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







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EDITORIAL COMMENT

Reasons for Lack of Improvement in Treatment With Evidence-Based Therapies in Heart Failure*

Biykem Bozkurt, MD, PhD

Despite increased awareness with recommendations in practice guidelines and performance measures, we have not made much progress in the proportion of patients treated with guideline-directed medical therapy (GDMT) or patients achieving target doses of heart failure (HF) medications in the last 2 decades (1-6) (Figure 1A). Specifically, rates for angiotensin-converting enzyme (ACE) inhibitor, angiotensin receptor blocker (ARB), beta-blocker (BB), or mineralocorticoid receptor antagonist (MRA) use have not increased in the last 18 years according to population-based HF registries (Figure 1A) (1-6). There are several reasons hypothesized for this lack of improvement, including provider aversion and inertia, patient intolerance and side effects, lack of payer or insurance coverage, and data and cost limitations. For ease of the reader, in this editorial, these are categorized into 3 ability domains as patient-, provider-, and payer-related issues (4) (Figure 1B).

Improvement in the proportions of patients treated with GDMT (except for a very modest increase in the uptake of treatment with ARB with neprilysin inhibitor [ARNi]—a relatively new medication for HF)—or maximum doses of GDMT among 2,588 treatment-eligible patients over 12 months (3) (Figures 1A and 1C). In more than two-thirds of the patients, there were no medication changes despite suboptimal doses. There were higher rates of discontinuation or dose decreases (12%) for ACE inhibitors/ARBs than an initiation or dose increase (7%) at 12 months, resulting in lower rates of ACE inhibitor/ARB use at 12 months compared with baseline (Figure 1C). Dose initiation or increase rates were modest for other medications and were comparable to discontinuations, resulting in an overall net flat effect, except for a modest increase in ARNi use (7%) at 12 months. Patients were treated with any of the following: ARNi, BB, or MRA; and <1% of patients were previously treated with target doses of ARNi, BB, and MRA at baseline and 12 months (3).

The CHAMP-HF (Change the Management of Patients With Heart Failure) registry enrolled nearly 5,000 patients with chronic systolic HF, who did not require heart transplantation, left ventricular assist

Tam metin

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
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