

University of Groningen

Cardiovascular effects of non-cardiovascular drugs in heart failure

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DOI:
[10.33612/diss.132706675](https://doi.org/10.33612/diss.132706675)

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

Publication date:
2020

[Link to publication in University of Groningen/UMCG research database](#)

Citation for published version (APA):
Yurista, S. (2020). *Cardiovascular effects of non-cardiovascular drugs in heart failure*. University of Groningen. <https://doi.org/10.33612/diss.132706675>

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Effects of sodium–glucose co-transporter 2 inhibition with empaglifozin on renal structure and function in non-diabetic rats with left ventricular dysfunction after myocardial infarction

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ABSTRACT

Background: The use of sodium–glucose co-transporter 2 inhibitors (SGLT2i) is currently expanding to cardiovascular risk reduction in non-diabetic subjects, but renal (side-)effects are less well studied in this setting.

Methods: Male non-diabetic Sprague-Dawley rats underwent permanent coronary artery ligation to induce MI, or sham surgery. Rats received chow containing empagliflozin (EMPA) (30 mg/kg/day) or control chow. Renal function and electrolyte balance were measured in metabolic cages. Histological and molecular markers of kidney injury, parameters of phosphate homeostasis and bone resorption were also assessed.

Results: EMPA resulted in a two-fold increase in diuresis, without evidence for plasma volume contraction or impediments in renal function in both sham and MI animals. EMPA increased plasma magnesium levels, while the levels of glucose and other major electrolytes were comparable among the groups. Urinary protein excretion was similar in all treatment groups and no histomorphological alterations were identified in the kidney. Accordingly, molecular markers for cellular injury, fibrosis, inflammation and oxidative stress in renal tissue were comparable between groups. EMPA resulted in a slight increase in circulating phosphate and PTH levels without activating FGF23-Klotho axis in the kidney and bone mineral resorption, measured with CTX-1, was not increased.

Conclusions: EMPA exerts profound diuretic effects without compromising renal structure and function or causing significant electrolyte imbalance in a non-diabetic setting. The slight increase in circulating phosphate and PTH after EMPA treatment was not associated with evidence for increased bone mineral resorption suggesting that EMPA does not affect bone health.

Keywords: Sodium–glucose co-transporter 2 inhibitors • Diabetes • Heart failure • Renal function

INTRODUCTION

Sodium glucose co-transporter 2 inhibitors (SGLT2i) reduce cardiovascular (CV) events and prevent heart failure (HF) hospitalizations when given to diabetic subjects with either established CV disease or with multiple risk factors for CV disease [1–3]. Furthermore, SGLT2i also appear to slow the progression of diabetic kidney disease in patients with type 2 diabetes mellitus (T2D) [4–6]. On the other hand, it has also been suggested that SGLT2-inhibitors promote bone mineral reabsorption and may increase fracture risk in these patients [7–9].

The substantial reduction in HF hospitalizations observed in patients with T2D have led to the hypothesis that these drugs could also be beneficial in non-diabetic patients. Indeed, we and others have recently demonstrated that the SGLT2i EMPA improves cardiac function and ameliorates cardiac remodelling in non-diabetic animals with HF after a large myocardial infarction (MI) [10,11]. The effects of SGLT2i on clinical outcomes are also currently under investigation in several phase 3 clinical trials that include diabetic and non-diabetic HF patients (NCT03057977, NCT03057951, NCT03036124, NCT03619213). Recently, the Dapagliflozin And Prevention of Adverse-outcomes in Heart Failure (DAPA-HF, NCT03036124) trial revealed that the SGLT2i dapagliflozin reduced the risk of worsening heart failure or cardiovascular mortality, regardless of the presence or absence of diabetes [12]. Importantly, dapagliflozin did not appear to compromise renal function [12].

SGLT2i have potent diuretic properties, which may partially explain their beneficial effects on HF-hospitalisations [13,14]. Nevertheless, the diuretic effects of SGLT2i could also limit the use of SGLT2i in HF patients without T2D. While diuretics are recommended to alleviate congestion in HF patients, their side effects include plasma volume contraction, electrolyte imbalance, renal dysfunction and even kidney injury [15,16]. As renal dysfunction is strongly associated with impaired outcome in HF patients [16–18], careful titration of diuretics is of paramount importance in HF [16]. While the effects of SGLT2i on renal function are well described in patients with T2D, little is known about the renal effects of SGLT2i in the non-diabetic context. We therefore aimed to determine the effects of empagliflozin (EMPA) on renal structure and function in non-diabetic rats with LV dysfunction after MI, as this is a realistic clinical scenario if not now, likely in the near future.

METHODS

The current analysis represents a renal substudy of a recently published article [10]. We refer to this publication for detailed description of the methods and the cardiometabolic effects.

Animals

The study was performed in male non-diabetic Sprague-Dawley rats weighing 250-280 g (Envigo, The Netherlands). Animals were fed ad libitum and housed conventionally in groups of two to four rats with 12:12 h light-dark cycles. The study was approved by the local Animal Ethics committee (IvD number 16487-02-001) and we followed ARRIVE guidelines when reporting this study [19].

Myocardial infarction surgery

Rats were randomized to HF or sham surgery under isoflurane anaesthesia. HF was induced by permanent ligation of the proximal portion of the left coronary artery as previously described [20,21]. Sham operated rats underwent the same procedure but without coronary ligation.

Investigational drug

Empagliflozin (BI 10773) was mixed with standard rat chow (R/M-H V1534-70, Ssniff, Germany) in a final concentration of 200 mg/kg intended to reach an average dose of 30 mg/kg/day.

