Iatrogenic Decompensated Heart Failure

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Published online: 21 February 2020
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Abstract
Purpose of Review To provide an overview of the potential iatrogenic causes of acute decompensated heart failure (AHF) and an evidence-based management strategy to address this.

Recent Findings As the heart failure (HF) population continues to age and become burdened with greater comorbidities and polypharmacy, patients become more susceptible to the iatrogenic precipitants of HF. The following clinical scenarios are familiar to clinicians, but the sequelae to AHF are often unanticipated: HF medications withdrawn during an intercurrent illness and not restarted, cardiotoxic therapy prescribed for cancer without timely and regular monitoring of left ventricular function, excessive intravenous fluids administered for sepsis or postoperatively, a blood transfusion volume not adjusted for body weight, iatrogenic anaemia that goes unnoticed or an inappropriate type of pacemaker implanted in a patient with underlying left ventricular systolic dysfunction.

Summary Iatrogenic decompensated HF is a phenomenon that is infrequently documented in the literature but increasingly confronted by clinicians of all specialties. It is associated with a high mortality and morbidity rate. By having greater awareness of these triggers, iatrogenic AHF should be one that is prevented rather than managed when it occurs.

Keywords Iatrogenic · Decompensated heart failure · Pharmacotherapy · Fluid management · High-output heart failure · Pacemaker

Introduction

One of the major challenges of managing acute decompensated heart failure (AHF) is identifying and addressing the precipitating factors, which are often multifactorial. The European Society of Cardiology (ESC) guidelines for heart failure (HF) emphasize on recognizing intrinsic cardiovascular triggers (such as acute coronary syndrome, arrhythmias and hypertension) and extrinsic insults such as infection and respiratory and renal dysfunction [1]. However, what is less described but seen increasingly more commonly in daily practice are precipitants related to inadvertent harm from acts of commission or omission by physicians, or directly by a form of medical therapy, which we collectively refer to as iatrogenic decompensated HF (IAHF). Little is known of its prevalence, and this type of data is not collected in the annual UK National Heart Failure Audit which analysed over 58,000 AHF hospitalisations [2]. An observational study in 1996 found that iatrogenesis accounted for 7% of HF admissions, and was associated with higher mortality and longer hospital stays compared with non-iatrogenic causes [3] though, this difference in mortality rate could have very likely been confounded by other comorbidities, additional medications or the presence of infection. With an aging population burdened with increasing comorbidities and polypharmacy combined with newer medications and technology, these seemingly innocuous therapies may unknowingly decompensate the delicate neurohormonal balance in these patients; hence, the current prevalence of IAHF is likely to be higher. An overview of these precipitants and its management implications is discussed under four major categories: pharmacotherapy, fluid management, high-output HF and pacemaker devices summarized in Table 1.
Pharmacotherapy

Withholding and Delaying HF Medications

It is well-established that in patients with HF with reduced ejection fraction (HFrEF), renin-angiotensin-aldosterone system inhibitors (RAASis), e.g. ACE inhibitors (ACEi) and angiotensin-receptor blockers (ARBs), beta-blockers, mineralocorticoid-receptor antagonists (MRAs), the more recent combination sacubitril/valsartan, and sodium-glucose transport protein 2 inhibitors (regardless of diabetes status) markedly improve survival and reduce HF hospitalizations against placebo [4, 5]. The delay in starting, inappropriate discontinuation or failure to restart these prognostically vital medications can put these patients at risk of acute decompensation of stable chronic heart failure and sometimes cause haemodynamic deterioration.

RAASi is often misunderstood as a nephrotoxic drug. Introduction of the UK electronic acute kidney injury alert (AKI e-alert) system has exacerbated this anxiety, and a reflex cessation of RAASi amongst hospital and community practitioners occurs when a small serum urea or creatinine (sCr) rise is seen [6]. RAASi induces renal efferent arterial vasodilatation, and a resultant fall in intra-glomerular pressure is expected, reflected by an initial sCr rise and a decline in glomerular filtration rate (GFR) in the first 2 weeks. Moreover, GFR is dependent on blood pressure (BP). In HF patients who frequently have chronic kidney disease (CKD) and hypertension, the BP range for intra-renal autoregulation becomes narrower, so a small drop in BP can lead to a modest fall in GFR through RAASi-mediated vasodilation rather than intrinsic kidney injury [7]. New national guidance recommends withholding RAASi only if sCr increases by > 30% or potassium level ≥ 6.0 [8••]. It reminds physicians that an AKI e-alert does not automatically mean withdrawal of RAASi but instead, stimulate inquiry into other potential causes that may include AHF itself. AHF can lead to a disproportionate rise in urea via anti-diuretic hormone release, raised renal interstitial pressures in systemic venous congestion or functional ureteric obstruction from tense ascites [9, 10]. A study of > 16,000 patients found that discontinuation of RAASi in HFrEF patients is associated with higher mortality and re-admission rates at 30 and 90 days and 1 year [11]. In fact, RAASi offers the greatest mortality reduction in HFrEF patients with the worse renal function at baseline [12]. Therefore even with moderate-severe renal dysfunction, there is consensus to continue RAASi if benefits outweigh risks [8••].

