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Systematic review

# Efficacy of implantable haemodynamic monitoring in heart failure across ranges of ejection fraction: a systematic review and meta-analysis

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**ABSTRACT**

**Aims** We conducted a meta-analysis of randomised controlled trials (RCTs) of implantable haemodynamic monitoring (IHM)-guided care.

**Methods** PubMed and Ovid MEDLINE were searched for RCTs of IHM in patients with heart failure (HF). Outcomes were examined in total (first and recurrent) event analyses.

**Results** Five trials comparing IHM-guided care with standard care alone were identified and included 2710 patients across ejection fraction (EF) ranges. Data were available for 628 patients (23.2%) with heart failure with preserved ejection fraction (HFpEF) (EF  $\geq 50\%$ ) and 2023 patients (74.6%) with heart failure with a reduced ejection fraction (HFrEF) (EF  $< 50\%$ ). Chronicle, CardioMEMS and HeartPOD IHMs were used. In all patients, regardless of EF, IHM-guided care reduced total HF hospitalisations (HR 0.74, 95% CI 0.66 to 0.82) and total worsening HF events (HR 0.74, 95% CI 0.66 to 0.84). In patients with HFrEF, IHM-guided care reduced total worsening HF events (HR 0.75, 95% CI 0.66 to 0.86). The effect of IHM-guided care on total worsening HF events in patients with HFpEF was uncertain (fixed-effect model: HR 0.72, 95% CI 0.59 to 0.88; random-effects model: HR 0.60, 95% CI 0.32 to 1.14). IHM-guided care did not reduce mortality (HR 0.92, 95% CI 0.71 to 1.20). IHM-guided care reduced all-cause mortality and total worsening HF events (HR 0.80, 95% CI 0.72 to 0.88).

**Conclusions** In patients with HF across all EFs, IHM-guided care reduced total HF hospitalisations and worsening HF events. This benefit was consistent in patients with HFrEF but not consistent in HFpEF. Further trials with pre-specified analyses of patients with an EF of  $\geq 50\%$  are required.

**PROSPERO registration number** CRD42021253905.

**INTRODUCTION**

Remote monitoring using implanted devices may provide useful information about the natural history of congestion leading to decompensation in people with heart failure (HF). Early identification of increasing cardiopulmonary pressures and intervention to reduce these might decrease the risk of subsequent HF hospitalisation. Based on the CardioMEMS Heart Sensor Allows Monitoring of Pressure to Improve Outcomes in New York Heart Association Class III Heart Failure Patients (CHAMPION) trial, one implantable pulmonary

**WHAT IS ALREADY KNOWN ON THIS TOPIC**

⇒ Implantable haemodynamic monitoring (IHM) to guide treatment has a class IIb recommendation in the 2021 European Society of Cardiology Heart Failure Guidelines.

**WHAT THIS STUDY ADDS**

⇒ This is the first meta-analysis to examine the effectiveness of IHM-guided care across a range of ejection fractions (EFs), combining data from the five randomised trials that investigated IHM-guided treatment, including a pre-COVID-19 sensitivity analysis from the recent Hemodynamic-Guided Management of Heart Failure (GUIDE-HF) trial. This study demonstrates that IHM-guided treatment was effective at reducing worsening heart failure (HF) events in patients with an EF of  $< 50\%$ ; it is uncertain if patients with HF with preserved EF receive the same benefit.

**HOW THIS STUDY MIGHT AFFECT RESEARCH, PRACTICE OR POLICY**

⇒ This study indicates that future trials should focus on people with an EF of  $\geq 50\%$  with pre-specified analyses to confirm the effectiveness of IHM-guided care in this population. IHM-guided treatment can be considered as a strategy in patients with an EF of  $< 50\%$ .

artery pressure monitor (CardioMEMS, Abbott, Illinois, USA) received a class IIb recommendation to reduce HF hospitalisations in the 2021 European Society of Cardiology Heart Failure guidelines.<sup>1</sup> In the Hemodynamic-Guided Management of Heart Failure (GUIDE-HF) trial, the largest randomised controlled trial (RCT) to compare implantable haemodynamic monitoring (IHM) with standard care, IHM-guided care did not reduce HF hospitalisations overall, but sensitivity analyses suggested a modest benefit before the COVID-19 pandemic had an impact on patient management.<sup>2</sup>

No previous meta-analysis has included data from both the GUIDE-HF and the Left Atrial Pressure Monitoring to Optimize Heart Failure Therapy (LAPTOP-HF) study, the latter of which was an international RCT that reported its findings in 2016 (table 1). Additionally, despite IHM randomised



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**Table 1** Randomised controlled trials of IHM-guided HF management compared with standard care

| Trial      | First author and year                        | Design, country                                    | Primary efficacy endpoint  | Numbers enrolled | EF                        | NYHA class | Previous HF event                                       | Follow-up |
|------------|--|--|--|------------------|---------------------------|------------|---|-----------|
| COMPASS-HF | Bourge <i>et al.</i> , <sup>16</sup> 2008    | Single-blind,* multicentre RCT; USA                | HF hospitalisation and ED and urgent clinic visit for intravenous therapy (included hypovolaemic events) | 274              | No EF inclusion criterion | III-IV     | ≤6 months (or ED visit)                                 | 6 months  |
| CHAMPION   | Abraham <i>et al.</i> , <sup>18</sup> 2011   | Single-blind,* multicentre RCT; USA                | HF hospitalisation   | 550              | No EF inclusion criterion | III        | ≤12 months  | 6 months  |
| REDUCE-HF  | Adamson <i>et al.</i> , <sup>10</sup> 2011   | Single-blind,* multicentre RCT; USA                | HF hospitalisation and ED and urgent clinic visit for intravenous therapy                                | 400              | No EF inclusion criterion | II-III     | ≤12 months  | 12 months |
| LAPTOP-HF  | Abraham <i>et al.</i> , <sup>17</sup> 2016   | Multicentre RCT (no blinding); USA and New Zealand | HF hospitalisation and complications from HF therapy   | 486              | No EF inclusion criterion | III        | ≤12 months (or BNP ≥400 pg/mL or NT-proBNP ≥1500 pg/mL) | 12 months |
| GUIDE-HF   | Lindenfeld <i>et al.</i> , <sup>2</sup> 2021 | Single-blind,* multicentre RCT; USA and Canada     | All-cause mortality and HF hospitalisation and ED and urgent clinic visit for intravenous therapy        | 1000             | No EF inclusion criterion | II – IV    | ≤12 months (or BNP ≥250 pg/mL or NT-proBNP ≥1000 pg/mL) | 12 months |