Experimental protocol

As described previously, this study represents a renal-oriented sub study of a recently published cohort of non-diabetic rats with LV dysfunction after MI [10]. Rats were treated with EMPA or chow starting either 2 days before surgery (EMPA-early) or 2 weeks after surgery (EMPA-late). After 10 weeks of treatment with EMPA or vehicle, rats were anesthetized, blood was drawn (either anti-coagulated with EDTA or sodium heparin) and the organs were rapidly excised. Kidney tissues were sectioned transversally and processed for immunohistochemistry or snap-frozen for molecular analysis. Rats with an infarct size of less than 15% were excluded from analysis.

Metabolic cage

Two weeks before termination, rats were placed in metabolic cages to monitor 24-h water and food intake and 24-h urine collection, as previously described [10]. After 24-h, blood samples were drawn from tail vein and plasma was collected. Plasma and urine were stored at -80 °C for later analysis.

Histology

The kidney was sectioned transversally, fixed by immersion in 4% buffered formaldehyde solutions for 48 h (Klinipath, The Netherlands) and subsequently embedded in paraffin according to standard procedures. Paraffin sections were stained with periodic acid Schiff (PAS), as described before [22]. Mesangial matrix expansion, focal glomerulosclerosis and

interstitial fibrosis were evaluated histomorphologically by an experienced renal pathologist (HvG).

Blood measurements

Upon sacrifice, 8 ml of blood was drawn from the abdominal aorta and urine was obtained from the bladder. Routine plasma biochemistries (urea, creatinine, phosphorus, magnesium, calcium, uric acid) were determined by Roche COBAS (Roche Diagnostics, Germany). Plasma parathyroid hormone (PTH) concentration was analyzed using Immutopics intact PTH ELISA kit 60-2500 (San Clemente, USA) according to the manufacturer's protocol. Fibroblast growth factor 23 (FGF-23) level was measured in plasma using LSBio intact FGF-23 ELISA kit according to the manufacturer's protocol (Seattle, USA). Cross-linked C-terminal telopeptide of type I collagen (CTX-1) concentration in plasma was measured using RatLapsTM EIA kit (IDS, Boldon, UK).

Urine measurements

Routine urinalysis (total protein, creatinine and phosphorus) were determined on the Roche COBAS (Roche Diagnostics, Germany). Creatinine clearance, as an estimation for glomerular filtration rate, was calculated from 24 h urinary and plasma creatinine levels, as previously described [10,22].

Quantitative real-time PCR

RNA was extracted from the kidney using TRIzol reagent (Invitrogen Corp., USA) and cDNA was synthesized by QuantiTect Reverse Transcription Kit (Qiagen, The Netherlands), as previously described [10], following manufacturer's protocol. Relative gene expressions were determined by a qRT-PCR (Bio-Rad Lab, The Netherlands). 36B4 reference gene was used to correct all measured mRNA expression. Primer sequences can be found in the supplementary *Table S1*.

Statistical analysis

Data are presented as means \pm standard errors of the mean (SEM). To compare normally distributed parameters, One-way analysis of variance (ANOVA) followed by Tukey's Post Hoc test was used. When data were not normally distributed, a non-parametric Kruskal-Wallis test followed by a Mann-Whitney U test with correction for multiple comparisons was used. To compare EMPA and vehicle treatment independent of treatment allocation, an independent t-test or a Mann-Whitney U test was used, where appropriate. Differences were considered significant at $p < 0.05$. IBM SPSS Statistics for Windows, Version 23.0 (IBM Corp, USA) was used to perform all statistical analysis.

RESULTS

A total of 140 rats were randomized to MI or sham surgery, 47 rats died during the surgical procedure and 20 rats with an infarct < 15% of the left ventricle were excluded from analysis, leaving a total of 73 rats for the current analysis. The final group sizes were 8 for the sham-vehicle (Sham-Veh) group, 19 for sham-EMPA group, 22 for MI-vehicle (MI-Veh) group, and 24 for MI-EMPA group.

TABLE 1. Cardiovascular characteristics.

