



University of Dundee

SGLT2-inhibitors; more than just glycosuria and diuresis

Fathi, Amir; Vickneson, Keeran; Singh, Jagdeep S

Published in: **Heart Failure Reviews**

DOI: 10.1007/s10741-020-10038-w

Publication date: 2020

Document Version Publisher's PDF, also known as Version of record

Link to publication in Discovery Research Portal

Citation for published version (APA): Fathi, A., Vickneson, K., & Singh, J. S. (2020). SGLT2-inhibitors; more than just glycosuria and diuresis. *Heart* Failure Reviews. https://doi.org/10.1007/s10741-020-10038-w

General rights

Copyright and moral rights for the publications made accessible in Discovery Research Portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

- Users may download and print one copy of any publication from Discovery Research Portal for the purpose of private study or research.
- You may not further distribute the material or use it for any profit-making activity or commercial gain.
 You may freely distribute the URL identifying the publication in the public portal.

Take down policy

If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.



SGLT2-inhibitors; more than just glycosuria and diuresis

Amir Fathi¹ · Keeran Vickneson² · Jagdeep S. Singh^{2,3}

Accepted: 5 October 2020 © The Author(s) 2020

Abstract

Heart failure (HF) continues to be a serious public health challenge despite significant advancements in therapeutics and is often complicated by multiple other comorbidities. Of particular concern is type 2 diabetes mellitus (T2DM) which not only amplifies the risk, but also limits the treatment options available to patients. The sodium-glucose linked cotransporter subtype 2 (SGLT2)-inhibitor class, which was initially developed as a treatment for T2DM, has shown great promise in reducing cardiovascular risk, particularly around HF outcomes – regardless of diabetes status.

There are ongoing efforts to elucidate the true mechanism of action of this novel drug class. Its primary mechanism of inducing glycosuria and diuresis from receptor blockade in the renal nephron seems unlikely to be responsible for the rapid and striking benefits seen in clinical trials. Early mechanistic work around conventional therapeutic targets seem to be inconclusive. There are some emerging theories around its effect on myocardial energetics and calcium balance as well as on renal physiology. In this review, we discuss some of the cutting-edge hypotheses and concepts currently being explored around this drug class in an attempt better understand the molecular mechanics of this novel agent.

Keywords SGLT2-inhibitors \cdot Renal disease \cdot Heart failure \cdot Calcium handling \cdot Myocardial energetics \cdot Ventricular remodelling

Introduction

Heart failure (HF) is a growing public health concern with 8.5 million people expected to be living with this condition in the year 2030 at a staggering cost of \$70 billion in the United States alone [1]. HF not only shortens life, but also reduces its quality. There have been considerable advancements in the management of this disease using a variety of neurohormonal modulators and, more recently, with device therapy. Nevertheless, HF remains a challenge to treat particularly because

Amir Fathi and Keeran Vickneson contributed equally to this work.

Jagdeep S. Singh jxsingh@dundee.ac.uk

- ¹ Department of Neuroanaesthesia and Critical Care, National Hospital for Neurology and Neurosurgery, University College London, London, UK
- ² Division of Molecular and Clinical Medicine, School of Medicine, University of Dundee, Dundee, UK
- ³ Department of Cardiology, The Edinburgh Heart Center, Royal Infirmary of Edinburgh, 51 Little France Crescent, Edinburgh EH16 4SA, UK

it is frequently associated with other co-morbidities such as hypertension, type 2 diabetes mellitus (T2DM), ischaemic heart disease and renal impairment. Ischaemic heart disease is the commonest cause of heart failure and T2DM amplifies that risk further. Patients with ischaemic cardiomyopathy and T2DM face a doubling of mortality risk compared to patients without T2DM. Indeed, T2DM increases mortality risk regardless of the underlying aetiology of heart failure [2].

Interestingly, there appears to be a bidirectional relationship between T2DM and HF; not only can T2DM cause HF, but it can also be the consequence of it [3]. This underscores the importance of optimal diabetes control in patients with HF. Although there is a multitude of efficacious medications for T2DM, they do not appear to reduce the risk of cardiovascular (CV) outcomes and in the context of HF—some of these agents may even be harmful. A meta-analysis involving 95,000 individuals showed a 42% increased risk of incident HF with the use of thiazolidinediones in patients with T2DM while DPPIV-inhibitors increased that risk by 25% [4]. Another meta-analysis of large HF trials showed patients with HF using insulin had a 27% increased risk of all-cause mortality [5]. These data highlight the complexity, and potential dangers, of managing T2DM and HF concomitantly. A novel class of anti-diabetes therapy known as the sodium glucose-linked cotransporter subtype 2 (SGLT2) inhibitor has shown potential in addressing this area of urgent unmet need, not only by improving glycaemia but also preserving renal function and reducing hard HF outcomes as well. In this review, we discuss the evidence behind the striking CV benefits and unpick some of the unique characteristics of this drug class that could potentially herald a new generation of targeted agents in the treatment of HF.

SGLT2 inhibitors – what it says on the tin

Canagliflozin, dapagliflozin, empagliflozin and ertugliflozin are four currently available agents of the SGLT2-inhibitor class. All four are indicated for T2DM, while dapagliflozin recently received FDA approval for HF, with or without T2DM [6]. As the name implies, SGLT2-inhibitors work by inhibiting the sodium-glucose cotransporter subtype-2, located in the S1 and S2 segments of the proximal convoluted tubule (PCT) of the kidney. SGLT2 is a low affinity high capacity transporter ($K_m 2 \text{ mM}$) responsible for approximately 90% reabsorption of filtered plasma

glucose, and the remainder is reabsorbed by SGLT1 [7]. Potent SGLT2-inhibition prevents the reabsorption of filtered glucose as well as sodium, resulting in glucosuria and natriuresis. (Fig. 1) Other pleotropic benefits of SGLT2-inhibition include weight loss (1.8 to 2.7 kg), reductions in blood pressure (systolic blood pressure (BP): 1.0–2.6 mmHg; diastolic BP 0.7–2.2 mmHg) [9], without increases in heart rate and a low potential of inducing hypoglycaemia [10].

Unlike other antidiabetic agents, the glucose lowering effect of SGLT2 inhibitors is independent of pancreatic beta-cell function and insulin sensitivity – an ideal property in the context of T2DM disease progression [11] and metabolic milieu of the T2DM phenotype [12]. This insulinindependent fall in plasma glucose reduces insulin requirements and induces a rise in the glucagon-to-insulin ratio, shifting metabolism towards a catabolic state [13]. The caloric loss coupled with increased lipolysis (from catabolism) are responsible for the sustained steady-state weight loss seen with SGLT2-inhibition. (Fig. 2) Body composition studies have shown approximately two-thirds of sustained weight loss is attributable to loss of body fat [14].



Fig. 1 Normal renal tubular resorption of glucose. The diagram also identifies the site at which SGLT2-inhibitors act [8]. Abbreviations: PCT-proximal convoluted tubules; SGLT-sodium-glucose cotransporter



Fig. 2 Overview of the effects of SGLT2-inhibitor therapy. Abbreviations: CV-cardiovascular; EPO-erythropoietin; Hb-haemoglobin; Hct-haematocrit; NHE-1- sodium-hydrogen exchanger subtype-1;

PCT-proximal convoluted tubule (of renal nephron); SGLT-sodiumglucose linked cotransporter; TG-tubuloglomerular

In addition, glucosuria and natriuresis simulate an osmotic diuretic effect resulting in reduced plasma volume. Filtered glucose that is not reabsorbed increases tubular fluid osmolarity, flattening the osmotic gradient between tubular fluid and the interstitium thereby reducing electrolyte-free water reabsorption. This is expected to provide some relief on cardiac filling volume and blood pressure which in turn results in reduced preload and afterload respectively. (Fig. 2) Unlike conventional sodium-driven diuretics, SGLT2-inhibititor-mediated free water excretion is expected to be more efficient in relieving signs and symptoms of interstitial congestion without adversely compromising intravascular fluid status or causing reflex sympathetic activation [15, 16]. The recently published RECEDE-CHF trial confirmed that in a cohort of patients with established HF on long-term diuretic therapy, the addition of empagliflozin increased urine output by over 500 ml/day and more than half of that volume was in the form of electrolyte-free water clearance [17].

These are the primary mechanisms of action of the SGLT2-inhibitor class that have been purported to be responsible for its beneficial effects, however as our understanding of this drug class evolves there is a recognition that there may be other novel pathways at play – we explore these pathways in the later sections of this article.

Cardiovascular outcome trials – the breakthrough

In 2015, the Empagliflozin Cardiovascular Outcome Event Trial in Type 2 Diabetes Mellitus Patients (EMPA-REG OUTCOME) was the first FDA-mandated cardiovascular outcome trial (CVOT) of the class. It studied 7020 individuals with T2DM, randomised to empagliflozin or placebo over a mean follow-up period of 3.1 years [18]. There were striking reductions in 3-point major adverse cardiovascular events (MACE); (HR 0.86; 95% CI 0.74-0.99), all-cause mortality (HR 0.68; 95% CI 0.57-0.82), CV mortality (HR 0.62; 95% CI 0.49–0.77) and HF hospitalisation (HR 0.65; 95% CI 0.50–0.85) [18]. Further analyses revealed the CV benefit with empagliflozin was independent of CV comorbidity burden [19, 20] and renal function [21]. Although the most promising effects of the drug were around HF outcomes, only 10% of the cohort had a diagnosis of HF prior to randomisation and, importantly, there was no formal confirmation / characterisation of HF (e.g. by measuring natriuretic peptides or performing echocardiography). Another unexpected finding was the rapidity of benefit, with the HF hospitalisation and CV mortality curves diverging within weeks of initiation of therapy.

The Canagliflozin Cardiovascular Assessment Study (CANVAS) and Dapagliflozin Effect on Cardiovascular Events-Thrombolysis in Myocardial Infarction 58 (DECLARE-TIMI 58) trials were the next two CVOTs to establish CV safety and efficacy of canagliflozin and dapagliflozin respectively. Both studies utilised a broader inclusion criteria of patients, which included those at high risk of developing CV disease as well as patients with already established CV disease. The majority of participants in CANVAS had established CV disease, while the reverse was true for DECLARE-TIMI 58. Both had a similar proportion of patients with a history of HF to EMPA-REG OUTCOME. Canagliflozin significantly reduced 3-point MACE (HR 0.86; 95% CI 0.75-0.97) and HF hospitalisation (HR 0.67; 95% CI 0.52-0.87), but there was no significant reduction in mortality [22]. Dapagliflozin had no effect on 3-point MACE (HR 0.93; 95% CI 0.84-1.03) or CV death (HR 0.98; 95% CI 0.82-1.17) but there was still significantly lower hospitalisations for HF (HR 0.73; 95% CI 0.61-0.88) [23]. In post-hoc analysis of DECLARE-TIMI 58 where participants were stratified by ejection fraction, dapagliflozin reduced risk of CV death and HF hospitalisation to a greater extent in patients with reduced ejection fraction (EF) [24], however these findings are to be interpreted with caution given the small proportion of patients and lack of robust characterisation of HF. A meta-analysis of all three CVOTs showed an overall 14% reduction in MACE, 23% reduction in composite HF hospitalisation and CV death and 45% reduction in renal disease progression [25]. These findings have also been reproduced in large, real-world clinical studies [26, 27].

