Study Rationale and Design of a Study of Empagliflozin’s Effects in Patients With Type 2 Diabetes Mellitus and Coronary ARtery Disease: The EMPA-CARD Randomized Controlled Trial

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Study protocol

Keywords: SGLT2 inhibitor, Empagliflozin, Randomized controlled trial, Coronary artery disease, Inflammation, Type 2 diabetes mellitus
Abstract

Background: Recent trials have revealed that sodium-glucose co-transporter 2 inhibitors (SGLT2-i) are effective against hyperglycemia and also reduce micro- and macro-vascular complications in patients with type 2 diabetes mellitus (T2DM). Most of the beneficial cardiovascular effects have been investigated in patients with heart failure and coronary artery disease (CAD). Yet, no human study has been conducted to investigate the mechanisms underlying these clinically beneficial effects in patients with CAD. Accordingly, the EMPA-CARD trial was designed to focus on the molecular effects of empagliflozin in patients with T2DM and CAD.

Methods: In this multicenter, triple-blind randomized controlled trial, patients with documented known T2DM and CAD will be recruited. They will be randomized on a 1:1 ratio and assigned into two groups of empagliflozin 10 mg/daily and placebo. The primary endpoint is the effect of empagliflozin on changes of plasma interleukin 6 (IL-6) after 26 weeks of treatment. The secondary endpoints will consist of changes in other inflammatory biomarkers (Interleukin 1-beta and high-sensitive C-reactive protein), markers of oxidative stress, platelet function, and glycemic status.

Discussion: The EMPA-CARD trial mainly tests the hypothesis that SGLT2 inhibition by empagliflozin may improve inflammatory status measured as reduction in inflammatory biomarkers in patients with T2DM and CAD. The results will provide information about the underlying mechanisms of SGLT2 inhibition that mediate the beneficial effects of this medication on clinical outcomes.


Introduction

Diabetes mellitus (DM) is a chronic disease that is characterized by high blood glucose levels due to the inability to produce adequate amounts of insulin and/or resistance to the hormone's actions (1). Type 2 diabetes mellitus (T2DM) is a complex metabolic condition and, as the most common type of diabetes, it affects some 37% of the population in western and middle-eastern societies leading to long-term complications (2). The major complications of this disease fall into two general categories of microvascular and macrovascular involvements, which can consequently lead to cardiovascular, neurological, ophthalmological, and renal damage (3). Cardiovascular complications are the leading cause of death in the affected population. Atherosclerosis develops more rapidly in these patients and increases the incidence of a heart attack two to three times (4). The management methods require multifaceted strategies, including regular physical activity, proper diet, and medications to control hyperglycemia, hypertension, inflammation, and dyslipidemia (5, 6). Among various medications available to treat diabetes, one of the newest ones is sodium-glucose co-transport 2 inhibitors (SGLT2-i) including empagliflozin, dapagliflozin, and canagliflozin. Empagliflozin is a selective inhibitor of the SGLT2 co-transporter expressed in the proximal tubule of nephrons. Inhibition of SGLT2 reduces
hyperglycemia by decreasing glucose reabsorption in the kidneys leading to increased urinary glucose excretion, reductions in blood glucose and HbA1c (7). The EMPA-REG OUTCOME trial and its sub-studies revealed the additional beneficial effects of this medication versus standard treatment protocols in patients with T2DM and heart failure. These effects include a reduction in Major Adverse Cardiovascular Events (MACE), blood pressure, urinary albumin exertion, and improvements in lipid profile and vascular resistance (8, 9). Some hypotheses have been advanced to explain the beneficial cardiovascular effects of this agent including hemodynamic changes due to decreased plasma volume following glycosuria-induced diuresis, changes in heart fuel consumption, improvements in overall metabolism by a reduction of inflammation, and direct effects on the cardiac tissue; however, the underlying mechanisms remain obscure (10, 11). Many factors contribute to the physiopathology of cardiovascular complications in patients with T2DM including an increase in pro-inflammatory factors, imbalance in oxidative stress and a hyper-coagulation state with abnormal platelet function (12). In this regard, animal models and in-vitro studies have been performed to evaluate the effects of empagliflozin on inflammatory state and platelet function. The results showed a reduction in oxidative stress, pro-inflammatory phenotype, and glucotoxicity, but platelet function remained unchanged (13, 14). To date, no human model randomized trial has been performed to evaluate the inflammation and platelet dysfunction hypothesis in T2DM patients with coronary artery disease (CAD). Hence, the main objective of this study is to test the hypothesis that empagliflozin improves the inflammatory state in T2DM patients with documented known CAD.