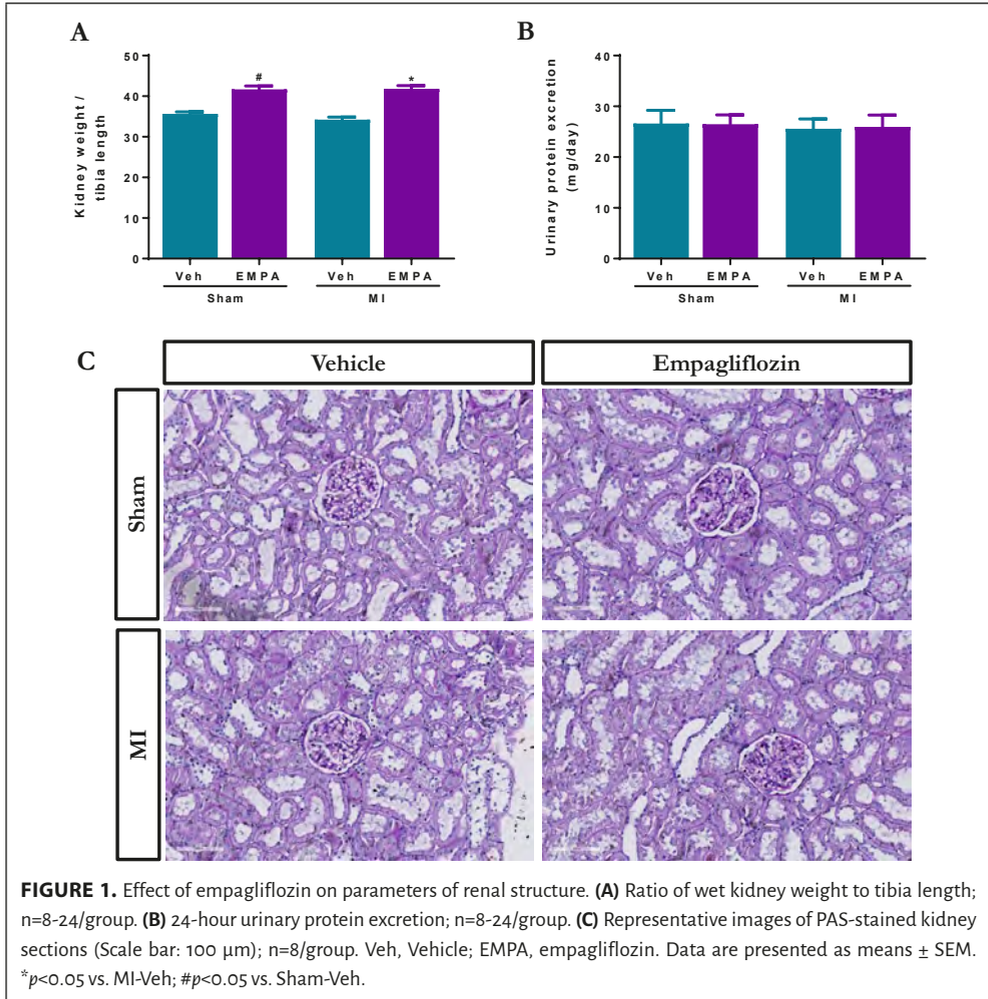
Parameters	Sham-Veh	Sham-EMPA	MI-Veh	MI-EMPA
MI size (%)	0	0	31.3 ± 1.5	33.5 ± 2.0
VWs/TL (mg/mm)	30.8 ± 0.4	30.7 ± 0.6	33.9 ± 0.5#	30.7 ± 0.6*
LVEF (%)	73 ± 3	75 ± 2	43 ± 2#	54 ± 2*
% Fibrosis	4.8 ± 1.7	3.95 ± 0.9	19.5 ± 2.5#	10.1 ± 0.6*
Cardiomyocyte CSA (µm ²)	447.9 ± 158.3	450.4 ± 103.3	814.4 ± 49.0#	629.1 ± 28.6*
SBP (mmHg)	114 ± 4	116 ± 4	118 ± 4	109 ± 9
DBP (mmHg)	81 ± 4	82 ± 4	85 ± 3	78 ± 8
ANP mRNA (fold change)	1.00 ± 0.18	1.33 ± 0.24	6.51 ± 0.84#	3.67 ± 0.61*
β-MHC/α-MHC mRNA ratio (fold change)	1.00 ± 0.18	1.04 ± 0.07	1.98 ± 0.18#	1.16 ± 0.07*

Data are presented as means ± SEM. **p* < 0.05 vs. MI-Veh; #*p* < 0.05 vs. Sham-Veh.

Veh, Vehicle; EMPA, empagliflozin; MI, myocardial infarction; VW/TL, ventricular weight/tibia length; LVEF, LV ejection fraction, CSA, cross-sectional area; SBP, systolic blood pressure; DBP, diastolic blood pressure; ANP, atrial natriuretic peptides; β-MHC, myosin heavy chain isoform beta; α-MHC, myosin heavy chain isoform alpha.