Cardiovascular Outcomes Following Ertugliflozin Treatment in Type 2 Diabetes Mellitus Participants With Vascular Disease (VERTIS CV) is the most recent CVOT to investigate effects of ertugliflozin in over 8000 patients with established CV disease [28]. The inclusion criteria used was very similar to EMPA-REG OUTCOME, with a larger proportion of patients with HF at baseline in VERTIS CV (23.1% vs 10.1%). Ertugliflozin failed to demonstrate superiority in the primary outcome of 3-point MACE (HR 0.97 95% CI 0.85-1.11) or any of the key secondary outcomes; CV death (HR 0.92 95% CI 0.77-1.11) and renal disease progression (HR 0.81 95% CI 0.63—1.04). However, a 30% reduction in risk of hospitalisation for HF was seen with ertugliflozin, consistent with the class effect seen in previous CVOTs. The discordant results in CV death or renal disease progression, especially in a secondary prevention cohort is surprising given the benefits seen in other CVOTs; further analysis is awaited.

Findings from these trails led the leading cardiac (and diabetes) societies to recommend the use of SGLT2-inhibitors in patients with coronary artery disease with T2DM to reduce the risk of future CV events [29, 30].

HF with reduced ejection fraction – SGLT2-inhibitors are fast becoming a must have drug

The Dapagliflozin and Prevention of Adverse outcomes in Heart Failure (DAPA-HF) trial was the first to prospectively investigate the benefits of dapagliflozin in a HF population with reduced ejection fraction (HFrEF) with or without diabetes. Dapagliflozin demonstrated a remarkable reduction in composite CV death or worsening HF (defined as hospitalisation or urgent visit for HF) (HR 0.74 [95% CI 0.65-0.85]), with each component being reduced by 18% and 30% respectively [31]. Patients also experienced less symptoms of HF in the dapagliflozin arm compared to conventional HF therapy as evidenced by a significant and clinically meaningful improvement in Kansas City Cardiomyopathy Questionnaire score. Some noteworthy aspects of the study include: 1) majority of the participants were on well-established background HF therapy (ACEi/ARB/ ARNI 94%, BB 96%, MRA 71%), which further underscores the added, incremental benefit of dapagliflozin on CV outcomes; 2) CV benefit manifested early and was maintained throughout course of the study; 3) short median follow-up period of 18 months as compared to other key HF therapies (with the exception of beta-blockers) and 4) pre-specified analyses showed CV benefit was consistent, regardless of diabetes status at baseline and there were no safety concerns regarding hypoglycaemia or adverse volume depletion in non-diabetic HF patients [31].

Empagliflozin Outcome Trial in Patients with Chronic Heart Failure with Reduced Ejection Fraction (EMPEROR-Reduced) was designed to be more ambitious than DAPA-HF, recruiting a HF population that had on average more severe systolic dysfunction (mean EF 27% vs. 31%; N-terminal pro B-type natriuretic peptide (NT-proBNP) 1907 vs. 1437) [32]. After a median follow-up of 16 months, the incidence of primary outcome of CV death or hospitalisation for HF was significantly reduced by empagliflozin therapy (HR 0.75; 95% CI 0.65–0.86]), largely driven by lower HF hospitalisations. [32]. The lack of signal in CV mortality (HR 0.92; 95% CI 0.75–1.12) was somewhat surprising. Perhaps, in patients with more advanced HF, the ability of empagliflozin to further reduce mortality risk may have been a step too far.

The true CV benefit with SGLT2-inhibition could be benchmarked against the only other novel therapy shown to reduce mortality in a well-treated HF population—the angiotensin receptor-neprilysin inhibitor sacubitril-valsartan, which works by augmenting the natriuretic peptide system and simultaneously inhibiting the renin–angiotensin–aldosterone system [33]. Dapagliflozin and empagliflozin both showed greater relative and absolute risk reduction in primary composite outcome of CV death or HF hospitalisation compared to sacubitril-valsartan (PARADIGM-HF) in comparable cohorts of HF patients [34]. The benefit seen with SGLT2-inhibitors was primarily driven by superior reductions in HF hospitalisation. In fact, the effect size seen in the SGLT2-inhibitor trials may have been undermined by the relatively older and sicker (larger proportion of NYHA III/ IV) cohort and a significantly shorter median follow-up time.

HF with preserved ejection fraction – are SGLT2-inhibitors the holy grail?

HF with preserved ejection fraction (HFpEF) makes up approximately half of the HF population. Both HFrEF and HFpEF have similar prognoses but very different pathophysiology [35]. HFrEF is due to impaired emptying of the left ventricle (LV) from systolic dysfunction, perpetuated by systemic neurohormonal activation. HFpEF on the other hand is largely due to LV diastolic dysfunction, either passively from increased myocardial stiffness or actively from impaired relaxation [36]. The lack of efficacy seen with ACE-inhibitors/ARBs [37-39], MRAs [40] and sacubitril / valsartan [41] in HFpEF suggest that excessive loading is not a key pathogenic mechanism or adverse adaptation in this HF phenotype. Increasing evidence points to abnormalities at the cellular level (inflammation, myocardial energetics, calcium handling and extracellular matrix composition) as potential targets for HFpEF management [42]. As we will discuss in the subsequent sections, SGLT2-inhibitors have unique properties that could specifically address these molecular changes particularly around calcium handling and myocardial energetics.

Several large dedicated HFpEF studies; Empagliflozin Outcome Trial in Patients with Chronic Heart Failure with Preserved Ejection Fraction (EMPEROR-Preserved) (NCT03057951) and Dapagliflozin Evaluation to Improve the Lives of Patients with Preserved Ejection Fraction Heart Failure (DELIVER) (NCT03619213), are nearing completion and will characterize SGLT2-inhibitor benefit in this cohort.

Cardiorenal syndrome – an unmet need in HF

The heart and kidney are inextricably linked, where direct or indirect effects of one dysfunctional organ can initiate and perpetuate the combined disorder of both organs, and is often referred to as the cardiorenal syndrome [43]. Up to 60% of patients with HF have co-morbid chronic kidney disease (CKD), thus increasing their risk of mortality compared with patients with just HF alone [44, 45]. Besides cardioprotection, the renal protective effects of SGLT2-inhibitors are derived from multiple converging pathways related to its primary mechanism of action; natriuresis and glucosuria.

In patients with T2DM, the increased filtered glucose elicits upregulation of SGLT1 and SGLT2 expression in

the proximal convoluted tubule of the nephron. As a consequence of the increased glucose (and sodium) absorption by these transporters in the proximal part of the nephron, there is reduced sodium delivery distally to the macula densa causing afferent renal arteriolar dilatation and hyperfiltration. Excessive hydrostatic damage to the glomerulus from hyperfiltration is thought to be the main driver for diabetic nephropathy [46]. By inhibiting sodium resorption in the proximal nephron thereby increasing distal delivery, SGLT2inhibitors can restore tubuloglomerular feedback and normalise renal blood flow. The preferential effect of SGLT2inhibition on the afferent, rather than efferent arteriole, may explain why renal function returns to baseline after an initial dip in estimated glomerular filtration rate (eGFR), with stability over time (positive eGFR slope) [47, 48]. In comparison, RAAS-inhibitors slow renal disease progression but eGFR either stabilises after the initial dip or slowly declines over a similar period (negative eGFR slope) [49, 50]. In addition, cardiorenal syndrome severely limits the available options of HF medications (ACE-inhibitors/ARBs, MRAs and ARNIs), all of which cumulatively increase risk of acute kidney injury (AKI) and hyperkalaemia in a dose-dependent manner. As Lam et al. point out, cardiorenal syndrome remains one of the key unmet needs in the management of HF, which could be directly addressed by SGLT2-inhibitors [51].

Following consistently positive effects on renal endpoints in early trials, the Evaluation of the Effects of Canagliflozin on Renal and Cardiovascular Outcomes in Participants with Diabetic Nephropathy (CREDENCE) trial was the first dedicated renal endpoint study in patients with diabetic kidney disease, using canagliflozin. Of the 4401 patients, all had a diagnosis of chronic kidney disease, 50.4% also had established CV disease and 16% had baseline HF [52]. Canagliflozin delayed renal disease progression (HR 0.60; 95% CI 0.48–0.76) and reduced risk of end-stage kidney disease (HR 0.68; 95% CI 0.54–0.86) [52]. Risk of hospitalisation from HF was also significantly reduced by 39%, consistent with results from previous CVOTs. There was no increase in incidence of AKI or hyperkalaemia with SGLT2 inhibition.

A Study to Evaluate the Effect of Dapagliflozin on Renal Outcomes and Cardiovascular Mortality in Patients With Chronic Kidney Disease (DAPA-CKD) provides further affirmation for SGLT2-inhibitor use in both diabetic and non-diabetic CKD aetiologies. Dapagliflozin reduced the risk of worsening renal function or death from kidney failure by 44%, hospitalisation for HF or CV death by 29% and all-cause mortality by 31% (all p < 0.05) – regardless of T2DM status [53]. Results from EMPA-KIDNEY are keenly awaited.

Are SGLT2-inhibitors a diabetes drug or a drug for HF and CKD?

Despite multiple guideline recommended disease-modifying therapies, patients with HF continue to have a poor prognosis [54]. Results from recent CV and renal outcome trials are very promising highlighting SGLT2-inhibitors' potential to treat T2DM, CV disease, HF and CKD—conditions that are invariably linked by the common CV risk factor profile [18, 22, 23, 52]. It has also become evidently clear that the cardiorenal benefits of SGLT2-inhibitors are independent of its modest reductions in conventional risk factors (HbA1c, blood pressure, cholesterol), T2DM status and renal function.

The question remains: Should cardiologists and nephrologists be proactively prescribing SGLT2-inhibitors to all patients with CV and renal disease?

From a CV perspective, there have now been analyses of three patient groups; 1) primary prevention cohort (part of CANVAS and DECLARE TIMI-58), 2) secondary prevention cohort (EMPA-REG OUTCOMES, VERTIS CV, part of CANVAS and DECLARE TIMI-58) and 3) HF cohort (DAPA-HF and EMPEROR-Reduced). As the CV risk profile increases, so does the apparent 'efficacy' of SGLT2-inhibition on CV mortality and HF hospitalisation. However, as seen in EMPEROR-Reduced, there is a signal that there may be an upper limit of HF severity beyond which the mortality benefits of SGLT2-inhibition plateaus, whilst morbidity benefits (reduced hospitalisations) remain robust throughout the entire spectrum of disease risk and severity.

So far only canagliflozin has received regulatory approval for use in patients with T2DM and CKD following data from the CREDENCE trial [55]. In patients with non-diabetic nephropathy, the answer is slowly emerging, especially following very encouraging results from DAPA-CKD, resulting in it being granted fast-track designation by the FDA. [56]. This highlights the urgent clinical need to slow down CKD progression and improve quality of life as well as life expectancy in this expanding cohort. Current evidence indicate that the renoprotective benefits conferred may just be the most striking effect of this drug class thus far—perhaps as a consequence of the very limited treatment options available for CKD, unlike in HF.