**Methods And Experimental Design**

2.1. **Study design and Intervention**

EMPA-CARD is a multicenter triple-blind randomized placebo-controlled trial designed to evaluate the effect of treatment with empagliflozin (10 mg once daily) versus placebo for 26 weeks, in addition to standard care, in patients with T2DM and coronary artery disease.

2.1.1. **Study objectives:**

The primary objective of the EMPA-CARD trial is to evaluate the effect of empagliflozin on the inflammatory state in patients with T2DM and coronary artery disease after 26 weeks of treatment (Figure 1). Other objectives which will be investigated as the trial's sub-studies are shown in Table 1.

2.1.2. **Ethics considerations:**

The EMPA-CARD trial is conducting in accordance with the principles of the declaration of Helsinki and all subsequent revisions. The study was approved by the ethics committee of Zanjan university of Medical Sciences. All patients provided with written informed consent prior to the recruitment. Moreover, the study protocol was prospectively registered on the Iranian Registry of Clinical Trials (www.IRCT.ir, Identifier: IRCT20190412043247N2).
2.1.3. **Trial population and eligibility assessment**

Our goal is to randomize 100 participants from registries of 4 clinical centers (i.e., Mousavi hospital (coronary angiography registry), Vali-e-Asr hospital, and two cardiology clinics of Zanjan University of Medical Sciences) in Zanjan, Iran. The pooled population of the clinics is 8000 patients, whose records were extracted from registries at the pre-recruitment phase. Patients aged between 45 and 75 years old with T2DM and known coronary artery disease (established by coronary angiography) who are on background standard anti-ischemic and anti-diabetic therapy with an HbA$_{1c}$ between 6.5 and 9 were eligible for inclusion. The type and dose of background anti-diabetic medications should be constant for at least 3 months prior to recruitment. Patients’ left ventricular ejection fraction (LVEF) must be higher than 40%. Moreover, they must not be receiving anti-oxidant, anticoagulant, or antiplatelet medications or supplements (except for aspirin at 80 mg/d) for at least 3 months prior to recruitment. Detailed information for inclusion and exclusion criteria is provided in Table 2.

2.1.4. **Randomization and follow-up**

The randomization method employed in this study is stratified randomization. During the screening process, eligible patients are stratified by gender (male and female), age (45-54, 55-64, and 65-75 years old), and HbA$_{1c}$ (6.5-7.9% and 8-9%) and assigned into one of the two arms of the study (A or B). The randomization sequence is created using Winpepi software (version 11.6). All participants will be followed up by phone calls every 4 weeks for assuring the proper medication usage, and for premature discontinuation or development of any adverse events. At the end of weeks 2, 12, and 26, patients are instructed to attend clinic visits for complete checkup and physical examination. Patients are also advised to return all used and unused medications at the end of week 26. The study’s follow-up and procedures are summarized in Table 3.