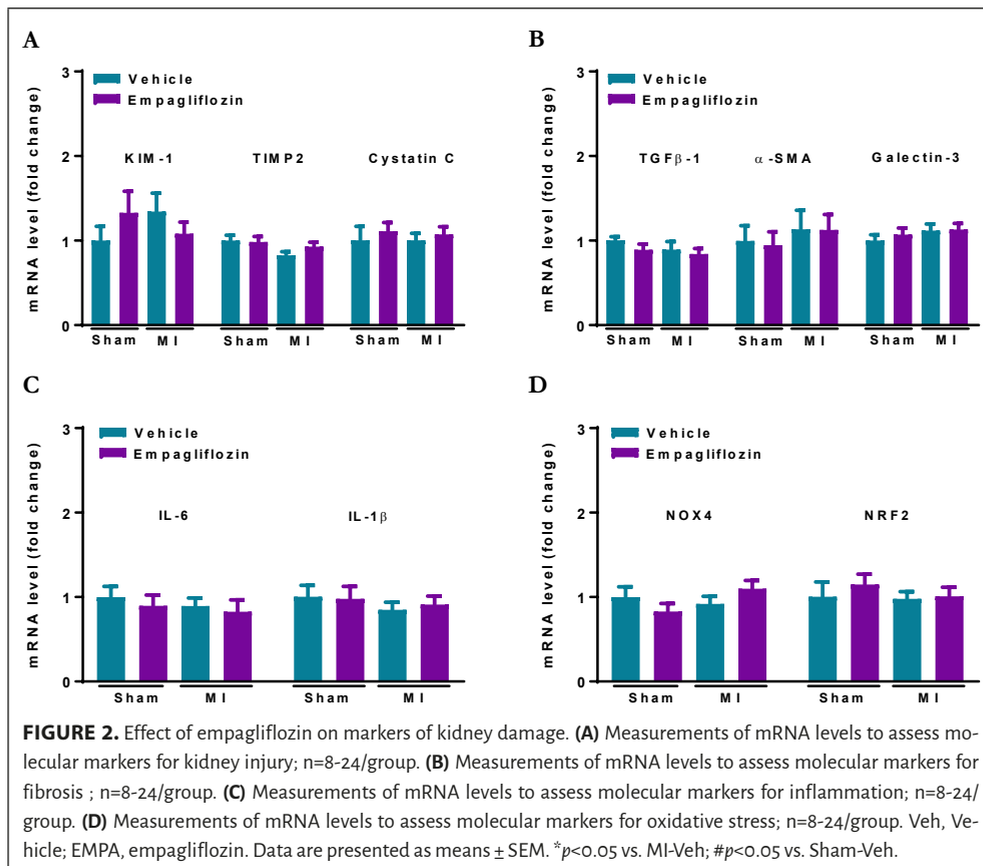
General and cardiac effects of EMPA

A detailed description of the effects of EMPA on food and fluid intake as well as cardiac structure and function in this cohort have been recently published [10], as a reference they are depicted in *Table 1*. Food intake was comparable between vehicle and EMPA treated groups (*Table 2*), resulting in an average daily intake of 30 mg/kg/day of EMPA. While the size of the MI was comparable between the vehicle and the EMPA treatment groups, EMPA resulted in a marked improved cardiac function and attenuated echocardiographic and histological indices for cardiac remodelling (*Table 1*).



Effects of EMPA on renal structure

To investigate the effect of EMPA on the renal structure, wet weight of the kidney and 24-h protein excretion were measured and kidney sections stained with PAS were analysed. The relative wet kidney weight was slightly increased in sham and MI animals treated with EMPA compared to vehicle-treated rats (*Figure 1A*). Daily protein excretion did not differ among the groups, indicating that EMPA did not cause proteinuria (*Figure 1B*). Furthermore, histomorphological changes were not observed in sham and MI animals treated with EMPA or vehicle (*Figure 1C*).



To assess molecular markers for kidney damage, mRNA expression of markers for kidney injury, fibrosis, inflammation and oxidative stress were determined. The cellular injury markers kidney injury molecule-1 (KIM-1), tissue inhibitor of metalloproteinases 2 (TIMP2) and Cystatin C, that is used as a marker to estimate GFR, as well as the kidney fibrosis markers transforming growth factor beta-1 (TGF- β 1), alpha-smooth muscle actin (α -SMA) and Galectin-3 were comparable between groups (Figure 2A-B). Moreover, the inflammatory markers interleukin 6 (IL-6) and interleukin 1 beta (IL-1 β) and the oxidative stress markers NADPH oxidase 4 (NOX4) and the nuclear factor (erythroid-derived 2)-like 2 (NRF2) were also comparable (Figure 2C-D). Taken together, our results indicate that the small increase in kidney weight observed in our cohort was not associated with evidence of structural damage to the kidney. The increase in kidney weight is probably caused by non-pathological fluid accumulation that was removed by alcohol solutions in dehydration step.

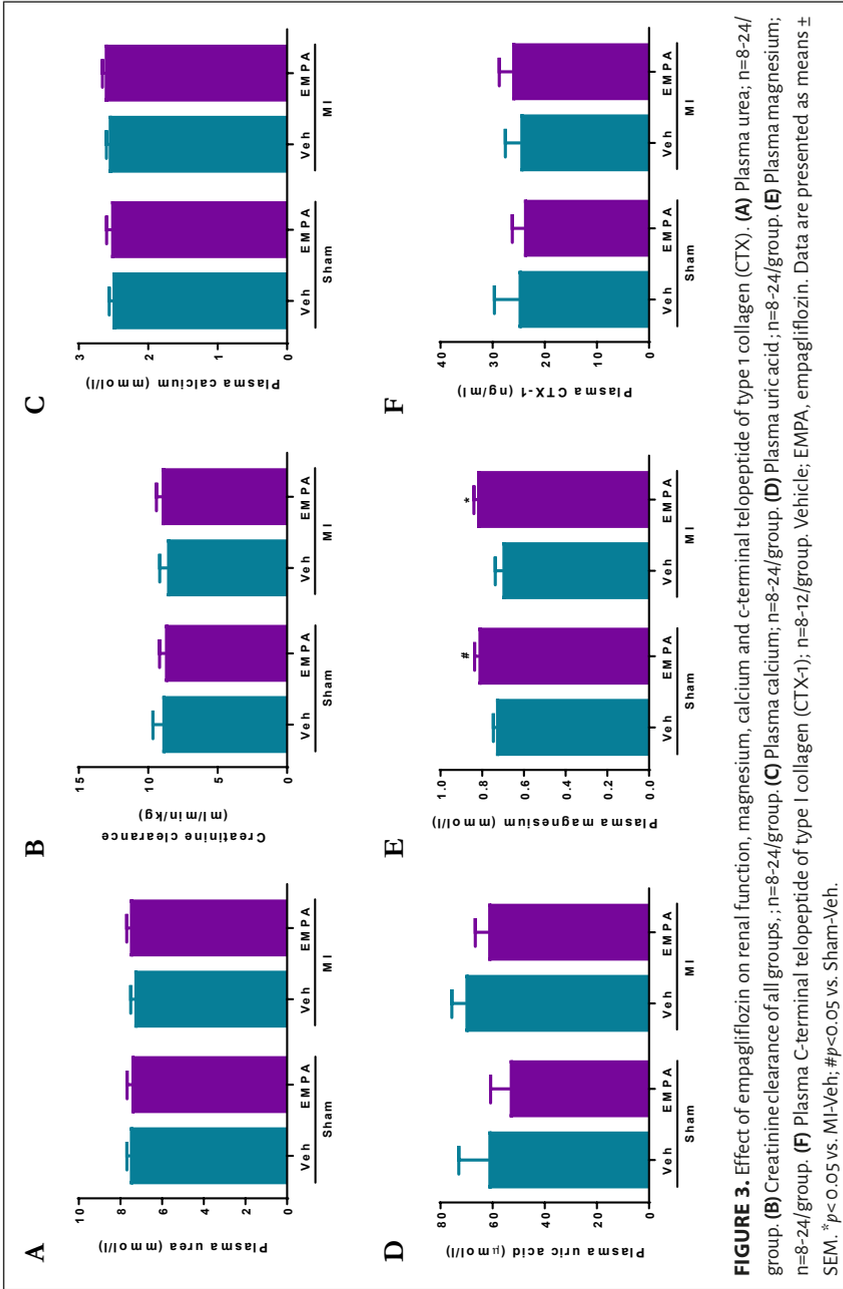
Effects of EMPA on electrolytes and renal function