Mechanistic trials in SGLT2-inhibitors—more questions than answers

A characteristic feature of the pathophysiology of HF is the activation of compensatory pathways; however, sustained activation of the neurohormonal system (sympathetic nervous system and renin–angiotensin–aldosterone system) results in maladaptive remodelling of the ventricles and myocardial injury, which perpetuate the disease state. Being able to reverse this remodelling is an important determinant of long-term severity of HF and mortality [57]. Previous HF therapies have all unequivocally exhibited reverse remodelling effects on the failing heart [58–60]. It was therefore sensible that the main premise behind the first batch of mechanistic studies of SGLT2-inhibitors were based on the hypothesis that they too could induce reverse remodelling owing to the diuretic and BP lowering properties, which should improve ventricular loading in the dysfunctional ventricle.

The Research Into the Effect of SGLT2 Inhibition on Left Ventricular Remodeling in Patients With Heart Failure and Diabetes Mellitus (REFORM) trial was the first to try to determine the mechanistic effect of SGLT2-inhibitors specifically in the HF population. At 1 year, no significant difference in LV remodelling (LV end systolic volume, end diastolic volume, LV mass index or EF), as assessed by cardiac magnetic resonance imaging was seen with dapagliflozin therapy [61]. The absence of effect on remodelling was unexpected given the striking improvements in HF outcomes. There are two possible reasons why dapagliflozin therapy was neutral on LV remodelling. First, the majority of patients in the REFORM trial were on ACEi/ARBs (89%), beta-blockers (82%) and MRA (41%), which have potent effects on ventricular loading. The addition of an SGLT2-inhibitor, with only modest diuretic and BP effects, would thus confer little incremental benefit on haemodynamic load and parameters of LV remodelling. Similar conclusions can also be drawn from the DEFINE-HF Trial, where dapagliflozin did not significantly reduce NT-proBNP levels, a biomarker of ventricular end diastolic pressure, over 12 weeks, as compared to placebo [62]. Second, improvement in LV remodelling seen with SGLT2-inhibitors appear to usually occur in the early stages of the disease spectrum. Post-hoc exploratory analyses from the REFORM trial suggest dapagliflozin reduced LV volumes and indexed LV mass in patients with LVEF \geq 45%, with no interaction seen in other key secondary outcomes [63]. These findings are consistent with improvements in HF outcomes in the CVOTs, in which a large proportion of patients probably had early, occult HF. Other mechanistic trials such as the EMPA-HEART and DAPA-LVH trials support this hypothesis, demonstrating reductions in LV mass in patients with T2DM and LV hypertrophy without LV dysfunction or symptoms of HF [64].

Putting everything together, there is a definite clinical benefit from SGLT2-inhibitor therapy particularly in the context of hard HF outcomes, however the mechanism of that effect remains unclear (Table 1). The glucose-lowering and diuretic effects from its primary mode of action on the nephron, although important, simply cannot explain the speed and magnitude of benefit seen in the large outcome

Trial Name	Drug	Duration (median)	Cohort	Primary outcome	Key secondary outcomes
Cardiovascular Outcome Trials EMPA-REG Outcome [18]	Empagliflozin vs. placebo	3.1 years	n= 7020; T2DM with established CVD	3P-MACE (HR 0.86; 95% CI 0.74-0.99)	 - 3P-MACE + hospitalisation for UA (HR 0.89; 95% CI 0.78 to 1.01) - CV death (HR 0.62; 95% CI 0.49 to 0.77) - HHF (HR 0.65; 95% CI 0.50 to 0.85) - CV death/HHF (HR 0.66; 95% CI 0.55 to 0.79) - Death from any cause (HR 0.68; 95% CI 0.57 to 0.82)
CANVAS [22]	Canagliflozin vs. placebo	2.4 years	n=9734; Poorly controlled T2DM plus i) age 30+ and history of symptomatic atherosclerotic CVD or ii) age 50+ and high risk of CVD	3P-MACE (HR 0.86; 95% CI 0.75–0.97)	 - CV death (hazard ratio, 0.87; 95% CI 0.72 to 1.06) - Progression of albuminuria (30% increase) (HR 0.73; 95% CI 0.67 to 0.79) - CV death/HHF (HR 0.78; 95% CI 0.67 to 0.91)
DECLARE-TIMI 58 [23]	Dapagliflozin vs. placebo	4.2 years	n = 17,160; age 40 + with T2DM and either his- tory or high risk of atherosclerotic CV events	3P-MACE (HR 0.93; 95% CI 0.84-1.03) CV death/HHF (HR 0.83; 95% CI 0.73-0.95)	 ->40% reduction in eGFR/new ESRD/renal death/CV death (HR 0.76; 95% CI 0.67–0.87) - Death from any cause (HR 0.93; 95% CI 0.82–1.04)
VERTIS CV [28] Heart Failure Outcome Trials	Ertugliflozin vs. placebo	6.1 years	n= 8246; T2DM and established ASCVD	3P-MACE (HR 0.97; 95% CI 0.85–1.11)	 - CV death/HHF (HR 0.88; 95% CI 0.75–1.03) - HHF (HR 0.70; 95% CI 0.54–0.90) - Progression of renal disease (HR 0.81; 95% CI 0.63–1.04)
DAPA-HF [31]	Dapagliflozin vs. placebo	1.5 years	n=4744; HFrEF (LVEF < 40%); NTproBNP> 400–600 (depending on criteria); with or without T2DM	Time to first occurrence of CV death/ HHF/ urgent HF visit (HR 0.74; 95% CI 0.65–0.85)	- CV Death/HHF (HR 0.75; 95% CI 0.65-0.85) - ≥50% sustained reduction in eGFR/ reaching ESRD/renal death (HR 0.71; 95% CI 0.44-1.16) - KCCQ (HR 1.18; 95% CI 1.11- 1.26) - Death from any cause (HR 0.83; 95% CI 0.71-0.97)

Table 1 (continued)					
Trial Name	Drug	Duration (median)	Cohort	Primary outcome	Key secondary outcomes
EMPEROR-Reduced [32]	Empagliflozin vs. placebo	16 months	n= 3730; HFrEF (LVEF $\leq 40\%$; NYHA II-IV); NTproBNP> 600–5000 (specific criteria based on diagnosis of AF and EF); with or without diabetes	Time to first occurrence of CV death/ HHF (HR 0.75; 95% CI 0.65–0.86)	 CV death (HR 0.92; 95% CI 0.75-1.12) First HHF (HR 0.69; 95% CI 0.59-0.81) HHF (HR 0.70; 95% CI 0.58-0.85) HHF (HR 0.70; 95% CI 0.58-0.85) Decline in eGFR (1.73 ml/ min/1.73m²/year slower decline in treatment arm; 95% CI 1.10-2.37 Death from any cause (HR 0.92; 95% CI 0.77 to 1.10)
DELIVER [65] (Currently recruiting – est comple- tion June 2021)	Dapagliflozin vs. placebo	2.75 years	n=6100; HFpEF (LVEF>40%); Elevated NT-proBNP; Ambulatory and hospitalised patients	Time to first occurrence of CV death/ HHF/ urgent HF visit	 - KCCQ - Worsening NYHA class - Total number of CV death or HHF - Time to death from any cause
EMPEROR-Preserved [66] (est completion Nov 2020)	Empagliflozin vs. placebo	3.2 years	n≈5988; HFpEF (LVEF>40%) + structural heart disease; NTproBNP> 300; with or without diabetes	Time to first occurrence of CV death/ HHF	 CV death HHF All-cause hospitalisation Change in KCCQ RRT or sustained reduction of ≥ 40% eGFR All-cause mortality Onset of DM
Mechanisuc / Biomarker Triais REFORM [61]	Dapagliflozin vs. placebo	12 months	n=58; T2DM; stable HFrEF (LVEF $\leq 45\%$) NYHA class I-III; on furosem- ide ≤ 80 mg or other loop diuretics; eGFR ≥ 45	Change in indexed LVESV (2.49 mL/m2; 95% CI -6.30 to 11.28)	 LVMi (2.5 g/m2; 95% CI -3.95 to 8.95) LVEF (0.69%; 95% CI -3.32 to 4.69) LVEDV (indexed: 3.9 mL/m2; 95% CI -7.05 to 14.85) Weight (-2.26 kg; 95% CI -4.83 to 0.31) Weight (-2.26 kg; 95% CI -14.51 to 2.11) SBP (-4.7 mmHg; 95% CI -14.51 to 5.11) Haematorrit (2.89%; 95% CI 1.14 to 4.64) Loop diuretic dose (-29.06 mg; 95% CI -42.17 to -15.95)

Table 1 (continued)					
Trial Name	Drug	Duration (median)	Cohort	Primary outcome	Key secondary outcomes
RECEDE-CHF [17]	Empagliflozin vs. placebo	6 weeks	n= 23; T2DM and HF (NYHA II-III; LVEF < 50%); eGFR ≥ 45 ml/ min/1.73m ² ; median NT-proBNP 2381; on furosemide or equivalent loop diuretic therapy	Mean change in 24-h urinary vol- ume at 6 weeks (545 ml, 95% CI 136–954)	 - 24-h urinary sodium excretion (7.85 mmol/L; 95% CI -2.43 to 6.73) - Electrolyte-free water clearance (312 ml; 95% CI 26-598) - Fractional clearance of sodium (0.11%; 95% CI -0.22 to 0.44) - Change in urine creatinine (-1.66 mmol/L; 95% CI -3.07 to -0.25) - Change in NTproBNP (283.4 pg/ml; 95% CI -835.8 to 1402.3)
DAPA-LVH [67]	Dapagliflozin vs. placebo	12 months	n=66; T2DM; BMI≥23; BP<145/90 mmHg; LV hypertrophy	Change in indexed LVM (-2.82 g; 95% CI -5.13 to -0.51)	 LVMi (-0.20 g/m2; 95% CI -1.21 to 0.80) LVEF (0.79%; 95% CI -1.14 to 2.72) EDV (-1.59mLS; 95% CI -7.06 to 3.87) ESV (-1.12mLs; 95% CI -3.50 to 1.25) Ambulatory SBP (-3.63 mmHg; 95% CI -6.44 to -0.82) Ambulatory SBP (-3.63 mmHg; 95% CI -948.13 to -271.28) Weight (-3.77 kg; 95% CI -4.92 to -2.61) Haematocrit (2.90%; 95% CI 1.84 to 3.96) NT-proBNP (-103.68 pg/mL; 95% CI -326.90 to 119.54)
EMPA HEART Cardio-Link 6 [64]	Empagliflozin vs. placebo	6 months	n=97; T2DM; established CVD (MI≥6 months, or coronary revas- cularization≥2 months)	Change in indexed LVM (-3.35 g/ m2; 95% CI -5.9 to -0.81)	- LVEDV (-0.9 mL/m2; 95% CI -8.5 to -6.7) - ESV (-2.2 mL/m2; 95% CI -7.3 to 2.8) - LVEF (2.2%; 95% CI -0.2 to 4.7) - NT-proBNP (7.4 pg/mL; 95% CI - 10.3, 25.1)

Table 1 (continued)

