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Title: Cardiopulmonary functional capacity: Another piece of the puzzle of SGLT2 inhibition in heart failure?

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The glifozins have become an integral part of the treatment of heart failure [1]. Following the results of the DAPA-HF trial [2] which were soon followed by EMPEROR-Reduced trial [3], they firmly established themselves as a core therapy for heart failure with reduced ejection fraction, even upending decades of sequential treatment initiation algorithms in guidelines. Ever since the glifozins were shown to be beneficial in patients with type 2 diabetes, there has been immense interest in their mechanism of action and this has only intensified since the discovery that they improved outcomes in patients with heart failure with multiple studies speculating on potential mechanisms of action that underlie the benefits in heart failure [4]. It is possible that there is no single mechanism as the benefit of these drugs also extends to patients with chronic kidney disease as well as patients with type 2 diabetes. It is difficult to imagine that in every disease area there is only one single mechanism of action. Indeed, secondary analysis of many of these trials suggest multiple benefits for patients. One of the major findings from the trials of glifozins and heart failure with reduced ejection fraction is a very consistent benefit on symptoms and health-related quality of life [5,6]. This is as important if not more important often to patients than reductions in morbidity and mortality. Understanding why patients feel better may also help us understand the mechanisms that lead to a reduction in morbidity and mortality that are observed in the randomised trials. There have been other trials that have used health-related quality of life as an outcome [7, 8,9]. These have been very consistent with the randomised trials that examined morbidity and mortality outcomes. A consistent finding of all of these trials is an improvement in health-related quality of life in all domains. The Kansas City Cardiomyopathy Questionnaire (KCCQ) is commonly used to assess health-related quality of life. This questionnaire is a patient reported metric that quantifies symptoms, physical function, quality of life, and social function over the prior 2 weeks. In DAPA-HF dapagliflozin caused more patients to improve their KCCQ score and less to suffer a deterioration in their score [6]. One remaining question has been how does dapagliflozin do this? In this issue of the Journal Palau and colleagues present the results of a small, randomised trial looking at the effect of dapagliflozin on functional capacity as measured by cardiopulmonary exercise testing [10].
In their study patients were randomised either to dapagliflozin or placebo (with 45 patients in each group) and in this double-blinded trial patients who had heart failure with a reduced ejection fraction underwent maximal cardiopulmonary exercise testing at baseline, 1 and 3 months. The primary outcome of the trial was a change in peak VO₂ at 1 and 3 months. Several secondary endpoints were also examined, these included changes at 1 and 3 months in the 6-minute walk test distance and health related quality of life measured by the Minnesota Living with Heart Failure Questionnaire. Echocardiographic parameters (left ventricular ejection fraction, E/e’ and left ventricular and atrial volumes) were also measured to look for signs of cardiac remodelling. The cohort that they enrolled was very representative of patients with heart failure with reduced ejection fraction. However, there was one characteristic that merits special attention. This was an extremely well treated population. Of the randomized patients, 89% of were receiving sacubitril/valsartan, 91% were receiving a beta blocker and 74% of patients were receiving a mineralocorticoid receptor antagonist. This is much better than many clinical trials and registries of patients with heart failure with reduced ejection fraction. Despite this excellent baseline therapy, the investigators reported that peak VO₂ increased in patients who were randomised to dapagliflozin at 1-month and at 3 months. There was no difference in 6-minute walk test distance or with the health-related quality of life questionnaire. There was no statistically significant difference in echocardiographic parameters either. At first sight this may seem to be at odds with the results of trials like DEFINE-HF [7] or the sub studies of the large randomised trials of SGLT2 inhibitors which have shown a benefit on health-related quality of life [5,6]. However, this sample was relatively small and the time period over which the change was examined was also fairly short in comparison to other trials or analyses. Although neither the 6-minute walk distance or scores from the Minnesota Living with Heart Failure Questionnaire were statistically significantly different between the two groups the change and each of these parameters was in favour of dapagliflozin. As might be expected...
in a trial of such short duration there was no difference in structural measures of cardiac function such as ejection fraction or volumes or indeed filling pressures is measured by E/e’.

Previous studies of remodelling in patients with heart failure have required longer periods of follow up to show any effect on cardiac structure [11]. How important is the improvement in peak VO₂ that we have seen in this trial? An analysis of the HF-ACTION trial which collected information and peak VO₂ in 1620 patients with heart failure with reduced ejection fraction, reported that a 6% increase in peak VO₂, when adjusted for other significant predictors was associated with a 5% lower risk of the composite of all-cause mortality or all-cause hospitalisation [12]. There was 8% lower risk of cardiovascular mortality or heart failure hospitalisation associated with this 6% increase of peak VO₂ at 3 months. Therefore peak VO₂ is clearly related to outcomes in this population. In the study by Palau et al the baseline peak VO₂ was 13.4 mL/kg/min and at 3-months this was 13.7 mL/kg/min. The significant difference between the dapagliflozin and placebo groups arose because the peak VO² fell from 12.9 to 12.7 mL/kg/min on average. This led to a greater difference between the two groups at 3 months compared to baseline and dapagliflozin appeared to preserve and improve peak VO₂ in a short period of time. This preservation phenomenon also follows the pattern that we have seen with other biological parameter such as renal function and health related quality of life where less people deteriorated with dapagliflozin. Returning to the question of how do glifozins work in heart failure, the findings of Palau et al take us further forward in our understanding of the many potential mechanisms that may be at play. While the findings do not complete the puzzle, they add an important piece to the picture.
References


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