EMPA resulted an increased in sodium and glucose excretion as well as a twofold increase in urine production (Table 2). Fluid intake was also twofold higher in EMPA-treated sham and MI groups and there was no evidence for plasma volume contraction as evidenced by comparable haematocrit levels (Table 2) and plasma urea concentrations (Figure 3A). Similarly, creatinine clearance was comparable between the groups (Figure 3B). No changes were observed in plasma concentrations of glucose, sodium and potassium (Table 2). Plasma calcium and uric acid (Figure 3C and 3D) were also comparable between EMPA and vehicle-treated groups. EMPA has been shown to increase magnesium levels in patients with diabetes [23] and we also detected a small increase in plasma magnesium levels in the EMPA-treated sham and MI groups (Figure 3E).

TABLE 2. General characteristics of rats with LV dysfunction and sham-operated animals.

Parameters	Sham-Veh	Sham-EMPA	MI-Veh	MI-EMPA
Water intake (ml /24h)	33.7 ± 0.9	59.8 ± 1.3#	31.9 ± 0.6	63.4 ± 1.3*
Food intake (g/24 h)	32.9 ± 0.3	33.3 ± 0.3	32.9 ± 0.3	33.6 ± 0.2
Urine Production (ml/24 h)	13.56 ± 1.16	32.79 ± 1.48#	14.35 ± 0.66	34.09 ± 1.40*
Plasma glucose (mmol/l)	13.72 ± 1.47	12.31 ± 0.90	13.72 ± 0.57	12.62 ± 0.87
Plasma sodium (mmol/l)	138.83 ± 1.05	138.88 ± 0.34	139.30 ± 0.36	140.00 ± 0.47
Plasma potassium (mmol/l)	5.08 ± 0.25	4.80 ± 0.12	4.83 ± 0.07	4.79 ± 0.08
Glucose excretion (mmol/day)	0.01 ± 0.01	8.98 ± 0.84#	0.01 ± 0.00	11.07 ± 0.92*
Sodium excretion (mmol/day)	1.93 ± 0.08	2.87 ± 0.15#	1.85 ± 0.15	3.13 ± 0.17*
Hematocrit (L/L)	45.85 ± 1.39	48.81 ± 0.90	48.42 ± 1.05	49.16 ± 0.71
Insulin/glucagon ratio	4.15 ± 0.77	1.65 ± 0.22#	4.42 ± 0.78	1.66 ± 0.10*

Data are presented as means ± SEM. * $p < 0.05$ vs. MI-Veh; # $p < 0.05$ vs. Sham-Veh.