The rationale follows for beta-blockers: the benefits are higher with increasing severity of HFrEF but with greatest risk when stopped. This is usually due to anxiety about its negative inotropic effects or in context of hypotension [13]. Unless there is severe hypotension, ESC recommends continuing beta-blockers at a reduced dose [1]. Besides, beta-blockers cannot be the reason for AHF in patients maintained on a long-term steady dose [14]. Discontinuation can trigger rebound tachycardia, ventricular arrhythmia and aggravate angina [15, 16]. This is associated with higher mortality and rehospitalization rates [13]. Conversely, continuing beta-blockers in AHF correlates with lower mortality and admission rates [13, 17].

Even planned withdrawal of HF medications can lead to AHF in apparently asymptomatic chronic HF. In a prospective randomized trial of patients with stable dilated cardiomyopathy, MRA, beta-blockers and RAASi were sequentially weaned off. However, even with a controlled withdrawal of these medications, it resulted in a substantial fall in the left ventricular ejection fraction in 40% of patients, which was detected within 8 weeks [18]. This emphasizes the point that HF medications should only be temporarily withheld when absolutely necessary.

Cardiotoxicity and Adverse Drug Reactions

Direct myocardial toxicity and fluctuations in afterload and preload are common mechanisms behind drugs implicated in AHF. An established phenomenon is cancer therapy–related cardiac dysfunction, whereby left ventricular ejection fraction (LVEF) falls by > 10 to < 53%, manifesting acutely or delayed, even as long as 20 years after completion of therapy [19]. Anthracyclines (e.g. doxorubicin) can cause dose-dependent cardiotoxicity that is generally irreversible, whereas biologically targeted drugs (e.g. trastuzumab for HER-2 positive breast cancer) can induce cardiotoxicity which is usually reversible upon prompt drug cessation or initiation of HF medications [20, 21]. According to ESC guidelines, a baseline echocardiogram is recommended before initiation of such therapy irrespective of the clinical history. Low-risk patients with a normal baseline echocardiogram and no clinical risk factors for HF should undergo echocardiography every 4 cycles of trastuzumab or after an equivalent dose of 200 mg/m².

Table 1 Summary of potential iatrogenic causes for AHF

| Pharmacotherapy | Withholding HF medications |
|-----------------|---------------------------|
|                  | Delays in initiating HF medications |
|                  | Cardiotoxicity |
|                  | Adverse drug reactions |
| Fluid management | Excessive intravenous fluid |
|                  | Under-diuresis |
|                  | Transfusion-associated circulatory overload |
|                  | Dehydration |
| High-output HF   | Arterio-venous fistula |
|                  | Anaemia |
| Pacemaker-related HF | Pacing-induced LV systolic dysfunction |
|                  | Pacemaker wire–related tricuspid regurgitation |
|                  | Pacemaker syndrome |
anthracycline, while surveillance should be more frequent in high-risk patients. After completion of treatment, follow-up scans should be arranged at 1 and 5 years, and sometimes continued longer depending on the patient’s initial risk stratification. If clinical HF develops during or following treatment, it should be treated according to current ESC guidelines for HF with the initiation of cardio-protective medications such as ACEi and beta-blockers, usually prescribed together. Interruption or continuation of cancer treatment will depend on the risks versus benefits, determined by factors such as the degree of left ventricular systolic dysfunction (LVSD), cancer prognosis and responsiveness to the cancer therapy [1, 19, 22].