2.1.5. **Blinding**

Empagliflozin and placebo tablets with the same color, shape, and package with the different code combination numbers are used for groups A and B. After enrollment, a code is assigned to each patient and will be used until the end of this study. Patients, researchers and healthcare providers who gather information and assess the outcomes and analyze the data of the study are blinded to the assigned treatment (empagliflozin or placebo) such that all patients will be evaluated under the unique assigned code and the treatment groups of A or B. In case of a need for un-blinding, the research council of Zanjan University of Medical Sciences will be notified by the chief investigator to discuss the terms of un-blinding. The un-blinded treatment list is held by the Zanjan University Medical Documentation Council which is not involved in the study. If the un-blinding is approved by the research council, the Documentation Council will be notified and ultimately the patient will be sent to an external clinical visit for further assessment and follow-up; meanwhile the chief investigator and research council will remain blinded.

2.2. **Outcomes**
To assess the outcomes of the study and sub-studies, patients undergo blood sampling, electrocardiography (ECG), and 2D trans-thoracic echocardiography (Table 3).

2.2.1. Primary outcome

The primary outcome of this study is the changes in plasma interleukin 6 levels after 26 weeks of the treatment between the two study arms.

2.2.2. Secondary outcomes for the primary objective

The secondary outcomes for testing the hypothesis of the study's primary objective are categorized into 4 divisions including 1) Changes in additional plasma inflammatory biomarker levels (high-sensitive C-reactive protein (hs-CRP) and plasma interleukin 1-beta (IL-1b)); 2) Changes in oxidative stress status (lymphocytic reactive oxygen species levels (ROS), plasma levels of malondialdehyde (MDA), carbonyl level, total antioxidant capacity (T-AOC), reduced glutathione level (GSHr), catalase enzyme (CAT) activity, and superoxide dismutase enzyme (SOD) activity); 3) Changes in platelet function (CD62-P expression on the platelet surface); and 4) Changes in glycemic status (Fasting Blood Glucose (FBS), HbA1C, and Homeostatic Model Assessment for Insulin Resistance (HOMA-IR)).

2.2.3. Additional secondary outcomes

The additional secondary outcomes will be considered for the EMPA-CARD sub-studies including 1) Changes in cardiac biomarkers (Serum high-sensitive Troponin I (hs-Troponin-I), Brain Natriuretic Peptide (BNP) and N-terminal pro-Brain Natriuretic Peptide (NT-proBNP) levels); 2) Changes in Echocardiographic parameters (left ventricular systolic and diastolic function and right ventricular function); 3) Changes in hematopoietic status and renal function (blood hematocrit, hemoglobin, and serum erythropoietin, and urine microalbuminuria levels); 4) Changes in lipid profile (serum total cholesterol, Low-Density Lipoprotein (LDL) cholesterol, High-Density Lipoprotein (HDL) cholesterol, and triglyceride levels); and 5) Changes in electrocardiographic parameters (PR interval duration, QRS complex duration, QT interval duration, ST-segment deviation, and T wave alternans).

All the above secondary outcomes will be assessed as the changes from baseline to week 26 of treatment in each study arm. Measurements of the biomarkers will be performed in the biotechnology laboratory of the Zanjan University of Medical Sciences and the Mousavi Hospital’s laboratory. The detailed information about the mentioned outcomes is presented in Table 4.

2.3. Biobank

For the measurements and analysis of some outcomes, a biobank is established in which the separated plasma/serum and urine samples are allocated in separate micro-tubes and stored in an ultra-low temperature freezer with the patient’s dedicated code. Based on the regulations of the local data protection agency, after the measurements, all the remaining samples will be anonymized and the
biobank will be discontinued. All patients are informed in the consent form about the establishment of the biobank.

2.4. Study monitoring

Monitoring the study and data source verification is the responsibility of the Research Council of Zanjan University of Medical Sciences, Iran. All the forms used in this study, including the consent form, trial procedure manual, trial violation form, adverse events form, and outcomes’ related checklists, were evaluated and validated by a team of clinical trial monitors and the above Council. Recruitment visits are conducted under direct observation of the team. Human subject de-identification is an issue which is considered as local/regional and national requirements among all presentations and publications. To fulfill that requirement, appropriate measurements such as encoding or deletion are enforced. All of the study investigators are informed to follow the trial procedure manual. Also, any deviation from the protocol by any investigator will be recorded into the trial violation form designed by the Ethics Committee of the Zanjan University of Medical Sciences. All of the study forms are in the Persian language.