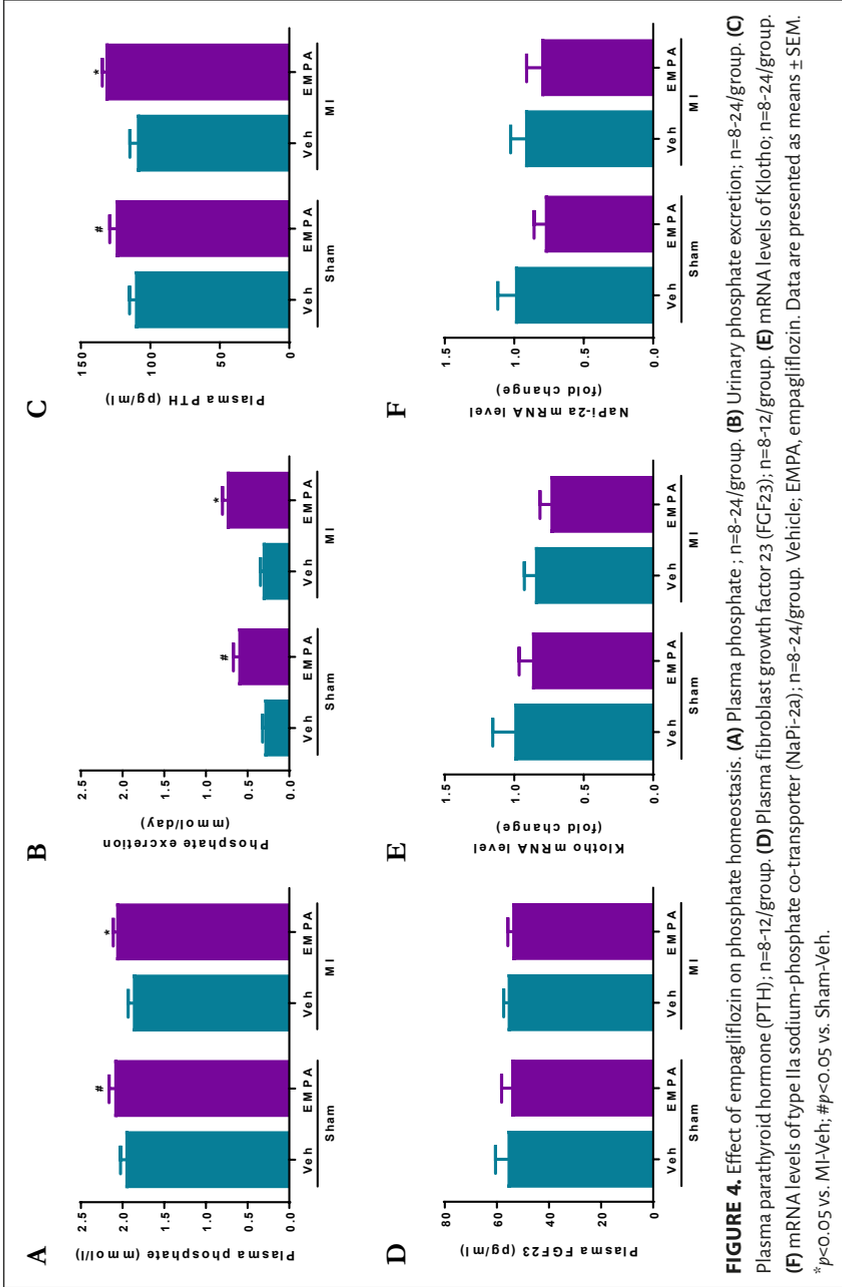


FIGURE 4. Effect of empagliflozin on phosphate homeostasis. **(A)** Plasma phosphate ; n=8-24/group. **(B)** Urinary phosphate excretion; n=8-24/group. **(C)** Plasma parathyroid hormone (PTH); n=8-12/group. **(D)** Plasma fibroblast growth factor 23 (FGF23); n=8-12/group. **(E)** mRNA levels of Klotho; n=8-24/group. **(F)** mRNA levels of type IIa sodium-phosphate co-transporter (NaPi-2a); n=8-24/group. Vehicle; EMPA, empagliflozin. Data are presented as means \pm SEM. ^{*}p<0.05 vs. MI-Veh; [#]p<0.05 vs. Sham-Veh.

Effects of EMPA on phosphate homeostasis

Phosphate homeostasis is regulated by a complex process involving intestinal, renal and bone handling [24]. Plasma phosphate plays an essential role on bone health [25]. It has been suggested that SGLT2i stimulate diabetes-related bone resorption and can increase the fracture-risk in patients with diabetes [7,26]. To assess whether this occurs in non-diabetic rats with HF, we evaluated several parameters of phosphate homeostasis namely plasma levels of PTH and FGF23 as well as the renal expression of Klotho and renal type 2a sodium-phosphate co-transporter (NaPi-2a).

EMPA increased plasma phosphate levels by 8% (*Figure 4A*) and increased the urinary excretion of phosphate (*Figure 4B*). Furthermore, EMPA increased circulating PTH levels (*Figure 4C*). Plasma levels of FGF23 were not affected by EMPA (*Figure 4D*), nor was kidney expression of klotho (*Figure 4E*) and NaPi-2a (*Figure 4F*) different in sham and MI groups. In addition, EMPA had no effect on bone mineral resorption marker CTX-1 in both sham and MI animals. These findings suggest that EMPA marginally increases plasma phosphate and PTH levels without affecting FGF23-klotho axis thus does not affect bone resorption in a non-diabetic setting.

DISCUSSION

We previously discovered that the anti-diabetic drug empagliflozin (EMPA) improves cardiac function in non-diabetic rats with HF after MI, suggesting that it may also benefit non-diabetic HF patients. In the current analysis we aimed to determine the renal effects of EMPA in non-diabetic HF and were able to demonstrate that (1) the beneficial effects of EMPA on cardiac function and remodelling were not offset by detrimental consequences on renal structure and function. (2) The marked diuretic effects of EMPA were not associated with severe plasma volume contraction, nor did EMPA alter plasma levels of glucose and (most) electrolytes. (3) Structural, functional and molecular analyses did not provide any suggestion for EMPA-induced (long term) renal damage in rats with or without HF. (4) EMPA increased circulating magnesium levels. (5) EMPA increased circulating phosphate and PTH levels without activating the FGF23-klotho axis. Accordingly, EMPA did not promote bone mineral resorption. These findings suggest that the beneficial effect of EMPA in non-diabetic HF are not offset by renal side effects or an increased fracture risk.

Renal dysfunction is common in HF [27], and worsening renal function (WRF) has been associated with mortality and HF hospitalisations [17]. Moreover, HF has been associated with elevated galectin-3 that play a central role in both heart and kidney fibrosis [28,29]. Congestion is one of the clinical hallmarks of HF and congestion can also promote renal dysfunction [30,31]. HF patients are often treated with diuretics to prevent or treat congestion, but little is known about their effect on prognosis [16,32]. In high doses, diuretics

activate the renin-angiotensin-aldosterone system (RAAS) and may thereby promote HF progression [33,34]. Furthermore, diuretics can cause plasma volume contraction, worsen renal function and can cause various electrolyte disturbances including hypokalemia, hypomagnesemia, hypocalcemia, hyponatremia, and hyperuricemia [35–37]. While mineralocorticoid receptor antagonists (MRAs) have mild diuretic effects and do improve prognosis in HF with reduced ejection fraction (HFrEF) [38], hyperkalemia and WRF are common side effects of these drugs as well [39]. Our findings showed that the potent diuretic effects of EMPA are not accompanied by renal dysfunction and electrolyte imbalance are reassuring, and SGLT2i may offer a safe opportunity to alleviate congestion.