 $\underline{\textcircled{O}}$ Springer

Trial Name	Drug	Duration (median)	Cohort	Primary outcome	Key secondary outcomes
DEFINE-HF [62]	Dapagliflozin vs. placebo	12 weeks	n= 263; HFrEF (LVEF ≤ 40%, and NYHA class II-III) with or without T2DM	i) average of 6- and 12-week mean NT-proBNP (HR 0.95; 95% CI 0.84 to 1.08) ii) \geq 5-point increase in average of 6- and 12-week KCCQ-OS or \geq 20% reduction in average of 6- and 12-week NT-proBNP (OR 1.8, 95% CI 1.03 to 3.06)	 KCCQ 6 weeks (OR 1.8; 95% CI 1.04 to 3.12) and 12 weeks (OR 1.7; 95% CI 0.98 to 3.1) ≥ 20% reduction NT-proBNP at 6 weeks (OR 1.1; 95% CI 0.6 to 1.9) and 12 weeks (OR 1.9; 95% CI 1.09 to 3.31) ≥ 20% reduction in BNP at 6 weeks (OR 1.5; 95% CI 0.9 to 2.7) and 12 weeks (OR 2.0; 95% CI 1.1 to 3.4) 6 MWT—adjusted distance at week 12 (304 m vs 301 m) Mean weight reduction of -1.33 kg without T2DM Dapa vs. placebo (95% CI -2.41 to -0.23 kg) HHF or urgent HF visits (HR 0.84; 95% CI 0.35 to 1.97)
EMPA-TROPISM [68] (est completion Dec 2020)	Empagliflozin vs. placebo	6 months	n=84; stable HFrEF (LVEF <50%) NYHA II-III); with or without T2DM	Changes in LVESV/LVEDV	LVEF index - VO ₂ Consumption - 6MWT - KCCQ-12
Renal Outcome Trials CREDENCE [52]	Canagliflozin vs. placebo	2.6 years	n= 4401; T2DM; CKD (eGFR 30 to < 90 ml/ minute/1.73 m ²); albuminuria (UACR > 300 to 5000); on ACEi/ARB therapy	ESRD/ serum creatinine ×2 baseline (HR (30 + days)/ renal or CV death (HR 0.70; 95% CI 0.59 to 0.82)	 CV death/HHF (HR 0.69; 95% CI 0.57 to 0.83) CV death/ MI/ stroke (HR 0.80; 95% CI 0.67 to 0.95) HHF (HR 0.61; 95% CI 0.47 to 0.80) CV death (HR 0.78; 95% CI 0.61 to 1.00) CV death from any cause (HR 0.83; 95% CI 0.68 to 1.02) CV death/ MI/ stroke/ hospitalization from from 0.40 (HR 0.74; 95% CI 0.63 to 0.80)
DAPA-CKD [69]	Dapagliflozin vs. placebo	2.4 years	n= 4304; with or without diabetes; eGFR \geq 25 and \leq 75 ml/min/1.73m ² ; UACR \geq 200 or \leq 5000 mg/g; maximum tolerated daily dose of ACEi or ARB	≥50% decline in eGFR/reaching ESRD/CV death/renal death (HR 0.61; 95% CI 0.51–0.72)	 HHF/ CV death (HR 0.71; 95% CI 0.55–0.92) Death from any cause (HR 0.69; 95% CI 0.53–0.88) ≥ 50% decline in eGFR/reaching ESRD/renal death (HR 0.56; 95% CI 0.45–0.68)

continued)	
-	
able	

Trial Name	Drug	Duration (median)	Cohort	Primary outcome	Key secondary outcomes
EMPA-Kidney [70] (est completion June 2022)	Empagliflozin vs. placebo	3.1 years	n≈6000; CKD+risk of kidney disease pro- gression (depending on criteria); g on ACEi or ARB therapy	Time to first occurrence of: (i) Kidney disease progression (ESRD, sustained decline in eGFR to <10 mL/min/1.73m ² , renal death, decline of $\geq 40\%$ in eGFR) or (ii) Cardiovascular death	Time to: - HHF or CV death - All-cause hospitalisations - Death from any cause - First occurrence of kidney disease progression - CV death - CV death or ESRD
3P-MACE, Composite of: CV De Stage Renal Disease; KCCQ, Kan cle; CVD, cardiovascular disease;	ath; non-fatal MI; non-fatal S sas City Cardiomyopathy Qu LVEDV, LV end-diastolic vc	Stroke); ASC estionnaire; (olume; ESV,]	VD, atherosclerotic cardiovascular dis 5MWT, 6-min walk test; ITT, intentior End-systolic volume; EDV, end diastol	ease; UA, unstable angina; LVEF, left n to treat analysis; LVSD, left ventriculi lic volume; BIA, Bioelectrical impeden	ventricle ejection fraction; ESRD, end ar systolic dysfunction; LV, left ventri- ce Analysis; CPET, Cardio-pulmonary

Exercise Testing; LVM, LV mass; CMRI, cardiac magnetic resonance imaging; LVMi, BSA indexed LVM; CKD, chronic kidney disease; ACEi, angiotensin-converting-enzyme inhibitor; ARB

angiotensin-receptor blocker; UACR, urine albumin-to-creatinine ratio; RRT, renal replacement therapy

trials. Early mechanistic trials on conventional markers of LV remodelling have been inconclusive, raising the possibility that SGLT2-inhibitors act in a novel way that is not reflected by changes in LV remodelling as we currently understand them. In the following sections we explore new and emerging hypotheses into the molecular changes brought about by SGLT2-inhibition and their potential role in treating HF.

Calcium and the cardiomyocyte

The ionic balance within the cardiomyocyte is finely modulated by a variety of ion pumps on the cell surface membrane which interact via multiple overlapping pathways. Calcium (Ca^{2+}), is a key ion involved in excitation–contraction coupling, cardiac rhythmicity, also acting as a second messenger in regulating gene transcription for myocyte hypertrophy and other pathological remodelling pathways [71].

Action potentials generated at the cell surface trigger the opening of voltage-dependent L-type Ca^{2+} channels facilitating Ca^{2+} influx into the cardiomyocyte where it then binds to ryanodine R2 receptors (RyR2) on the sarcoplasmic reticulm (SR) to release intrasarcoplasmic Ca^{2+} . This process (called calcium-induced calcium release) amplifies the cytoplasmic Ca^{2+} content, which then bind to cardiac troponin, triggering muscle contraction. At the end of systole, the majority of Ca^{2+} ions are recycled back to the SR for storage via the SR Ca^{2+} -ATPase (SERCA2a), while the rest are extruded extracellularly via the sodium-calcium exchanger (NCX) [72] (Fig. 3).

HF and T2DM cause structural and functional changes to these pumps, and in doing so, disrupt the finely-balanced calcium homeostasis. There is increased activity of phospholamban which inhibits SERCA2a resulting in reduced Ca²⁺ reuptake into SR. This prolongs the relaxation phase (causing diastolic dysfunction) and reduces SR Ca²⁺ content available for use in the next contraction cycle (cauing systolic dysfunction). Likewise, phosphorylation of RyR2 pumps cause diastolic leakage of Ca²⁺ from the SR further reducing its storage capacity and increasing arrhythmogenicity due to increased cytosolic Ca^{2+} during diastole [73]. Additionally, there is also increased expression of SGLT1 and the sodium-hydrogen exchanger 1 (NHE1) on the cardiomyocyte cell membrane which increases intracytoplasmic sodium (Na⁺) concentrations [74, 75]. This reduces the efficiency of NCX pumps on the cell surface as well as those on mitochondria (which require low intracytoplasmic Na⁺ concentrations), resulting in increased cytoplasmic Ca²⁺ and reduced mitochondrial Ca²⁺ leading to myocyte hypertrophy, increased oxidative stress and accelerated cell death [76, 77] (Fig. 3).

SGLT2 is not expressed on the heart [78], yet there is strong pre-clinical evidence that SGLT2-inhibitors

Fig. 3 Schematic representation of sodium and calcium ion balance in the cardiomyocyte. Figure 2a: Ion balance in a healthy heart. Figure 2b: Abnormal calcium and sodium balance as a consequence of heart failure and type 2 diabetes Abbreviations: Ca²⁺-calcium; K⁺-potassium; LTCa-L-type calcium channel; Na⁺-sodium; Na⁺/K⁺ ATPase-sodium–potassium adenosine triphosphatase pump; NCX-sodium-calcium

exchanger; NHE-1-sodium-hydrogen exchanger subtype-1; PLB-phospholamban; ROS-reactive oxygen species; RyR2-ryanodine receptor 2; SERCA2a- sarcoplasmic reticulum calcium adenosine triphosphatase subtype 2a; SGLT-1-sodium-glucose cotransporter subtype 1; Blue spheres: calcium ions; Orange cubes: sodium ions; Green sphere: glucose molecule: Pink hexagon: potassium ion influence Ca^{2+} handling by modulating intracytoplasmic Na⁺ in the cardiomyocyte. This is supported by evidence that empagliflozin improves SERCA2a efficiency [79] and reduces RyR2-dependent Ca²⁺ leak [80, 81]. Additionally, empagliflozin may also preserve mitochondrial function and reduce oxidative damage [82]. SGLT2-inhibitors achieve this by inhibiting NHE1 (SGLT2-inhibitors inactivate NHE1 by binding to its Na⁺ binding site) and possibly SGLT1 as well (all SGLT2-inhibitors have intrinsic SGLT1 blocking ability – albeit to different degrees) [77, 83, 84].

Interestingly, T2DM is not the only trigger for SGLT1 upregulation in the heart; there is a 31% increase in SGLT1 expression seen in obese HF patients *without* T2DM [75]. Furthermore, preclinical data on empagliflozin shows its effects on myocyte Ca²⁺ and Na⁺ is independent of extracellular glucose levels and SGLT2 activity [85]. These findings could possibly explain the equal benefit of dapagliflozin in patients with or without T2DM in the DAPA-HF and EMPEROR-Reduced trials. Nevertheless, the only way to truly determine the effect of SGLT2-inhibiton on myocardial ionic homeostasis is by performing a dedicated clinical trial.

Powering the heart

The healthy heart utilises a variety of fuels to power its function. Under normal conditions, mitochondrial oxidative phosphorylation accounts for 95% of myocardial energy demands, whilst only 5% is met by glycolysis and TCA cycle. Between 70%—90% of the ATP generated by the heart is derived from free fatty acids (FFA) and the remainder from glucose and other substrates such as ketones, branch chain amino acids and lactate [86]. The heart can rapidly alter its fuel mix between these substrates depending upon the workload, perfusion and substrate bioavailability [87]. There are, however, important differences between these fuels; FFA requires more oxygen per ATP molecule produced compared to the more oxygen efficient glucose. However, each molecule of the energy dense FFA produces more ATP than that of glucose. Interestingly, ketones are more efficient than both, producing more energy with less oxygen compared to glucose and FFA respectively - leading to it being dubbed as a 'super fuel' [88].

FFAs are the primary energy source for the heart under most conditions, although its proportion of the overall fuel mix changes situationally. For example in the post prandial state, with higher circulating glucose and insulin, there is increased glucose uptake through myocardial GLUT proteins [86]. Following exercise when levels are raised, more lactate is used for energy production. Similarly, when ketone levels are raised it becomes the preferred fuel; and the heart has been shown to be able to oxidise more ketones per unit of mass than any other organ [89]. In contrast, during high intensity exercise (physiological hypoxemia) or ischaemia (pathological) there is a shift toward the more oxygenefficient glucose and glycolytic ATP production [88].