2.5. Adverse event recording

Any adverse events, related or unrelated, whether reported by the patient or discovered by the investigators during the study follow-up (until week 28) will be recorded in the adverse events form. At the same time, the Research Council of the Zanjan University of Medical Sciences will be notified at the time of the recording. The severity and the importance of the adverse events will be evaluated by a physician. The standard adverse event form is presented in the supplementary material.

2.6. Potential confounding factors

According to the nature of the variables in primary and secondary outcomes, two potential major study confounding factors were predicted: 1) patients’ nutritional status and 2) patients’ physical activity status. To evaluate the impact of these possible confounding factors, the food frequency questionnaire (FFQ) and the physical activity questionnaire (FAQ) are used. The forms are filled at the day of recruitment. Also, patients are advised not to change their routine diet or physical activity during the study.

2.7. Statistical considerations

2.7.1. Sample size calculations

The sample size was estimated based on the changes in plasma IL6 level as a primary outcome of treatment. According to a previous study, the SGLT2-i dapagliflozin reduced the mean plasma IL-6 level from 6.6 to 5.8 (pg/mL) with the standard deviation (SD) of 8.9 (15). The minimal important difference (MID) of the IL-6 level was calculated 4.45 (pg/mL) using 0.5×SD (the distributional-based method) (16). With a study power of 80%, an alpha level of 0.05%, and MID of 4.45, the required sample size was
calculated to be 41 patients in each arm. Considering a 20% dropout during the study, the sample size was rounded up to 50 participants in each arm. The sample size was calculated using the G-power software (version 3.1.9.2).

2.7.2. Statistical analysis plan

We will use the means ± standard deviations (SD) and frequencies and percentages n (%) to interpret the results. The noninferiority margin and 95% confidence interval (CI) will be estimated for primary outcomes. The noninferiority will be accepted if the upper 95% CI for the relative risk (RR) lies below the margin of noninferiority. An intention-to-treat analysis will be used to examine the effect of missing data. Superiority analysis will be performed at $\alpha=0.025$ level. The interaction effects will be examined using graphical and statistical tests. We will consider the probability of 0.001 for the interim analysis and 0.025 for the main analysis. We will perform Bayesian analysis of the covariance test to examine the effects of intervention on IL6. All statistical tests will be carried out in the in Rv.4 environment. The primary data of the FFQ will be evaluated by using the Nutritionist software version 4 (N4).

2.7. Current status

At present, the mentioned sites are actively recruiting patients. The pre-recruitment phase (evaluation and extraction of 8000 patients’ medical records from hospitals and clinics registries) was started on November 11, 2019. The patients’ recruitment phase (the first patient who was consented and randomized) was started on June 29, 2020 and until November 9, 2020, 91 patients were randomized. The end date of expected recruitment may be extended due to the COVID-19 pandemic but it is expected that the recruitment phase will ends by the end of December 2020.

Discussion

The EMPA-CARD randomized placebo-controlled clinical trial seeks to investigate the mechanisms underlying the cardiovascular effects of empagliflozin in patients with T2DM who have proven CAD at the same time.

