Essential components in natriuretic peptide-guided management of heart failure: an intervention synthesis

Jason Oke, Alison Clements, Julie McLellan, Clare Bankhead, Clare J Taylor, Graeme Spence, Amitava Banerjee, Rafael Perera

ABSTRACT

Aim To identify the key components of natriuretic peptide (NP)-guided treatment interventions which reduced hospitalisation in patients with heart failure (HF).

Methods and results We extracted detailed information on the components of interventions from studies of NP-guided treatment of HF identified in a previous systematic review. We used meta-regression techniques to assess univariate associations between components and the strength of the reduction in HF hospitalisations and all-cause mortality. A Bayesian meta-analysis approach was used to re-estimate study-level effects in order to identify the study with the most effective NP-guided monitoring intervention. Finally, we examined the intervention options common to the studies in which the 95% credible interval excluded no effect. We identified eight components of NP-guided treatment from ten studies. Univariate comparisons produced mainly equivocal results, but single trial choice and common components analysis led to similar conclusions. Using a predefined treatment protocol, setting a stringent NP target (N-terminal pro-B-type natriuretic peptide of 1000 pg/mL or B-type natriuretic peptide 100 pg/mL) and including a relative target were potential key components to reducing HF hospitalisations using NP-guided therapy.

Conclusion This analysis provides a description of the key components of NP-guided treatment which could help policy makers develop specific recommendations for HF management. Our research suggests that NP-guided interventions could be simplified, but more research in relevant health settings, such as primary care, is required.

INTRODUCTION

B-type natriuretic peptide (BNP) and the biologically inactive N-terminal fragment (NT-proBNP) are cardiac hormones collectively known as natriuretic peptides (NPs) that are synthesised and secreted in the ventricular myocardium. Serum concentrations are related to left ventricular filling pressures, and can be useful to discriminate between heart failure (HF) and other causes of breathlessness. Concentrations are higher in untreated or decompensated HF and fall after treatment. Therefore, NPs are a useful marker of cardiac function and treatment response.

Monitoring NP concentrations may guide decisions about when to alter the dose of medication, but evidence on the benefit of guiding therapy using plasma concentrations of NP has been conflicting. The most recent systematic review of 18 randomised controlled trials of NP-guided treatment of HF reported no evidence on reduction in all-cause mortality nor HF mortality, but did find evidence that NP-guided treatment reduced the number of hospital admissions. As with many systematic reviews, these studies implemented closely related but not identical versions of NP-guided treatment. Therefore, it is not clear which version should be implemented in clinical practice, where in the patient pathway the test should be conducted or which features of the intervention are critical for effectiveness. Current methods to guide the transition from research evidence to clinical practice are poorly developed, and clinical guidance is often not covered in the systematic review literature. One methodological approach to overcome the challenges of moving from evidence to practice is to use
an intervention synthesis as described by Glasziou et al.\textsuperscript{4} This consists of three broad approaches: (1) choosing a single trial based on criteria such as feasibility, cost and effectiveness; (2) common components hybrid which extracts and recombines components based on frequency and importance; and (3) model-guided synthesis, an approach which models the mechanisms of action to assess the importance of the intervention components. The single trial-based choice and common components hybrid method is viable for a wide range of systematic reviews, whereas the model-guided synthesis is probably only possible for large systematic reviews.\textsuperscript{4}

The aim of this study was to determine the effective intervention components which were associated with a reduction in hospitalisations for patients with HF. This could enable policy makers and practitioners to determine specific recommendations for monitoring HF.

**METHODS**

Our intervention synthesis builds on a systematic review of the effectiveness of NP-guided monitoring for HF.\textsuperscript{6} In brief, the systematic review included trials comparing management guided by serial BNP or NT-proBNP with usual care. The primary outcome in the original review was all-cause mortality with secondary outcomes of HF mortality, HF admissions, adverse events, costs and quality of life. For our intervention synthesis, we used HF hospitalisations as the primary outcome and all-cause mortality as a secondary outcome. All-cause mortality was considered as a secondary outcome because in the original systematic review NP-guided therapy was not associated with a significant reduction in all-cause mortality, whereas it was associated with a significant reduction in HF hospitalisations.

