to be useful in this regard [13,14]. Park et al. [14] demonstrated the utility of LV-GLS as a prognostic indicator in patients with acute HF throughout all ranges of LVEF. They demonstrated that the rate of all-cause death was well stratified according to GLS, but not to LVEF [14]. With regard to the specific etiology of HF, LV-GLS worse than −14% was an independent predictor of all-cause mortality and hospitalization for HF in acute MI with LVEF > 40% [15]. In patients receiving potentially cardiotoxic chemotherapy, baseline and follow-up measurement of LV-GLS are recommended because LV-GLS can detect early subclinical LV dysfunction in these patients [16].

Right ventricular strain
The RV has a complex structure that causes difficulty in the estimation of systolic function. The pathophysiology of RV dysfunction is also complex, because RV function is influenced by a wide range of factors, such as volume status, pulmonary vascular resistance, and LV function [17]. Therefore, many studies have failed to show the predictive value of RV function in patients with HF [18]. RV strain has recently become one of the most actively investigated topics in the measurement of RV function [19]. Iacoviello et al. [20] reported that RV-GLS and RV free wall strain were independent predictors of all-cause mortality in patients with chronic stable HF. The most recent study, including 1,824 patients with acute HF, showed that the group with decreased biventricular strain had the poorest prognosis. In addition, poorer RV-GLS was an independent predictor of all-cause mortality in acute HF patients without pulmonary hypertension [21]. Limitations, such as intervendor variability, the impact of age and sex, load dependency, and variable cutoff value, make GLS difficult to incorporate into daily practice; however, accumulating evidence may result in more universal usage of GLS in the future [13,16,22].

CT and CMR
Cardiac CT can be used to evaluate the coronary arteries in HF patients with low to intermediate pretest probability of CAD or equivocal noninvasive stress test results. CMR has the unique advantage of myocardial tissue characterization, which helps to determine the etiology of various cardiomyopathies and estimate myocardial viability. However, the role of imaging modalities other than transthoracic echocardiography was restricted in the 2016 European Society of Cardiology (ESC) guidelines to estimation of the etiology in HF patients [1].

With recent progress in CMR techniques, additional information about myocardial tissue has become available. Pre- and post-contrast T1 time and extracellular volume (ECV) are known to be related to myocardial tissue composition [23]. Mascherbauer et al. [24] reported a negative correlation between post-contrast T1 time and E/e’ in 61 HFpEF patients with diagnoses confirmed by right heart catheterization. They also showed that short post-contrast T1 time (< 388.3 ms) was an independent predictor of HF hospitalization and cardiovascular mortality [24]. In a prospective cohort study of 117 non-ischemic dilated cardiomyopathy (NIDCM) patients, ECV was a better prognostic marker of cardiac events than the presence of mid-wall late gadolinium enhancement (LGE) [25]. However, the difference in administration, distribution, and excretion of gadolinium can cause potential variability of post-contrast T1 time and ECV and, therefore, native T1 has emerged as a new alternative method [26]. In 637 NIDCM patients, native T1 was an independent predictor of all-cause and HF composite endpoint, while post-contrast T1 was not [27]. Furthermore, efforts are being made to use CT for tissue characterization. Prior studies have shown that CT could be used to evaluate myocardial delayed enhancement for non-ischemic cardiomyopathy as well.
as MI, through similar contrast kinetics of iodine to gadolinium. Moreover, the recently developed dual-energy technique could improve the image quality of CT for evaluation of myocardial delayed enhancement through monochromatic imaging. Mapping of iodine distribution within the myocardium using the dual-energy technique could improve diagnostic accuracy for evaluation of myocardial perfusion and ECV fraction [28-30]. In patients with HF, myocardial scarring detected by CT is similar to the presence of LGE by CMR [31]. Despite limitations including radiation exposure and a shortage of experience, CT may supersede CMR in the future due to its versatility, especially in the field of ischemic cardiomyopathy [32].

**Novel biomarkers and genetic testing**

**Biomarkers**

As there is a considerable body of evidence for NP biomarkers, they have already been incorporated into the American expert consensus for HF [33]. In the European guidelines, NPs are essential elements for the diagnosis of HFrEF and acute HF [1].

There are increasing numbers of novel biomarkers. The biomarkers for myocardial injury and fibrosis, such as soluble suppression of tumorigenicity-2 (ST2), galectin-3, and high-sensitivity cardiac troponin, can be used for additive risk stratification in HF patients [33, 34]. Soluble ST2, a marker for cardiac stress and fibrosis, showed good performance for additive risk stratification of death in acute HF in an individual patient-based meta-analysis [35]. In addition, a smaller decrease in soluble ST2 48 hours after presentation was an independent predictor of 1-year mortality in patients with acute HF who visited the emergency department [36]. The value of soluble ST2 as a prognostic indicator was also demonstrated in ambulatory HF patients [37]. Notably, soluble ST2 can be useful in patients with renal insufficiency because it is not influenced by renal function [37, 38]. Galectin-3 is a marker of inflammation and fibrosis. It can also provide prognostic information in patients with acute or chronic HF and identify patients who can benefit from medical therapy [34,39]. In addition, galectin-3 can also be a potential treatment target for fibrosis. Several galectin-3 inhibitors are being developed in experimental studies [39]. In addition, other new biomarkers, such as procalcitonin, mid-regional pro-atrial NP, and growth differentiation factor-15, are also under investigation [34,40]. Individual novel biomarkers may not be useful as prognostic indicators, but combinations of these biomarkers can show better performance in the prediction of mortality [41].

Nevertheless, the results of studies regarding biomarker-guided therapy were variable [42]. NP-guided pharmacological therapy reduced all-cause mortality and HF hospitalization in a meta-analysis of 2,686 patients with chronic HF [43]. However, Guiding Evidence Based Therapy Using Biomarker Intensified Treatment in Heart Failure (GUIDE-IT), the most recent multi-center randomized controlled trial (RCT), yielded disappointing results. In that study, NT-proBNP-guided therapy did not show significant benefits compared with usual care in high-risk HFrEF patients [44]. There were a number of possible reasons for this failure, including: (1) the patients allocated to the biomarker-guided group in the clinical trials already had a low level of NP because they generally received strict guideline-directed medical therapy (GDMT) in advance; (2) the additional treatment in patients with high NP levels was limited to increases or addition of diuretics; and (3) it was not clear how much low NP level is needed to improve the patient outcomes. Therefore, it would be erroneous to conclude that biomarker-guided therapy is useless. GDMT has not been used in real-world situations due to the gap between guidelines and clinical practice [45,46]. Thus, biomarker-guided therapy may still have room for improvement.

**Genetic testing**

Inherited cardiomyopathies account for a small portion of HF cases. However, such cases cannot be ignored because the diagnosis, clinical manifestations, and treatment are different from those of other types of HF. For example, a patient with Anderson-Fabry disease can show systemic manifestations, including chronic kidney disease with proteinuria and stroke. The diagnosis requires confirmation by a specific test for enzyme activity or genetic testing and should be treated with enzyme replacement therapy [47]. Accordingly, genetic testing provides accurate information regarding the etiology of HF, potentially followed by a specific therapy and better clinical outcome [48,49].
Recently, the Heart Failure Society of America published new guidelines for genetic testing in patients diagnosed with genetic cardiomyopathy [50]. The guidelines were developed according to evidence about the validity and influence of genetic testing on the prognosis of diseases. Briefly, genetic testing is recommended in patients with hypertrophic cardiomyopathy (HCM), NIDCM, arrhythmogenic right ventricular cardiomyopathy, cardiomyopathy with other extracardiac manifestations, and LV non-compaction. Careful family history taking, phenotypic screening, and genetic counseling are recommended for family members of patients with genetically confirmed inherited cardiomyopathies.

In addition, a new classification method of genetic cardiomyopathy was proposed. Traditionally, the type of cardiomyopathy was defined by morphology, i.e., dilated, hypertrophic, and restricted cardiomyopathy. The European and American guidelines utilized complex classification methods because a wide variety of mutations as well as non-genetic cardiomyopathy can show similar findings [51,52]. Recent progress in testing methods, such as next-generation sequencing, revealed that several genetic mutations are responsible for single inherited cardiomyopathy and that there are many subclinical mutation carriers in the population. Therefore, a new classification incorporating these factors was required. The MOGE(S) classification includes morphofunctional phenotype (M), involved organ system (O), genetic inheritance pattern (G), etiology (E), and functional status (S) of the disease. This new classification integrates the phenotype, genotype, and function of an individual patient. Therefore, it has advantages regarding standardization and expandability of nomenclature [53].

**PHARMACOLOGICAL TREATMENT**

**Conventional pharmacological treatment**

Pharmacological treatment of HF has been evolving through increased understanding of its pathophysiology and the development of new drugs (Table 1 [54-89], Fig. 3). Before the 1980s, treatment of HF largely depended on bed rest and fluid restriction, and only digitalis and diuretics were prescribed for HF patients. As decreased LV contractility was thought to be the main cause of HF, there were many attempts to improve contractility by administration of inotropic drugs to HF patients. However, most clinical trials of inotropic agents were stopped prematurely because of their poor results [90]. From the mid-1970s, vasodilators have been used to increase cardiac efficiency by reducing afterload, and the first large RCT of vasodilator treatment in HF was published in 1986. In this study, vasodilator treatment improved mortality, but subsequent larger studies suggested that it was not beneficial for the long-term survival of HF patients [54].