This shift in metabolism to favour glucose over FFA is a rapid response mechanism to acute changes, however, just like neurohormonal activation in the context of ventricular loading, if left unchecked it can result in 'metabolic remodelling' leading to cardiomyocyte dysfunction. Indeed, it is the most consistent metabolic change seen in animal and clinical studies of HF. A chronic overreliance on glucose results in suppression of genes related to FFA beta-oxidation pathways including their upstream regulators such as PPARalpha [90]. Downregulation of FFA oxidation capacity causes intracytoplasmic FFA accumulation, resulting in cardiac steatosis and lipotoxicity from reactive oxygen species [91]. Additionally, there will also be a large energy deficit due to the inability of glucose to produce nearly as much ATP as FFA oxidation. Together, this metabolic remodelling results in increased cellular toxicity and an energy deficit, culminating in cardiomyocyte dysfunction [90] (Fig. 4).

On the other hand, insulin resistance and increased proinflammatory cytokinesis (TNF and IL-6) which are hallmarks of T2DM, also induce lipolysis thereby increasing FFA delivery to the cardiomyocyte and worsening lipotoxicity [92]. As a further complication, insulin resistance reduces glucose delivery into the cardiomyocyte from insulin-sensitive GLUT4 (predominant route of glucose entry into cardiomyocyte) and GLUT8, forcing the heart to rely on the lower capacity (and insulin-independent) GLUT1 and SGLT1 routes. This exacerbates the energy deficit already present from lipotoxicity, thereby accelerating cardiomyocyte dysfunction [90, 93].

Given the effects of the various fuel substrates on cardiac efficiency and potentially even cellular dysfunction, optimizing myocardial energetics could be a key therapeutic target for patients with HF and T2DM. SGLT2-inhibitors are uniquely poised to fill this area of need as they have consistently shown to increase plasma ketone levels by increased hepatic synthesis from lipolysis and reduced renal loss [94]. (Fig. 2) Indeed, a recent porcine model of post infarct HF found metabolic remodelling with reduced FFA metabolism and anaerobic glycolysis resulted in a significant energy deficit. However, treatment with empagliflozin was able to switch myocardial metabolism away from anaerobic glycolysis toward utilisation of ketones, FFA and branch chain amino acids. The improved energetics resulted in higher myocardial ATP content and reduced adverse LV remodelling, enhanced LV function and reduced neurohormonal activation at 2 months post infarct [89]. Clinically, Nielsen and colleagues showed that an infusion of ketones bodies in patients with HF resulted in increased cardiac output and improved EF with an associated increase in heart rate and myocardial oxygen consumption. This was a small, proof of

◄Fig. 4 Schematic representation of myocardial energy consumption. Figure 3a: Myocardial energetics in healthy heart under resting conditions. Figure 3b: Myocardial energetics as a consequence of heart failure and type 2 diabetes. Figure 3c: Changes in myocardial energetics with the use of SGLT2-inhibitor therapy in patients with heart failure and type 2 diabetes Abbreviations: ATP-adenosine triphosphate; BCAA-branch chain amino acid; FFA-free fatty acid; HFheart failure; ROS-reactive oxygen species; SGLT2- sodium-glucose cotransporter subtype 2; T2DM- type 2 diabetes mellitus

concept trial in HF patients without T2DM where measurements were taken acutely following ketone infusion, administered in low dose insulinemic euglycaemic clamp conditions, acheiving fairly high levels of ketonemia [95]. It is yet unclear what the effects of a more modest, chronic ketonemia, as seen in SGLT2-inhibition, will be. There are ongoing efforts to determine the clinical effect of SGLT2-inhibition on myocardial energetics in the context of HF [96]. These studies will help determine if the future of managing HF will include metabolic modulation to improve myocardial energetics just as neurohormonal modulation to improve ventricular loading forms the backbone of HF therapeutics today.

SGLT2-inhibitor induced erythropoiesis

SGLT2-inhibitors increase haemoglobin and haematocrit levels. Initially these observations were attributed solely to haemoconcentration from its diuretic effect [97]. We now know that although there is an initial rise in urinary volume, this plateaus after a few weeks but haematocrit continues to rise well beyond that, suggesting a different mechanism driving this effect [98]. There is strong evidence demonstrating SGLT2-inhibition inducing erythropoiesis; a 12-week study reported increased haematocrit, haemoglobin, reticulocyte count and erythropoietin (EPO) with dapagliflozin [99], with similar results in a trial using empagliflozin in patients with T2DM and ishcaemic heart disease [100].

The mechanism behind these observations have yet to be elucidated, however there are a few hypotheses. EPO is synthesized in the renal cortex by EPO-producing fibroblasts. Patients with T2DM have increased filtration of glucose resulting in upregulation of SGLT1 and SGLT2 in the PCT to increase glucose resorption capacity, however this is a net energy-consuming process. The resulting relative cortical hypoxia and increased oxidative stress from higher energy demands of these transporters in the renal cortex causes the cortical fibroblasts to undergo transformation into myofibroblasts which no longer produce EPO. SGLT2-inhibitors block these transporters, thereby reducing energy demands—the cortical injury is reduced, the transformation reverses and EPO production capacity is restored. Additionally, SGLT2inhibition increases sodium delivery to the distal portions of the nephron which causes upregulation of medullary sodium transporters in the loop of henle and terminal nephron resulting in relative medullary hypoxia, which stimulates erythropoiesis. It has also been suggested that increased plasma ketone levels, which are associated with SGLT2-inhibitor use, directly contribute to EPO synthesis [100, 101].

There is evidence that erythropoiesis-stimulating agents (in the non-HF population) have beneficial cardiac effects including increased survival following ischaemia-reperfusion injury, reduced apoptosis, increased angiogenesis and improved myocardial contractility [102–104]. However, previous trials using various erythropoiesis-stimulating agents in patients with HF have been underwhelming and in the case of darbepoetin alfa, there was no effect on HF outcomes, but an increased risk of thromboembolic events [105]. Indeed, this was also an initial concern with SGLT2inhibitor therapy [106] but that has not been borne out with more recent trials and real-world data [18, 22, 23, 107]. In fact, a mediation analysis of the EMPA-REG OUTCOME showed changes in haemoglobin and haematocrit were responsible for approximately 50% of the risk reduction in CV death in that trial [108]. Perhaps SGLT2-inhibitors induce a more physiological effect, by restoring favourable renal physiology, which is salutary to the heart compared to exogenous stimulation of erythropoiesis by other agents.

Modulation of the sympathetic nervous system

The sympathetic nervous system is closely linked to the pathophysiology of HF and T2DM. Hypoperfusion from HF results in sympathetic activation, however persistently elevated sympathetic activity results in ventricular remodelling that perpetuates HF [109]. Similarly, insulin resistance in T2DM results in hyperinsulinemia which increases sympathetic tone through effects on central nervous outflow, baroreceptor reflex sensitivity and alteration of noradrenaline metabolism. However, persistently increased sympathetic activity results in insulin resistance from chronic skeletal muscle beta-adrenergic receptor activation and muscle hypoperfusion from persistent alpha-adrenergic vasoconstriction within skeletal muscles [110].

Interestingly, SGLT2-inhibitors have also shown potential in modulating sympathetic activity, thereby breaking the vicious cycle of chronic sympathetic activation in patients with T2DM and HF. Large clinical trials thus far have shown no reflex increase in heart rate following BP reduction from SGLT2-inhibition; a surrogate marker for sympathetic blockade. Mechanistic studies have supported these findings by demonstrating lower plasma metanephrines [89] and reduced noradrenaline and tyrosine hydroxylase (rate-limiting enzyme in catecholamine synthesis) activity in animal models [111]. Others have also noted that ketone bodies (which are increased with SGLT2-inhibitor therapy) attenuate sympathetic tone by suppressing G protein-coupled receptor 41 (GPR41) which is widely distributed in sympathetic ganglia [112]. These effects may also be responsible for lower sudden cardiac deaths in patients on SGLT2-inhibiton due to lower arrhythmia potential.

SGLT2 inhibitors – a new paradigm

The pathophysiology of HF involves 2 overarching themes – abnormal loading conditions on the heart (the failing pump) and dysfunctional mechanics at the cellular level (the failing cardiomyocyte) [113]. Insofar as the treatment of chronic HF is concerned, the benefit derived from current HF therapies are 'mechanical' due to favourable effects on ventricular loading. For the majority of patients, in spite of optimized loading conditions, HF disease progression continues unabated due to ongoing cellular dysfunction. We propose that SGLT2-inhibitors are uniquely poised to address this important but frequently overlooked and poorly understood aspect of HF.

Nevertheless, there may be a differential effect depending on where the patient is on the HF disease spectrum. In the early stages, with adequate cellular functional reserve, optimizing the molecular milieu within the cardiomyocyte with SGLT2-inhibitor therapy (i.e. improved calcium handling, efficient myocardial energetics, optimized oxygen delivery etc.) along with modest improvements in ventricular loading (i.e. diuresis and reduced BP) confers a large benefit. However, that benefit gradually wanes as cellular dysfunction, and indeed, cellular death occurs. In such advanced circumstances, improving haemodynamic loading becomes key, with the heavy lifting being done by neurohormonal modulators and SGLT2-inhibitors playing an important, but secondary role. If this hypothesis is true, then SGLT2inhibitors will very likely be the first therapeutic agent to be beneficial in patients with HFpEF where cardiomyocyte dysfunction predominates - we will have to wait and see if this is borne out in the large clinical trials currently underway.

Conclusions

Some have dubbed treating HF as the "last great battle in the war on cardiovascular disease" [113]. Despite great advances in treatment options, managing HF in the modern era remains an uphill battle. SGLT2-inhibitors have demonstrated great promise in the prevention and treatment of HF and CKD. As more clinical data are collected, its therapeutic potential is being realised well beyond its initial intended use as a diabetes drug. With a clearer understanding of these molecular mechanisms, we will be able to fully harness its true potential and perhaps even pave the way for a new era of molecular therapeutic agents in this fight against HF. Author contributions: AF and KV performed the literature search and drafted the manuscript; JSS conceptualized the review, critically revised and edited the manuscript.

Funding: JSS was funded by the European Foundation for the Study of Diabetes, Clinical Diabetes Research Program in Macrovascular Complications of Diabetes.