We contacted authors to obtain information regarding the interventions evaluated as part of the systematic review.\textsuperscript{6} For the purpose of the intervention synthesis, we went back to the original trial publications and attempted to gather as much detail on the components of intervention as possible. Two members of the team (AC and CB) extracted and coded the component information.

First, we examined univariate associations between each component and size of the relative risk (RR) reduction by comparing effect estimates in subcategories of each component. We compared the RRs of hospitalisation and all-cause mortality across component subgroups, using meta-regression to test for equality.\textsuperscript{9} We presented the estimated ratio of RRs with 95% CIs for each component in a figure. For defining stringency of the target, we used the lowest threshold reported in the included studies and defined stringency as 100 pg/mL (or equivalent units) for BNP and 1000 pg/mL for NT-proBNP. We converted targets in units of pmol/L for NT-proBNP using an online calculator\textsuperscript{10} and assumed that ng/L and pg/mL were equivalent (ie, 100 ng/L is the same as 100 pg/mL). We report thresholds in units of pg/mL because the majority of the studies in the review reported NP in this unit. Studies with higher targets were classed as not stringent. We looked at the frequency of monitoring across the whole length of the study (ie, titration and control phase). Studies were deemed to be ‘more frequent’ monitoring if at any time scheduled visits were at intervals less than every 3 months.

We then re-estimated the study-level RRs with 95% credible intervals using a Bayesian meta-analysis approach (see online supplementary file 1). This was to account for biases in which smaller studies report larger effect sizes.\textsuperscript{11}

We first identified the trial with the largest RR reduction (single-trial choice) following re-estimation and assessed this version of the intervention. We then separated the trials into studies that appeared to have the ‘more effective’ versions of the intervention using the criteria of the 95% credible interval. The ‘more effective’ studies had a 95% credible interval which excluded ‘no difference’ (RR=1), whereas in the ‘less effective’ studies the 95% credible interval crossed the line of ‘no difference’. Component options were selected for the new composite intervention based on which of the component choices

| Component | Description | Categorisation (where applicable) |
|-----------|-------------|----------------------------------|
| 1. Setting | Where the majority of the patients’ management took place, that is, specialist outpatient at ambulatory clinics. | 1. Specialist heart failure or cardiology clinic. 2. Non-specified outpatient or other. |
| 2. Telephone contact | Was telephone contact with a healthcare professional available to patients? | 1. Yes. 2. None specified. |
| 3. Education | Did the studies incorporate education for self-management? | 1. Yes. 2. None specified. |
| 4. Treatment protocols | Did the studies use predefined rules/ algorithms for the up titration of medication? | 1. Predefined. 2. Investigator judgement. |
| 5. Targets | Was the target NP level based on an absolute level or were relative changes to baseline considered? | 1. Absolute. 2. Incorporate relative changes. |
| 6. Stringency | How aggressive was the NP target? | 1. Less stringent. 2. Stringent. |
| 7. Trigger | Was a relative value set for treatment changes (eg, reduction by 30%)? | 1. Yes. 2. No. |
| 8. Monitoring frequency | What was the most frequent rate of monitoring (either in titration or control phase)? | 1. More frequent than every 3 months. 2. Every 3 months or longer. |

*Threshold for less/more stringent was 100 pg/ml (or equivalent units) for BNP and 1000 pg/mL for NT-proBNP.

NP, natriuretic peptide.
RESULTS
The review found 3394 references and identified 18 studies suitable for consideration. One further study has been published since the review was finished.12 HF hospitalisation rates were reported in 11 studies: Troughton et al,3 Jourdain et al,13 Anguita et al,14 Berger et al,15 Krupicka et al,16 Lainchbury et al,17 Karlström et al,18 Januzzi et al,19 Schou et al,20 Skvortsoy 201521 and Felker et al.12 Mortality outcomes were reported in six additional studies: Pfisterer et al,22 Beck-da-Silva et al,23 Persson et al,24 Eurlings et al,25 Shah et al26 and Shochat et al.27 Hospitalisation outcomes but not mortality were reported in Januzzi et al,19 and hence 16 studies were included in the secondary analysis. A flow chart showing the selection process can be found in online supplementary file 1.

We identified eight components of NP-guided treatment from the original ten studies (table 1). An evidence summary table that shows detailed descriptions of the specific components used for each study can be found in online supplementary table S1.