Since the 1980s, HF has been understood as a neurohormonal disease, and physicians attempted to improve cardiac function by blocking the renin-angiotensin-aldosterone system and sympathetic activation. ACEI was reported to reduce mortality and hospitalization in HFrEF patients [91], and β-blockers also improved LV function and reduced mortality and hospitalization rates [92]. Unlike ACEI, β-blockers do not have a class effect, and evidence of beneficial effects in treatment of HF have been reported only for bisoprolol, sustained-release metoprolol, carvedilol, and nebivolol [93,94]. Subsequently, additional prognostic benefits were reported when MRA was combined with ACEI and β-blocker therapy [95]. Angiotensin receptor blockers also showed beneficial effects on prognosis in patients with HFrEF and has been used as an alternative treatment option when ACEI cannot be used [96]. Combination therapy targeting the neurohormonal system significantly improved the prognosis of patients with HFrEF compared to the use of vasodilator and inotropic agents and is still the mainstay of pharmacological treatment of HF [94].

In the 2000s, several new medications were introduced for treatment of HF. Ivabradine is an inhibitor that acts on the If channel of the sinoatrial node, which reduces heart rate independently of β-blocker use. In patients with HFrEF, ivabradine improves the clinical outcome in cases with resting heart rate > 70 bpm despite adequate β-blocker therapy [55]. Tolvaptan is an oral vasopressin-2 receptor antagonist with excellent diuresis activity and is expected to be beneficial in decongestion of acute heart failure (AHF) patients [97]. As many AHF patients have renal dysfunction, resistance to diuretics, and electrolyte imbalance such as hyponatremia, tolvaptan seemed to be more beneficial in these patients [98]. However, tolvaptan failed to show a posi-
Table 1. Landmark pharmacological and non-pharmacological studies of heart failure

| Topic | Study | Year | Author/Group | Patients | No. | Intervention | Comparator | Primary end-point | Outcome | Important secondary outcome | FU duration |
|-------|-------|------|-------------|----------|-----|--------------|------------|-------------------|---------|----------------------------|-------------|
| Drug  | V-HeFT [54] | 1986 | Cohn et al. | LVEF < 45%, DCMP | 642 | ISDN/hydralazine | Prazocin or placebo | All-cause death | RRR 0.34 (0.04–0.54; vs. placebo, at 2 yr) | 2.3 yr (mean) |
|       | CONSENSUS [57] | 1987 | CONSENSUS Trial Study group | NYHA IV | 253 | Enalapril | Placebo | All-cause death | RRR 0.40 at 6 mon (p = 0.002) | RRR for mortality at 1 yr 0.31 (p = 0.001) | 188 day (mean) |
|       | SOLVD [58] | 1991 | SOLVD Investigators et al. | LVEF ≤ 35%, NYHA II–IV | 2,569 | Enalapril | Placebo | All-cause death | RRR 0.16 (0.05–0.26) | RRR for all-cause death or HF hospitalization: 0.26 (0.18–0.34) | 4 yr |
|       | USCP [59] | 1996 | Packer et al. | LVEF ≤ 35% | 1,094 | Carvedilol | Placebo | All-cause death | RRR 0.65 (0.39–0.80) | RRR for CV death or hospitalization: 0.38 (0.18–0.53) | 12 mon |
|       | RALES [60] | 1999 | Pitt et al. | LVEF ≤ 35%, NYHA III–IV | 1,663 | Spironolactone | Placebo | All-cause death | RR 0.70 (0.60–0.82) | CV death or hospitalization RR: 0.68 (0.59–0.78) | 2 yr |
|       | ATLAS [61] | 1999 | Packer et al. | LVEF ≤ 30%, NYHA II–IV | 3,164 | High dose lisinopril | Low dose lisinopril | All-cause death | HR 0.92 (0.82–1.03) | HR for all-cause death or hospitalization for any cause: 0.88 (0.82–0.96) | 45.7 mon (median) |
|       | CHARM [62] | 2003 | Granger et al. | LVEF ≤ 40%, NYHA II–IV | 2,028 | Candesartan | Placebo | CV death, HF hospitalization | HR 0.70 (0.60–0.81) | HR for CV death: 0.80 (0.66–0.96) | 33.7 mon (median) |
|       | SHIFT [55] | 2010 | Swedberg et al. | LVEF ≤ 35%, NYHA II–IV | 6,558 | Ivabradine | Placebo | HF death or hospitalization | HR 0.82 (0.75–0.90) | HR for HF hospitalization: 0.74 (0.66–0.83) | 22.9 mon (median) |
|       | PARADIGM-HF [56] | 2014 | McMurray et al. | LVEF ≤ 40%, NYHA II–IV | 8,399 | Valsartan/sacubitril | Enalapril | CV death or hospitalization for HF | HR 0.80 (0.73–0.87) | HR for all-cause death: 0.84 (0.76–0.93) | 27 mon (median) |
|       | EMPA-REG OUTCOME [63] | 2015 | Zinman et al. | T2DM at high risk for CV events | 7,020 | Empagliflozin | Placebo | CV death, nonfatal MI, nonfatal stroke | HR 0.89 (0.74–0.99) | HR for all-cause death: 0.68 (0.57–0.82) | 3.1 yr (median) |
|       | LEADER [64] | 2016 | Marso et al. | T2DM at high risk for CV events | 9,340 | Liraglutide | Placebo | CV death, nonfatal MI, nonfatal stroke | HR 0.87 (0.78–0.97) | HR for all-cause death: 0.85 (0.74–0.97) | 3.8 yr (median) |