An emerging hypothesis is that SGLT2i may directly inhibit the Na⁺/H⁺ exchanger isoform 1 (NHE1) in the myocardium and NHE isoform 3 (NHE3) in the kidney [40–43]. It is thought that the suppression of NHE1 by SGLT2i reduces cardiac injury, hypertrophy and fibrosis as well as reduce in the risk of cardiovascular death and hospitalization for heart failure [40,41], and AMPK might also mediate indirect SGLT2i–NHE interactions in the heart [44]. Whereas the suppression of NHE3, which is increased in HF, leads to the inhibition of proximal tubular reabsorption of sodium and thereby decrease intravascular volume and cardiac wall stress [40,41,45]. Thus, inhibition of NHE1 and NHE3 may be beneficial to prevent and/or treat heart failure.

The considerable increase in diuresis observed with SGLT2i suggests that these drugs could be classified as diuretics. A study in patients with T2D demonstrated that SGLT2i dapagliflozin may have a diuretic-like effect beyond the glycemic control [46]. The mechanisms of SGLT2i mediated diuresis are, however, very different from classical diuretics prescribed to HF patients. SGLT2i function as osmotic diuretics and reduce interstitial fluid volume without causing major changes to plasma volume and sodium levels, whereas the reverse is true for loop diuretics [47,48]. Another difference between SGLT2i and classical diuretics is that SGLT2i promote uric acid excretion and may therefore prevent gout [49]. Nevertheless, the cardiovascular benefits of SGLT2i in high risk exceed the benefits of classical diuretics, making it highly unlikely that these benefits are merely explained by enhanced diuresis [50]. Future studies should better map the interplay and interactions of joint use of thiazides and loop diuretics and SGLT2i.

The US Food and Drug Administration (FDA) has raised the concern for acute kidney injury (AKI) associated with SGLT2i canagliflozin and dapagliflozin use [51]. However, a longitudinal analysis from two large health system registries, Mount Sinai chronic kidney disease registry and the Geisinger Health System cohort, demonstrated that SGLT2i use is not associated with an increased risk of AKI in patients with T2D [52]. Furthermore, a study by Cahn *et al.* confirmed that SGLT2i do not increase risk for AKI compared with DPP4 inhibitors among patients with T2D [53]. DAPA-HF trial revealed that the beneficial effects of SGLT2i dapagliflozin was not related to any adverse events on renal function [12]. Our

findings support and extend this observation as we found no evidence of renal damage, by either functional (plasma urea, creatinine clearance) or structural (proteinuria, renal histopathological analysis) renal damage after EMPA treatment. Similarly, we have performed a comprehensive screening of the relevant markers for kidney injury, fibrosis, inflammation and oxidative stress, and no significant differences were observed among the various groups, providing molecular proof that there are no damaging processes occurring in kidneys of rats treated with SGLT2i.

3 SGLT2i increases circulating Mg^{2+} levels in individuals with T2D and therefore it has been proposed that the increase in circulating Mg^{2+} could at least partially underlie the cardio-beneficial effects of SGLT2i [23,54,55]. Mg^{2+} homeostasis is maintained by the interaction of intestine, bone and kidneys [56,57]. Several hormones have been suggested to affect the Mg^{2+} balance. Studies in rats showed that Mg^{2+} reabsorptive capacity in the distal segment of the kidney is increased after glucagon and PTH infusion [58,59]. Insulin causes a shift of magnesium from the extracellular to the intracellular space, resulting in a decrease in plasma magnesium and an increase in erythrocyte magnesium content in both healthy non-diabetics and diabetic individuals [60,61]. In our study, we observed that EMPA reduced insulin/glucagon ratio (Table 2) and increased PTH levels in plasma (Figure 4C). Building upon these observations, our data suggest that increases in plasma magnesium levels are most probably explained by the effects of EMPA on insulin, glucagon, and PTH.

Our observation demonstrated that EMPA caused a slight but significant increase in serum phosphate and PTH concentrations, and is in line with similar observations in patients with or without diabetes [62–64]. The increase in PTH levels has raised the concern that SGLT2i may promote bone mineral resorption and increase fracture risk [62]. The putative mechanisms are sought in the fact that SGLT2i increase tubular sodium concentrations and increase renal phosphate reabsorption through enhanced activity of sodium-phosphate cotransporter (NaPi) [62]. The resultant increase in plasma phosphate levels will then trigger a reactive increase in PTH secretion by the parathyroid gland and the secretion of FGF23 by osteocytes. PTH and FGF-23 promote bone resorption and reduce bone mineralisation, respectively, and are both implicated in osteoporosis [24,65]. Furthermore, both factors suppress the activity of NaPi-2a and NaPi-2c in the kidney which promotes urinary phosphate excretion [66]. A study by Weir *et al.* demonstrated that in patients with T2D receiving SGLT2i canagliflozin versus placebo, minimal increases in serum phosphate and magnesium were within normal limits and has no clinical relevance [67]. While we did observe an increase in plasma phosphate and PTH levels, the observed changes were within physiological range [68]. Furthermore, calcium levels were unaltered and we did not find evidence for enhanced bone resorption. Plasma FGF23 and renal expression of NaPi-2a were also unaltered, indicating that EMPA did not result in the activation of the FGF-23-Klotho axis. In addition, no effect on plasma CTX-1 was noted in both vehicle and empa-treated animals.