\*Patients but not investigators were blinded to haemodynamic data.  
BNP, brain natriuretic peptide; CHAMPION, CardioMEMS Heart Sensor Allows Monitoring of Pressure to Improve Outcomes in New York Heart Association Class III Heart Failure Patients; COMPASS-HF, Chronicle Offers Management to Patients with Advanced Signs and Symptoms of Heart Failure; ED, emergency department; EF, ejection fraction; GUIDE-HF, Hemodynamic-Guided Management of Heart Failure; HF, heart failure; IHM, implantable haemodynamic monitor; LAPTOP-HF, Left Atrial Pressure Monitoring to Optimize Heart Failure Therapy; NT-proBNP, N-terminal pro-B-type natriuretic peptide; NYHA, New York Heart Association; RCT, randomised controlled trial; REDUCE-HF, Reducing Events in Patients with Chronic Heart Failure.

trials recruiting patients across a range of ejection fractions (EFs), no meta-analysis reported the effect of IHM on the reduction of HF hospitalisations and related events in subgroups of

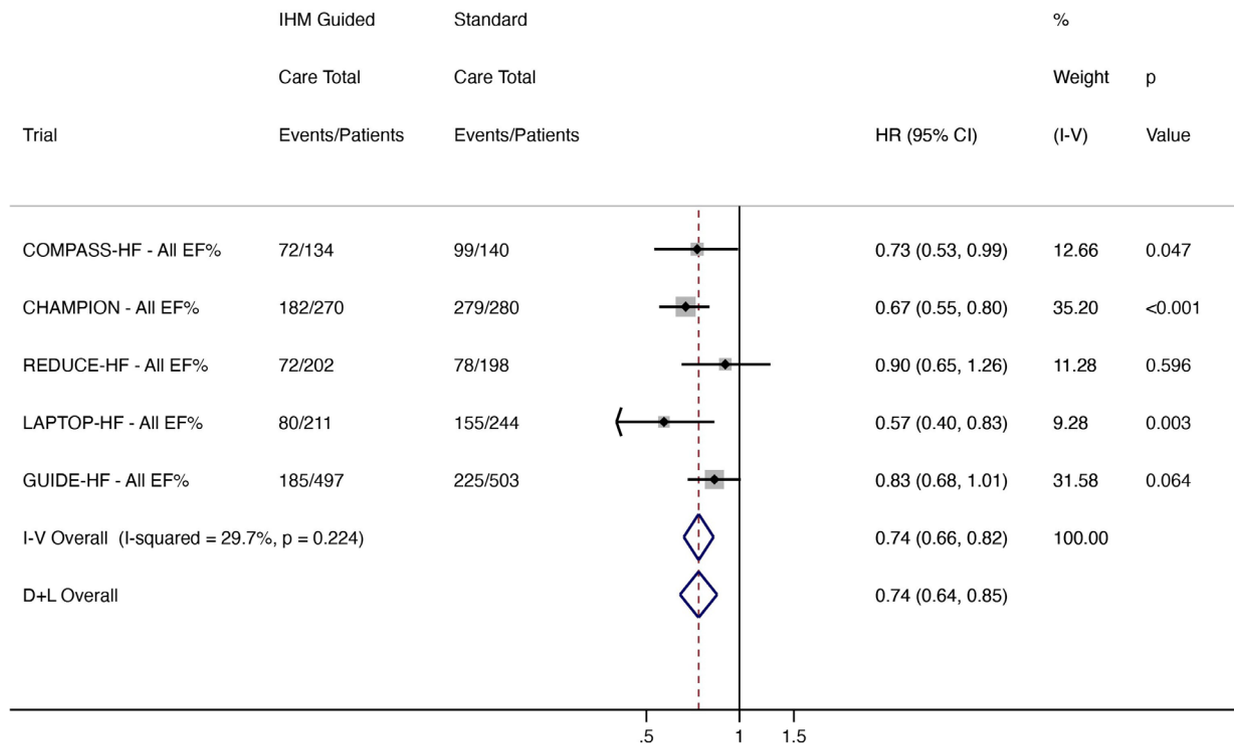
patients with HF with preserved ejection fraction (HFpEF) or heart failure with a reduced ejection fraction (HFrEF). This type of monitoring is of particular interest in patients with HFpEF,

**Table 2** Key baseline characteristics of patients enrolled in randomised controlled trials of IHM-guided HF management compared with standard care

|                         | COMPASS-HF   | CHAMPION   | REDUCE-HF  | LAPTOP-HF  | GUIDE-HF   |
|-------------------------|--|--|--|--|--|
| Age (years)             | IHM arm: 58±14<br>Control arm: 58±13                                       | IHM arm: 61±13<br>Control arm: 62±13                 | IHM arm: 55±15<br>Control arm: 55±15                                       | IHM and control arms: 62±12  | IHM arm: 71 (64-76)<br>Control arm: 70 (64-77)   |
| Sex (male, %)           | IHM arm: 66<br>Control arm: 64   | IHM arm: 72<br>Control arm: 73                       | IHM arm: 70<br>Control arm: 67   | IHM and control arms: 75   | IHM arm: 62<br>Control arm: 63   |
| Race (%)                | IHM arm<br>▶ White: 73<br>Control arm<br>▶ White: 80                       | IHM arm<br>▶ White: 73<br>Control arm<br>▶ White: 73 | IHM arm<br>▶ White: 55<br>Control arm<br>▶ White: 58                       | n/r  | IHM arm<br>▶ White: 81<br>Control arm<br>▶ White: 80   |
| EF (n=participants)     | ≥50%: 70<br><50%: 204  | ≥50%: 66<br>≥40–49%: 53<br>≤40%: 456<br><40%: 430    | IHM and control arms: mean<br>EF 23%±7                                     | IHM and control arms, EF >35%:<br>121<br>IHM and control arms, EF ≤35%:<br>365<br>Mean EF 30%±15 | >50%: 321<br>≤50%: 679   |
| NYHA class (%)          | IHM arm:<br>▶ III: 84<br>▶ IV: 16<br>Control arm:<br>▶ III: 87<br>▶ IV: 13 | IHM arm: III only<br>Control arm: III only           | IHM arm:<br>▶ II: 53<br>▶ III: 47<br>Control arm:<br>▶ II: 45<br>▶ III: 55 | IHM arm: III only<br>Control arm: III only   | IHM arm:<br>▶ II: 29<br>▶ III: 65<br>▶ IV: 6<br>Control arm:<br>▶ II: 30<br>▶ III: 65<br>▶ IV: 5 |
| Ischaemic aetiology (%) | IHM arm: 47<br>Control arm: 44   | IHM arm: 59<br>Control arm: 62                       | IHM arm: 45<br>Control arm: 44   | IHM and control arms: 46   | IHM arm: 42<br>Control arm: 38   |
| Diuretics (%)           | IHM arm: 93<br>Control arm: 99   | IHM arm: 92<br>Control arm: 92                       | IHM arm: 93<br>Control arm: 97   | n/r  | IHM arm: 95<br>Control arm: 95   |
| ACEi/ARB (%)            | IHM arm: 85<br>Control arm: 81   | IHM arm: 76<br>Control arm: 79                       | IHM arm: 92<br>Control arm: 92   | n/r  | IHM arm*: 64<br>Control arm*: 64   |
| Beta blockers (%)       | IHM arm: 83<br>Control arm: 81   | IHM arm: 90<br>Control arm: 91                       | IHM arm: 96<br>Control arm: 96   | n/r  | IHM arm: 89<br>Control arm: 88   |

\*ARNI/ACEi/ARB.