Components and univariate associations with hospitalisation and all-cause mortality outcomes
Component 1: setting
Twelve of the studies were based in specialist HF or cardiology clinics. In four studies, the descriptions were ambiguous or unclear, and so were classed as non-specific or other.

Component 2: telephone contact
Only three studies in total included telephone contact as part of the intervention. In the Berger et al study, the multidisciplinary care and NT-proBNP-guided intensive management arms included telephone contact with an HF nurse. Similarly, the patients enrolled in the Schou et al study had free daily access to telephone consultations with an HF nurse who was supervised by cardiologists.

Component 3: education
Six studies included some form of an educational component to their intervention. In some studies, education was aimed at promoting adherence to medication, whereas in others education was broader and promoted enhancement of self-management and lifestyle.

Component 4: treatment protocols
Eight studies used predefined (but different) treatment protocols which laid out specific instructions for
uptitratin of therapy. Six studies used investiga-

tor judgement to modify therapy in response to NP moni-

toring. The Januzzi et al 19 study justified this choice by

i stating that ‘it was believed that such an approach would

confound the concept of standard HF, which does not
typically rely on such care’. In two of the studies, it was

ot possible to ascertain if a predefined or investigator
judgement approach was used.

Component 5: NP targets
Nine studies set goals based on absolute target NP levels,
although these varied in stringency. Six studies included
a relative NP target based on baseline NP level. For
example, Skvortsov et al 21 set a goal of getting NT-proBNP
below 1000 pg/mL or 50% lower than the baseline
NT-proBNP level. Schou et al 20 did not set a target.

Component 6: target stringency
In all but one study, a single target was set. Karlström
et al 18 defined BNP targets of 150 ng/L in patients <75
years and 300 ng/L (BNP) target for patients over 75.
Five studies set ‘stringent’ targets and ten studies set ‘less
stringent’ targets. As Schou et al 20 did not set a target, we
did not include this study in this comparison.

Component 7: NP trigger
None of the studies made no mention of using a specific
trigger value of NP and set only an overall target level
NP. Seven studies were classed as having set ‘Relative’
triggers to change therapy. For example, the Berger et
al study used an increase or decrease of 30% to opti-
mise therapy. Similarly the Schou et al 20 study set a refer-
ce change value of 30% increase from baseline to take
action and mandated completion of a clinical checklist in
the absence of symptoms or signs of congestion.

Component 8: monitoring frequency
Twelve studies monitored NP more frequently than every
3 months and four studies (Anguita et al, 14 Jourdain et al,
13 Krupicka et al, 16 and Lainchbury et al 17) set moni-
toring intervals at 3 months or longer (figure 1).

Single-trial-based choice
The Skvortsov et al 21 study had the largest RR reduction in
hospital admissions and all-cause mortality using standard
meta-analysis methods (see online supplementary file 1),
but this was a fairly small study (N=76) and the effect
may be optimistic (a small study effect). The Bayesian

Table 2  HF hospitalisation: Presence or absence of component options split by studies with study-specific credible intervals
excluding the null effect (RR=1) (lower part of the table) and studies with credible intervals not excluding the null effect (upper
part of the table)

| Component options for new composite intervention are selected based on the most common option in the ‘more effective’ studies. NA, not available; NP, natriuretic peptide; RR, relative risk. |
|---|---|---|---|---|---|---|---|
| Set in heart failure clinic | Telephone contact | Educational component | Predefined treatment protocol | Incorporated relative NP target | Stringent target | Relative NP trigger value | Monitoring more frequently (<every 3 months) |
| Anguita et al 14 | ✔ | ✗ | ✗ | ✗ | ✔ | ✗ | ✔ |
| Felker et al 12 | ✔ | ✗ | ✗ | ✗ | ✔ | ✗ | ✔ |
| Karlström et al 18 | ✔ | ✗ | ✗ | ✗ | ✔ | ✗ | ✔ |
| Lainchbury et al 17 | ✗ | ✗ | ✔ | ✗ | ✗ | ✗ | ✔ |
| Schou et al 20 | ✔ | ✔ | ✔ | ✔ | ✔ | ✔ | ✔ |
| Proportion | 0.80 | 0.20 | 0.6 | 0.40 | 0 | 0.5 | 0.20 | 0.80 |
| Studies excluding, RR=1 | | | | | | | |
| Berger et al 15 | ✔ | ✔ | ✔ | ✔ | ✗ | ✗ | ✔ |
| Januzzi et al 19 | ✔ | ✗ | ✗ | ✗ | ✗ | ✗ | ✗ |
| Jourdain et al 13 | ✗ | ✗ | ✔ | ✗ | ✗ | ✗ | ✗ |
| Krupicka et al 16 | ✔ | ✗ | ✗ | ✗ | ✔ | ✗ | ✗ |
| Skvortsov et al 21 | ✔ | ✗ | ✗ | ✗ | ✔ | ✗ | ✗ |
| Troughton et al 3 | ✔ | ✗ | ✗ | ✗ | ✔ | ✗ | ✗ |
| Proportion | 0.833 | 0.167 | 0.167 | 0.6 | 0.33 | 0.667 | 0.167 | 0.5 |