An estimated 50% of HF patients aged ≥ 65 suffer from at least 5 comorbidities. This results in greater exposure to other drugs such as steroids for chronic obstructive pulmonary disease, non-steroidal anti-inflammatory drugs (NSAIDs) for osteoarthritis and pioglitazone for type 2 diabetes mellitus (T2DM) [23]. These medications can increase fluid retention, afterload or preload, and potentially destabilize the neurohormonal and haemodynamics balance in HF. Steroids have a propensity to cause hypertension and fluid retention associated with a dose-dependent risk of precipitating or worsening HF [24]. A similar effect is seen in NSAIDs mediated by the inhibition of cyclo-oxygenase (COX) to trigger sodium and water retention. One study demonstrated a ten-fold increased risk of AHF over 72 months of NSAID use [25]. Indeed COX-2 inhibitors can also lead to a dose-dependent increased risk of HF hospitalization, and is discouraged in patients with ischaemic heart disease by ESC guidance [26, 27]. Fluid retention is also seen in alpha1-blockers (e.g. alfuzosin and doxazocin) and the oral hypoglycaemic agent, thiazolidinedione (e.g. pioglitazone). The former is associated with a two-fold risk of HF compared with other anti-hypertensives, while the latter is found to exacerbate existing HF and increase risk of de novo HF [28, 29]. Thus, these familiar drugs which are often prescribed in HF patients for other morbidities, e.g. osteoarthritis and T2DM, should be substituted for safer alternative.

Another comorbidity that frequently coexists is atrial fibrillation. For rate control, non-dihydropyridine calcium channel blockers should be avoided in HFrEF due to their negative inotropic activity and association with AHF [30]. For rhythm control, class I antiarrhythmics, e.g. flecainide, can significantly depress LV function and was found to precipitate AHF in patients with baseline LVSD in the CAST trial [31]. Another black-box warning is dronedarone (class III antiarrhythm). Yet, fluid management will only be as accurate as the clinician. Jugular venous pressure can be subjective and chest radiographic signs of HF can be non-specific [39, 40]. To complicate matters, elderly patients can present with subclinical congestion [41]. Hence, we frequently see the counterintuitive dilemma of treating with diuretics and intravenous fluids together. One study found 11% inpatients with AHF (without sepsis or bleeding) were given intravenous fluids (median 1000 ml) alongside diuretics, presumably to counterbalance any harm from over-diuresis. This increases LV filling pressures and prolongs the AHF episode [42]. Another fear to address is the initial rise in sCr with

**Fluid Management**

**Excessive Intravenous Fluid and Under-Diuresis**

The first Surviving Sepsis Campaign initiative recommended rapid intravenous fluid administration of up to 30 ml/kg to restore BP from third-space fluid losses [34]. However, this strategy is unsafe in patients with HF and sepsis. Achieving a mean arterial pressure (MAP) of 65 mmHg may not be realistic in some HFrEF patients who have lower than average baselines, and pushing more fluids to aim for 30 ml/kg can precipitate pulmonary oedema. This is supported by a retrospective study of patients with septic shock where intravenous fluid of 30 ml/kg as an early goal-directed therapy was associated with higher rates of fluid overload, which resulted in increased use of diuretics and thoracocentesis for pleural effusions [35]. This approach can even delay the anticipation and initiation of vasopressors or even mechanical circulatory support. The impact on survival between an initial liberal fluid strategy and restrictive fluids with early use of vasopressors in the setting of septic shock is still uncertain and of interest; CLOVERS is an ongoing multicentre randomized study investigating this question [36]. For now, international guidelines recommend an initial fluid challenge of 250-ml bolus over 30 min up to 500 ml with a low threshold to consider noradrenaline or dobutamine support on intensive care [37].

Similarly, judicious fluid management is imperative in surgical patients during the intraoperative and postoperative period. Apart from the obvious consequence of pulmonary oedema, liberal intravenous fluid is also associated with bowel oedema which can lead to ileus, poor intestinal absorption and bacterial translocation [38]. This can potentially delay the transition to oral hydration and delay the re-initiation and absorption of vital anti-heart failure medications, of which most are not available in intravenous form. Above all, the key is regular clinical assessment of fluid status.

Yet, fluid management will only be as accurate as the clinical evaluation of fluid status which can be challenging and contradictory between clinicians. Jugular venous pressure can be subjective and chest radiographic signs of HF can be non-specific [39, 40]. To complicate matters, elderly patients can present with subclinical congestion [41]. Hence, we frequently see the counterintuitive dilemma of treating with diuretics and intravenous fluids together. One study found 11% inpatients with AHF (without sepsis or bleeding) were given intravenous fluids (median 1000 ml) alongside diuretics, presumably to counterbalance any harm from over-diuresis. This increases LV filling pressures and prolongs the AHF episode [42]. Another fear to address is the initial rise in sCr with
intravenous diuretics, swaying physicians to hold back on the furosemide dose, resulting in under-diuresis. The DOSE-AHF reassures us that high-dose intravenous furosemide versus low-dose regimen shows no differences in renal function over 60 days; rather, the former strategy offers greater diuresis without any significant renal impairment [43]. Patients with fluid overload should have diuretics up-titrated to achieve euvolaemia and not stopped or reduced prematurely based on renal function alone.