ACEi, ACE inhibitor; ARB, angiotensin receptor blocker; ARNI, angiotensin receptor blocker-nepilysin inhibitor; CHAMPION, CardioMEMS Heart Sensor Allows Monitoring of Pressure to Improve Outcomes in New York Heart Association Class III Heart Failure Patients; COMPASS-HF, Chronicle Offers Management to Patients with Advanced Signs and Symptoms of Heart Failure; EF, ejection fraction; GUIDE-HF, Hemodynamic-Guided Management of Heart Failure; HF, heart failure; IHM, implantable haemodynamic monitoring; LAPTOP-HF, Left Atrial Pressure Monitoring to Optimize Heart Failure Therapy; n/r, not reported; NYHA, New York Heart Association; REDUCE-HF, Reducing Events in Patients with Chronic Heart Failure.



**Figure 1** Total (first and recurrent) HF hospitalisations in all patients regardless of EF. CHAMPION, CardioMEMS Heart Sensor Allows Monitoring of Pressure to Improve Outcomes in New York Heart Association Class III Heart Failure Patients; COMPASS-HF, Chronicle Offers Management to Patients with Advanced Signs and Symptoms of Heart Failure; D+L, DerSimonian and Laird; EF, ejection fraction; GUIDE-HF, Hemodynamic-Guided Management of Heart Failure; HF, heart failure; IHM, implantable haemodynamic monitor; I-V, inverse variance; LAPTOP-HF, Left Atrial Pressure Monitoring to Optimize Heart Failure Therapy; REDUCE-HF, Reducing Events in Patients with Chronic Heart Failure.

in whom evidence-based treatment options are limited. Therefore, in this meta-analysis of all randomised trials using IHM, we investigated whether treatment guided by such monitoring reduced the risk of total (first and recurrent) HF hospitalisations, total worsening HF events (HF hospitalisation and emergency department (ED) and urgent clinic visit for intravenous HF therapy) and all-cause mortality, when compared with standard care, in patients with HF across a range of EFs.

## METHODS

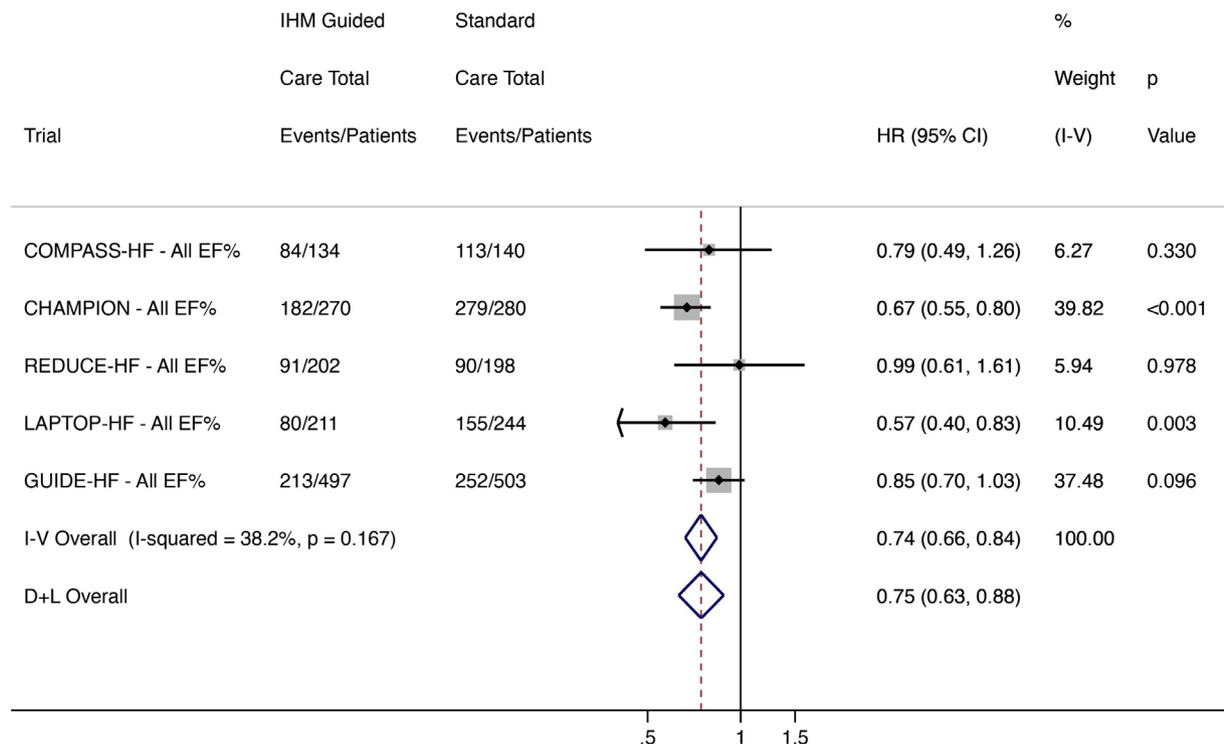
### Search strategy and data extraction

A systematic review of RCTs in patients with HF was performed, comparing IHM-guided care versus standard therapy. This meta-analysis was registered on PROSPERO (CRD42021253905). The Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines were followed to conduct the literature search, data extraction and reporting. A PRISMA checklist is included in online supplemental appendix, table S1). Bias was assessed using the Revised Cochrane Risk-of-Bias Tool for Randomised Trials<sup>3</sup> (online supplemental appendix, table S2). Searches were performed on public databases (PubMed and Ovid MEDLINE) between 1 May 2020 and 5 September 2022 using the terms “heart failure AND implantable AND haemodynamic monitoring” and “left atrial pressure AND heart failure AND monitoring” and “pulmonary artery pressure monitoring AND heart failure”. All studies published up to 5 September 2022 were eligible. No restriction was placed on study size, language or country of publication. Titles and abstracts were screened based on pre-specified inclusion criteria using the population, intervention, comparator, outcomes and study framework:

- Population: patients with HF.