Component options for new composite intervention are selected based on the most common option in the ‘more effective’ studies. NA, not available; NP, natriuretic peptide; RR, relative risk.
Table 3  All-cause mortality: presence or absence of component options split by studies with study-specific credible intervals excluding the null effect (RR=1) (lower part of the table) and studies with credible intervals not excluding the null effect (upper part of the table)

| Set in heart failure clinic | Telephone contact | Educational component | Predefined treatment protocol | Incorporated relative NP target | Stringent target | Relative NP trigger value | Monitoring more frequently (<every 3 months) | Proportion |
|-----------------------------|-------------------|-----------------------|-----------------------------|--------------------------------|------------------|------------------------|---------------------------------------------|-----------|
| Studies not excluding, RR=1 |                   |                       |                             |                                |                  |                        |                                             |           |
| Anguita et al14              | ✔                 | ✗                     | ✗                           | ✗                              | ✔                | ✗                      | ✗                                          | 0.778     |
| Beck-da-Silva et al23        | ✔                 | ✗                     | ✗                           | ✗                              | ✔                | ✗                      | ✗                                          | 0.333     |
| Berger et al15               | ✔                 | ✔                     | ✔                           | ✔                              | ✗                | ✔                      | ✗                                          | 0.555     |
| Karlström et al18            | ✔                 | ✗                     | ✗                           | ✔                              | ✗                | ✗                      | ✗                                          | 0.5      |
| Krupicka et al16             | ✔                 | ✗                     | ?                           | ✔                              | ✗                | ✔                      | ✗                                          | 0.375     |
| Lainchbury et al17           | ✗                 | ✗                     | ✔                           | ✗                              | ✗                | ✔                      | ✗                                          | 0.25      |
| Persson et al24              | ✗                 | ✗                     | ✗                           | ✔                              | ✗                | ✔                      | ✗                                          | 0.555     |
| Shochat et al27              | ✗                 | ✗                     | ✗                           | ✔                              | ✗                | ✔                      | ✗                                          | 0.778     |
| Shah et al26                 | ✔                 | ✗                     | ✗                           | ✔                              | ✗                | ✔                      | ✗                                          | 0.714     |

| Studies excluding, RR=1      |                   |                       |                             |                                |                  |                        |                                             |           |
| Eurlings et al25             | ✔                 | ✗                     | ✗                           | ✔                              | ✗                | ✔                      | ✗                                          | 0.714     |
| Felker et al12               | ✔                 | ✗                     | ✗                           | ✔                              | ✗                | ✔                      | ✗                                          | 0.142     |
| Jourdain et al33             | ✗                 | ✗                     | ✗                           | ✔                              | ✗                | ✔                      | ✗                                          | 0.667     |
| Pfisterer et al22            | ✔                 | ✗                     | ✗                           | ✔                              | ✗                | ✔                      | ✗                                          | 0.429     |
| Skvortsov et al21            | ✔                 | ✗                     | ✗                           | ✔                              | ✗                | ✔                      | ✗                                          | 0.286     |
| Troughton et al3             | ✔                 | ✗                     | ✗                           | ✔                              | ✗                | ✔                      | ✗                                          | 0.857     |

| Proposed new composite      |                   |                       |                             |                                |                  |                        |                                             |           |
| Intervention                | ✗                 | ✗                     | ✗                           | ✔                              | ✗                | ✔                      | ✗                                          |           |

Component options for the new composite intervention are selected based on the most common option in the ‘more effective’ studies. NA, not available; NP, natriuretic peptide; RR, relative risk.