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| Topic | Study | Year | Author | Patients | No. | Intervention | Comparator | Primary endpoint | Outcome | Important secondary outcome | FU duration |
|-------|-------|------|--------|----------|-----|-------------|------------|-----------------|---------|---------------------------|------------|
| CANVAS [65] | 2017 | Neal et al. | T2DM at high risk for CV events | 10,142 | Canagliflozin | Placebo | CV death, nonfatal MI, nonfatal stroke | HR 0.86 (0.75–0.97)<sup>a</sup> | HR for CV death: 0.87 (0.72–1.06)<sup>b</sup> | HR for amputation: 1.97 (1.41–2.75)<sup>c</sup> | 188 wk (mean) |
| CANTOS [66] | 2017 | Ridker et al. | History of MI patients and elevated hsCRP | 10,061 | Canakinumab | Placebo | CV death, nonfatal MI, nonfatal stroke | HR 0.85 (0.74–0.98)<sup>a</sup> | Increase fatal infection or sepsis (0.31 per 100 person/yr vs. 0.18 per 100 person/yr, p = 0.02)<sup>c</sup> | 3.7 yr (median) |
| DECLARE-TIMI 58 [67] | 2018 | Wiviott et al. | T2DM with multiple risk factors for CV disease or known CV disease | 17,160 | Dapagliflozin | Placebo | CV death, MI, ischemic stroke | HR 0.93 (0.84–1.03)<sup>a</sup> | HR for CV death and HF hospitalization: 0.83 (0.73–0.95)<sup>a</sup> | 4.2 yr (median) |
| ATTR-ACT | 2018 | Maurer et al. | TTR cardiac amyloidosis with HF | 441 | Tafamidis | Placebo | All-cause death | HR 0.70 (0.51–0.96)<sup>a</sup> | 30 mon |
| ICD MADIT [69] | 1996 | Moss et al. | Prior MI (3 wk), LVEF ≤ 35%, NYHA I–III, asymptomatic NSVT or inducible VT | 196 | ICD | Medical therapy | All-cause death | HR 0.46 (0.26–0.82)<sup>a</sup> | 27 mon (mean) |
| MUSTT [70] | 1999 | Buxton et al. | CAD, LVEF ≤ 40%, asymptomatic NSVT and inducible sustained VT | 704 | Antiarrhythmic drugs or ICD | Medical therapy | Cardiac arrest, arrhythmic death | HR 0.73 (0.53–0.99)<sup>a</sup> | HR for ICD vs. no ICD: 0.24 (0.13–0.45)<sup>a</sup> | 39 mon (median) |
| MADIT II [71] | 2002 | Moss et al. | Prior MI (≥ 1 mon), LVEF ≤ 35% | 1,232 | ICD | Medical therapy | All-cause death | HR 0.69 (0.51–0.93)<sup>a</sup> | 20 mon (mean) |
| Topic       | Study       | Year | Author              | Patients | No.  | Intervention | Comparator | Primary end-point                  | Outcome                        | Important secondary outcome                | FU duration |
|------------|-------------|------|---------------------|----------|------|--------------|------------|-----------------------------------|----------------------------------|-------------------------------------------|--------------|
| DEFINITE   | Kadish et al. | 2004 | NIDCM, LVEF ≤ 36%   | 229      | ICD  | Medical therapy | All-cause death | HR 0.65 (0.40–1.06)  | HR for arrhythmic SCD: 0.20 (0.66–0.71) | 29 mon (mean) |
| DINAMIT    | Hohnloser et al. | 2004 | Recent MI (6 to 40 day), LVEF ≤ 35%, NIDCM | 674      | ICD  | Medical therapy | All-cause death | HR 1.68 (0.76–1.55)  | HR for arrhythmic death: 0.42 (0.22–0.83) | 30 mon (mean) |
| SCD-HeFT   | Bardy et al.  | 2005 | NYHA II–III, LVEF ≤ 35% | 2,521    | Amiodarone ICD | Medical therapy | All-cause death | Amiodarone HR 1.06 (0.86–1.30) | ICD HR 0.77 (0.62–0.96) | 46 mon (median) |
| DANISH     | Kober et al.  | 2016 | NYHA II–III, or CRT candidate with IV, LVEF ≤ 35%, NIDCM | 1,116    | ICD (including CRT if indicated) | Medical therapy (including CRT if indicated) | All-cause death | HR 0.87 (0.68–1.12)  | HR for sudden death: 0.50 (0.31–0.82)     | 68 mon (median) |
| VEST       | Olgin et al.  | 2018 | Recent MI, LVEF ≤ 35% | 2,302    | Wearable ICD | Medical therapy | Sudden death or death from ventricular tachyarrhythmia | RR 0.67 (0.37–1.21)  |                                      |                                      |
| CRT        | Bristow et al. | 2004 | NYHA III–IV, LVEF ≤ 35%, QRSd ≥ 120 ms, sinus | 1,520    | CRT-P, CRT-D | Medical therapy | All-cause death, hospitalization | CRT-P HR 0.81 (0.69–0.96) | CRT-D HR 0.80 (0.68–0.99) | Medical 11.9 mon CRT-P 16.2 mon CRT-D 15.7 mon |
| CARE-HF    | Cleland et al. | 2005 | NYHA III–IV, LVEF ≤ 35%, QRSd ≥ 120 ms, sinus, LVESD ≥ 30 mm | 813      | CRT  | Medical therapy | All-cause death, hospitalization for CV event | HR 0.63 (0.51–0.77)  | HR for all-cause death: CRT-P 0.76 (0.58–1.01) | 29 mon (mean) |
| REVERSE    | Linde et al.  | 2008 | NYHA I–II, LVEF ≤ 40%, QRSd ≥ 120 ms, sinus, LVEDD ≥ 55 mm | 610      | CRT-on, CRT-off | HF clinical composite response (worse) | Worsened primary end-point: 16% vs. 21% (p = 0.10) | HR for HF hospitalization: 0.47 (p = 0.03) | 12 mon  |
| Topic     | Study        | Year | Author          | Patients               | No.  | Intervention                      | Comparator | Primary endpoint                              | Outcome                           | Important secondary outcome       | FU duration |
|-----------|--------------|------|-----------------|------------------------|------|----------------------------------|------------|---------------------------------------------|-----------------------------------|----------------------------------|--------------|
| MADIT-CRT | [80]         | 2009 | Moss et al.     | NYHA I–II, LVEF ≤ 35%, QRSd ≥ 130 ms, sinus | 1,820 | CRT-D                           | ICD        | All-cause death or HF                     | HR 0.66 (0.52–0.84)\(^a\)       | HR for all-cause death: 1.00 (0.69–1.44)\(^b\) | 2.4 yr (mean) |
| RAFT      | [81]         | 2010 | Tang et al.     | NYHA II–III, LVEF ≤ 30%, QRSd ≥ 120 ms, sinus or AF | 1,798 | CRT-D                           | ICD        | All-cause death or HF hospitalization     | HR 0.75 (0.64–0.87)\(^a\)       | HR for all-cause death: 0.75 (0.62–0.91) | 40 mon (mean) |
| VAD       | REMATCH      | 2001 | Rose et al.     | NYHA IV, LVEF ≤ 25%, ineligible for HT | 129  | Axial pulsatile flow LVAD (HeartMate) | Medical therapy | All-cause death                           | RR 0.52 (0.34–0.78)\(^a\)       | RR for serious adverse event: 2.35 (1.86–2.93)c | NA           |
| HMII-BTT  | [82]         | 2007 | Miller et al.   | NYHA IV, awaiting HT  | 333  | Axial continuous flow LVAD (HeartMate II) | Status at 180 day (HT, recovery, on mechanical support) | Survival during support: 68% at 6 mon | Improved NYHA functional class and quality of life at 3 mon\(^a\) | Continuous 126 day (median) |
| HMII-DT   | [84]         | 2009 | Slaughter et al.| NYHA IIIB–IV, LVEF ≤ 25%, ineligible for HT | 200  | Axial continuous flow LVAD (HeartMate II) | Axial pulsatile flow LVAD (HeartMate II | Disabling stroke or device removal due to malfunction at 2 yr | HR 0.38 (0.27–0.54)\(^a\)       | Less infection, right HF, respiratory failure, renal failure, rehospitalization\(^c\) | Continuous 1.7 yr Pulsatile 0.6 yr (median) |
| ENDURANCE | [85]         | 2017 | Rogers et al.   | NYHA IIIB–IV, LVEF ≤ 25%, ineligible for HT | 446  | Centrifugal continuous flow LVAD (HeartWare) | Axial continuous flow LVAD (HeartMate II) | Disabling stroke or device removal due to malfunction at 2 yr | Noninferior (p = 0.01 for noninferiority)\(^b\) | More stroke (29.7% vs. 12.1%, p < 0.001)c | NA (2 yr) |
| MOMENTUM 3| [86]         | 2017 | Mehra et al.    | Advanced HF patients who are ineligible or waiting for HT | 294  | Centrifugal continuous flow LVAD (HeartMate III) | Axial continuous flow LVAD (HeartMate II) | Disabling stroke or device removal due to malfunction at 6 mon | Noninferior (p < 0.001 for noninferiority)\(^b\) | No pump thrombosis (6% vs. 10.3%, p < 0.001) | NA (6 mon) |
| CABG      | STICH [87]   | 2016 | Velazquez et al.| LVEF ≤ 35%, CAD amenable to CABG | 1,212 | CABG                            | Medical therapy | All-cause death                           | HR 0.86 (0.72–1.04)\(^b\)       | HR for all-cause death or hospitalization due to CV cause: 0.74 (0.64–0.85)\(^a\) | NA           |
### Table 1. Continued

| Topic | Study | Year | Author | Patients | No. | Intervention | Comparator | Primary end-point | Outcome | Important secondary outcome | FU duration |
|-------|-------|------|--------|----------|-----|--------------|------------|-------------------|---------|-----------------------------|-------------|
| HT    | SCHEDULE | 2014 | Andreassen et al. | De novo HT recipient | 115 | Everolimus with low dose cyclosporin | Standard dose cyclosporin | GFR at 12 mon | 79.8 mL/min/1.73 m² vs. 61.5 mL/min/1.73 m² (p < 0.001)<sup>a</sup> | Less cardiac allograft vasculopathy (50.0% vs. 64.6%, p = 0.003)<sup>a</sup> | NA (12 mon) |
|       | PROCEED II | 2015 | Ardehali et al. | HT recipient | 130 | Organ Care System | Standard cold storage | Patient and graft survival at 30 day | 94% vs. 97% (p = 0.45)<sup>b</sup> | NA | |

FU, follow-up; V-HeFT, Vasodilator-Heart Failure Trial I; LVEF, left ventricular ejection fraction; DCMP, dilated cardiomyopathy; ISDN, isosorbide dinitrate; RRR, relative risk reduction; CONSENSUS, Cooperative North Scandinavian Enalapril Survival Study; NYHA, New York Heart Association; SOLVD, Studies of Left Ventricular Dysfunction; HF, heart failure; USCP, U.S. Carvedilol Program; CV, cardiovascular; RALES, Randomized Aldactone Evaluation Study; RR, relative risk; ATLAS, Assessment of Treatment with Lisinopril and Survival; MADIT, Multicenter Automatic Defibrillator Implantation Trial; HR, hazard ratio; CHARM, Candesartan in Heart failure-Assessment of mortality and Morbidity; SHF, Systolic Heart failure treatment with the If inhibitor ivabradine Trial; PARADIGM-HF, Prospective Comparison of ARNI with ACEI to Determine Impact on Global Mortality and Morbidity in Heart Failure; EMPA-REG OUTCOME, Empagliflozin Cardiovascular Outcome Event Trial in Type 2 Diabetes Mellitus; T2DM, type 2 diabetes mellitus; MI, myocardial infarction; LEADER, Liraglutide Effect and Action in Diabetes: Evaluation of Cardiovascular Outcome Results; CANVAS, Canagliflozin Cardiovascular Assessment Study; CANTOS, Canakinumab Anti-inflammatory Thrombosis Outcome Study; hsCRP, high-sensitivity C-reactive protein; DECLARE-TIMI 58, Dapagliflozin Effect on Cardiovascular Events Thrombolysis in Myocardial Infarction 58; ATHER-HEFT, Transhyretin Amyloidosis Cardiomyopathy Clinical Trial; TTR, transthyretin; ICD, implantoable cardioverter-defibrillator; MADIT, Multicenter Automatic Defibrillator Implantation Trial; NSVT, nonsustained ventricular tachycardia; VT, ventricular tachycardia; MUSTT, Multicenter Unstained Tachycardia Trial; CAD, coronary artery disease; DEFINITE, Defibrillators in Non-Ischemic Cardiomyopathy Treatment Evaluation; NIDCM, nonischemic dilated cardiomyopathy; SCD, sudden cardiac death; DINAMIT, Defibrillator in Acute Myocardial Infarction Trial; SCD-HeFT, Sudden Cardiac Death in Heart Failure Trial; DANISH, Danish Study to Assess the Efficacy of ICDs in Patients with Non-ischemic Systolic Heart Failure on Mortality; CRT, cardiac resynchronization therapy; VEST, Vest Prevention of Early Sudden Death Trial; COMPANION, Comparison of Medical Therapy, Pacing, and Defibrillation in Heart Failure; QRSD, QRS duration; CRT-P, cardiac resynchronization therapy without defibrillator; CRT-D, cardiac resynchronization therapy with defibrillator; CARE-HF, Cardiac Resynchronization in Heart Failure; IVEED, left ventricular end-systolic dimension; REVERSE, Resynchronization reVerses Remodeling in Systolic left ventricular dysfunction; RAFT, Resynchronization–Defibrillation for Ambulatory Heart Failure Trial; AF, atrial fibrillation; HT, heart transplantation; VAD, ventricular assisted device; REMATCH, Randomized Evaluation of Mechanical Assistance for the Treatment of Congestive Heart Failure; LVAD, left ventricular assist device; HMII-BTT, HeartMate II bridge to transplant; HMII-DT, HeartMate II destination therapy; ENDURANCE, The HeartWare™ Ventricular Assist System as Destination Therapy of Advanced Heart Failure; NA, not available; MOMENTUM 3, Multicenter Study of MagLev Technology in Patients Undergoing Mechanical Circulatory Support Therapy with HeartMate 3; CABG, coronary artery bypass surgery; STICH, Surgical Treatment for Ischemic Heart Failure; SCHEDULE, Scandinavian Heart Transplant Everolimus De novo study with Early Calcineurin Inhibitors Avoidance; GFR, glomerular filtration rate; PROCEED II, ex vivo perfusion of donor hearts for human heart transplantation.