Most of the recent published clinical trials have focused on the effects of empagliflozin in patients with heart failure in which the superiority of empagliflozin has been reported. The recent meta-analysis with the pooled population of 8,474 patients (DAPA-HF and EMPEROR-Reduced population) showed that treatment with an SGLT2-i was associated with a significant reduction in cardiovascular death (pooled hazard ratio of 0.86) and re-hospitalization due to heart failure complications as well as improvements in renal function (17). However, limited data are available regarding the beneficial effects of empagliflozin in patients with established CAD. The EMPA-HEART Cardiolink-6 trial showed that 6 months of treatment with empagliflozin significantly reduced left ventricular mass index (LVMi) in patients with T2DM and CAD (18). In this study we are exploring molecular mechanisms and cellular changes that potentially mediate these beneficial effects. Plasma levels of IL-6, a pro-inflammatory cytokine which also has anti-inflammatory properties promotes and regulates inflammatory processes in patients with T2DM, and
changes in its level is considered as the primary outcome of this study. The rationale of choosing this biomarker as a leading factor for investigation among other candidates is that IL-6 and tumor necrosis factor-alpha (TNF-α) appear to have the highest impact on the promotion of CAD in patients with diabetes mellitus (19). Moreover, the other factors measured in this study are either directly or indirectly affected by IL-6. It has been shown that IL-6 reduces oxidative stress and apoptosis of pancreatic β-cell by promoting autophagy, an adaptive response to cellular stress. Nevertheless, it seems that the inflammation and fatty acid oxidation, which is also promoted by IL-6, have a greater impact on increasing the oxidative stress in patients with T2DM. This could enhance the rate and magnitude of coronary atherosclerosis and subsequent CAD (20, 21). Although it is uncertain whether empagliflozin has beneficial effects in patients with CAD, the EMPA-CARD trial is the first study designed to provide a better understanding of the effects of this medication on inflammation in these patients.

Declarations

Acknowledgment:

We would like to express our thanks to Dr. Hossein Chiti, Dr. Ahmad Jalilvand, Mr. Ehsan Janzamin, Ms. Zeinab Darvishian, Dr. Atieh Asgari, and Dr. Homa Taheri for their fruitful consultations and for providing us with resources. We thank Dr. Abidi Pharmaceutical Company® for providing us with the medication and placebo and for sponsoring the study. Last but not least, we thank all the patients and collaborators for their patience and cooperation through the study.

Ethics approval and consent to participate

The EMPA-CARD trial is conducting in accordance with the principles of the declaration of Helsinki and all subsequent revisions. The study was approved by the ethics committee of Zanjan university of Medical Sciences. All patients provided with written informed consent prior to the recruitment. Moreover, the study protocol was prospectively registered on the Iranian Registry of Clinical Trials (www.IRCT.ir, Identifier: IRCT20190412043247N2).

Consent for publication:

Not applicable.

Funding:

The EMPA-CARD study is funded by Dr. Abidi Pharmaceutical Company® and Zanjan University of Medical Sciences (Grant Code: 1602001000).

Role of Dr. Adibi Pharmaceutical Company:

The company supplied the medication and placebo for the study. The company had no role in the development of the protocol, process of the study, or preparation of this manuscript.
**Conflict of Interest:**

The authors contributing to the study have nothing relevant to disclose. Only the study authors are responsible for conducting and reporting the final contents of the trial.

**Data and materials availability:**

The data/information supporting this study is available from the corresponding author upon request.

**Authors’ contribution:**

Sepehr Gohari: Concepts, Design and study’s main idea, Literature search, Main investigator and outcome assessor, Manuscript preparation, patients follow up.

Tara Reshadmanesh: Design, outcome assessor, literature search, manuscript preparation.

Hadi Khodabandehloo: Design, literature search, Revision, Concepts.

Mojtaba Fathi: Design, Concepts, Revision.

Hassan Ahangar: Design, Concepts, Revision, Study coordination, Definition of intellectual content.


Faramarz Ismail-Beigi: Design, Revision.

Samin Ghanbari: Outcome assessor and patient follow up.

Mohsen Dadashi: Outcome assessor and patient follow up.

Muhammad Javad Muhammadi: Outcome assessor and patient follow up.

Sheida Gohari: statistical analysis, Manuscript editing.

Saeid Ghaffari: Design.