These results are in contrast with observations by Thraikill *et al.* who studied a model for diabetic bone disease [7]. Our findings are also in line with recent evidence that demonstrated that SGLT2i do not result in a net increase in fracture risk [69,70]. Taken together, our data suggest that EMPA does not affect the bone mineral resorption in a non-diabetic context.

STUDY LIMITATIONS

Despite strengths related to the direct measures of renal structure, function and other physiological parameters, our study does have limitations. In addition to the inherent limitations associated with our experimental heart failure model and differences among species, there are a number of specific limitations to our study. First, in accordance with the previous studies, our post-MI HF model did not induce renal dysfunction in rats [71,72]. Our findings may therefore not reflect the effects of EMPA in patients with both cardiac and renal dysfunction. Second, we compared EMPA with placebo in this study and therefore cannot extrapolate whether there will be a meaningful interaction with other HF drugs, including diuretics. Third, our study was not focussed on bone health and we did not perform direct measurements of bone mineral density. Nevertheless, our study does provide compelling evidence suggesting that the cardiac benefits of EMPA are not associated with detrimental renal consequences or impediments in bone health. Another potential limitation of our study was the exclusion of 20 rats with small infarctions because their cardiac function is close to normal. As a sensitivity analysis we performed all analysis both with and without rats with small infarctions, and no differences in the outcomes were observed (data not shown). Further research is clearly required to ascertain whether the improvement of cardiac function was associated with diuretic properties of EMPA treatment, or, improvement of cardiac function caused the diuresis.

CONCLUSION

SGLT2 inhibition with EMPA exerts profound diuretic effects without compromising renal structure and function or causing significant electrolyte imbalance. The slight increase in circulating phosphate and PTH after EMPA treatment was not associated with evidence for increased bone mineral resorption suggesting that EMPA does not affect bone health.

ACKNOWLEDGEMENTS

We acknowledge Boehringer Ingelheim for supplying empagliflozin and control chow, and thank Dr. Eric Mayoux for support and expert advice throughout the project. We thank Janny Takens for expert technical assistance and advice.

FUNDING

Dr. Yurista is supported by a grant from the Indonesia Endowment Fund for Education (LPDP No. 20150722083422). Dr. de Boer is supported by the Netherlands Heart Foundation (CVON DOSIS, grant 2014-40, CVON SHE-PREDICTS-HF, grant 2017-21, and CVON RED-CVD, grant 2017-11); and the Innovational Research Incentives Scheme program of the Netherlands Organization for Scientific Research (NWO VIDI, grant 917.13.350). Dr. Westenbrink is supported by The Netherlands Organisation for Scientific Research (NWO VENI, grant 016.176.147).

COMPLIANCE AND ETHICAL STANDARD

Ethics approval and consent to participate

The study was approved by the University of Groningen Ethical Committee on Animal Experimentation (IvD number: 16487-02-001). The investigation conforms to the Guide for the Care and Use of Laboratory Animals published by the US National Institutes of Health and in accordance with national regulations.

Competing interest

SRY, HHWS, HvG, JLH, HJLH, LMM, SUO and BDW do not report conflicts of interest relative to this report. The UMCG, which employs all authors, has received research grants and/or fees from AstraZeneca, Abbott, Bristol-Myers Squibb, Novartis, Roche, Trevena, and ThermoFisher GmbH. Dr. de Boer received personal fees from MandalMed Inc, Novartis, and Servier.

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SUPPLEMENTARY MATERIAL

Supplementary Table

TABLE S1. List of primers used in this study.

Genes	Forward primer (5'-3')	Reverse primer (5'-3')
KIM-1	AGAGAGAGCAGGACACAGGCTT	ACCCGTGGTAGTCCCAAACA
TIMP2	TGGACGTTGGAGGAAAGAAG	TGTCCCAGGGCACAATAAAG
Cystatin C	AGCGAGTACAACAAGGGCAGCAAC	TTGTCAGGGTGTGTGTGCCTTCC
TGF- β 1	AAGAAGTCACCCGCGTGCTA	TGTGTGATGTCTTTGGTTTTGTCA
α -SMA	CATCATGCGTCTGGACTTGG	TCACGCTCAGCAGTAGTCAC
Galectin-3	CCCGCTCAATGAGAACAAC	ACCGCAACCTTGAAGTGGTC
IL-6	AGCGATGATGCACTGTCAGA	TAGCACACTAGGTTTGCCGA
IL-1 β	TGTGATGAAAGACGGCACACC	GGGAACTGTGCAGACTCAAC
NOX4	ATGTTGGGCCTAGGATTGTGT	AAGTTCAGGGCGTTCACCAA
NRF2	ATCCAGACAGACACCAGTGGATC	GGCAGTGAAGACTGAACTTTCA
36B4	GTTGCCTCAGTGCCTCACTC	GCAGCCGCAAATGCAGATGG
Klotho	CGTGAATGAGGCTCTGAAAGC	GAGCGGTCACTAAGCGAATACG
NaPi-IIa	CCAACATCGGGACCTCTGTC	CCACCACGGATGTTGAAGGA
ANP	ATGGGCTCCTTCTCCATCAC	TCTACCGGCATCTTCTCCTC
α -MHC	GACAACTCCTCCGCTTTGG	AAGATCACCCGGGACTTCTC
β -MHC	TCTGGAGGCCTTTGGCAATG	GATGCCAACTTTCCTGTTGC