<sup>a</sup>Positive result.

<sup>b</sup>Neutral results.

<sup>c</sup>Serious adverse event.
Figure 3. Progression of heart failure treatment: medications, devices, and transplantation. HT, heart transplantation; TAH, total artificial heart; V-HeFT, Vasodilator-Heart Failure Trial I; CONSENSUS, Cooperative North Scandinavian Enalapril Survival Study; SOLVD, Studies of Left Ventricular Dysfunction; USCP, U.S. Carvedilol Program; RALES, Randomized Aldactone Evaluation Study; ATLAS, Assessment of Treatment with Lisinopril and Survival; MADIT, Multicenter Automatic Defibrillator Implantation Trial; MUSTT, Multicenter Unstained Tachycardia Trial; VAD, ventricular assisted device; MMF, mycophenolate mofetil; CHARM, Candesartan in Heart failure-Assessment of moRtality and Morbidity; SHIFT, Systolic Heart failure treatment with the I2 inhibitor ivabradine Trial; PARADIGM-HF, Prospective Comparison of ARNI with ACEI to Determine Impact on Global Mortality and Morbidity in HF; EMPA-REG, Empagliflozin Cardiovascular Outcome Event Trial in Type 2 Diabetes Mellitus; CANVAS, Canagliflozin Cardiovascular Assessment Study; LEADER, Liraglutide Effect and Action in Diabetes: Evaluation of Cardiovascular Outcome Results; CANTOS, Canakinumab Anti-inflammatory Thrombosis Outcome Study; ATTR-ACT, Transthyretin Amyloidosis Cardiomyopathy Clinical Trial; DECLARE, Dapagliflozin Effect on Cardiovascular Event Results; DEFINITE, Defibrillators in Non-Ischemic Cardiomyopathy Treatment Evaluation; DINAMIT, Defibrillator in Acute Myocardial Infarction Trial; COMPANION, Comparison of Medical Therapy, Pacing, and Defibrillation in Heart Failure; SCHEDULE, Scandinavian Heart Transplant Everolimus De novo study with Early Calcineurin Inhibitors Avoidance; PROCEED II, ex vivo perfusion of donor hearts for human heart transplantation; ENDURANCE, The HeartWare Ventricular Assist System as Destination Therapy of Advanced Heart Failure; MOMENTUM 3, Multicenter Study of MagLev Technology in Patients Undergoing Mechanical Circulatory Support Therapy with HeartMate 3; ICD, implantable cardioverter-defibrillator; CRT, cardiac resynchronization therapy.
tive result in several clinical trials in AHF, and further studies are needed, therefore, to determine its applicability in the treatment of HF [99].

The Prospective Comparison of ARNI with ACEI to Determine Impact on Global Mortality and Morbidity in HF (PARADIGM-HF) study published in 2014 demonstrated the clinical efficacy of angiotensin receptor-neprilysin inhibitor (ARNI) in HFrEF [56]. ARNI is a combination of valsartan and the neprilysin inhibitor, sacubitril. Neprilysin is an endopeptidase that degrades several endogenous vasoactive peptides, including NPs, bradykinin, and adrenomedullin. Sacubitril increases the concentrations of these peptides by inhibiting neprilysin, and peptides antagonize the opposite action of neurohormonal overactivation [100]. Experimental studies have suggested that simultaneous inhibition of the renin-angiotensin system and neprilysin can more effectively decrease neurohormonal activation, which aggravates HF. As the combination of ACEI and sacubitril increased serious angioedema, valsartan and sacubitril constituted ARNI. In a large double-blind RCT, ARNI was superior to ACEI alone in reducing the risk of death and hospitalization for HFrEF patients [56]. ARNI is already being used for HFrEF patients as a standard practice, and clinical trials are underway regarding the role of ARNI in HFP EF patients and post-MI patients.

While there have been steady developments regarding pharmacological treatment for use in patients with HFrEF, no drugs have shown clear mortality benefits in patients with HFP EF. At present, drug therapy in HFP EF is focused on controlling symptoms and treating risk factors and comorbidities [12].

**Emerging pharmacological treatment**

**SGLT-2 inhibitor and glucagon-like peptide 1 agonist**

The SGLT-2 inhibitors represent a novel class of antihyperglycemic agents that increase urinary excretion of glucose in the renal tubules [101]. Empagliflozin showed good outcomes in large RCT (Empagliflozin Cardiovascular Outcome Event Trial in Type 2 Diabetes Mellitus Patients [EMPA-REG OUTCOM E]). Empagliflozin reduced HF hospitalization and cardiovascular death in patients with type 2 diabetes, with a consistent benefit in patients with HF. The serious adverse event rate of empagliflozin was similar, but the rate of genital infection was higher than placebo [63]. Canagliflozin also reduced the risk of cardiovascular death and hospitalization due to HF in patients with type 2 diabetes and elevated risk of cardiovascular disease (Canagliflozin Cardiovascular Assessment Study [CANVAS] trial). However, administration of canagliflozin increased the incidence of volume depletion, fracture, and amputation compared with placebo [65]. In addition, dapagliflozin also reduced the rates of cardiovascular death and hospitalization for HF in a recent RCT (Dapagliflozin Effect on Cardiovascular Events-Thrombolysis in Myocardial Infarction 58 [DECLARE-TIMI 58]) [67].

SGLT-2 inhibitors induce glycosuria and diuresis, which can be expected to reduce blood pressure, improve glycemic control, result in weight loss, and improve insulin sensitivity [102]. It has been reported that SGLT-2 inhibitors have cardioprotective effects by improving cardiac metabolism. A study in a murine model showed that empagliflozin increases cardiac adenosine triphosphate (ATP) production by activating cardiac oxidation of glucose and fatty acids [103], although the precise underlying mechanism is not fully understood yet.

Large-scale cardiovascular outcome studies using other SGLT-2 inhibitors are underway [104], and a multinational observational study, including South Korea, is currently in progress [105,106]. Moreover, the benefit of SGLT-2 inhibitors may persist regardless of the presence of diabetes. Empagliflozin significantly reduced the rate of cardiac deterioration in HF without diabetes in a murine model [107]. Human clinical studies on the effects of SGLT-2 inhibitors in HF without diabetes are also underway; DAPA-HF (NCT03036124) for dapagliflozin, EMPEROR-Reduced (NCT03057977) and EMPEROR-Preserved (NCT03057951) for empagliflozin [108].

Liraglutide, a glucagon-like peptide 1 analog, showed decreased all-cause mortality and cardiovascular death compared with placebo in a large-scale RCT [64]. As the U.S. Food and Drug Administration requires cardiovascular safety data for any new antidiabetic medications before approval, novel antidiabetic medications with additional cardiovascular benefits are likely to be developed in the future as well.
New drugs in specific cardiomyopathy fields

New drugs that act on cardiac myosin have been developed and tested for efficacy in specific diseases. Mavacamten, which acts as an inhibitor of cardiac myosin ATPase and reduces cardiac contractility, is under investigation in patients with obstructive HCM [109]. PIONEER-HCM (NCT02842244) is a phase 2 trial of mavacamten. In a pilot study performed in 11 patients with symptomatic obstructive HCM, significant decreases in both post-exercise peak and resting LV outflow tract gradient were observed in patients with mavacamten treatment [110]. With these positive results, a phase 3 study of EXPLORER-HCM is currently underway (NCT03470545).

In contrast to mavacamten, omecamtiv mecarbil (OM) is a selective cardiac myosin activator that increases myocardial systolic function. When administered to patients with AHF, the clinical effect of OM in relieving dyspnea was not clear (Acute Treatment With Omecamtiv Mecarbil to Increase Contractility in Acute Heart Failure [ATOMIC-AHF]) [111]. However, patients with chronic HF showed positive results after OM treatment with increased cardiac function, decreased ventricular dimension, and decreased serum NT-proBNP level (Chronic Oral Study of Myosin Activation to Increase Contractility in Heart Failure [COSMIC-HF]) [112]. The Global Approach to Lowering Adverse Cardiac Outcomes Through Improving Contractility in Heart Failure (GALACTIC-HF; NCT02929329) trial is currently underway to determine the clinical role of OM in comparison with placebo when added to current HF standard medication in patients with chronic HF [113].