‘shrinkage’ analysis, which re-estimates study-level effects, pulled back the Skvortsov et al21 study from an observed RR (95% CI) of 0.25 (0.09 to 0.67) to 0.54 (0.26 to 0.85) for hospitalisation and from 0.35 (0.12 to 0.98) to 0.78 (0.58 to 0.96) for all-cause mortality. Following re-estimation, the Skvortsov et al21 study remained the study with the largest reduction in both hospital admissions and all-cause mortality, but shrinkage analysis had the effect of making it comparable with Januzzi et al19, Jourdain et al33 and Berger et al15 (figure 2).

Patients in the Skvortsov et al21 study were managed in a specialist HF clinic. The intervention did not include telephone contact or any specialist education component. At each visit, NP measurement was considered in conjunction with clinical assessment, and specific instructions were given on the basis of clinical symptoms and NP trends. Target NP levels were stringent (NT-proBNP 1000 pg/mL), but relative changes from baseline (>50% reduction) were also included. Outpatient monitoring was monthly in controlled patients and every 2 weeks in those whose clinical condition appeared to be deteriorating. A notable feature of the Skvortsov et al21 study is that, unlike the other studies, the majority of the patients recruited into the study were New York Heart Association (NYHA) stage 4 (see online supplementary file 1).

Common components approach
After re-estimation using the Bayesian approach, in the hospitalisation outcome, six studies had credible intervals excluding RR of 1 (the ‘more effective’ interventions) and five studies had credible intervals that crossed RR of 1 (considered ‘less-effective’ versions of the interventions). For all-cause mortality, seven studies had credible intervals that excluded 1 and the remaining nine had credible intervals that crossed 1. No component options were exclusive to the more effective studies (ie, occurred...
in all of the more effective studies and in none of the less effective studies). Applying the common components approach to the hospitalisation outcome (table 2), our proposed new composite intervention would be based in a specialist clinic, would not include an education component or provide telephone contact but would use a predefined treatment protocol, aim to reduce NP to a stringent target (<1000 pg/mL), and also consider relative targets (eg, 50% reduction from baseline) but not include a relative trigger (ie, increasing doses based on short-term changes in NP). Applying the same method to the all-cause mortality outcomes (table 3), our composite intervention would be comprised of a pre-defined treatment protocol, a relative and stringent target with more frequent monitoring. Monitoring frequency was considered equivocal for the hospitalisation outcome.

Component options based on all-cause mortality were the same as for the hospitalisation outcome, with the exception of monitoring frequency. For all-cause mortality, the studies that monitored more frequently than every 3 months outnumbered the studies monitoring less frequently. Only the Skvortsov et al study included all of these specific component options.

**DISCUSSION**

In our study we have attempted to identify the key components of NP-guided monitoring for patients with chronic HF using an intervention synthesis approach. While a comprehensive systematic review and meta-analysis had shown that the use of NP can reduce the risk of hospitalisation, specific guidance on which form of the intervention could be adopted in practice was missing. Our analyses have helped reveal which components are perhaps unnecessary and which components may be essential in reducing admissions for patients with HF.

The Bayesian meta-analytic approach had the effect of reducing heterogeneity in the all-cause mortality outcome, suggesting that most of the observed variation is attributable to sampling variability as opposed to ‘real’ treatment effect modification. For HF hospitalisation, the heterogeneity was preserved after the Bayes analysis, suggesting the observed differences in intervention efficacy may be real. While univariate analyses of individual components produced mainly equivocal results, the single trial-based choice and common components analysis led to a similar conclusion; using a predefined treatment protocol and setting a stringent target alongside a relative target are potentially key components in NP-guided therapy reducing HF hospitalisations.

Current guidelines from the European Society of Cardiology (ESC) highlight the residual uncertainty about the value of NP-guided therapy and make no specific recommendations for monitoring. The National Institute for Health and Care Excellence suggests considering specialist monitoring of patients in whom uptitration is problematic or who have been admitted to the hospital. A recent individual patient data meta-analysis reported that the benefit of NP-guided therapy is confined to patients with heart failure with reduced ejection fraction (HFrEF), and although comorbidities were associated with mortality outcomes this was not the case for HF-related hospitalisations. Similarly, a meta-regression found limited evidence of effect modification around therapy types, NYHA class, and having hypertension or diabetes, but length of follow-up and age were associated with HF-related hospitalisations.