- Intervention: IHM-guided care.
- Comparator: standard care.
- Outcomes of interest.
  - HF hospitalisation.
  - Worsening HF events (HF hospitalisation and ED and urgent clinic visits for intravenous HF therapy).
  - All-cause mortality.
  - All-cause mortality and HF hospitalisation.
  - All-cause mortality and worsening HF events.
- Study design: RCTs.

Full-text articles of original trial reports and published articles with retrospective analyses of the RCTs were included. Data presented at conferences were included if the presentation was available and verifiable by the researchers. Hazard ratios (HRs) and incidence rate ratios (IRRs) for endpoints were recorded. IRRs are approximations of HRs and were included as the effect estimate if HRs were not available as has previously been reported.<sup>4–6</sup> If either HR or IRR was not reported in the literature, the IRR was calculated using event numbers and study cohort time at risk. Ninety-five per cent CIs were calculated if only a p value and effect estimate were reported.<sup>7</sup> Numbers of patients in EF subgroups and their numbers of events were calculated from available data where necessary. Two researchers (JPC and MMYL) independently extracted and analysed the data. Results were compared and differences resolved by consensus with opinion from a third author. All authors reviewed the analysis results and contributed to drafting the report. If data were not available, the original study authors were contacted and data were requested.



**Figure 2** Total (first and recurrent) worsening HF events (HF hospitalisation and ED and urgent clinic visit for intravenous HF therapy) in all patients regardless of EF. CHAMPION, CardioMEMS Heart Sensor Allows Monitoring of Pressure to Improve Outcomes in New York Heart Association Class III Heart Failure Patients; COMPASS-HF, Chronicle Offers Management to Patients with Advanced Signs and Symptoms of Heart Failure; D+L, DerSimonian and Laird; ED, emergency department; EF, ejection fraction; GUIDE-HF, Hemodynamic-Guided Management of Heart Failure; HF, heart failure; IHM, implantable haemodynamic monitor; I-V, inverse variance; LAPTOP-HF, Left Atrial Pressure Monitoring to Optimize Heart Failure Therapy; REDUCE-HF, Reducing Events in Patients with Chronic Heart Failure.

### Statistical analysis

Statistical analyses were performed using Stata17. As the trials investigated three devices across different decades we used a random effects (DerSimonian and Laird (D+L)<sup>8</sup>) model so that differences in the studies' designs and cohorts would be accounted for within the analysis. We performed sensitivity analyses of each meta-analysis using fixed-effect models. Only the result of the fixed-effect model in patients with HFpEF is reported, as the other fixed-effect analyses were consistent with the reported random effect models.  $I^2$  statistic for percentage heterogeneity was computed with corresponding p values. Forest plots graphically report the pooled effect size estimates, the degree of heterogeneity and the weighted contribution each study made to the analyses. All outcomes were examined in total events (first and recurrent) analyses.

### Definitions of HFpEF and HFrEF

HFpEF was defined as HF with an EF of  $\geq 50\%$  in keeping with the 2021 European Society of Cardiology Heart Failure guidelines<sup>1</sup> and the recently proposed universal definition of HF.<sup>9</sup> HFrEF was defined as HF with an EF of  $< 50\%$ , with the inclusion of patients with heart failure with mildly reduced ejection fraction (HFmrEF, EF 41%–49%) as well as patients with an EF of  $\leq 40\%$ .<sup>19</sup> There were insufficient data available to further subclassify the trial cohorts into a HFmrEF subgroup. Only Chronicle Offers Management to Patients with Advanced Signs and Symptoms of Heart Failure (COMPASS-HF) pre-specified an analysis of patients with an EF of  $\geq 50\%$ . While EF was not an inclusion criterion for the Reducing Events in Patients with Chronic Heart Failure (REDUCE-HF) trial, patients included in

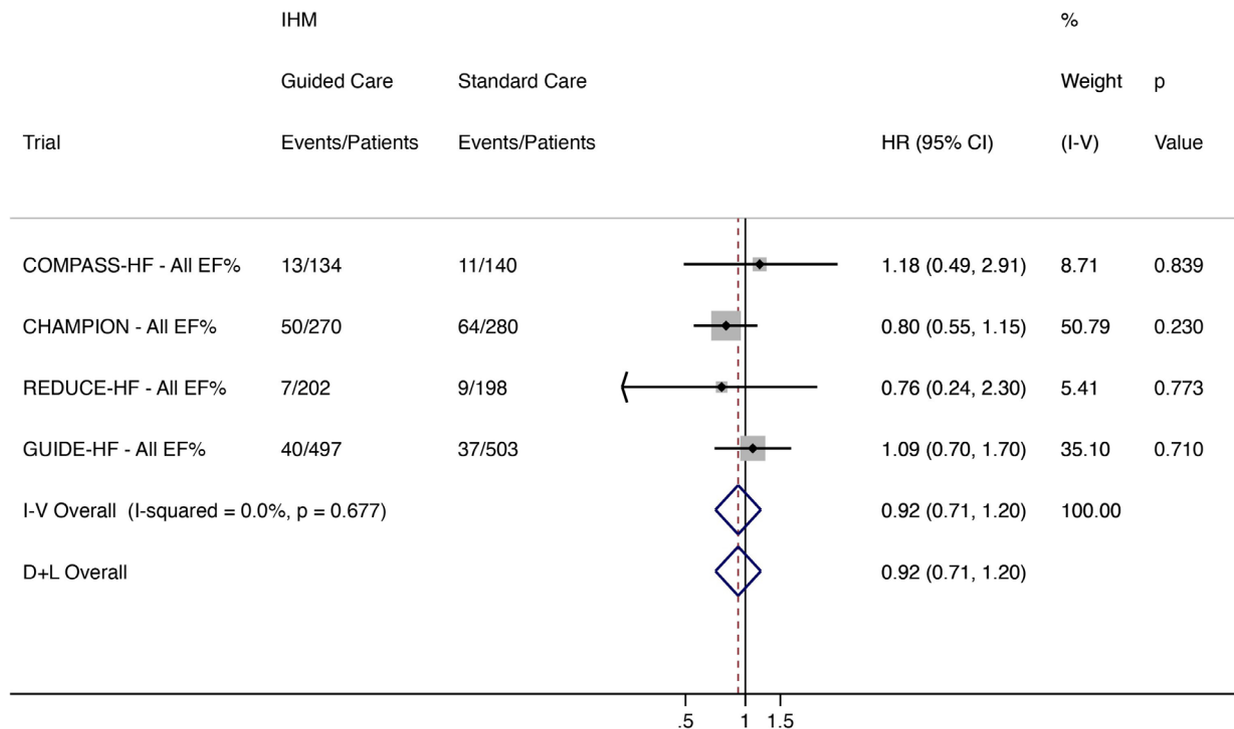
that trial had severely impaired systolic function with a mean ( $\pm$ SD) EF of  $23\% \pm 7\%$ .<sup>10</sup> Patients from the REDUCE-HF trial were therefore included in the HFpEF (EF  $< 50\%$ ) analysis. Similarly, patients from LAPTOP-HF, in whom the mean EF was  $30\% \pm 15\%$ <sup>11</sup> were included in the HFpEF (EF  $< 50\%$ ) analysis.