The Transthyretin Amyloidosis Cardiomyopathy Clinical Trial (ATTR-ACT) showed remarkable results in transthyretin amyloid cardiomyopathy. Fibrillogensis in amyloid cardiomyopathy occurs when the tetrameric structure of the transthyretin protein dissociates into intermediates, which misassemble into amyloid fibrils. Tafamidis binds to the thyroxine-binding sites of transthyretin with high affinity and selectivity, and inhibits dissociation of tetramers into monomers. In this multicenter, double-blind RCT, tafamidis showed reductions in all-cause mortality and cardiovascular-related hospitalization rates and reduced the decline in functional capacity compared with placebo [68].

Anti-inflammatory therapy

Various cytokines have been shown to play important roles in determining cardiac function under pathophysiological conditions. Several cytokines, including tumor necrosis factor α, transforming growth factor β, and interleukins (ILs), such as IL-1, IL-4, IL-6, IL-8, and IL-18, are involved in the development of various inflammatory cardiac pathologies. There have been many clinical trials to improve cardiac pathology by blocking these cytokines, but most have failed to demonstrate clinical efficacy [114].

Anti-inflammatory therapy using canakinumab, a monoclonal antibody targeting IL-1β, led to a significantly lower rate of recurrent cardiovascular events in patients with previous MI compared to placebo. However, because canakinumab caused serious infectious complications, the all-cause mortality rate was not different from the placebo group (Canakinumab Anti-inflammatory Thrombosis Outcome Study [CANTOS] trial) [66]. The IL-1 receptor antagonist, anakinra, is another potential candidate for anti-inflammatory therapy. Administration of anakinra showed improvement of peak VO₂ in recently decompensated systolic HF patients (Recently Decompensated Heart Failure Anakinra Response Trial [REDHART]) [115], but did not lead to changes in HF-pEF patients ( Decompensated Heart Failure Anakinra Response Trial 2 [DHART2]) [116]. It is unclear whether anakinra is effective in preventing cardiac remodeling after acute MI, but a multicenter, double-blind placebo-controlled clinical trial is currently underway (Virginia Commonwealth University-Anakinra Remodeling Trial 3 [VCU-ART3] trial; NCT01950299) [117].

Gene therapy

Cardiac gene therapy, involving the production of proteins with curative efficacy by transferring specific exogenous genes, was proposed as an important alternative therapeutic approach [118]. The major targets of gene therapy are the β-adrenergic system, angiogenesis, cytoprotection, and stem cell homing, among which research on Ca²⁺ cycling protein is a representative project. The sarcoplasmic/endoplasmic reticulum Ca²⁺-ATPase (SERCA2a) regulates the contraction and relaxation of myocardial cells by transporting calcium into the sarcoplasmic reticulum from the cytosol. Regardless of the etiology, it has been demonstrated that SERCA2a is
deficient in experimental models of HF, and correction of SERCA2a deficiency can improve calcium influx and cardiac function. However, a recent study showed that administration of SERCA2a does not improve the clinical course of HFrEF patients [119]. Despite disappointing results, gene therapy still has potential and further studies are required. One of the major obstacles to gene therapy is the delivery method of the therapeutic materials into the target cells. As intravenous injection did not show sufficient effect to transduce the myocardium, intracoronary injection, myocardial injection, and pericardial injection have been suggested according to the condition of the patient, the type of vector, and the target gene. Further advances in vectors and delivery methods will be essential for the clinical application of gene therapy [120].

**Stem cell therapy**

As there is no alternative way to regenerate or replace damaged cardiomyocytes, there has been a great deal of interest in the development of stem cell therapy [121]. Numerous studies yielded optimistic results using stem cells to improve myocardial function and ventricular remodeling, but the results were inconsistent [122]. Human pluripotent stem cells (hPSCs) have emerged to replace embryonic stem cells, which maintain the similarity to embryonic stem cells but without the ethical issues or risks of rejection [123]. The efficiency of differentiation from hPSCs to cardiomyocytes and bioengineering technology to improve the therapeutic effects of hPSC-derived cardiomyocytes have improved over the past several decades. However, cellular heterogeneity, immaturity, arrhythmogenicity, and tumorigenicity are problems that remain to be resolved [123].

**DEVICES AND NON-SURGICAL INTERVENTIONS**

**Implantable cardioverter-defibrillator & cardiac resynchronization therapy**

Many pioneering landmark trials from the 1990s have confirmed the efficacy of implantable cardioverter-defibrillator (ICD) and cardiac resynchronization therapy (CRT) for improving cardiovascular outcome in HF patients (Table 1, Fig. 3). Current ESC guidelines generally suggest ICD for primary prevention in symptomatic HF patients with LVEF ≤ 35%, despite > 3 months of optimal medical therapy [1]. The detailed indications are slightly different according to the specific cardiomyopathy in patients (Table 2) [1,124,125]. For example, HCM with a high risk of sudden cardiac death, dilated cardiomyopathy due to lamin A/C (LMNA) mutation, cardiac sarcoidosis with unexplained syncope, or myocardial scarring seen on CMR, are indications for ICD regardless of LVEF [126]. However, the Danish Study to Assess the Efficacy of ICDs in Patients with Non-ischemic Systolic Heart Failure on Mortality (DANISH) trial, which

| Table 2. Current class I indications of cardiac implantable electronic devices in patients with heart failure |
|---------------------------------------------------------------------------------------------------|
| **ICD for secondary prevention**                          | **ICD for primary prevention**                             | **CRT**                                                  |
| ACC/AHA (2013) [125]                                       | NIDCM or ICM at least 40 days post-MI on chronic GDMT with (1) LVEF ≤ 35% and NYHA class II or III symptom (I-A) or (2) LVEF ≤ 30% and NYHA class I symptom (I-B) | Sinus rhythm with LVEF ≤ 35% on GDMT and LBBB with QRSd ≥ 150 ms and NYHA class III (I-A) or ambulatory IV (I-A) or II (I-B) |
| ESC (2016) [1]                                             | Recovery from ventricular arrhythmia with hemodynamic instability (IA) | Symptomatic HF (NYHA II–III) with LVEF ≤ 35% despite ≥ 3 months of OMT in ICM (IA) or NIDCM (IB) | Symptomatic HF with sinus rhythm and LVEF ≤ 35% despite OMT with LBBB with QRSd ≥ 150 ms (I-A) or 130–149 ms (I-B) |

ICD, implantable cardioverter-defibrillator; CRT, cardiac resynchronization therapy; ACC/AHA, American College of Cardiology/American Heart Association; NIDCM, nonischemic dilated cardiomyopathy; ICM, ischemic cardiomyopathy; MI, myocardial infarction; GDMT, goal-directed medical therapy; LVEF, left ventricular ejection fraction; NYHA, New York Heart Association; LBBB, left bundle branch block; QRSd, QRS duration; ESC, European Society of Cardiology; HF, heart failure; OMT, optimal medical therapy.
included 1,116 nonischemic cardiomyopathy patients, showed no mortality benefit with prophylactic ICD implantation [75], which reflects the different risk of sudden cardiac death in nonischemic cardiomyopathy and ischemic cardiomyopathy. However, as more than half of the patients received optimal medical therapy and CRT in both groups, this may have influenced the lack of significant results associated with prophylactic ICD. Therefore, the role of prophylactic ICD implantation to reduce mortality in HF patients may be reestablished with the further development of HF therapy. The application of wearable ICD in patients with recent MI and LVEF ≤ 35% did not show significant differences compared with GDMT, with regard to rates of sudden death or death from ventricular tachyarrhythmia at 90 days [76].

The generally accepted class I indications for CRT are restricted to symptomatic patients with LVEF ≤ 35%, despite optimal medical therapy, who have left bundle branch block and QRS duration ≥ 130 ms [127]. In Korea, there has been a rapid increase in the implantation of cardiac implantable electronic devices (Fig. 4).

Recent studies have focused on determining ways to achieve a high response rate in patients with CRT implants. Based on LGE detected by CMR, which can directly visualize myocardial scarring, patients receiving LV lead placement away from LGE showed better clinical outcome [128]. In the Targeted Left Ventricular Lead Placement to Guide Cardiac Resynchronization Therapy (TARGET) study, 220 patients were randomly assigned to two groups with or without targeted LV lead deployment using speckle-tracking 2-dimensional radial strain measured by transthoracic echocardiography. The targeted LV lead placement group showed a higher portion of CRT responders and lower rate of combined endpoint compared with the control group [129]. The combination of multimodality imaging, including nuclear imaging and radial strain, demonstrated a higher response rate compared with the control group, but the clinical outcomes were similar between the two groups [130].