Cocco et al also did examine why only some of the many NP-guided trials demonstrated encouraging trends. They concluded that the different approaches used in the studies with particular respect to patient selection (acute vs chronic forms of HF) and the choice of NP target were important. Some of the studies examined, set targets that were too high to reduce cardiac events, which is consistent with our finding that stringency of target may be associated with reducing hospitalisation. Finally, they observed that target NP values were chosen empirically, as opposed to being evidence-based, and these levels varied significantly. As per the ESC position they also concluded that NP-guided therapy usually led to intensification of therapy, and this could potentially explain reductions in HF-related hospitalisations. We could not rule this out with the data available to us.

The results of this analysis can be generalised to a diverse patient group; all of the studies included people of older age (up to 85 years of age), while some but not all of the studies recruited patients with heart failure with preserved ejection fraction (HFrEF) as well as HFrEF. There were also patients represented from a wide spectrum of chronic HF, including asymptomatic and symptomatic forms of the disease, but not those with significant comorbidities, awaiting heart surgery or low life expectancy, who were excluded. Finally, the interventions were for the most part well described, and as part of the initial systematic review detailed information on the interventions used in the trials was obtained.

There are a number of limitations to our analysis. The initial systematic review only identified 10 studies that reported HF hospitalisation rates and 15 with all-cause mortality outcomes. Even with the addition of the most recent trial, this may be too few to be able to identify all of the ‘active’ components of most interventions. The small sample size is likely to have limited our ‘statistical power’ to detect all but the largest treatment covariate interactions. The studies that have evaluated the use of NP-guided monitoring in patients with HF are heterogeneous for factors other than the intervention components examined in this study. Therefore, a more nuanced analysis that is able to adjust for the varying demographic and clinical characteristic of the participants, outcome evaluation and assay techniques may be required to determine the optimal approach to NP-guided monitoring. This is probably only possible with a comprehensive individual patient data meta-analysis including all of the relevant studies.
By doing multiple comparisons in the univariate analyses, we run the risk that a statistical difference could occur by chance. In fact, only 1 of the 16 comparisons we made was statistically significant at the 5% level, and it therefore seems more likely that with this number of studies we are more likely to be at risk of making a type 2 error and missing real differences. In addition, the outcome does not take account of how long these hospital stays were and this could also have been important. In all of the studies contributing hospitalisation data, NP was managed in ambulatory clinics by HF specialists, and therefore does not provide any evidence that NP-guided monitoring if implemented in a primary care setting could reduce hospitalisation to the same degree. Berger et al warned that ‘deployment of their intervention using ambulatory HF specialists and home nurses may not be feasible in all health care systems, and further adaptation of this approach will be necessary in different settings’. Our analysis did not extend to considering monitoring with BNP versus NT-proBNP because in the initial review a stratified analysis comparing NT-proBNP with BNP with respect to all-cause and HF mortality, all-cause and HF admissions and quality of life showed no differences. However, there is evidence to suggest that BNP immunoassays exhibit greater between-method variability than NT-proBNP assays (coefficient of variation (CV) for BNP=13.0%, CV for NT-proBNP=8.7%), suggesting that BNP immunoassays are affected by large systematic differences in analytical performance. On this evidence alone, one would favour NT-proBNP if NP-guided monitoring included setting a stringent target.

Lastly, adverse events were reported in only a few studies, and we have not been able to assess whether any specific version of the intervention is associated with more or fewer problems.

In this analysis we have attempted to provide information about the critical features of NP-guided monitoring of chronic HF to reduce hospitalisation. While there are tools to aid the evaluation of complex interventions in trials, when it comes to meta-analysis we are reductionist and tend to focus on simple questions which may not translate to clinical practice. The approach used in this paper takes account of the complexity of NP-guided management (or therapy) as an intervention and attempts to provide practical recommendations for monitoring. While the findings of this review were unable to unequivocally determine an optimal intervention, a future intervention synthesis incorporating more data may provide clear procedural details of the essential elements of NP-guided monitoring. Further research to evaluate the clinical and cost-effectiveness of any implemented NP-guided HF monitoring within different healthcare systems is also needed.