### Efficacy endpoints

The primary efficacy endpoints for each of the included trials were examined in total (first and recurrent) event analyses comparing the effect of IHM-guided care with standard care alone. These endpoints were as follows:

- ▶ COMPASS-HF and REDUCE-HF: total worsening HF events. HF hospitalisations for less than 24 hours were included in the composite endpoint in REDUCE-HF.
- ▶ LAPTOP-HF: total HF hospitalisation and complications of HF treatment such as hypotension and acute renal failure.
- ▶ CHAMPION: total HF hospitalisation.
- ▶ GUIDE-HF: all-cause mortality and total worsening HF events.

Recurrent events were analysed in a negative binomial regression model in COMPASS-HF and an Andersen-Gill model in CHAMPION, REDUCE-HF, LAPTOP-HF and GUIDE-HF. As these methods yield very similar results in simulations<sup>12</sup> and trial datasets,<sup>13</sup> they were used in the meta-analysis. Meta-analysis was performed for (1) total HF hospitalisations, (2) total worsening HF events (HF hospitalisation and ED visit and urgent clinic visit for intravenous HF therapy), (3) all-cause mortality, (4) all-cause mortality and total HF hospitalisations, and (5) all-cause mortality and total worsening HF events. Only hospitalisations for greater than 24 hours in REDUCE-HF were included



**Figure 3** All-cause mortality in all patients regardless of EF for (COMPASS-HF, CHAMPION, REDUCE-HF and GUIDE-HF). CHAMPION, CardioMEMS Heart Sensor Allows Monitoring of Pressure to Improve Outcomes in New York Heart Association Class III Heart Failure Patients; COMPASS-HF, Chronicle Offers Management to Patients with Advanced Signs and Symptoms of Heart Failure; D+L, DerSimonian and Laird; EF, ejection fraction; GUIDE-HF, Hemodynamic-Guided Management of Heart Failure; HF, heart failure; IHM, implantable haemodynamic monitor; I-V, inverse variance; LAPTOP-HF, Left Atrial Pressure Monitoring to Optimize Heart Failure Therapy; REDUCE-HF, Reducing Events in Patients with Chronic Heart Failure.

in the total HF hospitalisation analysis (aforementioned item 1). All-cause mortality data from COMPASS-HF, CHAMPION, REDUCE-HF and GUIDE-HF were pooled. The GUIDE-HF main trial results were published in 2021, followed by an analysis examining the impact of the COVID-19 pandemic on that trial's event rates.<sup>14</sup> We performed a sensitivity analysis examining the rate of HF events including the pre-COVID event rates and the results from the other four included trials.

## RESULTS

### Literature review and search strategy

A total of 1373 articles were identified by searching electronic databases. Two further articles<sup>11 15</sup> were found by hand searching references and internet searches. A PRISMA flowchart outlines the search process and identification of relevant articles (online supplemental appendix, figure S1). Five RCTs were identified (table 1): COMPASS-HF,<sup>16</sup> CHAMPION,<sup>17 18</sup> REDUCE-HF,<sup>10</sup> LAPTOP-HF<sup>11 15</sup> and GUIDE-HF.<sup>2</sup> The 18-month results for the CHAMPION trial were used in this meta-analysis. The HR for HF hospitalisation at 1 year was reported for 455 of the 486 randomised in LAPTOP-HF by the lead investigator at the annual meeting of the Heart Failure Association of the European Society of Cardiology in 2017.<sup>15</sup>

### Trial characteristics

COMPASS-HF, CHAMPION and REDUCE-HF were conducted in the USA. LAPTOP-HF was conducted in the USA and New Zealand. GUIDE-HF was conducted in the USA and Canada. The COMPASS-HF and REDUCE-HF studies evaluated the Chronicle pressure sensor (Medtronic, Minnesota, USA). The CardioMEMS device (Abbott, Illinois, USA) was investigated in CHAMPION and GUIDE-HF. The

HeartPOD device (St. Jude, Minnesota, USA) was investigated in LAPTOP-HF (table 1). The main trial characteristics are summarised in table 2.

All participants underwent implantation of haemodynamic monitors and were randomised to receive HF care guided by haemodynamic data or receive standard care. COMPASS-HF, CHAMPION, REDUCE-HF and GUIDE-HF were single-blind studies where investigators, but not patients, had access to the treatment group haemodynamic data. Patients were unaware of their randomised assignment groups in these four trials. LAPTOP-HF had no blinding (ie, patients and investigators were aware of the intervention assignment groups).