**Role of AF ablation in HF**

AF is the most common type of arrhythmia in HF populations, and it can cause deterioration of LV function as well as symptoms of HF [131]. Rate control and anticoagulation are the mainstays of AF treatment. The 2016 ESC guidelines specify that a rhythm control strategy using amiodarone or AF ablation may be considered in chronic AF patients with HFrEF who are symptomatic despite optimal medical therapy (Class IIb) [1]. However, a recently published RCT showed a beneficial effect of AF ablation in HFrEF patients compared with pharmacological control (Catheter Ablation for Atrial Fibrillation with Heart Failure [CASTLE-AF] trial). In this RCT, patients who received AF ablation therapy showed lower rates of all-cause mortality, worsening or hospitalization for HF, and cardiovascular death after about 3 years [132]. Another RCT (Ablation versus Amiodarone for Treatment of Atrial Fibrillation in Patients With Congestive Heart Failure and an Implanted ICD/CRTD [AATAC] trial) showed that AF ablation was superior to amiodarone treatment with regard to maintenance of sinus rhythm, unplanned hospitalization, and mortality [133]. However, the CASTLE-AF trial included a highly selected population (398 of 3,013 screened patients), and a beneficial effect of AF ablation was observed in patients with young age (< 65 years), NYHA functional class II, LVEF ≥ 25%, and without diabetes [132]. The results of the Catheter Ablation Versus Anti-arrhythmic Drug Therapy for Atrial Fibrillation (CABANA) trial (NCT00911508), which enrolled 2,204 patients ≥ 65 years or with more than one risk factor for stroke, did not show a beneficial effect of AF...
Ablation in patients with high risk of stroke. The composite primary endpoints consisting of death, disabling stroke, serious bleeding, or cardiac arrest at 5 years were similar between ablation and drug therapy groups (hazard ratio, 0.86; 95% confidence interval, 0.65 to 1.15 for intention-to-treat analysis) [134]. If a patient has HF and symptomatic AF but is not a candidate for AF ablation or has failed to respond to this treatment, atrioventricular junction ablation followed by CRT may be a useful treatment option. Despite the small size of the study population, the Ablate and Pace in Atrial Fibrillation plus Cardiac Resynchronization Therapy (APAF-CRT) trial showed a decreased rate of hospitalization for HF and improved quality of life in patients undergoing atrioventricular junction ablation and CRT [135].

Remote monitoring
Remote monitoring has become one of the most active fields in the management of HF. Due to the importance of volume status in HF patients, there have been efforts to estimate and use hemodynamics as a guide for treatment. The CardioMEMS (St. Jude Medical, St. Paul, MN, USA) Heart Sensor Allows Monitoring of Pressure to Improve Outcomes in NYHA Class III Heart Failure Patients (CHAMPION) trial, a prospective RCT enrolling 550 symptomatic patients with chronic HF regardless of LVEF, showed that hemodynamic monitoring with a wireless implantable pulmonary artery pressure monitoring system (CardioMEMS) could significantly reduce HF hospitalization rate [136]. In the subgroup of 445 HFrEF patients on GDMT in the CHAMPION study, pulmonary artery pressure-guided management also reduced HF hospitalization and mortality rates [137]. Based on the previous results, CardioMEMS was approved by the U.S. Food and Drug Administration in 2014 and was included in the 2016 European guideline (Class IIb) [1].

Large amounts of information are available from patients with a preexisting cardiac implantable electronic device, including heart rate, lead profile, battery status and arrhythmic events. In the INfluence of home moniToring on mortality and morbidity in heart failure patients with IMpaired LEft ventricular function (IN-TIME) trial, without the need for additional invasive procedures, 664 patients with LVEF ≤ 35% implanted with ICD or CRT were randomized into two groups with or without telemonitoring, and a significantly lower rate of clinical composite endpoint was observed in the telemonitoring group at 1 year [138]. Remote monitoring also decreased the time to clinical decision and the length of hospital stay for cardiovascular hospitalization in the Clinical Evaluation of Remote Notification to Reduce Time to Clinical Decision (CONNECT) trial [139]. However, a meta-analysis including nine RCTs of remote monitoring showed no additional benefit with regard to survival or patient safety [140].

The diagnostics included in implantable devices show good predictive capability for impending HF decompensation using device-specific algorithms, such as OptiVol (Medtronic, Minneapolis, MN, USA) [141]. However, the OptiLink HF study did not show a significant difference in mortality according to alerts regarding changes in intrathoracic impedance reflecting patients’ fluid status [142]. The Multisensor Chronic Evaluation in Ambulatory Heart Failure Patients (MultiSENSE) study showed that HeartLogic multisensory index and alert algorithm (Boston Scientific, Marlborough, MA, USA) can be a good predictor of HF decompensation [143]. Based on these findings, the Multiple Cardiac Sensors for Management of Heart Failure (MANAGE-HF) trial in which patients were randomized according to whether HeartLogic alerts were turned on or off is currently ongoing (NCT03237858). As methodologies for remote monitoring using implantable devices can be extended, we expect better results in the near future.

Percutaneous correction of functional mitral regurgitation
Functional or secondary mitral regurgitation (MR), frequently accompanied by HFrEF, is a meaningful predictor of mortality after adjusting for clinical, echocardiographic, or laboratory variables [144]. However, surgical treatment of moderate ischemic MR in addition to coronary artery bypass surgery failed to show LV reverse remodeling and mortality improvement [145]. As the benefit of surgery for functional MR is questionable, the American guidelines for valvular heart disease published in 2014 recommended mitral valve surgery for secondary MR only in patients with symptomatic severe MR or moderate MR undergoing other cardiac surgery [146]. Percutaneous approaches to correct secondary MR in HF patients are actively studied because
the devices for percutaneous treatment of MR have been advanced throughout repair, annuloplasty, and replacement of the mitral valve [147], and the surgical risk of HFrEF combined with MR is high. A recent RCT for severe secondary MR comparing percutaneous mitral valve repair using MitraClip with medical therapy versus medical therapy alone (Percutaneous Repair with the MitraClip Device for Severe Functional/Secondary Mitral Regurgitation [MITRA-FR]) showed similar mortality and HF hospitalization rates [148]. On the other hand, in the Cardiovascular Outcomes Assessment of the MitraClip Percutaneous Therapy for Heart Failure Patients with Functional Mitral Regurgitation (COAPT) trial performed in 614 symptomatic patients with HF and moderate-to-severe secondary MR despite maximal medical therapy, the addition of percutaneous mitral repair with MitraClip to medical therapy was associated with lower rates of HF hospitalization and all-cause mortality than medical therapy alone within 2 years of follow-up [149].

Other percutaneous approaches to improve the outcomes of functional MR are also under investigation. The outcomes with the Cardioband system, a device for percutaneous mitral annuloplasty, were reported recently. Most patients showed moderate or less residual MR and improved symptoms at 1 year [150]. The Annular Reduction for Transcatheter Treatment of Insufficient Mitral ValvE (ACTIVE) randomized trial is recruiting patients to evaluate the efficacy of the Cardioband system along with optimal medical therapy (NCT03016975). Various devices for percutaneous mitral valve replacement are also under investigation but are still at the level of early feasibility studies at present [151,152]. Moreover, Neochord, which was developed for transapical repair of MR with artificial chordae, is also the subject of an RCT in comparison with surgical mitral valve repair in degenerative MR (NCT02803957).

Other interventions: inter-atrial shunting, vagus nerve stimulation, and others

Several forward-looking device therapies are under development. The Reduce Elevated Left Atrial Pressure in Patients With Heart Failure (REDUCE LAP-HF) I study is a phase II RCT of an interatrial shunt in symptomatic HF patients with LVEF \( \geq 40\% \) and elevated exercise pulmonary capillary wedge pressure. The device significantly reduced post-exercise pulmonary capillary wedge pressure at 1 month [153] and showed similar safety outcome at 1 year [154]. As autonomic imbalance is important in the pathophysiology of HF, vagus nerve stimulation is thought to be a potential treatment target [155]. Recent phase II (NEural Cardiac TherApY foR Heart Failure [NECTAR-HF]) [156,157] and phase III RCTs (Increase of Vagal Tone in Heart Failure [INO-VATE-HF] [158] for symptomatic HF with LV dysfunction reported results of vagus nerve stimulation but failed to show significant decreases in LV end-systolic diameter or improvement of clinical outcomes.

Although renal denervation has been suggested as an alternative treatment option for resistant hypertension, the results of RCTs were disappointing [159,160]. However, another RCT indicated the possibility of revival. In the Renal Denervation With the Symplicity SpyralTM Multi-electrode Renal Denervation System in Patients With Uncontrolled Hypertension in the Absence of Antihypertensive Medications (SPYRAL HTN-OFF MED) trial, the renal denervation group showed a decrease of about 5 mmHg in systolic and diastolic blood pressure with 24-hour ambulatory blood pressure monitoring after 3 months, while the sham group showed no significant changes in blood pressure [161]. Two new RCTs for renal denervation have recently begun: endovascular ultrasound renal denervation to treat hypertension (RADIANCE-HTN SOLO; NCT02649426) and SPYRAL HTN-ON MED (NCT02493975). As hypertension is one of the most important etiologies of HF, renal denervation may be a promising technology if these trials succeed.