There is growing evidence that the use of N-terminal pro-B-type natriuretic peptide (BNP) and ultrasonography can help in fluid assessment. With serial BNPs, a falling level correlates well with a trend towards euvolaemia [44]. Echocardiography can reveal an inferior vena cava dilatation with little collapsibility to steer physicians away from further fluid administration, while lung ultrasound (LUS) can reveal bilateral diffuse vertical hyperechoic reverberations known as B-lines which can identify subclinical pulmonary oedema. The presence of ≥ 15 B-lines correlates with NT-proBNP > 1000 pg/ml and E/e’ ratio > 15 [45]. This may even lead to improved patient outcomes as demonstrated in a small randomized trial, LUS-HF, which found that recently discharged HF patients who were assessed using LUS during follow-up were associated with a significantly reduced rate of HF re-admissions [46]. Clinicians should thus integrate these newer technologies with physical examination for a more accurate judgement of fluid status.

**Transfusion-Related Circulatory Overload**

Heart Failure, CKD and hypertension may be associated risk factors with transfusion-related circulatory overload (TACO) which manifests as pulmonary oedema within 6 h of transfusion and is associated with raised BP, BNP and a high mortality [47]. Small frail elderly adults are theoretically at greater risk of fluid overload if they receive a greater volume of blood transfusion than necessary. An average unit of packed red blood cells is 280 ml, and a dose of 4 ml/kg generally raises haemoglobin by 10 g/l [48]. Thus, an 80-year-old lady weighing 50 kg will only require 200 ml to raise haemoglobin by 10 g/l. However, if a whole unit is transfused and there is underlying HF, AHF may quickly follow. Although a Cochrane review indicated insufficient evidence, premedication with furosemide before blood transfusion remains common practice [49]. In contrast, volume depletion may precipitate AHF in the presence of LV outflow tract obstruction (LVOTO). A reduced LV volume and tachycardia can augment the intraventricular gradient and diastolic pressures precipitating pulmonary oedema. These are mainly reported through case series. For example, one case reported a patient with hypertrophic cardiomyopathy who developed significant LVOTO and pulmonary oedema after being kept nil by mouth and having received light sedation. This subsequently improved with intravenous fluids and beta-blockers [50]. Interestingly, mid-cavity LVOTO can also occur in up to 25% patients with Takotsubo cardiomyopathy predisposed by its basal myocardial hyperkinesia, excessive sympathetic stimulation and, in extreme cases, cardiogenic shock can ensue when inotropic agents are used [51, 52]. Overall, too much or too little intravenous fluids can trigger AHF if we overlook the clinical context.

**High-Output Heart Failure**

**Anaemia**

Severe anaemia can also worsen LVOTO as described above, but it is better known for unmasking myocardial ischaemia and causing high-output heart failure [53]. Anaemia can promote renal nitric oxide synthase activity. This causes peripheral vasodilation, neurohormonal activation, driving cardiac output (CO) above 8 l/min which over time leads to LV hypertrophy and dilation. All this can culminate to high-output HF [54, 55]. Anaemia is an independent predictor of HF mortality, yet a comorbidity that can go unnoticed [56]. It can be iatrogenic from surgical blood loss, post-catheterization haematuria or venepunctures, accelerated by anticoagulation or antiplatelets, and in turn, exacerbate a pre-existing anaemia of chronic disease from hepatic upregulation or antiplatelets, and in turn, exacerbate a pre-existing anaemia of chronic disease from hepatic upregulation [57]. Ferritin and transferrin saturation should therefore be routinely checked in HF. This can guide the prescription of intravenous iron which reduces HF hospitalization and improves exercise capacity, and is recommended over blood transfusion given the TACO risk [48, 58].

**Arterio-venous Fistula**

Less commonly, the low systemic vascular resistance from a systemic arterio-venous shunt can similarly activate the systemic neurohormonal system. The degree of CO augmentation depends on the shunt size and magnitude of flow. Nephrologists should be wary that dialysis patients are at further risk of HF due to the additive effects of co-existing anaemia [55]. An arterio-venous fistula can be created inadvertently during vascular access e.g. in angiography or temporary pacing. A case report described a patient with AHF due to an iatrogenic femoral arterio-venous fistula from cannulation, and was found to have a CO > 16 l/min which was resolved after shunt closure [59]. Cardiologists should therefore be vigilant of this rare complication.