Compliance with ethical standards

Conflict of interest JSS has received financial and non-financial support from Boehringer Ingelheim, Astra Zeneca and Merck Sharp and Dohme for work that is relevant but unrelated to this article. AF and KV report no conflict of interest.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References

- Heidenreich PA, Albert NM, Allen LA, Bluemke DA, Butler J, Fonarow GC et al (2013) Forecasting the impact of heart failure in the United States: a policy statement from the American Heart Association. Circ Heart Fail 6(3):606–619. https://doi. org/10.1161/HHF.0b013e318291329a
- Cubbon RM, Adams B, Rajwani A, Mercer BN, Patel PA, Gherardi G et al (2013) Diabetes mellitus is associated with adverse prognosis in chronic heart failure of ischaemic and non-ischaemic aetiology. Diab Vasc Dis Res 10(4):330–336. https://doi.org/10.1177/1479164112471064
- Andersson C, Norgaard ML, Hansen PR, Fosbøl EL, Schmiegelow M, Weeke P et al (2010) Heart failure severity, as determined by loop diuretic dosages, predicts the risk of developing diabetes after myocardial infarction: a nationwide cohort study. Eur J Heart Fail 12(12):1333–1338. https://doi.org/10.1093/ eurjhf/hfq160
- Udell JA, Cavender MA, Bhatt DL, Chatterjee S, Farkouh ME, Scirica BM (2015) Glucose-lowering drugs or strategies and cardiovascular outcomes in patients with or at risk for type 2 diabetes: a meta-analysis of randomised controlled trials. Lancet Diabetes Endocrinol 3(5):356–366. https://doi.org/10.1016/ s2213-8587(15)00044-3
- Cosmi F, Shen L, Magnoli M, Abraham WT, Anand IS, Cleland JG et al (2018) Treatment with insulin is associated with worse outcome in patients with chronic heart failure and diabetes. Eur J Heart Fail 20(5):888–895. https://doi.org/10.1002/ ejhf.1146
- 6. U.S. Food and Drug Administration. FDA approves new treatment for a type of heart failure. https://www.fda.gov/

news-events/press-announcements/fda-approves-newtreatment-type-heart-failure. Accessed 10 June 2020

- Wright EM, Hirayama BA, Loo DF et al (2007) Active sugar transport in health and disease. J Intern Med 261(1):32–43. https://doi.org/10.1111/j.1365-2796.2006.01746.x
- Singh JS, Fathi A, Vickneson K, Mordi I, Mohan M, Houston JG et al (2016) Research into the effect Of SGLT2 inhibition on left ventricular remodelling in patients with heart failure and diabetes mellitus (REFORM) trial rationale and design. Cardiovasc Diabetol 15:97. https://doi.org/10.1186/s12933-016-0419-0
- Mazidi M, Rezaie P, Gao HK, Kengne AP (2017) Effect of sodium-glucose cotransport-2 inhibitors on blood pressure in people with type 2 diabetes mellitus: A systematic review and meta-analysis of 43 randomized control trials with 22 528 patients. J Am Heart Assoc 6(6). https://doi.org/10.1161/ jaha.116.004007
- Toyama T, Neuen BL, Jun M, Ohkuma T, Neal B, Jardine MJ et al (2019) Effect of SGLT2 inhibitors on cardiovascular, renal and safety outcomes in patients with type 2 diabetes mellitus and chronic kidney disease: A systematic review and meta-analysis. Diabetes Obes Metab 21(5):1237–1250. https://doi.org/10.1111/ dom.13648
- Nauck MA (2014) Update on developments with SGLT2 inhibitors in the management of type 2 diabetes. Drug Des Devel Ther 8:1335–1380. https://doi.org/10.2147/dddt.s50773
- Ferrannini E, Baldi S, Frascerra S, Astiarraga B, Heise T, Bizzotto R et al (2016) Shift to Fatty Substrate Utilization in Response to Sodium-Glucose Cotransporter 2 Inhibition in Subjects Without Diabetes and Patients With Type 2 Diabetes. Diabetes 65(5):1190–1195. https://doi.org/10.2337/db15-1356
- Alatrach M, Laichuthai N, Martinez R, Agyin C, Ali AM, Al-Jobori H et al (2020) Evidence Against an Important Role of Plasma Insulin and Glucagon Concentrations in the Increase in EGP Caused by SGLT2 Inhibitors. Diabetes 69(4):681–688. https://doi.org/10.2337/db19-0770
- 14. Cefalu WT, Leiter LA, Yoon KH, Arias P, Niskanen L, Xie J et al (2013) Efficacy and safety of canagliflozin versus glimepiride in patients with type 2 diabetes inadequately controlled with metformin (CANTATA-SU): 52 week results from a randomised, double-blind, phase 3 non-inferiority trial. Lancet 382(9896):941–950. https://doi.org/10.1016/ s0140-6736(13)60683-2
- Hallow KM, Helmlinger G, Greasley PJ, McMurray JJV, Boulton DW (2018) Why do SGLT2 inhibitors reduce heart failure hospitalization? A differential volume regulation hypothesis. Diabetes Obes Metab 20(3):479–487. https://doi.org/10.1111/dom.13126
- Staels B (2017) Cardiovascular Protection by Sodium Glucose Cotransporter 2 Inhibitors: Potential Mechanisms. Am J Med 130(6s):S30–S39. https://doi.org/10.1016/j.amjmed.2017.04.009
- Mordi NA, Mordi IR, Singh JS, McCrimmon RJ, Struthers AD, Lang CC et al (2020) Renal and Cardiovascular Effects of SGLT2 Inhibition in Combination with Loop Diuretics in Patients with Type 2 Diabetes and Chronic Heart Failure: The RECEDE-CHF Trial. Circulation. https://doi.org/10.1161/ circulationaha.120.048739
- Zinman B, Wanner C, Lachin JM, Fitchett D, Bluhmki E, Hantel S et al (2015) Empagliflozin, Cardiovascular Outcomes, and Mortality in Type 2 Diabetes. N Engl J Med 373(22):2117–2128. https://doi.org/10.1056/NEJMoa1504720
- Fitchett D, Inzucchi SE, Cannon CP, McGuire DK, Scirica BM, Johansen OE et al (2019) Empagliflozin Reduced Mortality and Hospitalization for Heart Failure Across the Spectrum of Cardiovascular Risk in the EMPA-REG OUTCOME Trial. Circulation 139(11):1384–1395. https://doi.org/10.1161/ circulationaha.118.037778

- 20. Fitchett D, Butler J, van de Borne P, Zinman B, Lachin JM, Wanner C et al (2018) Effects of empagliflozin on risk for cardiovascular death and heart failure hospitalization across the spectrum of heart failure risk in the EMPA-REG OUTCOME(R) trial. Eur Heart J 39(5):363–370. https://doi.org/10.1093/eurheartj/ehx511
- Wanner C, Lachin JM, Inzucchi SE, Fitchett D, Mattheus M, George J et al (2018) Empagliflozin and Clinical Outcomes in Patients With Type 2 Diabetes Mellitus, Established Cardiovascular Disease, and Chronic Kidney Disease. Circulation 137(2):119–129. https://doi.org/10.1161/ circulationaha.117.028268
- 22. Neal B, Perkovic V, Mahaffey KW, de Zeeuw D, Fulcher G, Erondu N et al (2017) Canagliflozin and Cardiovascular and Renal Events in Type 2 Diabetes. N Engl J Med 377(7):644–657. https://doi.org/10.1056/NEJMoa1611925
- Wiviott SD, Raz I, Bonaca MP, Mosenzon O, Kato ET, Cahn A et al (2019) Dapagliflozin and Cardiovascular Outcomes in Type 2 Diabetes. N Engl J Med 380(4):347–357. https://doi. org/10.1056/NEJMoa1812389
- 24. Kato ET, Silverman MG, Mosenzon O, Zelniker TA, Cahn A, Furtado RHM et al (2019) Effect of Dapagliflozin on Heart Failure and Mortality in Type 2 Diabetes Mellitus. Circulation 139(22):2528–2536. https://doi.org/10.1161/ circulationaha.119.040130
- Zelniker TA, Wiviott SD, Raz I, Im K, Goodrich EL, Bonaca MP et al (2019) SGLT2 inhibitors for primary and secondary prevention of cardiovascular and renal outcomes in type 2 diabetes: a systematic review and meta-analysis of cardiovascular outcome trials. Lancet 393(10166):31–39. https://doi.org/10.1016/ s0140-6736(18)32590-x
- 26. Kosiborod M, Lam CSP, Kohsaka S, Kim DJ, Karasik A, Shaw J et al (2018) Cardiovascular Events Associated With SGLT-2 Inhibitors Versus Other Glucose-Lowering Drugs: The CVD-REAL 2 Study. J Am Coll Cardiol 71(23):2628–2639. https://doi.org/10.1016/j.jacc.2018.03.009
- 27. Patorno E, Pawar A, Franklin JM, Najafzadeh M, Déruaz-Luyet A, Brodovicz KG et al (2019) Empagliflozin and the Risk of Heart Failure Hospitalization in Routine Clinical Care. Circulation 139(25):2822–2830. https://doi.org/10.1161/ circulationaha.118.039177
- Cannon C, McGuire D, Cherney D. Results of the eValuation of ERTugliflozin EffIcacy and Safety CardioVascular Outcomes Trial (Vertis CV). 80th American Diabetes Association Scientific Sessions; 2020 June 16. Online Symposium
- Cosentino F, Grant PJ, Aboyans V, Bailey CJ, Ceriello A, Delgado V et al (2020) 2019 ESC Guidelines on diabetes, pre-diabetes, and cardiovascular diseases developed in collaboration with the EASD. Eur Heart J 41(2):255–323. https://doi.org/10.1093/ eurheartj/ehz486
- Davies MJ, D'Alessio DA, Fradkin J, Kernan WN, Mathieu C, Mingrone G et al (2018) Management of Hyperglycemia in Type 2 Diabetes, 2018 A Consensus Report by the American Diabetes Association (ADA) and the European Association for the Study of Diabetes (EASD). Diabetes Care 41(12):2669–2701. https:// doi.org/10.2337/dci18-0033
- McMurray JJV, Solomon SD, Inzucchi SE, Kober L, Kosiborod MN, Martinez FA et al (2019) Dapagliflozin in Patients with Heart Failure and Reduced Ejection Fraction. N Engl J Med 381(21):1995–2008. https://doi.org/10.1056/NEJMoa1911303
- 32. Packer M, Anker SD, Butler J, Filippatos G, Pocock SJ, Carson P et al (2020) Cardiovascular and Renal Outcomes with Empagliflozin in Heart Failure. N Engl J Med. https://doi.org/10.1056/ NEJMoa2022190
- 33. Singh JSS, Burrell LM, Cherif M, Squire IB, Clark AL, Lang CC et al (2017) Sacubitril/valsartan: beyond natriuretic peptides.