Acknowledgements We would like to thank Paul Glassziou for his help in developing the analysis.

Contributors RP conceived and designed the research idea, CB, JM and AC acquired, extracted and coded the data for analysis. RP, CB, JM, AC and JO met regularly to refine the analysis plan and interpret the data. JO and GS performed the statistical analysis. JO drafted the manuscript. CJT and AB provided clinical input to the project and made critical revisions to the manuscript. All authors gave final approval of the manuscript.

Funding This work is supported by funding from the National Institute for Health Research (NIHR) School for Primary Care Research (project reference number 156) and the Programme Grants for Applied Research (ref: RP-PG-1210-12003). The research was partially supported by the National Institute for Health Research (NIHR) Oxford Biomedical Research Centre Programme. The views expressed are those of the author(s) and not necessarily those of the NHS, the NIHR or the Department of Health.

Competing interests None declared.

Patient consent Not required.

Provenance and peer review Not commissioned; externally peer reviewed.

Open access This is an open access article distributed in accordance with the Creative Commons Attribution 4.0 Unported (CC BY 4.0) license, which permits others to copy, redistribute, remix, transform and build upon this work for any purpose, provided the original work is properly cited, a link to the licence is given, and indication of whether changes were made. See: http://creativecommons.org/licenses/by/4.0/.

REFERENCES

1. Weber M, Hamm C. Role of B-type natriuretic peptide (BNP) and NT-proBNP in clinical routine. Heart 2006;92:843–9.
2. Roberts E, Ludman AJ, Dzworyński K, et al. The diagnostic accuracy of the natriuretic peptides in heart failure: systematic review and diagnostic meta-analysis in the acute care setting. BMJ 2015;350:h910–h910.
3. Troughton RW, Frampton CM, Yardley TG, et al. Treatment of heart failure guided by plasma amino-terminal brain natriuretic peptide (N-BNP) concentrations. Lancet 2000;355:1126–30.
4. Vasan RS. Expanding indications for natriuretic peptides: importance of better new (research) protocols. Am Heart J 2004;148:743–6.
5. Porapakkham P, Porapakkham P, Zimmet H. B-type natriuretic peptide-guided therapy. Arch Intern Med 2010;170:507.
6. McLellan J, Heneghan CJ, et al. Cochrane Heart Group. B-type natriuretic peptide-guided treatment for heart failure, Perera R, Cochrane Database of Systematic Reviews. 135. Chichester, UK: John Wiley & Sons, Ltd. 2016.
7. Hoffmann TC, Glassziou PP, Boutron I, et al. Better reporting of interventions: template for intervention description and replication (TIDieR) checklist and guide. BMJ 2014;348:g1687.
8. Glassziou PP, Chalmers I, Green S, et al. Intervention synthesis: a missing link between a systematic review and practical treatment(s). PLoS Med 2014;11:e1001690.
9. Borenstein M, Higgins JP. Meta-analysis and subgroups. Prev Sci 2013;14:134–43.
10. Unitilsa.com. 2017. NT-proBNP (N-terminal pro B-type natriuretic peptide) http://unitilsa.com/node/163 (cited 22 May 2017).
11. Rücker G, Carpenter JR, Schwarzer G. Detecting and adjusting for small-study effects in meta-analysis. Biom J 2011;53:351–68.
12. Felker GM, Anstrom KJ, Adams KF, et al. Effect of natriuretic peptide-guided therapy on hospitalization or cardiovascular mortality in high-risk patients with heart failure and reduced ejection fraction: a randomized clinical trial. JAMA 2017;318:713.
13. Jourdain P, Jondeau G, Furcht F, et al. Plasma brain natriuretic peptide-guided therapy to improve outcome in heart failure: the STARS-BNP Multicenter Study. J Am Coll Cardiol 2007;49:1733–9.
14. Anguita M, Esteban F, Castillo JC, et al. Utilidad de los valores del péptido natriurético cerebral frente al control clínico habitual para la monitorización del tratamiento en pacientes con insuficiencia cardiaca. Medicina Clinica 2010;135:435–40.
15. Berger R, Moore D, Peter S, et al. N-terminal pro-B-type natriuretic peptide-guided, intensive patient management in addition to multidisciplinary care in chronic heart failure 3-arms, prospective, randomized pilot study. J Am Coll Cardiol 2010;55:645–53.
16. Krupicka J, Janota T, Hradec J. Optimization of heart failure therapy guided by plasma BNP concentrations. European Society of Cardiology 2010.
17. Lainchbury JG, Troughton RW, Strangman KM, et al. N-terminal pro-B-type natriuretic peptide-guided treatment for chronic heart failure: results from the BATTLESCARRED (NT-proBNP-Assisted Treatment To Lessen Serial Cardiac Readmissions and Death) trial. J Am Coll Cardiol 2009;55:53–60.
18. Karlström P, Alehagen U, Boman K, et al. Brain natriuretic peptide-guided treatment does not improve morbidity and mortality in extensively treated patients with chronic heart failure: responders to treatment have a significantly better outcome. *Eur J Heart Fail* 2011;13:1096–103.

