

INPLASY PROTOCOL

To cite: Yang et al. The Efficacy and Safety of LCZ696 in CKD Patients: A Systematic Review and Meta-Analysis. Inplasy protocol 202240045. doi: 10.37766/inplasy2022.4.0045

Received: 07 April 2022

Published: 08 April 2022

Corresponding author:
Xu Jinsheng

xjs5766@126.com

Author Affiliation:
Departments of Nephrology,
The Fourth Hospital of Hebei
Medical University,
Shijiazhuang, PR China.

Support: 20577701D.

Review Stage at time of this submission: Completed but not published.

Conflicts of interest:
None declared.

The Efficacy and Safety of LCZ696 in CKD Patients: A Systematic Review and Meta-Analysis

Yang, XY¹; Jin, JJ²; Cheng, MJ³; Li, YJ⁴; Li, YZ⁵; Zhao, YF⁶; Bai, YL⁷; Xu, JS⁸.

Review question / Objective: To assess the efficacy and safety of LCZ696 compared to ACEI/ARBs in patients with CKD(eGFR<60ml/min/1.73 m²), we carried out the meta-analysis.

Condition being studied: CKD is a global health problem that costs health-care systems a bunch of prices, and it has a high prevalence of between 11% and 13% around the world. Many large clinical trials have shown the advantages of LCZ696 for patients with heart failure, but the effect of it on people who have CKD has not been tested in a great number of studies.

Information sources: Two reviewers independently conducted comprehensive searches until January 15(th), 2022 in the widely used medical databases, including PubMed, the Cochrane Library and Embase.

INPLASY registration number: This protocol was registered with the International Platform of Registered Systematic Review and Meta-Analysis Protocols (INPLASY) on 08 April 2022 and was last updated on 08 April 2022 (registration number INPLASY202240045).

INTRODUCTION

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patients with heart failure, but the effect of it on people who have CKD has not been tested in a great number of studies.

METHODS

Participant or population: Randomized controlled trials that included patients with CKD (eGFR < 60 ml/min/1.73 m²).

Intervention: Sacubitril valsartan (LCZ696).

Comparator: ACEI/ARBs.

Study designs to be included: Randomized controlled trials.

Eligibility criteria: (I) The type of trial was RCT; (II) The subjects contained CKD (eGFR < 60 ml/min/1.73 m²) patients; (III) The experimental group was LCZ696 and the control group was ACEI/ARB; (IV) All studies had data about renal function and renal outcomes.

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Main outcome(s): The change of Scr; The change of eGFR; The incidence of ESRD; The incidence of cardiovascular events; The incidence of hyperkalemia; The incidence of hypotension.

Quality assessment / Risk of bias analysis: We used the Cochrane Collaboration's tool for assessing risk of bias to evaluate the bias of the trials presented.

Strategy of data synthesis: All statistical analysis in this paper were performed on Review Manager 5.3. Because the outcomes of the study were dichotomous data, the OR with 95% CI was computed. The heterogeneity of studies was assessed with the Cochran Q test, I² statistic and forest maps. Heterogeneity was low when I² was less than 25%, moderate when I² was between 25% and 50% and high when I² was greater than 50%. If there was

heterogeneity in the outcomes, the reasons of heterogeneity were analyzed via meta regression, subgroup analysis, and sensitivity analysis.

Subgroup analysis: We carried out subgroup analysis according to intervention, reduction percentage in eGFR or age.

Sensitivity analysis: We did not investigate publication bias due to only 4 eligible studies.

Country(ies) involved: China.

Keywords: LCZ696, CKD, efficacy, safety.

Contributions of each author:

Author 1 - Yang Xinyue.

Email: 18712922179@163.com

Author 2 - Jin Jingjing.

Author 3 - Cheng Meijuan.

Author 4 - Li Yajing.

Author 5 - Li Yuzhe.

Author 6 - Zhao Yunfeng.

Author 7 - Bai Yaling.

Author 8 - Xu Jinsheng.

Email: xjs5766@126.com