Heart 103(20):1569–1577. https://doi.org/10.1136/heartjnl-2017-311295

- McMurray JJV, Solomon SD, Docherty KF, Jhund PS et al (2020) The Dapagliflozin And Prevention of Adverse outcomes in Heart Failure trial (DAPA-HF) in context. Eur Heart J. https://doi. org/10.1093/eurheartj/ehz916
- Pfeffer MA, Shah AM, Borlaug BA (2019) Heart Failure With Preserved Ejection Fraction In Perspective. Circ Res 124(11):1598–1617. https://doi.org/10.1161/ circresaha.119.313572
- 36. Zile MR, Baicu CF, Gaasch WH et al (2004) Diastolic heart failure–abnormalities in active relaxation and passive stiffness of the left ventricle. N Engl J Med 350(19):1953–1959. https:// doi.org/10.1056/NEJMoa032566
- Yusuf S, Pfeffer MA, Swedberg K, Granger CB, Held P, McMurray JJ et al (2003) Effects of candesartan in patients with chronic heart failure and preserved left-ventricular ejection fraction: the CHARM-Preserved Trial. Lancet 362(9386):777–781. https://doi.org/10.1016/s0140-6736(03)14285-7
- Massie BM, Carson PE, McMurray JJ, Komajda M, McKelvie R, Zile MR et al (2008) Irbesartan in patients with heart failure and preserved ejection fraction. N Engl J Med 359(23):2456–2467. https://doi.org/10.1056/NEJMoa0805450
- Cleland JG, Tendera M, Adamus J, Freemantle N, Polonski L, Taylor J (2006) The perindopril in elderly people with chronic heart failure (PEP-CHF) study. Eur Heart J 27(19):2338–2345. https://doi.org/10.1093/eurheartj/ehl250
- 40. Pitt B, Pfeffer MA, Assmann SF, Boineau R, Anand IS, Claggett B et al (2014) Spironolactone for heart failure with preserved ejection fraction. N Engl J Med 370(15):1383–1392. https://doi. org/10.1056/NEJMoa1313731
- Solomon SD, McMurray JJV, Anand IS, Ge J, Lam CSP, Maggioni AP et al (2019) Angiotensin-Neprilysin Inhibition in Heart Failure with Preserved Ejection Fraction. N Engl J Med 381(17):1609–1620. https://doi.org/10.1056/NEJMoa1908655
- 42. Lam CSP, Voors AA, de Boer RA, Solomon SD, van Veldhuisen DJ et al (2018) Heart failure with preserved ejection fraction: from mechanisms to therapies. Eur Heart J 39(30):2780–2792. https://doi.org/10.1093/eurheartj/ehy301
- Ronco C, Haapio M, House AA, Anavekar N, Bellomo R et al (2008) Cardiorenal syndrome. J Am Coll Cardiol 52(19):1527– 1539. https://doi.org/10.1016/j.jacc.2008.07.051
- 44. Hillege HL, Nitsch D, Pfeffer MA, Swedberg K, McMurray JJ, Yusuf S et al (2006) Renal function as a predictor of outcome in a broad spectrum of patients with heart failure. Circulation 113(5):671–678. https://doi.org/10.1161/circulationaha.105.580506
- 45. Heywood JT, Fonarow GC, Costanzo MR, Mathur VS, Wigneswaran JR, Wynne J et al (2007) High prevalence of renal dysfunction and its impact on outcome in 118,465 patients hospitalized with acute decompensated heart failure: a report from the ADHERE database. J Card Fail 13(6):422–430. https://doi. org/10.1016/j.cardfail.2007.03.011
- 46. Cherney DZ, Perkins BA (2014) Sodium-glucose cotransporter 2 inhibition in type 1 diabetes: simultaneous glucose lowering and renal protection? Can J Diabetes 38(5):356–363. https://doi. org/10.1016/j.jcjd.2014.05.006
- Fioretto P, Zambon A, Rossato M, Busetto L, Vettor R et al (2016) SGLT2 Inhibitors and the Diabetic Kidney. Diabetes Care 39(Suppl 2):S165–S171. https://doi.org/10.2337/dcS15-3006
- Cherney DZI, Heerspink HJL, Frederich R, Maldonado M, Liu J, Pong A et al (2020) Effects of ertugliflozin on renal function over 104 weeks of treatment: a post hoc analysis of two randomised controlled trials. Diabetologia 63(6):1128–1140. https://doi.org/10.1007/s00125-020-05133-4

- Björck S, Mulec H, Johnsen SA, Nordén G, Aurell M et al (1992) Renal protective effect of enalapril in diabetic nephropathy. BMJ 304(6823):339–343. https://doi.org/10.1136/bmj.304.6823.339
- Barnett AH, Bain SC, Bouter P, Karlberg B, Madsbad S, Jervell J et al (2004) Angiotensin-receptor blockade versus convertingenzyme inhibition in type 2 diabetes and nephropathy. N Engl J Med 351(19):1952–1961. https://doi.org/10.1056/NEJMoa0422 74
- 51. Lam CSP, Chandramouli C, Ahooja V, Verma S et al (2019) SGLT-2 Inhibitors in Heart Failure: Current Management, Unmet Needs, and Therapeutic Prospects. J Am Heart Assoc 8(20):e013389. https://doi.org/10.1161/jaha.119.013389
- 52. Perkovic V, Jardine MJ, Neal B, Bompoint S, Heerspink HJL, Charytan DM et al (2019) Canagliflozin and Renal Outcomes in Type 2 Diabetes and Nephropathy. N Engl J Med 380(24):2295– 2306. https://doi.org/10.1056/NEJMoa1811744
- 53. European Society of Cardiology. DAPA-CKD trial meets primary endpoint in patients with chronic kidney disease. https:// www.escardio.org/The-ESC/Press-Office/Press-releases/DAPA. Accessed 09 Sep 2020
- Shah KS, Xu H, Matsouaka RA, Bhatt DL, Heidenreich PA, Hernandez AF et al (2017) Heart Failure With Preserved, Borderline, and Reduced Ejection Fraction: 5-Year Outcomes. J Am Coll Cardiol 70(20):2476–2486. https://doi.org/10.1016/j. jacc.2017.08.074
- 55. Johnson & Johnson. U.S. FDA Approves INVOKANA® (canagliflozin) to Treat Diabetic Kidney Disease (DKD) and Reduce the Risk of Hospitalization for Heart Failure in Patients with Type 2 Diabetes (T2D) and DKD. https://www.jnj.com/u-sfda-approves-invokana-canagliflozin-to-treat-diabetic-kidneydisease-dkd-and-reduce-the-risk-of-hospitalization-for-heartfailure-in-patients-with-type-2-diabetes-t2d-and-dkd. Accessed 18 June 2020
- 56. AstraZenaca. FDA grants Fast Track designation for Farxiga in chronic kidney disease. https://www.astrazeneca.com/mediacentre/press-releases/2019/fda-grants-fast-track-designationfor-farxiga-in-chronic-kidney-disease-27082019.html#:~:text= Astra Zenec a% 20tod ay% 20ann ounce d% 20that% 20the, chronic%20kidney%20disease%20(CKD). Accessed 09 Sep 2020
- 57. Kramer DG, Trikalinos TA, Kent DM, Antonopoulos GV, Konstam MA, Udelson JE et al (2010) Quantitative evaluation of drug or device effects on ventricular remodeling as predictors of therapeutic effects on mortality in patients with heart failure and reduced ejection fraction: a meta-analytic approach. J Am Coll Cardiol 56(5):392–406. https://doi.org/10.1016/j.jacc.2010.05.011
- Groenning BA, Nilsson JC, Sondergaard L, Fritz-Hansen T, Larsson HB, Hildebrandt PR et al (2000) Antiremodeling effects on the left ventricle during beta-blockade with metoprolol in the treatment of chronic heart failure. J Am Coll Cardiol 36(7):2072–2080
- 59. Cicoira M, Zanolla L, Rossi A, Golia G, Franceschini L, Brighetti G et al (2002) Long-term, dose-dependent effects of spironolactone on left ventricular function and exercise tolerance in patients with chronic heart failure. J Am Coll Cardiol 40(2):304–310. https://doi.org/10.1016/s0735-1097(02)01965-4
- 60. Greenberg B, Quinones MA, Koilpillai C, Limacher M, Shindler D, Benedict C et al (1995) Effects of long-term enalapril therapy on cardiac structure and function in patients with left ventricular dysfunction Results of the SOLVD echocardiography substudy. Circulation 91(10):2573–2581. https://doi.org/10.1161/01.cir.91.10.2573
- 61. Singh JSS, Mordi IR, Vickneson K, Fathi A, Donnan PT, Mohan M et al (2020) Dapagliflozin Versus Placebo on Left Ventricular Remodeling in Patients With Diabetes and Heart Failure: The

REFORM Trial. Diabetes Care 43(6):1356–1359. https://doi. org/10.2337/dc19-2187

- 62. Nassif ME, Windsor SL, Tang F, Khariton Y, Husain M, Inzucchi SE et al (2019) Dapagliflozin Effects on Biomarkers, Symptoms, and Functional Status in Patients With Heart Failure With Reduced Ejection Fraction: The DEFINE-HF Trial. Circulation 140(18):1463–1476. https://doi.org/10.1161/ circulationaha.119.042929
- 63. Singh JSS, Mordi I, Fathi A, Vickneson K, Donnan PT, Mohan M et al (2018) P905 Research into the effect of sodium-glucose linked transporter inhibition in left ventricular remodelling in patients with heart failure and diabetes mellitus. European Heart J 39(suppl_1). https://doi.org/10.1093/eurheartj/ ehy564.P905
- 64. Verma S, Mazer CD, Yan AT, Mason T, Garg V, Teoh H et al (2019) Effect of Empagliflozin on Left Ventricular Mass in Patients With Type 2 Diabetes Mellitus and Coronary Artery Disease: The EMPA-HEART CardioLink-6 Randomized Clinical Trial. Circulation 140(21):1693–1702. https://doi.org/10.1161/ circulationaha.119.042375
- NCT03619213. Dapagliflozin Evaluation to Improve the Lives of Patients With Preserved Ejection Fraction Heart Failure (DELIVER). https://clinicaltrials.gov/ct2/show/NCT03619213. Accessed 10 July 2020
- 66. NCT03057951. Empagliflozin Outcome Trial in Patients With Chronic Heart Failure With Preserved Ejection Fraction (EMPEROR-Preserved). https://clinicaltrials.gov/ct2/show/ NCT03057951. Accessed 10 July 2020
- 67. Brown AJM, Gandy S, McCrimmon R, Houston JG, Struthers AD, Lang CC et al (2020) A randomized controlled trial of dapagliflozin on left ventricular hypertrophy in people with type two diabetes: the DAPA-LVH trial. Eur Heart J. https://doi. org/10.1093/eurheartj/ehaa419
- NCT03485222. Are the "Cardiac Benefits" of Empagliflozin Independent of Its Hypoglycemic Activity? (ATRU-4). (EMPA-TROPISM). https://clinicaltrials.gov/ct2/show/NCT03485222. Accessed 10 July 2020
- NCT03036150. A Study to Evaluate the Effect of Dapagliflozin on Renal Outcomes and Cardiovascular Mortality in Patients With Chronic Kidney Disease (Dapa-CKD). https:// clinicaltrials.gov/ct2/show/NCT03036150. Accessed 10 July 2020
- NCT03594110. EMPA-KIDNEY (The Study of Heart and Kidney Protection With Empagliflozin). https://clinicaltrials.gov/ct2/ show/NCT03594110. Accessed 10 July 2020
- Goonasekera SA, Hammer K, Auger-Messier M, Bodi I, Chen X, Zhang H et al (2012) Decreased cardiac L-type Ca²+ channel activity induces hypertrophy and heart failure in mice. J Clin Invest 122(1):280–290. https://doi.org/10.1172/jci58227
- 72. Walweel K, Laver DR (2015) Mechanisms of SR calcium release in healthy and failing human hearts. Biophys Rev 7(1):33–41. https://doi.org/10.1007/s12551-014-0152-4
- Hasenfuss G (1998) Alterations of calcium-regulatory proteins in heart failure. Cardiovasc Res 37(2):279–289. https://doi. org/10.1016/s0008-6363(97)00277-0
- 74. Baartscheer A, Schumacher CA, van Borren MMGJ, Belterman CNW, Coronel R, Fiolet JWT et al (2003) Increased Na+/H+exchange activity is the cause of increased [Na+]i and underlies disturbed calcium handling in the rabbit pressure and volume overload heart failure model. Cardiovasc Res 57(4):1015–1024. https://doi.org/10.1016/s0008-6363(02)00809-x
- Lambert R, Srodulski S, Peng X, Margulies KB, Despa F, Despa S (2015) Intracellular Na+ concentration ([Na+]i) is elevated in diabetic hearts due to enhanced Na+-glucose cotransport. J Am Heart Assoc 4(9). https://doi.org/10.1161/jaha.115.002183