MECHANICAL CIRCULATORY SUPPORT AND HEART TRANSPLANTATION

Mechanical circulatory support

Left ventricular assist devices (LVAD) are rapidly being adopted for advanced HF treatment. These devices were initially used as a bridge to transplantation, but are now also commonly used as destination therapy. Advances in mechanical technology and surgical techniques have greatly increased the success rate and duration of ventricular assist devices. The HeartWare ventricular assist device (HVAD), which is a commercial LVAD using a centrifugal heart pump, showed a non-inferior outcome
in comparison with Heartmate II (The HeartWare™ Ventricular Assist System as Destination Therapy of Advanced Heart Failure [ENDURANCE]) [85]. This device is highly miniaturized to facilitate minimally invasive surgery and reduce surgical complications [162]. HVAD showed good clinical outcome and safety in the real-world registry data. Follow-up of 254 patients in a multicenter prospective registry study of patients transplanted with HVAD revealed a mean duration of support of 365 ± 280 days and success rates of 87% at 6 months, 85% at 1 year, and 73% at 3 years (post-market Registry to Evaluate the HeartWare Left Ventricular Assist System [ReVOLVE] study). During the follow-up period, 17% of the patients died, and the most common adverse event was bleeding (28%). In particular, the relatively high incidence of cerebrovascular accidents after HVAD implantation in previous studies was reduced to acceptable levels in the ReVOLVE study. Vigorous anticoagulation therapy and adequate blood pressure control play important roles in reducing the incidence of stroke [163]. There have also been improvements in minimally invasive techniques to reduce surgical complications (Fig. 5) [164].

Heartmate II, another LVAD, also showed acceptable clinical outcomes and safety for destination therapy [84] as well as use as a bridge to transplantation [83]. The newly developed Heartmate 3 is equipped with a fully magnetically levitated centrifugal flow pump, and showed a survival rate of 77.9% without serious complications at 2 years after device implantation (Multicenter Study of MagLev Technology in Patients Undergoing Mechanical Circulatory Support Therapy with HeartMate 3 [MOMENTUM 3]) [165]. This is an improvement compared with 56.4% for the previous version, Heartmate II. Although the survival rate is improving, many patients with Heartmate 3 implants still experience a range of complications, including bleeding, infection, stroke, right HF, and arrhythmias (Fig. 5) [86,165].

SynCardia, a sole total artificial heart (TAH) approved by the U.S. Food and Drug Administration, provides the most definitive treatment options for patients with biventricular failure who are not candidates for isolated LVAD placement [166]. The SynCardia system has recently developed a smaller 50 cc TAH that was designed to accommodate patients with low body surface area. This technical improvement should allow the device to be implanted in women and children, and it might be particularly useful in growing adolescents with palliated congenital heart disease [167].

Heart transplantation
Heart transplantation (HT) has become the standard treatment for selected patients with end-stage HF. Improvements in immunosuppressants, donor procurement, surgical techniques, and post-HT care have resulted in a substantial decrease in the incidence of acute allograft rejection, which had previously significantly limited survival of HT recipients. However, there are limitations to long-term allograft survival, including rejection, infection, coronary allograft vasculopathy, and malignancy (Fig. 6). Careful balance of immunosuppressive therapy and vigilant surveillance for complications can further improve long-term outcomes of HT recipients. Most transplant recipients have been treated with a combination of a calcineurin...
inhibitor, mycophenolate mofetil, and steroids [168]. To monitor the effects of immunosuppressive drugs and adjust the dose, physicians check the serum concentration of immunosuppressant. However, the serum concentration does not accurately reflect the degree of immunosuppression in a specific patient. Immune monitoring assay (Immuknow, Cylex, Columbia, MD, USA), a peripheral blood test, helps physicians to determine the degree of immunosuppression in patients by measuring the amount of ATP released from activated lymphocytes [169]. In a study to determine the efficacy of immune monitoring assay, patients with infectious complications had a low immune monitoring score, and some patients with rejection had a high score [170]. Further large-scale studies with more sophisticated measurements of immune monitoring methods are needed to achieve personalized immunosuppression.

Rejection diagnostic methods
Donor-derived cell-free DNA (dd-cfDNA) is an emerging noninvasive tool for diagnosis of rejection. The dd-cfDNA is detectable in both blood and urine of transplant recipients. After transplantation, dd-cfDNA rises to > 5% of total cfDNA and decreases sequentially to < 0.5% within 1 week. When a rejection event occurs, dd-cfDNA could increase up to 5-fold from the baseline value in the blood [171]. The dd-cfDNA is a potential candidate as a noninvasive tool for diagnosis of graft rejection, as the degree of dd-cfDNA elevation has been shown to be correlated with acute cellular rejection events, as determined by endomyocardial biopsy in early studies [172].

Expanding the donor pool
As the number of the donors is very small compared to patients requiring HT, there have been continuing efforts to expand the donor pool. To maximize the number of patients receiving HT, some transplantation centers now use extended criteria donor (ECD) hearts in high-risk recipients, and the outcomes seem to be acceptable. The general characteristics of ECD hearts are as follows: age > 50 years, female donor, heart from patients with cardiovascular death, hypertension, diabetes, elevation of cardiac troponin, LV systolic dysfunction (LVEF < 50%), and regional wall motion abnormalities. Moreover, the criteria for high-risk recipients are as follows: age > 65 years, renal insufficiency, peripheral artery disease, or poorly controlled diabetes. The ECD program has had little impact on the outcome of transplanted patients and seems to accomplish the purpose of expanding the donor pool [173].

The concept of donation after circulatory death (DCD) was introduced as part of the efforts to expand the donor pool. To minimize the damage due to ischemic time in DCD organs, trials to utilize 

\textit{ex vivo} perfusion systems were performed. The 

\textit{ex vivo} heart perfusion system maintains the heart in a beating and metabolically active state by supplying warm, oxygenated, and nutrient-enriched donor blood. Recently, 

\textit{ex vivo} perfusion systems have been reported to show non-inferiority with regard to outcome compared with standard cold storage methods, and further trials are currently underway [89].

**ACUTE HEART FAILURE**

AHF refers to rapid onset or worsening of symptoms and/or signs of HF. It is a life-threatening medical condition requiring urgent evaluation and treatment, typically leading to urgent hospital admission [1]. The
Table 3. Clinical characteristics and outcomes of acute heart failure registry

| Characteristic | KorAHF [46] | KorHF [175] | ATTEND [176] | ADHERE [177] | OPTIMIZE-HF [178] | EHFS II [179] | AHEAD Main [180,181] |
|----------------|-------------|-------------|-------------|-------------|-------------------|-------------|---------------------|
| Region         | Korea       | Korea       | Japan       | USA         | USA               | Europe      | Czech               |
| Recruitment period | Mar 2011– Feb 2014 | Jun 2004–2009 Apr | Apr 2007–Dec 2011 | Sep 2001–Jan 2004 | Mar 2003–Dec 2004 | Oct 2004–Aug 2005 | Sep 2006–Oct 2009 |
| No. of patients | 5,625       | 3,200       | 4,842       | 159,168     | 48,612            | 3,580       | 4,153               |
| Follow-up      | 5 years     | 5 years     | 180 day     | NA          | 60,90 day         | 3,12 months | 20 months (median) |

Demographics
- Age, yr, mean ± SD
  - KorAHF: 69 ± 15
  - KorHF: 68 ± 14
  - ATTEND: 73 ± 14
  - ADHERE: 72 ± 14
  - OPTIMIZE-HF: 73 ± 14
  - EHFS II: 70 ± 13
  - AHEAD Main: 72 ± 12
- Male sex, %
  - KorAHF: 53
  - KorHF: 50
  - ATTEND: 58
  - ADHERE: 48
  - OPTIMIZE-HF: 48
  - EHFS II: 61
  - AHEAD Main: 60

Co-morbidities, %
- Hypertension
  - KorAHF: 62
  - KorHF: 47
  - ATTEND: 69
  - ADHERE: 74
  - OPTIMIZE-HF: 71
  - EHFS II: 63
  - AHEAD Main: 73
- Diabetes
  - KorAHF: 40
  - KorHF: 31
  - ATTEND: 34
  - ADHERE: 44
  - OPTIMIZE-HF: 42
  - EHFS II: 33
  - AHEAD Main: 43
- Atrial fibrillation
  - KorAHF: 29
  - KorHF: NA
  - ATTEND: 36
  - ADHERE: 31
  - OPTIMIZE-HF: 31
  - EHFS II: 39
  - AHEAD Main: 27
- Chronic lung disease
  - KorAHF: 11
  - KorHF: 4
  - ATTEND: 10
  - ADHERE: 31
  - OPTIMIZE-HF: 28
  - EHFS II: 19
  - AHEAD Main: NA

Etiology, %
- Ischemic
  - KorAHF: 38
  - KorHF: 52
  - ATTEND: 31
  - ADHERE: 58
  - OPTIMIZE-HF: 46
  - EHFS II: 54
  - AHEAD Main: 56
- Hypertensive
  - KorAHF: 4
  - KorHF: NA
  - ATTEND: 18
  - ADHERE: NA
  - OPTIMIZE-HF: 23
  - EHFS II: 11c
  - AHEAD Main: 4