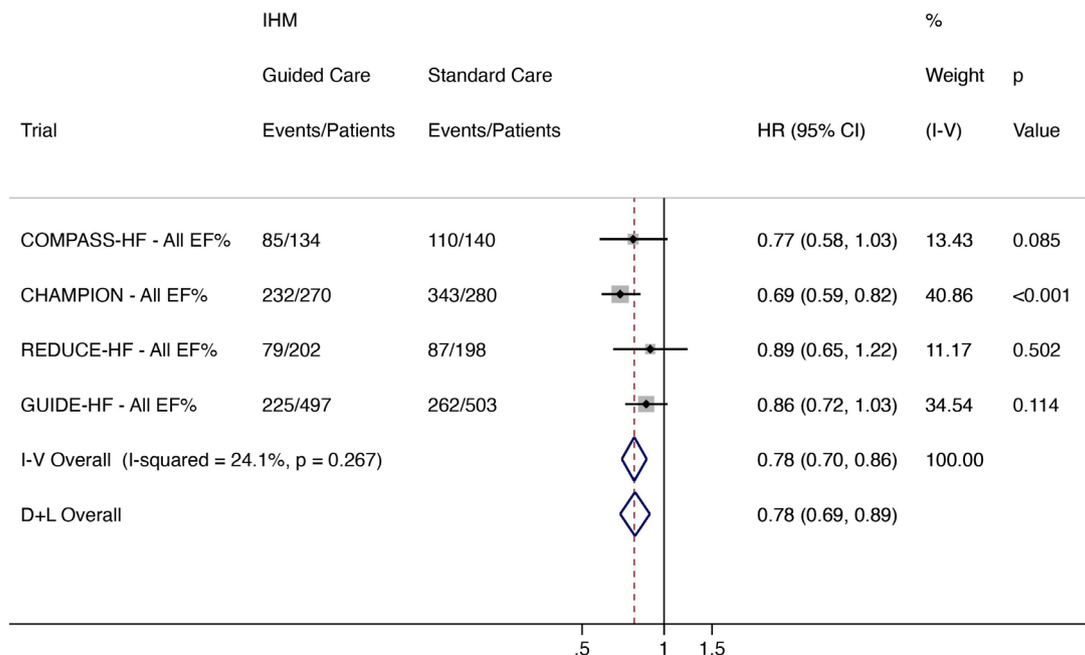
### Patients with HF regardless of EF

#### Total HF hospitalisations

There were 591 hospitalisations in 1314 patients receiving IHM-guided care compared with 836 events in 1365 standard care patients. HF hospitalisations were reduced in the IHM-guided care arm by 26% (HR 0.74, 95% CI 0.64 to 0.85; low heterogeneity ( $I^2$  29.7%)) (figure 1).

#### Total worsening HF events

There were 650 composite outcome events in 1314 patients receiving IHM-guided care and 889 events in 1365 standard care patients. IHM-guided care reduced total worsening HF events by 26% (HR 0.75, 95% CI 0.63 to 0.88; low heterogeneity ( $I^2$  38.2%)) (figure 2). In a sensitivity analysis of pre-COVID-19 event rates in the GUIDE-HF trial, IHM-guided care reduced HF events by 29% (HR 0.71, 95% CI 0.63 to 0.81; low heterogeneity ( $I^2$  2.9%)).



**Figure 4** Total (first and recurrent) HF hospitalisation and all-cause mortality in all patients regardless of EF (COMPASS-HF, CHAMPION, REDUCE-HF and GUIDE-HF). CHAMPION, CardioMEMS Heart Sensor Allows Monitoring of Pressure to Improve Outcomes in New York Heart Association Class III Heart Failure Patients; COMPASS-HF, Chronicle Offers Management to Patients with Advanced Signs and Symptoms of Heart Failure; D+L, DerSimonian and Laird; EF, ejection fraction; GUIDE-HF, Hemodynamic-Guided Management of Heart Failure; HF, heart failure; IHM, implantable haemodynamic monitor; I-V, inverse variance; LAPTOP-HF, Left Atrial Pressure Monitoring to Optimize Heart Failure Therapy; REDUCE-HF, Reducing Events in Patients with Chronic Heart Failure.

#### All-cause mortality

Mortality was reported in the COMPASS-HF, CHAMPION, REDUCE-HF and GUIDE-HF trials. Of 1103 patients, 110 (10.0%) in the IHM-guided care arm died compared with 121/1121 (10.8%) receiving standard care. IHM-guided care did not reduce all-cause mortality (HR 0.92, 95% CI 0.71 to 1.20; low heterogeneity ( $I^2$  0%)) (figure 3).

#### All-cause mortality and total HF hospitalisation

Data were available from COMPASS-HF, CHAMPION, REDUCE-HF and GUIDE-HF. A total of 621 events in 1103 patients occurred in the IHM arm and 802 events occurred in 1121 standard care patients. IHM-guided care, compared with standard care, reduced all-cause mortality and total HF hospitalisation by 22% (HR 0.78, 95% CI 0.69 to 0.89; low heterogeneity ( $I^2$  24.1%)) (figure 4).

#### All-cause mortality and total worsening HF events

A total of 680 events occurred in 1103 patients in the IHM arm and 855 events in 1121 standard care patients. IHM-guided care reduced all-cause mortality and worsening HF events by 20% (HR 0.81, 95% CI 0.69 to 0.94; moderate heterogeneity ( $I^2$  49.9%)). (online supplemental appendix, figure S2).

#### Patients with HFpEF (EF $\geq$ 50%)

Data were available from COMPASS-HF and CHAMPION for 136 patients with an EF of  $\geq$ 50%.<sup>19 20</sup> In the GUIDE-HF trial, a HR and event numbers were reported, but patient numbers for each randomised treatment group were not available.<sup>2</sup>

#### Total worsening HF events

A total of 186 events occurred in patients receiving IHM-guided treatment compared with 224 events in the standard care arm.

IHM-guided care reduced worsening HF when analysed using a fixed effect model (HR 0.72, 95% CI 0.59 to 0.88) but not a random-effects model (HR 0.60, 95% CI 0.32 to 1.14; high heterogeneity ( $I^2$  86.4%)) (figure 5).

#### Patients with HFrEF (EF <50%)

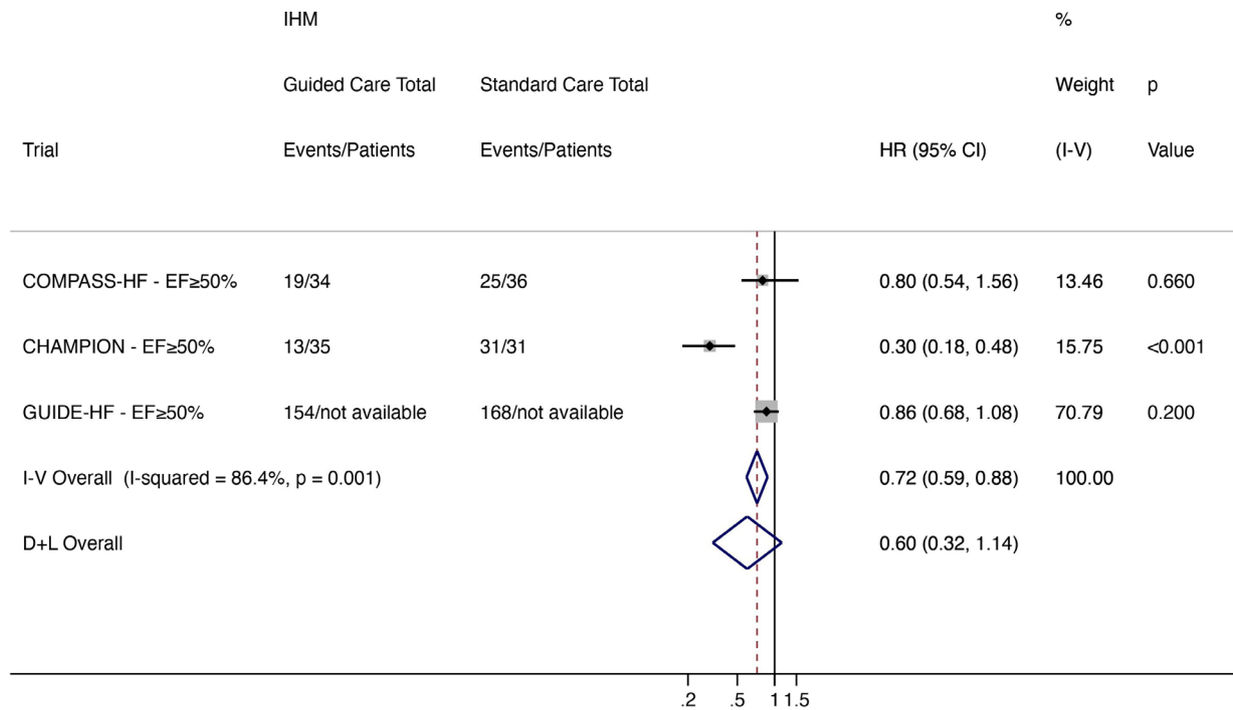
Data were available for 659 patients with an EF <50% from COMPASS-HF and LAPTOP-HF and for 856 patients with an EF  $\leq$ 40% from CHAMPION and REDUCE-HF. HRs and event numbers were reported for patients with an EF 40–50% and an EF <40% in GUIDE-HF but patient numbers were not available for each EF category in this trial.<sup>2</sup>