- Bay J, Kohlhaas M, Maack C et al (2013) Intracellular Na+ and cardiac metabolism. J Mol Cell Cardiol 61:20–27. https://doi. org/10.1016/j.yjmcc.2013.05.010
- 77. Uthman L, Baartscheer A, Schumacher CA, Fiolet JWT, Kuschma MC, Hollmann MW et al (2018) Direct cardiac actions of sodium glucose cotransporter 2 inhibitors target pathogenic mechanisms underlying heart failure in diabetic patients. Front Physiol 9. https://doi.org/10.3389/fphys.2018.01575
- Wright EM, Loo DD, Hirayama BA et al (2011) Biology of human sodium glucose transporters. Physiol Rev 91(2):733–794. https://doi.org/10.1152/physrev.00055.2009
- Hammoudi N, Jeong D, Singh R, Farhat A, Komajda M, Mayoux E et al (2017) Empagliflozin Improves Left Ventricular Diastolic Dysfunction in a Genetic Model of Type 2 Diabetes. Cardiovasc Drugs Ther 31(3):233–246. https://doi.org/10.1007/ s10557-017-6734-1
- Mustroph J, Wagemann O, Lücht CM, Trum M, Hammer KP, Sag CM et al (2018) Empagliflozin reduces Ca/calmodulin-dependent kinase II activity in isolated ventricular cardiomyocytes. ESC Heart Fail 5(4):642–648. https://doi.org/10.1002/ehf2.12336
- Lee TI, Chen YC, Lin YK, Chung CC, Lu YY, Kao YH et al (2019) Empagliflozin attenuates myocardial sodium and calcium dysregulation and reverses cardiac remodeling in streptozotocin-induced diabetic rats. Int J Mol Sci 20(7). https://doi. org/10.3390/ijms20071680
- Andreadou I, Efentakis P, Balafas E, Togliatto G, Davos CH, Varela A et al (2017) Empagliflozin Limits Myocardial Infarction in Vivo and Cell Death in Vitro: Role of STAT3, Mitochondria, and Redox Aspects. Front Physiol 8:1077. https://doi.org/10.3389/ fphys.2017.01077
- Uthman L, Baartscheer A, Bleijlevens B, Schumacher CA, Fiolet JWT, Koeman A et al (2018) Class effects of SGLT2 inhibitors in mouse cardiomyocytes and hearts: inhibition of Na+/H+ exchanger, lowering of cytosolic Na+ and vasodilation. Diabetologia 61(3):722–726. https://doi.org/10.1007/s00125-017-4509-7
- 84. Li Z, Agrawal V, Ramratnam M, Sharma RK, D'Auria S, Sincoular A et al (2019) Cardiac sodium-dependent glucose cotransporter 1 is a novel mediator of ischaemia/reperfusion injury. Cardiovasc Res 115(11):1646–1658. https://doi.org/10.1093/ cvr/cvz037
- 85. Baartscheer A, Schumacher CA, Wüst RCI, Fiolet JWT, Stienen GJM, Coronel R et al (2017) Empagliflozin decreases myocardial cytoplasmic Na+ through inhibition of the cardiac Na+/ H+ exchanger in rats and rabbits. Diabetologia 60(3):568–573. https://doi.org/10.1007/s00125-016-4134-x
- Doenst T, Nguyen TD, Abel ED (2013) Cardiac Metabolism in Heart Failure - Implications beyond ATP production. Circ Res 113(6):709–724. https://doi.org/10.1161/circresaha.113.300376
- Stanley WC, Recchia FA, Lopaschuk GD (2005) Myocardial substrate metabolism in the normal and failing heart. Physiol Rev 85(3):1093–1129. https://doi.org/10.1152/physrev.00006.2004
- Mudaliar S, Alloju S, Henry RR et al (2016) Can a Shift in Fuel Energetics Explain the Beneficial Cardiorenal Outcomes in the EMPA-REG OUTCOME Study? A Unifying Hypothesis Diabetes Care 39(7):1115–1122. https://doi.org/10.2337/dc16-0542
- Santos-Gallego CG, Requena-Ibanez JA, San Antonio R, Ishikawa K, Watanabe S, Picatoste B et al (2019) Empagliflozin Ameliorates Adverse Left Ventricular Remodeling in Nondiabetic Heart Failure by Enhancing Myocardial Energetics. J Am Coll Cardiol 73(15):1931–1944. https://doi.org/10.1016/j. jacc.2019.01.056
- Wende AR, Brahma MK, McGinnis GR, Young ME et al (2017) Metabolic Origins of Heart Failure. JACC Basic Transl Sci 2(3):297–310. https://doi.org/10.1016/j.jacbts.2016.11.009

- 91. Ritchie RH (2009) Evidence for a causal role of oxidative stress in the myocardial complications of insulin resistance. Heart Lung Circ 18(1):11–18. https://doi.org/10.1016/j.hlc.2008.11.003
- Yang Q, Vijayakumar A, Kahn BB (2018) Metabolites as regulators of insulin sensitivity and metabolism. Nat Rev Mol Cell Biol 19(10):654–672. https://doi.org/10.1038/s41580-018-0044-8
- Aerni-Flessner L, Abi-Jaoude M, Koenig A, Payne M, Hruz PW et al (2012) GLUT4, GLUT1, and GLUT8 are the dominant GLUT transcripts expressed in the murine left ventricle. Cardiovasc Diabetol 11:63. https://doi.org/10.1186/1475-2840-11-63
- 94. Mudaliar S, Polidori D, Zambrowicz B, Henry RR et al (2015) Sodium-Glucose Cotransporter Inhibitors: Effects on Renal and Intestinal Glucose Transport: From Bench to Bedside. Diabetes Care 38(12):2344–2353. https://doi.org/10.2337/dc15-0642
- 95. Nielsen R, Møller N, Gormsen LC, Tolbod LP, Hansson NH, Sorensen J et al (2019) Cardiovascular Effects of Treatment With the Ketone Body 3-Hydroxybutyrate in Chronic Heart Failure Patients. Circulation 139(18):2129–2141. https://doi. org/10.1161/circulationaha.118.036459
- 96. Åkerblom A, Oldgren J, Latva-Rasku A, Johansson L, Lisovskaja V, Karlsson C et al (2019) Effects of DAPAgliflozin on CAR-Diac substrate uptake, myocardial efficiency, and myocardial contractile work in type 2 diabetes patients-a description of the DAPACARD study. Ups J Med Sci 124(1):59–64. https://doi.org/10.1080/03009734.2018.1515281
- 97. Bailey CJ, Gross JL, Pieters A, Bastien A, List JF et al (2010) Effect of dapagliflozin in patients with type 2 diabetes who have inadequate glycaemic control with metformin: a randomised, double-blind, placebo-controlled trial. Lancet 375(9733):2223– 2233. https://doi.org/10.1016/s0140-6736(10)60407-2
- Sano M, Goto S (2019) Possible Mechanism of Hematocrit Elevation by Sodium Glucose Cotransporter 2 Inhibitors and Associated Beneficial Renal and Cardiovascular Effects. Circulation 139(17):1985–1987. https://doi.org/10.1161/ circulationaha.118.038881
- 99. Lambers Heerspink HJ, de Zeeuw D, Wie L, Leslie B, List J et al (2013) Dapagliflozin a glucose-regulating drug with diuretic properties in subjects with type 2 diabetes. Diabetes Obes Metab 15(9):853–862. https://doi.org/10.1111/dom.12127
- 100. Mazer CD, Hare GMT, Connelly PW, Gilbert RE, Shehata N, Quan A et al (2020) Effect of Empagliflozin on Erythropoietin Levels, Iron Stores, and Red Blood Cell Morphology in Patients With Type 2 Diabetes Mellitus and Coronary Artery Disease. Circulation 141(8):704–707. https://doi.org/10.1161/ circulationaha.119.044235
- 101. Morla L, Edwards A, Crambert G et al (2016) New insights into sodium transport regulation in the distal nephron: Role of G-protein coupled receptors. World J Biol Chem 7(1):44–63. https:// doi.org/10.4331/wjbc.v7.i1.44
- 102. Piuhola J, Kerkelä R, Keenan JI, Hampton MB, Richards AM, Pemberton CJ (2008) Direct cardiac actions of erythropoietin (EPO): effects on cardiac contractility, BNP secretion and ischaemia/reperfusion injury. Clin Sci (Lond) 114(4):293–304. https:// doi.org/10.1042/cs20070229

- 103. van der Meer P, Voors AA, Lipsic E, van Gilst WH, van Veldhuisen DJ et al (2004) Erythropoietin in cardiovascular diseases. Eur Heart J 25(4):285–291. https://doi.org/10.1016/j. ehj.2003.11.017
- 104. Sanchis-Gomar F, Garcia-Gimenez JL, Pareja-Galeano H, Romagnoli M, Perez-Quilis C, Lippi G et al (2014) Erythropoietin and the heart: physiological effects and the therapeutic perspective. Int J Cardiol 171(2):116–125. https://doi.org/10.1016/j. ijcard.2013.12.011
- 105. Swedberg K, Young JB, Anand IS, Cheng S, Desai AS, Diaz R et al (2013) Treatment of anemia with darbepoetin alfa in systolic heart failure. N Engl J Med 368(13):1210–1219. https://doi. org/10.1056/NEJMoa1214865
- 106. Wu JH, Foote C, Blomster J, Toyama T, Perkovic V, Sundström J et al (2016) Effects of sodium-glucose cotransporter-2 inhibitors on cardiovascular events, death, and major safety outcomes in adults with type 2 diabetes: a systematic review and meta-analysis. Lancet Diabetes Endocrinol 4(5):411–419. https://doi.org/10.1016/s2213-8587(16)00052-8
- 107. Kosiborod M, Birkeland KI, Cavender MA, Fu AZ, Wilding JP, Khunti K et al (2018) Rates of myocardial infarction and stroke in patients initiating treatment with SGLT2-inhibitors versus other glucose-lowering agents in real-world clinical practice: Results from the CVD-REAL study. Diabetes Obes Metab 20(8):1983–1987. https://doi.org/10.1111/dom.13299
- Inzucchi SE, Zinman B, Fitchett D, Wanner C, Ferrannini E, Schumacher M et al (2018) How Does Empagliflozin Reduce Cardiovascular Mortality? Insights From a Mediation Analysis of the EMPA-REG OUTCOME Trial. Diabetes Care 41(2):356– 363. https://doi.org/10.2337/dc17-1096
- 109. Florea VG, Cohn JN (2014) The autonomic nervous system and heart failure. Circ Res 114(11):1815–1826. https://doi. org/10.1161/circresaha.114.302589
- Perin PC, Maule S, Quadri R et al (2001) Sympathetic nervous system, diabetes, and hypertension. Clin Exp Hypertens 23(1–2):45–55. https://doi.org/10.1081/ceh-100001196
- 111. Herat LY, Magno AL, Rudnicka C, Hricova J, Carnagarin R, Ward NC et al (2020) SGLT2 Inhibitor-Induced Sympathoinhibition: A Novel Mechanism for Cardiorenal Protection. JACC Basic Transl Sci 5(2):169–179. https://doi.org/10.1016/j. jacbts.2019.11.007
- 112. Kimura I, Inoue D, Maeda T, Hara T, Ichimura A, Miyauchi S et al (2011) Short-chain fatty acids and ketones directly regulate sympathetic nervous system via G protein-coupled receptor 41 (GPR41). Proc Natl Acad Sci U S A 108(19):8030–8035. https://doi.org/10.1073/pnas.1016088108
- 113. Braunwald E (2013) Heart failure. JACC Heart Fail 1(1):1–20. https://doi.org/10.1016/j.jchf.2012.10.002

Publisher's Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.