19. Januzzi JL, Rehman SU, Mohammed AA, et al. Use of amino-terminal pro-B-type natriuretic peptide to guide outpatient therapy of patients with chronic left ventricular systolic dysfunction. *J Am Coll Cardiol* 2011;58:1881–9.

20. Schou M, Gustafsson F, Videbaek L, et al. Adding serial N-terminal pro brain natriuretic peptide measurements to optimal clinical management in outpatients with systolic heart failure: a multicentre randomized clinical trial (NorthStar monitoring study). *Eur J Heart Fail* 2013;15:818–27.

21. Кошкина ДЕ, Протасов ВН, Нарусов ОЮ. Терапия регресса сердечной недостаточности, 2015.

22. Pfisterer M, Buser P, Rickli H. BNP-guided vs symptom-guided heart failure therapy. *JAMA* 2009;301:383.

23. Beck-da-Silva L, de Bold A, Fraser M, et al. BNP-guided therapy not better than expert’s clinical assessment for beta-blocker titration in patients with heart failure. *Congest Heart Fail* 2005;11:248–55.

24. Persson H, Erntell H, Eriksson B, et al. Improved pharmacological therapy of chronic heart failure in primary care: a randomized Study of NT-proBNP Guided Management of Heart Failure—SIGNAL-HF (Swedish Intervention study—Guidelines and NT-proBNP Analyis in Heart Failure). *Eur J Heart Fail* 2010;12:1300–8.

25. Eurlings LW, van Pol PE, Kok WE, et al. Management of chronic heart failure guided by individual N-terminal pro-B-type natriuretic peptide guided therapy of chronic heart failure IMprove heart failure morbidity and mortality?). *J Am Coll Cardiol* 2010;56:2090–100.

26. Shah MR, Calif RM, Nohria A, et al. The STARBRITE trial: a randomized, pilot study of B-type natriuretic peptide-guided therapy in patients with advanced heart failure. *J Card Fail* 2011;17:615–21.

27. Shochat M, Shotan A, Dahan I. NT-proBNP-Guided preemptive treatment of outpatients with chronic heart failure followed in a outpatient clinic.Clinic. *J Card Fail* 2011;17:S56.

28. McMurray JJ, Adamopoulos S, Anker SD, et al. ESC Guidelines for the diagnosis and treatment of acute and chronic heart failure 2012. *Eur J Heart Fail* 2012;14:803–69.

29. National Institute for Health and Care Excellence. 2010.Chronic heart failure in adults:management https://www.nice.org.uk. 2010.

30. Brunner-La Rocca HP, Eurlings L, Richards AM, et al. Which heart failure patients profit from natriuretic peptide guided therapy? A meta-analysis from individual patient data of randomized trials. *Eur J Heart Fail* 2015;17:1252–61.

31. Savarese G, Trimarco B, Dellegrottaglie S, et al. Natriuretic peptide-guided therapy in chronic heart failure: a meta-analysis of 2,686 patients in 12 randomized trials. *PloS One* 2013;8:e68287.

32. Cocco G, Jerie P. Assessing the benefits of natriuretic peptides-guided therapy in chronic heart failure: a meta-analysis of 2,686 patients in 12 randomized trials. *Clin Chim Acta* 2012;414:112–9.

33. Campbell NC, Murray E, Darbyshire J, et al. Designing and evaluating complex interventions to improve health care. *BMJ* 2007;334:455–9.