#### Total worsening HF events

A total of 497 events occurred in the IHM-guided care arm compared with 681 events in the standard care arm. IHM-guided care reduced worsening HF by 25% when compared with standard care (HR 0.75, 95% CI 0.66 to 0.86; low heterogeneity ( $I^2$  6.7%)) (figure 6).

## DISCUSSION

The main results of this meta-analysis support the use of IHM in patients with symptomatic HF (irrespective of EF), demonstrating a 26% reduction in the risk of total worsening HF events, including hospital admission. This is the first meta-analysis to include total HF events from all IHM randomised trials, including LAPTOP-HF and the recently reported GUIDE-HF. We also report for the first time meta-analyses of the effectiveness of IHM-guided care in patients with HFrEF and HFpEF. The finding of a reduction in total worsening HF events in all patients regardless of EF was also present in patients with HFrEF, who comprised the majority of patients. The same benefit was not consistent in patients with HFpEF.



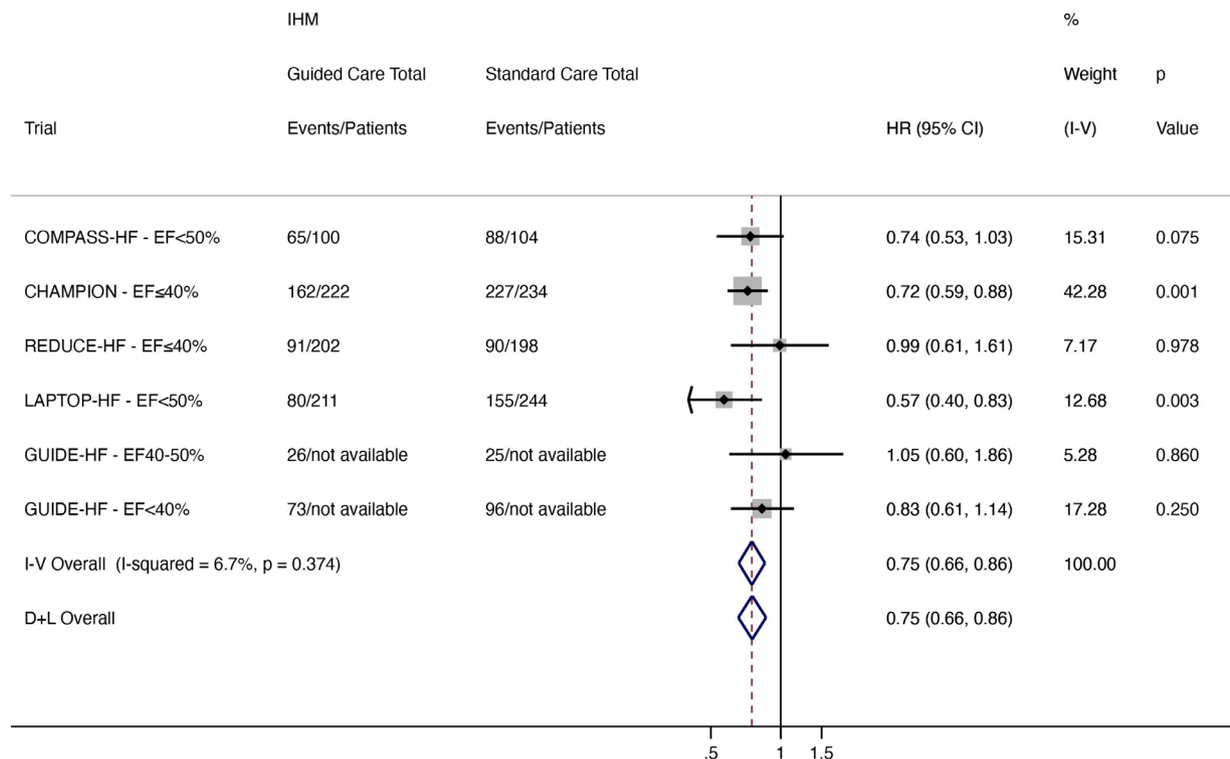
**Figure 5** Total (first and recurrent) worsening HF events (HF hospitalisation and ED and urgent clinic visit for intravenous HF therapy) in patients with HFpEF (EF  $\geq$ 50%). CHAMPION, CardioMEMS Heart Sensor Allows Monitoring of Pressure to Improve Outcomes in New York Heart Association Class III Heart Failure Patients; COMPASS-HF, Chronicle Offers Management to Patients with Advanced Signs and Symptoms of Heart Failure; D+L, DerSimonian and Laird; ED, emergency department; EF, ejection fraction; GUIDE-HF, Hemodynamic-Guided Management of Heart Failure; HF, heart failure; HFpEF, heart failure with preserved ejection fraction; IHM, implantable haemodynamic monitor; I-V, inverse variance; LAPTOP-HF, Left Atrial Pressure Monitoring to Optimize Heart Failure Therapy; REDUCE-HF, Reducing Events in Patients with Chronic Heart Failure.