**References**


Tables

Table 1. Trial objectives and measurement methods (measured at time = 0 and after 26 weeks of treatment)
<table>
<thead>
<tr>
<th>Objective</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary</td>
<td>1. To investigate the impact of empagliflozin on plasma IL-6 level in patients with T2DM patients and coronary artery disease (established by coronary angiography).</td>
</tr>
</tbody>
</table>
| Secondary | 1. Inflammatory biomarkers  
· Changes in serum hs-CRP levels.  
· Changes in plasma IL-1b levels.  
2. Oxidative stress status  
· Changes in lymphocytic reactive oxygen species.  
· Changes in plasma levels of malondialdehyde.  
· Changes in plasma carbonyl levels.  
· Changes in plasma antioxidant capacity.  
· Changes in plasma reduced glutathione levels.  
· Changes in catalase enzyme activity.  
· Changes in plasma superoxide dismutase enzyme activity.  
3. Platelet function  
· Changes in CD62-P expression on the platelet surface.  
4. Glycemic Status  
· Changes in fasting blood glucose levels.  
· Changes in HbA$_1c$ levels.  
· Changes in HOMA-IR.  |
| Additional secondary (Considered for substudies) | 1. Changes in cardiac biomarkers  
· Changes in serum hs-Troponin I levels.  
· Changes in serum BNP levels.  
· Changes in serum NT-proBNP levels.  
2. Echocardiographic Parameters  
· Changes in left ventricular systolic function.  
· Changes in left ventricular diastolic function.  
· Changes in right ventricular function.  
3. Hematopoietic status and renal function  
· Changes in serum erythropoietin levels.  |
· Changes in blood hematocrit levels.
· Changes in hemoglobin levels.
· Changes in Urine micro albuminuria levels.

4. Changes in lipid profile
· Changes in serum total cholesterol levels.
· Changes in serum LDL cholesterol levels.
· Changes in serum HDL cholesterol levels.
· Changes in serum triglyceride levels.

5. Electrocardiographic parameters
· Changes in PR interval duration.
· Changes in QRS complex duration.
· Changes in QT interval duration.
· Changes in ST segment deviation.
· Changes in T wave alternans.

IL-6: Interleukin 6, T2DM: Type 2 Diabetes Mellitus, hs-CRP: High sensitive C-Reactive Protein, IL-1b: Interleukin 1-beta, HbA$_1c$: Glycated Hemoglobin A$_1c$, HOMA-IR: Homeostatic Model Assessment for Insulin Resistance, hs-Troponin I: High Sensitive Troponin I, BNP: Brain Natriuretic Peptide, NT-proBNP: N-terminal pro-Brain Natriuretic Peptide, LDL cholesterol: Low Density Lipoprotein cholesterol, HDL cholesterol: High Density Lipoprotein cholesterol.

Table 2. List of inclusion and exclusion criteria for randomization
| **Inclusion Criteria** | 1. Age between 40-75 years  
2. HbA1c between 6.5 to 9  
3. Documented known diabetes mellitus type 2 under continuous fixed-dose anti-diabetic treatment for at least 3 months prior to the randomization  
4. Documented known coronary artery disease established by coronary angiography under continuous fixed-dose of anti-ischemic treatment for at least 3 months prior to the randomization  
5. BMI less than 40  
6. Fixed diet and physical activity  
7. eGFR greater than 45 ml/min/1.73m²  
8. Informed consent given in written form  
9. Resting heart rate between 60 to 100 b/min |
| **Exclusion Criteria** | 1. Pregnancy  
2. Heart Failure (NYHA class 3-4)  
3. Left ventricular ejection fraction <40%  
4. Use of anti-coagulant or anti-platelet drugs (except aspirin 80 mg/daily) for at least 3 months prior to the blood sampling  
5. Consumption of alcohol, continuous anti-inflammatory drugs (except aspirin 80mg/daily) or anti-oxidant supplement  
6. Use of pioglitazone  
7. History of allergic reaction to SGLT2-i medications  
8. History of SGLT2-i medication usage  
9. Gastrointestinal malabsorption disease  
10. History of CABG, ACS, TIA, CVA or PCI during past 3 month  
11. History or presence of malignancy  
12. Severe HTN  
   · Resting systolic blood pressure ≥ 180 mmHg  
   and/or  
   · Resting diastolic blood pressure ≥ 120 mmHg  
13. Anemia (Hb < 10 g/dL)  
14. Platelet count < 100000/µl |
15. History of infection during 1 one month prior to blood sampling
16. History of heart or lung transplant
17. Major psychiatric disorders
18. History of DKA
19. Elevated liver enzymes >3 times upper normal limit
20. Use of drugs which prolong QT interval
21. Presence of arrhythmia
22. Electrolyte imbalance
23. Patients with Pace Maker