**Pacemaker-Related HF**

A complication more familiar to cardiologists is pacing-induced cardiomyopathy defined by > 10% reduction in
LVEF after pacemaker implantation (having excluded other causes). There are several mechanisms behind this [60]. Firstly, the pacemaker wire may affect closure of the tricuspid valve which may cause haemodynamically significant tricuspid regurgitation. This is uncommon. Secondly, with VVI pacing, if the sinus node discharges faster than the programmed rate, the atria will contract against a closed atrioventricular valve causing atrioventricular dyssynchrony (known as pacemaker syndrome) which can precipitate AHF. This can be relieved by either lowering the pacing rate or upgrading to a dual-chamber pacemaker (e.g. DDD) which supports a more physiological AV sequential pacing [61]. However, dual-chamber pacing can also precipitate AHF in patients with underlying HF. The DAVID trial found that in patients with underlying LVSD, DDD pacing (especially when right ventricular apical pacing (RVP) exceeded 40%) led to more HF hospitalization or deaths compared with ventricular back-up pacing. This is thought to be driven by electromechanical dyssynchrony and subsequent maladaptive LV remodelling. Indeed, when DDD resulted in < 40% RVP, the HF hospitalization rate was lower [62]. The BLOCK-HF trial shed light on this. In patients with high-grade AV block and LVEF ≤ 50%, biventricular pacing was associated with a 10% absolute risk reduction in death and AHF hospitalization compared with RVP [63]. As a result, those patients with HFpEF regardless of severity that require anti-bradycardia pacing are recommended to have biventricular pacing de novo or as an upgrade [1, 61].

Conclusion

Iatrogenic decompensated HF is a phenomenon that is infrequently documented in the literature but increasingly confronted by clinicians of all specialties. It is associated with a high mortality and morbidity rate. As the HF population continues to age and become burdened with greater comorbidities and polypharmacy, patients become more susceptible to the iatrogenic precipitants of HF. Understandably, IAHF is sometimes unavoidable when concomitant renal dysfunction deteriorates, and the need to stop RAASi outweighs its benefit, or when an overarching concurrent illness develops, such as cancer which necessitates cardiotoxic chemotherapy. However, as long as clinicians are wary of the potential sequelae of AHF from these acts of commission or omission, we can anticipate and act promptly with timely monitoring of LV function and initiation of HF medications before AHF ensues. The already complex management of HF can be further complicated when a patient becomes acutely unwell with sepsis or significant anaemia. An effective yet simple approach would be regular clinical assessment integrated with the use of pulmonary ultrasound and biochemical markers such as BNP. Above all, by having greater awareness of these iatrogenic triggers, this phenomenon should be one that is prevented rather than managed when it occurs.

Compliance with Ethical Standards

Conflict of Interest The authors declare that they have no conflict of interest.

Human and Animal Rights and Informed Consent This article does not contain any studies with human or animal subjects performed by any of the authors.

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References

Papers of particular interest, published recently, have been highlighted as:

• Of major importance

1. Ponikowski P, Voors AA. 2016 ESC Guidelines for the diagnosis and treatment of acute and chronic heart failure: the Task Force for the diagnosis and treatment of acute and chronic heart failure of the European Society of Cardiology (ESC). Eur Heart J. 2016;37:2129–200.
2. National Heart Failure Audit (NHFA) April 2015 – March 2016. National Institute for Cardiovascular Outcomes Research (NICOR), Institute of Cardiovascular Science, University College London. 2016. http://www.ucl.ac.uk/nicor/audits/heartfailure/documents/annualreports/annual-report2015–6-v8.pdf. Accessed at 21 Sept 2019.
3. Rich M, Shah A, Vinson J, et al. Iatrogenic congestive heart failure in older adults: clinical course and prognosis. J Am Geriatr Soc. 1996;44:638–43.
4. Burnett H, Earley A, Voors AA, et al. Thirty years of evidence on the efficacy of drug treatments for chronic heart failure with reduced ejection fraction: a network meta-analysis. Circ Heart Fail. 2017;10:e003529.
5. McMurray JJV, Solomon SD, Inzucchi SE, et al. Dapagliflozin in patients with heart failure and reduced ejection fraction: the EMPEROR-Reduced Trial. J Am Coll Cardiol. 2019;73:1995–2008. https://doi.org/10.1016/j.jacc.2019.05.002.
6. Brar S, Ye F, James MT, et al. Association of angiotensin-converting enzyme inhibitor or angiotensin receptor blocker use with outcomes after acute kidney injury. JAMA Intern Med. 2018;178:1681.
7. Damman K, Navis G, Smilde TD, et al. Decreased cardiac output, venous congestion and the association with renal impairment in patients with cardiac dysfunction. Eur J Heart Fail. 2007;9:872–8.
8. Clark AL, Kalra PR, Petrie MC, Mark PB, Tomlinson LA, CRV T. Change in renal function associated with drug treatment in heart failure: national guidance. Heart. 2019;105:904–10 This article offers excellent guidance on the decision-making of continuation or withdrawal of RAAsIs in the face of changes in renal function.