Patients with an EF of  $<$ 50% have been shown to have higher resting intracardiac and pulmonary pressures than those with HFpEF, and in turn, patients with higher pressures are at greater risk of decompensation from even small rises in pressure.<sup>21–23</sup> The relative reduction of total worsening HF events with IHM monitoring observed in this meta-analysis was comparable with the magnitude of benefit found in patients with HFpEF treated with an angiotensin receptor blocker-nephrilysin inhibitor in the Prospective Comparison of ARNI With ACEI to Determine Impact on Global Mortality and Morbidity in Heart Failure (PARADIGM-HF) trial (sacubitril/valsartan reduced HF hospitalisations by 21% compared with enalapril) and the sodium glucose cotransporter 2 inhibitor dapagliflozin in the Dapagliflozin and Prevention of Adverse Outcomes in Heart Failure (DAPA-HF) trial (dapagliflozin reduced HF hospitalisations by 30% compared with placebo).<sup>24 25</sup>

Mortality data were not reported from LAPTOP-HF, and only 231 deaths were reported during the overall short average follow-up in the other four trials. With low numbers of deaths and limited follow-up periods, no IHM trial demonstrated a reduction in mortality with IHM-guided care. Accordingly, the 22% reduction in the composite endpoint all-cause mortality and HF hospitalisation with IHM-guided care observed in this analysis was driven by the favourable effect on hospitalisations. The rate of HF hospitalisations calculated for the standard care arms in individual trials in this analysis ranged from 42 per 100 patient-years in REDUCE-HF to an estimated 147 per 100 patient-years in COMPASS-HF. The reported rate of total HF hospitalisations in the standard care arm of GUIDE-HF, which investigated the only currently available IHM (CardioMEMS) in the most contemporary HF population, was 49.7 per 100 patient-years.<sup>2</sup> This was markedly higher than the composite

event rate for total HF hospitalisations and cardiovascular deaths in the placebo group of DAPA-HF (21.6 per 100 patient-years).<sup>26</sup> The substantially higher hospitalisation rates in the IHM trials highlight two considerations. First, that patients in these trials were highly selected and prognostically vulnerable. Second, rates of hospitalisation will differ between the healthcare setting in which the IHM trials were conducted (USA predominantly) and that of other diverse settings of care as indicated by event rates in more international contemporary HF trials. The effectiveness, including cost-effectiveness, of such a targeted intervention as IHM and the system of care required to deliver it may accordingly differ, depending on the setting and is an important consideration for future IHM studies (online supplemental appendix, table S3).

To date, the main source of information on the effect of IHM in patients with HFpEF has been the CHAMPION trial. In that trial, the rate of HF hospitalisation was 41 events per 100 patient-years in the IHM arm compared with 139 events per 100 patient-years in the standard care arm, giving a 70% relative risk reduction in HF hospitalisation among patients with an EF of  $\geq$ 50% when treatment was guided by IHM.<sup>19</sup> Again, the rate of HF hospitalisation was substantially higher in CHAMPION than observed in other contemporary trials of patients with HFpEF. In PARAGON-HF, the rate of HF hospitalisation and cardiovascular death in the valsartan group was 14.6 per 100 patient-years.<sup>27</sup> The reliability of the relative reduction for HF hospitalisation in patients with HFpEF reported in the CHAMPION trial is limited by the small number of patients (n=66) included in that analysis. Our new meta-analysis adds data on patients with HFpEF from COMPASS-HF and GUIDE-HF. With an additional 562 patients and 366 events, the fixed effect model demonstrated patients with HFpEF receiving IHM-guided



**Figure 6** Total (first and recurrent) worsening HF events (HF hospitalisation and ED and urgent clinic visit for intravenous HF therapy) in patients with HFrEF (EF<50%). CHAMPION, CardioMEMS Heart Sensor Allows Monitoring of Pressure to Improve Outcomes in New York Heart Association Class III Heart Failure Patients; COMPASS-HF, Chronicle Offers Management to Patients with Advanced Signs and Symptoms of Heart Failure; D+L, DerSimonian and Laird; ED, emergency department; EF, ejection fraction; GUIDE-HF, Hemodynamic-Guided Management of Heart Failure; HF, heart failure; HFpEF, heart failure with preserved ejection fraction; IHM, implantable haemodynamic monitor; I-V, inverse variance; LAPTOP-HF, Left Atrial Pressure Monitoring to Optimize Heart Failure Therapy; REDUCE-HF, Reducing Events in Patients with Chronic Heart Failure.

treatment had a 28% relative reduction in total worsening HF events. In the random-effects model, the average reduction was similar, but the CIs were wide, encompassing a potential 68% reduction to a 14% increase in such events with IHM-guided care. The difference in significance levels between models is in keeping with the high heterogeneity in the pooled sample. The effectiveness of IHM-guided treatment in patients with HFpEF remains uncertain and further trials are required with analyses in this population pre-specified in the study designs. On the available evidence, the patients who might benefit the most from IHM would have several characteristics, including a history of volume overload (as indicated by a recent HF hospitalisation), persisting symptoms and an EF of <50%.

This analysis has limitations. First, only two trials examined an IHM that is currently available (CardioMEMS), and three IHMs were examined over 18 years of investigation during which time the background management of patients with HF evolved with advancements in drug and device therapies.<sup>24 25 28–30</sup> Each IHM measured a different haemodynamic parameter. However, the IHM's haemodynamic measures were physiologically related (eg, ePAD (COMPASS-HF) provided a surrogate estimate for left atrial pressure (LAPTOP-HF)). Potential sources of bias exist. REDUCE-HF was terminated following concerns regarding 4-year pressure sensor failure in patients from other Chronicle device studies. A total of 400 patients from a recruitment target of 1300 patients had enrolled at the point of study termination. Consequently, REDUCE-HF was underpowered, with only 181 events reported compared with the 648 events expected. LAPTOP-HF was also terminated early after 1 year due to periprocedural safety concerns,<sup>11</sup> and mortality data from this

trial were not available. The meta-analysis effect estimates may have changed had both the REDUCE-HF and LAPTOP-HF trials achieved target recruitment. However, the inclusion of these studies in the meta-analysis reduced selection bias by including at least 1 year of follow-up data on clinically relevant outcomes from these RCTs. Based on patients in REDUCE-HF and LAPTOP-HF having a mean EF of  $23\% \pm 7\%$  and  $30\% \pm 15\%$ , respectively, both trials were included in the HFrEF analysis. The initial REDUCE-HF inclusion criteria also required participants to have an implantable cardioverter–defibrillator (ICD), favouring recruitment from a population with more severe HFrEF, the patient group in whom ICD implantation predominates. We cannot, however, completely exclude the possibility that some patients had EFs above these ranges. Individual cohort numbers were not available from all studies for all EF groups. We did not have individual participant level data to test the interaction between EF and IHM-guided care.

## CONCLUSIONS

IHM-guided treatment reduced total HF hospitalisation and worsening HF events. In subgroup analyses, patients with HFrEF appear to benefit from IHM-guided care, but whether the same benefit is present in patients with HFpEF remains uncertain. Further trials with pre-specified analyses of patients with an EF of  $\geq 50\%$  are required.

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