Table 3. SPIRIT flow diagram
<table>
<thead>
<tr>
<th>TIMEPOINT</th>
<th>Pre-recruitment</th>
<th>Baseline (Visit 1)</th>
<th>Visit 2</th>
<th>Visit 1</th>
<th>Visit 3</th>
<th>Follow up (Telephone visit 4)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>-</td>
<td>0</td>
<td>2±1 weeks after visit 0</td>
<td>12±2 weeks after visit 0</td>
<td>26±2 weeks after visit 0</td>
<td>28 weeks after visit 0</td>
</tr>
</tbody>
</table>

**ENROLMENT:**

- Patients’ medical records extraction: X
- Eligibility screen: X, X
- Informed consent: X
- Randomization: X
- Allocation: X

**INTERVENTIONS:**

- Empagliflozin 10mg (Intervention group)
- Placebo (Control group)

**ASSESSMENTS:**

- Demographic Data: X
- Vital signs: X, X, X, X, X
- Medical and drug history: X
- Concurrent medical conditions: X
- Concomitant medication: X, X, X, X, X
- Food frequency questionnaire: X
- Physical activity questionnaire: X
<table>
<thead>
<tr>
<th>Physical examination</th>
<th>X</th>
<th>X</th>
<th>X</th>
<th>X</th>
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</thead>
<tbody>
<tr>
<td>Drug container collection</td>
<td></td>
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<tr>
<td>Adverse events</td>
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<tr>
<td><strong>OUTCOME and Safety ASSESSMENTS</strong></td>
<td></td>
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<tr>
<td>Laboratory measurements (Outcomes)</td>
<td>X</td>
<td></td>
<td>X</td>
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<tr>
<td>2D-TTE</td>
<td>X</td>
<td></td>
<td>X</td>
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<tr>
<td>Electrocardiogram</td>
<td>X</td>
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<td>X</td>
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<tr>
<td>Laboratory measurements (Safety *)</td>
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</tbody>
</table>

* The safety measurements were done at the visit 0, visit 3 and whenever an adverse event occurs in a patient. The measurements included liver and renal function related laboratory measurements (Aspartate-Aminotransferase [AST], Alanine-Aminotransferase [ALT], Alkaline phosphatase [ALP], Creatinine [Cr] and Blood Urea Nitrogen [BUN]), Serum electrolyte measurements (Sodium [Na], Potassium [K]), Coagulative measurements (Prothrombin time [PT], Partial Thromboplastin Time [PTT]), Complete Blood Count (CBC), Urine Analysis (U/A), and Urine Culture (U/C) if the U/A become active and/or urogenital symptoms develop.