9. Mullens W, Abrahams Z, Skouri HN, et al. Elevated intrabdominal pressure in acute decompensated heart failure: a potential contributor to worsening renal function? J Am Coll Cardiol. 2008;51:300–6.

10. Testani JM, Cappola TP, Brensinger CM, et al. Interaction between loop diuretic-associated mortality and blood urea nitrogen concentration in chronic heart failure. J Am Coll Cardiol. 2011;58:375–82.

11. Gilstrap LG, Fonarow GC, Desai AS, et al. Initiation, continuation, or withdrawal of angiotensin-converting enzyme inhibitors/angiotensin receptor blockers and outcomes in patients hospitalized with heart failure with reduced ejection fraction. J Am Heart Assoc. 2017;6:e004675. https://doi.org/10.1161/JAHA.116.004675.

12. Clark H, Krum H, Hopper I. Worsening renal function during renin-angiotensin aldosterone system inhibitor initiation and long-term outcomes in patients with left ventricular systolic dysfunction. Eur J Heart Fail. 2014;16:41–8.

13. Jondeau G, Neuder Y, Eicher JC, et al. B-CONVINCED: Beta-blocker CONtinuation Vs. INterruption in patients with Congestive heart failure hospitalizeD for a decomposition episode. Eur Heart J. 2009;30:2186–92.

14. Prins KW, Neill JM, Tyler JO, Eckman PM, Duval S. Effects of beta-blocker withdrawal in acute decompensated heart failure: a systematic review and meta-analysis. J Am Coll Cardiol HF. 2015;3:647–53.

15. O’Brien ET. Beta-blockade withdrawal. Lancet. 1975;2:819.

16. Miller RR, Olson HG, Amsterdam EA, Mason DT. Propranolol withdrawal rebound phenomenon: exacerbation of coronary events after abrupt cessation of antianginal therapy. N Engl J Med. 1975;293:416–8.

17. Fonarow GC, Abraham WT, Albert NM, OPTIMIZE-HF Investigators and Coordinators, et al. Influence of beta-blocker continuation or withdrawal on outcomes in patients hospitalized with heart failure: findings from the OPTIMIZE-HF program. J Am Coll Cardiol. 2008;52:190–9.

18. Halliday BP, Wassell R, Lota AS, et al. Withdrawal of pharmacological treatment for heart failure in patients with recovered dilated cardiomyopathy (TRED-HF): an open-label, pilot, randomised trial. Lancet. 2018. https://doi.org/10.1016/S0140-6736(18)32484-X.

19. Plana JC, Galderisi M, Barac A, et al. Expert consensus for cancer treatments and cardiovascular toxicity developed under the auspices of the ESC Committee for Practice Guidelines. Eur Heart Journal. 2016;37:2768–801. https://doi.org/10.1093/eurheartj/ehw211.

20. Barry E, Alvarez JA, Scully RE, Miller TL, Lipschultz SE. Anthracycline-induced cardiotoxicity: course, pathophysiology, prevention and management. Expert Opin Pharmacother. 2007;8:1039–58. https://doi.org/10.1517/14656566.8.8.1039.

21. Smith I, et al. 2-year follow-up of trastuzumab after adjuvant chemotherapy in HER2-positive breast cancer: a randomized controlled trial. Lancet. 2007;369:29–36.

22. Zamorano JL, Lancellotti P, Munoz DR, et al. 2016 ESC Position Paper on cancer treatments and cardiovascular toxicity developed under the auspices of the ESC Committee for Practice Guidelines. Eur Heart Journal. 2016;37:2768–801. https://doi.org/10.1093/eurheartj/ehw211.

23. Saczynski JS, Go AS, Magid DJ, et al. Patterns of comorbidity in older adults with heart failure: the Research Network PRESERVE study. J Am Geriatr Soc. 2013;61:26–33.

24. Wei L, MacDonald TM, Walker BR. Taking glucocorticoids by prescription is associated with subsequent cardiovascular disease. Ann Intern Med. 2004;141:764–7.

