The Diuretic Effect of Sacubitril/Valsartan Might Be Clinically Relevant

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Recently, patients with heart failure have been prescribed a novel and innovative drug. Sacubitril/valsartan is a new drug modality that brings a 16% reduction in total mortality, a 20% reduction in cardiovascular mortality and 21% reduction in hospital admissions due to heart failure. The benefit is undoubtedly clinically relevant and the clinical trial which have shown such benefit have achieved an unprecedented statistical significance.1

The mechanism of action of sacubitril/valsartan combines the well-known vasodilatory effect of valsartan associated with the neutral endopeptidase (NEP) inhibition effect of sacubitril, which will ultimately result in increased serum levels of natriuretic peptides, increased action of endogenous natriuretic peptides in target tissues by prolonging its tissue half-life, and consequently increased vasodilatory, anti-proliferative and natriuretic effects.1

Although the current approach of replacing enalapril with sacubitril/valsartan might sound as a switch of vasodilators in patients with heart failure, the addition of natriuretic effect provided by sacubitril may in fact be the driving force of the clinical benefits. In favor of this concept we can make a few comments:

a. Hypotension, more frequently seen in sacubitril/valsartan than in the enalapril group, could possibly be associated with hypovolemia caused by the natriuretic effect of sacubitril;

b. Patients who received valsartan (160 mg twice daily) in the Val-HEFT trial2 did not show the same benefit on mortality or on hypotensive adverse events as those demonstrated in the PARADIGM-HF trial (sacubitril/valsartan 97/103 mg twice daily).

c. A post hoc analysis of data from the PARADIGM-HF study revealed that the increase in the mean dose of furosemide was smaller in the sacubitril/valsartan group compared with the enalapril group, and that the median dose of furosemide increased in the enalapril group, but not in the sacubitril/valsartan group.3

It is well known from observational studies and meta-analyses that increased doses of diuretics have been linked to worse prognosis in patients with heart failure. Despite inherent biases associated with observational studies, it is biologically plausible that diuretics are potentially harmful due to their hyperreninemic, vasoconstrictive and hypokalemic effects. One of the few clinical trials conducted on diuretics in patients with heart failure, the DOSE trial, have shown greater kidney toxicity associated with higher doses of furosemide. Diuretic dose reduction associated with sacubitril/valsartan therapy might be a desired secondary effect of this compound in patients with heart failure.4,5

In that sense, studies on diuretic withdrawal are mostly needed. The RBIC (REde Brasileira de Insuficiência Cardíaca – Heart Failure Brazilian Network) trial is under way and is intended to be the largest clinical trial ever conducted designed to assess the effects of diuretic withdrawal in ambulatory patients with heart failure.4 A subgroup of patients on sacubitril-valsartan will be compared with those on angiotensin-converting enzyme inhibitors/angiotensin receptor blocker for tolerance of diuretic withdrawal.

While no other data are available, it is reasonable to recommend closer attention to patients’ volume status and exercise a low threshold to decrease or even discontinue diuretics in heart failure patients on sacubitril/valsartan.

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