2D-TTE: 2-dimentional Trans-thoracic Echocardiography

Table 4. The detailed descriptions of selected outcomes
<table>
<thead>
<tr>
<th>Outcome</th>
<th>Method of Measurement</th>
<th>Mechanism</th>
<th>Time of Measurement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Changes in plasma IL-6 and 1-beta levels</td>
<td>Plasma IL-6 and IL-1b Elisa Kits</td>
<td>Key cytokines in development and maintenance of inflammation</td>
<td>Week 0 and 26</td>
</tr>
<tr>
<td>Changes in oxidative stress (Lymphocytic ROS, plasma MDA, Carbonyl, T-AOC, GSHr, CAT activity and SOD enzyme activity levels)</td>
<td>Flow Cytometry (DCFDA as dye), Spectrophotometry, Colorimetric and FRAP assays</td>
<td>ROS, MDA and Carbonyl Increase in inflammation and causes cell damage. GSHr, AOC, CAT activity and SOD activity increase in inflammation and reduce oxidative stress</td>
<td>Week 0 and 26</td>
</tr>
<tr>
<td>Changes in serum hs-CRP level</td>
<td>Turbidimetric assay</td>
<td>Circulating concentrations rise in response to inflammation</td>
<td>Week 0 and 26</td>
</tr>
<tr>
<td>Changes in platelet function</td>
<td>Flow Cytometry (MFI of CD62-P expression on platelet surface)</td>
<td>Platelet activity increases in DM and inflammation causing platelet dysfunction</td>
<td>Week 0 and 26</td>
</tr>
<tr>
<td>Changes in HbA₁c, FBS and Basal Insulin level</td>
<td>Enzymatic assay</td>
<td>Indicate the glycemic status which is reduce with anti-glycemic drugs</td>
<td>Week 0 and 26</td>
</tr>
<tr>
<td>Changes in HOMA-IR</td>
<td>fasting insulin (mU/mL) x fasting glucose (mg/dL)/405</td>
<td>A method used to quantify insulin resistance which increases in DM and insulin resistance conditions</td>
<td>Week 0 and 26</td>
</tr>
<tr>
<td>Changes in serum hs-Troponin I, BNP and NT-proBNP levels</td>
<td>Chemiluminescence assay</td>
<td>Specific cardiac biomarkers increase in cardiomyocyte damage and tension</td>
<td>Week 0 and 26</td>
</tr>
<tr>
<td>Changes in blood Hb, Hct and serum Erythropoietin</td>
<td>CBC and Serum Erythropoietin Elisa kit</td>
<td>Reflects the hematopoietic status</td>
<td>Week 0 and 26</td>
</tr>
<tr>
<td>Changes in urine microalbuminuria and albumin to creatinine ratio</td>
<td>Photometry and Enzymatic assays</td>
<td>Reflects the renal and glomerular function. May increase in DM as a microvascular adverse event.</td>
<td>Week 0 and 26</td>
</tr>
<tr>
<td>Changes in lipid profile (serum TG, Cholesterol, HDL,LDL cholesterol)</td>
<td>Enzymatic assay</td>
<td>Lipid disorders increase in DM which raises the cardiovascular complications that may decrease with empagliflozin treatment.</td>
<td>Week 0 and 26</td>
</tr>
<tr>
<td>Changes in echocardiography parameters</td>
<td>2D trans-Thoracic Echocardiography</td>
<td>Reflects left ventricular systolic and diastolic and right ventricular function</td>
<td>Week 0 and 26</td>
</tr>
</tbody>
</table>
Changes in electrocardiographic parameters | Electrocardiography | Reflects the electrophysiological activity of the heart | Week 0 and 26
---|---|---|---


**Figures**
Figure 1

Pathways contributing to the development of micro- and macro-vascular complications in patients with T2DM and insulin resistance (only coronary artery disease, retinopathy and nephropathy are illustrated).

*Indicates where the trial’s primary objective (potential molecular effects of empagliflozin on inflammatory state) will be assessed. IL-6: Interleukin 6. IL-1: Interleukin 1. ROS: Reactive Oxygen Species,
Figure 1

Pathways contributing to the development of micro- and macro-vascular complications in patients with T2DM and insulin resistance (only coronary artery disease, retinopathy and nephropathy are illustrated). *Indicates where the trial’s primary objective (potential molecular effects of empagliflozin on