25. Feenstra J, Heerdink ER, Grobbee DE, Stricker BH. Association of nonsteroidal anti-inflammatory drugs with first occurrence of heart failure and with relapsing heart failure: the Rotterdam Study. Arch Intern Med. 2002;162:265–70.

26. Gislason GH, Rasmussen JN, Abildstrom SZ, et al. Increased mortality and cardiovascular morbidity associated with use of nonsteroidal anti-inflammatory drugs in chronic heart failure. Arch Intern Med. 2009;169:141–9. https://doi.org/10.1001/archinternmed.2008.525.

27. Kristensen SD, Knuts J, Saraste A, et al. ESC/ESA Guidelines on non-cardiac surgery: cardiovascular assessment and management. Kardiol Pol. 2014;72:857–918.

28. Nesserhi FH. Doxazosin and congestive heart failure. J Am Coll Cardiol. 2001;38:1295–6.

29. Gerstein HC, Yusuf S, Bosch J, et al. Effect of rosiglitazone on the frequency of diabetes in patients with impaired glucose tolerance or impaired fasting glucose: a randomised controlled trial. Lancet. 2006;368:1096–105 [published correction appears in Lancet. 2006;368:1770].

30. Multicenter Diltiazem Postinfarction Trial Research Group. The effect of diltiazem on mortality and reinfarction after myocardial infarction: the Multicenter Diltiazem Postinfarction Trial Research Group. N Engl J Med. 1988;319:385–92.

31. Echt DS, Liebson PR, Mitchell LB, Peters RW, et al. Mortality and morbidity in patients receiving encaainide, flecaainide, or placebo: the Cardiac Arrhythmia Suppression Trial. N Engl J Med. 1991;324:781–8. https://doi.org/10.1056/NEJM199103213241201.

32. Kelm DJ, Perrin JT, Cartin-Ceba R, et al. Fluid overload in patients with severe sepsis and septic shock treated with early-goal directed therapy is associated with increased acute need for fluid-related medical interventions and hospital death. Shock. 2015;43(1):68–73. https://doi.org/10.1097/SHK.0000000000000268.

33. Swithin JG. Intravenous anaesthesia for the patient with left ventricular dysfunction. Semin Cardiothorac Vasc Anesth. 2006;10:43–8.

34. Levy MM, Evans LE, Rhodes A. The surviving sepsis campaign bundle: 2018 update. Crit Care Med. 2018;46(6):997–1000. https://doi.org/10.1097/ccm.0000000000003119.

35. Kelm DJ, Perrin JT, Cartin-Ceba R, et al. Fluid overload in patients with severe sepsis and septic shock treated with early-goal directed therapy is associated with increased acute need for fluid-related medical interventions and hospital death. Shock. 2015;43(1):68–73. https://doi.org/10.1097/SHK.0000000000000268.

36. Self WH, Semler MW, Bellomo R, et al. Liberal versus restrictive intravenous fluid therapy for early septic shock: rationale for a randomized trial. Ann Emerg Med. 2018;72:457–66.

37. Rhodes A, Evans LE, Alhazzani W, et al. Surviving sepsis campaign: international guidelines for management of sepsis and septic shock. Crit Care Med. 2017;45(3):486–552. https://doi.org/10.1097/ccm.0000000000002255.

38. Makaryus R, et al. Current concepts of fluid management in enhanced recovery pathways. Br J Anaesth. 2018;120(2):376–83. https://doi.org/10.1016/j.bja.2017.10.011.

39. Pellicori P, Clark AL. Newer technologies and jugular venous pressure. Br J Cardiol. 2014;21:146.

40. Gadsboll N, Holund-Carlson PF, Nielsen GG, et al. Symptoms and signs of heart failure in patients with myocardial infarction: reproducibility and relationship to chest X-ray, radionuclide ventriculography and right heart catheterization. Eur Heart J. 1989;10:1017–28.