Clinical status on admission
- De novo HF, %
  - KorAHF: 52
  - KorHF: 70
  - ATTEND: 64
  - ADHERE: 24
  - OPTIMIZE-HF: 12
  - EHFS II: 37
  - AHEAD Main: 58
- Lung congestion, %
  - KorAHF: 79 (edema)
  - KorHF: NA
  - ATTEND: 71 (rales)
  - ADHERE: 75 (edema)
  - OPTIMIZE-HF: 64 (rales)
  - EHFS II: NA
  - AHEAD Main: 18 (edema)
- Pulse rate, /min, mean ± SD
  - KorAHF: 93 ± 26
  - KorHF: 92 ± 26
  - ATTEND: 99 ± 29
  - ADHERE: NA
  - OPTIMIZE-HF: 87 ± 22
  - EHFS II: 95 (median)
  - AHEAD Main: 90 (median)
- SBP, mmHg, mean ± SD
  - KorAHF: 131 ± 30
  - KorHF: 131 ± 30
  - ATTEND: 146 ± 37
  - ADHERE: 144 ± 33
  - OPTIMIZE-HF: 143 ± 33
  - EHFS II: 135 (median)
  - AHEAD Main: 135 (median)
- LVEF < 40%, %
  - KorAHF: 55
  - KorHF: 74 (EF < 50%)
  - ATTEND: 53
  - ADHERE: 51
  - OPTIMIZE-HF: 49
  - EHFS II: 66 (EF < 45%)
  - AHEAD Main: 38 (EF < 30%)
- Creatinine, mg/dL, mean ± SD
  - KorAHF: 1.5 ± 1.5
  - KorHF: 1.5 ± 1.2
  - ATTEND: 1.4 ± 1.6
  - ADHERE: 1.8 ± 1.6
  - OPTIMIZE-HF: 1.8 ± 1.6
  - EHFS II: NA
  - AHEAD Main: 1.2 (median)

Pharmacologic treatment
- IV diuretics
  - KorAHF: 75
  - KorHF: 68
  - ATTEND: 76
  - ADHERE: 87
  - OPTIMIZE-HF: NA
  - EHFS II: 84
  - AHEAD Main: 84
- IV inotropes
  - KorAHF: 31
  - KorHF: 22
  - ATTEND: 19
  - ADHERE: 8
  - OPTIMIZE-HF: 7
  - EHFS II: < 29.8
  - AHEAD Main: NA
-IV vasodilators
  - KorAHF: 41
  - KorHF: 36
  - ATTEND: 78
  - ADHERE: 9
  - OPTIMIZE-HF: 14
  - EHFS II: 31
  - AHEAD Main: 19
The outcome of hospitalized HF patients in Korea has shown a modest improving trend over time, although treatment of AHF has not changed for several decades [3]. Clinical characteristics and outcomes of AHF registry according to different countries are summarized in Table 3 [46,174-181]. Many novel drugs have shown no clinical improvement [1]. This is because AHF syndrome is an event in the context of underlying HF, and not a disease entity per se [182]. Notably, the mid- or long-term outcome of AHF may not depend on treatment at the acute stage, but on the underlying disease status causing decompensation [46,183]. In addition, the classification of AHF is not clear [184-187]. Inadequate phenotyping is also responsible for the failure of treatments to improve outcomes in AHF. In this section, we focus on new treatment strategies and the attempt to reduce readmission to hospital (Table 4).

Diuretics are the mainstay of pharmacological treatment in AHF to improve symptoms [188]. Recently, the time-to-diuretics concept was proposed for AHF. A large prospective observational study showed a lower rate of in-hospital mortality in patients with door-to-diuretics time < 1 hour [189]. In contrast, another observational study failed to show any associations between clinical outcomes and short door-to-diuretics time [190]. Newly developed intravenous vasodilators failed to improve outcomes in AHF. Nesiritide, recombinant BNP with a vasodilating effect, did not show improvement in dyspnea and in death or rehospitalization but showed a significantly higher rate of hypotension in the Acute Study of Clinical Effectiveness of Nesiritide in Decompensated Heart Failure (ASCEND HF) RCT [191]. Serelaxin, i.e., recombinant human relaxin-2, improved dyspnea and 180-day mortality rate in the RELAXin in acute heart failure (RELAX-AHF) randomized trial [192], but did not show consistent results in the subsequent RELAX-AHF-2 study (NCT01870778) [193]. Ularitide did not improve clinical outcomes in the Trial of Ularitide Efficacy and Safety in Acute Heart Failure (TRUE-AHF) trial [194]. The inodilator, levosimendan, was associated with reduction of short- and long-term mortality rates compared with placebo in LV dysfunction patients with acute MI [195]. However, there was no mortality benefit of levosimendan in comparison with dobutamine [196]. Recently, the PIONEER-HF study showed encouraging results in HFrEF patients who were hospitalized...
for acute decompensated HF. The initiation of sacubitril-valsartan therapy led to a greater reduction of NT-proBNP concentration than enalapril therapy, with no significant difference in rate of adverse events, such as deteriorating renal function, hypotension, hyperkalemia, and angioedema [197]. However, the role of ARNI in AHF should be verified in larger prospective study.

As outlined above, the treatment strategy for AHF has not changed markedly over the last several years. However, the rate of readmission after an AHF episode

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Table 4. Recent evidence regarding mortality in treatment of acute heart failure

| Topic     | Study          | Year | Author                     | No.  | Intervention                          | Comparator | Finding                                                                 |
|-----------|----------------|------|----------------------------|------|---------------------------------------|------------|-------------------------------------------------------------------------|
| Diuretics | REALITY-AHF    | 2017 | Matsue et al. [189]         | 1,291| Door to diuretics time                | NA (prospective cohort) | Early treatment with IV loop diuretics (< 60 min) was associated with lower in-hospital mortality. |
|           | KorAHF         | 2018 | Park et al. [190]           | 5,625| Door to diuretics time                | NA (prospective cohort) | Door to diuretics time was not associated with clinical outcome. |
|           | EVEREST        | 2017 | Konstam et al. [99]         | 4,133| Tolvaptan                              | Placebo    | Tolvaptan did not show benefit in long-term mortality and composite of cardiovascular death and HF hospitalization. |
| Vasodilators | ASCEND-HF     | 2011 | O’Connor et al. [191]       | 7,141| Nesiritide                            | Placebo    | Nesiritide was not associated with change of HF rehospitalization and death within 30 days. |
|           | RELAX-AHF      | 2013 | Teerlink et al. [192]       | 1,161| Serelaxin                              | Placebo    | Serelaxin was associated with dyspnea relief and decrease in 180-day mortality. |
|           | RELAX-AHF-2    | 2017 | Abstract [193]              | 6,545| Serelaxin                              | Placebo    | Serelaxin did not showed significant difference in 180-day all-cause and cardiovascular mortality. |
|           | TRUE-AHF       | 2017 | Packer et al. [194]         | 2,157| Ulratitide                            | Placebo    | Ulratitide did not showed significant difference in cardiovascular death at a median follow-up of 15 months. |
| Inodilator | RUSSLAN        | 2002 | Moiseyev et al. [195]       | 504  | Levosimendan                          | Placebo    | Levosimendan was associated with reduction in 14- and 180-day mortality in patients with LV dysfunction due to AMI. |
|           | SURVIVE        | 2007 | Mebazaa et al. [196]        | 1,327| Levosimendan                          | Dobutamine | Levosimendan did not significantly reduce all-cause mortality at 180 days. |

REALITY-AHF, Registry Focused on Very Early Presentation and Treatment in Emergency Department of Acute Heart Failure; NA, not available; IV, intravenous; KorAHF, Korean Acute Heart Failure; EVEREST, Efficacy of Vasopressin Antagonism in Heart Failure Outcome Study With Tolvaptan; HF, heart failure; ASCEND-HF, Acute Study of Clinical Effectiveness of Nesiritide in Decompensated Heart Failure; RELAX-AHF, Trial of RELAXin in Acute Heart Failure; TRUE-AHF, Ularitide Efficacy and Safety in Acute Heart Failure; RUSSLAN, Randomised Study on Safety and effectiveness of Levosimendan in patients with left ventricular failure due to an Acute myocardial infarct; LV, left ventricle; AMI, acute myocardial infarction; SURVIVE, Survival of Patients With Acute Heart Failure in Need of Intravenous Inotropic Support.
is consistently high, reaching 20% and 50% at 1 and 6 months after discharge, respectively [198]. To reduce the healthcare costs associated with AHF, the hospital readmission reduction program was introduced in the USA, which awarded a penalty to hospitals with high 30-day readmission rates. However, the results have been disappointing. The 30-day and 1-year readmission rates decreased, while the mortality rate tended to increase [199]. In addition, the quality of care and clinical outcome were similar between hospitals with high and low risk-adjusted 30-day HF readmission rates [200], and the mortality rate was lower in the higher hospital-level 30-day episode payment [201]. It is possible that some hospitals attempted to improve the index rather than the true outcome by adopting methods, such as increased admission period, delaying readmission after 30 days, etc., resulting in an unintended increase in mortality rate [202]. Remote monitoring mentioned in the previous section can be utilized for the early detection of deterioration and the prevention of readmission in AHF patients.

**DIGITAL HEALTHCARE IN HEART FAILURE**

Digital healthcare has received much attention recently. As the amounts of information from diverse sources, such as electronic medical records, wearable devices, and genomic data, are increasing rapidly, artificial intelligence and machine learning are essential to collect, manage, and apply these data appropriately [203]. Ahmad et al. [204] reported that machine learning could improve prognostic prediction and phenotyping with different treatment responses in a large cohort of HF patients. The use of artificial intelligence and clinical decision support systems is not suitable for clinical application. However, it may help physicians to make decisions by organizing large amounts of data and building delicate prognostic models in the near future.

**CONCLUSIONS**

HF is becoming an increasingly important disease entity with the aging of society. According to its increasing prevalence, many new drugs and devices have been studied to improve clinical outcome in terms of mortality and quality of life in HF patients. Although few studies showed encouraging results, researchers are attempting to find subgroups in whom certain medications or devices could be most effective, new methods for better diagnosis and prediction of prognosis in HF patients, and new tools for treating HF.

**Conflict of interest**

No potential conflict of interest relevant to this article was reported.

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