41. Van Riet EE, Hoes AW, Limburg A, et al. Prevalence of unrecognized heart failure in older persons with shortness of breath on exertion. Eur J Heart Fail. 2014;16:772–7.
42. Bikdeli B, Strait K, Dharmarajan K, et al. Intravenous fluids in acute decompensated heart failure. JACC Heart Failure. 2015;3:127–33.
43. Felker GM, Lee KL, Bull DA, et al. Diuretic strategies in patients with acute decompensated heart failure. N Engl J Med. 2011;364:797–805.
44. Kazanegra R, Cheng V, Garcia A, Krishnaswamy P, Gardetto N, Clopton P, et al. A rapid test for B-type natriuretic peptide correlates with falling wedge pressures in patients treated for decompensated heart failure: a pilot study. J Card Fail. 2001;7:21–9.
45. Miglioranza MH, Gargani L, Sant’Anna RT, et al. Lung ultrasound for the evaluation of pulmonary congestion in outpatients: a comparison with clinical assessment, natriuretic peptides and echocardiography. JACC Cardiovasc Imaging. 2013;6(11):1141–51.
46. Rivas-Lasarte M, Maestro-Benedicto A, Fernandez-Martinez J, et al. Lung ultrasound may reduce heart failure hospitalizations: preliminary results from the LUS-HF Trial. Eur J Heart Fail. 2019. https://doi.org/10.1002/ejhf.1604.
47. Roubinian NH, Hendrickson JE, Triulzi DJ, et al. Contemporary risk factors and outcomes of transfusion-associated circulatory overload. Crit Care Med. 2018;46(4):577–85.
48. JPAC Joint United Kingdom (UK) Blood Transfusion and Tissue Transplantation Services Professional Advisory Committee. 4.13: Transfusion of blood components. Access Online- https://www.transfusionguidelines.org/transfusion-handbook/4-safe-transfusion-right-blood-right-patient-right-time-and-right-place/4-13-transfusion-of-blood-components. Accessed at 01/10/19.
49. Sarai M, Tejani AM. Loop diuretics for patients receiving blood transfusions. Cochrane Database Syst Rev. 2015;(2):CD010138. https://doi.org/10.1002/14651858.CD010138.pub2.
50. Arundepp K, Arunpreet K, Amrita D, et al. Try fluids for failing heart. Crit Care Med. 2013;41:346–7. https://doi.org/10.1097/01.ccm.0000440571.21226.8d.
51. Y-Hassan S, Tomvall P. Epidemiology, pathogenesis and management of takotsubo syndrome. Clin Auton Res. 2018;28(1):53–65. https://doi.org/10.1007/s10286-017-0465-z.
52. El Mahmoud R, Mansencal N, Pilliere R, et al. Prevalence and characteristics of left ventricular outflow tract obstruction in Takotsubo syndrome. Am Heart J. 2008;156:543–8.
53. Yokusoglu M, Nevruz O, Baysan O, et al. The altered autonomic nervous system activity in iron deficiency anemia. Tohoku J Exp Med. 2007;212:397–402.
54. Ni Z, Morcos S, Vaziri ND. Up-regulation of renal and vascular nitric oxide synthase in iron-deficiency anemia. Kidney Int. 1997;52:195–201.
55. Mehta PA, Dubrey SW. High output heart failure. Q J Med. 2009;102:235–41. https://doi.org/10.1093/qjmed/hcn147.
56. Cohen-Solal A, Damy T, Terbah M, et al. High prevalence of iron deficiency in patients with acute decompensated heart failure. Eur J Heart Fail. 2014;16(9):984–91.
57. Weiss G, Goodnough LT. Anemia of chronic disease. N Engl J Med. 2005;352:1011–23.
58. Jankowska EA, Tkaczyzsyn M, Suchocki T, et al. Effects of intravenous iron therapy in iron-deficient patients with systolic heart failure: a meta-analysis of randomized controlled trials. Eur J Heart Fail. 2016;18(7):786–95.
59. Sethi SS, Wiesner P, Green P, et al. An iatrogenic cause of a heart failure exacerbation. JACC. 2018;71:11.
60. Khurshid S, Epstein AE, Verdino RJ, et al. Incidence and predictors of right ventricular pacing-induced cardiomyopathy. Heart Rhythm. 2014;11(9):1619–25.
61. Brignole M, Auricchio A, Baron-Esquivias G, et al. ESC Guidelines on cardiac pacing and cardiac resynchronization therapy: the Task Force on cardiac pacing and resynchronization therapy of the European Society of Cardiology (ESC). Developed in collaboration with the European Heart Rhythm Association (EHRA). Eur Heart Journal. 2013;34:2281–329. https://doi.org/10.1093/eurheartj/eut206.
62. Wilkoff BL, Cook JR, Epstein AE, Dual Chamber and VVI Implantable Defibrillator Trial Investigators, et al. Dual-chamber pacing or ventricular backup pacing in patients with an implantable defibrillator: the Dual Chamber and VVI Implantable Defibrillator (DAVID) Trial. JAMA. 2002;288(24):3115–23.
63. Curtis AB, Worley SJ, Adamson PB, et al. Biventricular pacing for atrioventricular block and systolic dysfunction. N Engl J Med. 2013;368(17):1585–93. https://doi.org/10.1056/NEJMoa1210356.

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