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Diagnostic accuracy of natriuretic peptide screening for left ventricular systolic dysfunction in the community: systematic review and meta-analysis

Clare R. Goyder^{1*}, Andrea K. Roalfe¹, Nicholas R. Jones¹, Kathy S. Taylor¹, Charles D. Plumptre², Olivia James³, Thomas R. Fanshawe¹, F D Richard Hobbs¹ and Clare J. Taylor¹

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Abstract

Aims Heart failure (HF) is a global health burden and new strategies to achieve timely diagnosis and early intervention are urgently needed. Natriuretic peptide (NP) testing can be used to screen for left ventricular systolic dysfunction (LVSD), but evidence on test performance is mixed, and international HF guidelines differ in their recommendations. Our aim was to summarize the evidence on diagnostic accuracy of NP screening for LVSD in general and high-risk community populations and estimate optimal screening thresholds.

Methods We searched relevant databases up to August 2020 for studies with a screened community population of over 100 adults reporting NP performance to diagnose LVSD. Study inclusion, quality assessment, and data extraction were conducted independently and in duplicate. Diagnostic test meta-analysis used hierarchical summary receiver operating characteristic curves to obtain estimates of pooled accuracy to detect LVSD, with optimal thresholds obtained to maximize the sum of sensitivity and specificity.

Results Twenty-four studies were identified, involving 26 565 participants: eight studies in high-risk populations (at least one cardiovascular risk factor), 12 studies in general populations, and four in both high-risk and general populations combined. For detecting LVSD in screened high-risk populations with N-terminal prohormone brain natriuretic peptide (NT-proBNP), the pooled sensitivity was 0.87 [95% confidence interval (CI) 0.73–0.94] and specificity 0.84 (95% CI 0.55–0.96); for BNP, sensitivity was 0.75 (95% CI 0.65–0.83) and specificity 0.78 (95% CI 0.72–0.84). Heterogeneity between studies was high with variations in positivity threshold. Due to a paucity of high-risk studies that assessed NP performance at multiple thresholds, it was not possible to calculate optimal thresholds for LVSD screening in high-risk populations alone. To provide an indication of where the positivity threshold might lie, the pooled accuracy for LVSD screening in high-risk and general community populations were combined and gave an optimal cut-off of 311 pg/mL [sensitivity 0.74 (95% CI 0.53–0.88), specificity 0.85 (95% CI 0.68–0.93)] for NT-proBNP and 49 pg/mL [sensitivity 0.68 (95% CI 0.45–0.85), specificity 0.81 (0.67–0.90)] for BNP.

Conclusions Our findings suggest that in high-risk community populations NP screening may accurately detect LVSD, potentially providing an important opportunity for diagnosis and early intervention. Our study highlights an urgent need for further prospective studies, as well as an individual participant data meta-analysis, to more precisely evaluate diagnostic accuracy and identify optimal screening thresholds in specifically defined community-based populations to inform future guideline recommendations.

Keywords Biomarkers; Screening; Left ventricular systolic dysfunction; Natriuretic peptides

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Introduction

Approximately 40 million people worldwide are living with heart failure (HF), representing a major public health burden,^{1,2} but treatment can improve quality of life and survival and reduce hospitalizations.^{3,4} Detecting HF in the community, especially in the early stages, to achieve timely diagnosis is an urgent research priority.^{5–7} There is reliable evidence that treatment with renin–angiotensin–aldosterone system (RAAS) inhibitors, titrated to the appropriate dose, can limit progression from left ventricular systolic dysfunction (LVSD) to HF.^{8–11} Screening is one potential route to detect LVSD and provides an important opportunity for early intervention.

Echocardiography, and more recently cardiac magnetic resonance imaging (MRI), can accurately identify patients with LVSD, but these strategies are not feasible as general population screening tools due to poor cost-effectiveness. One potential screening strategy is an initial natriuretic peptide (NP) blood test with subsequent echocardiography or MRI performed only in patients with raised NP levels. There is evidence supporting the role of NP testing in the diagnosis of HF in symptomatic patients and limited evidence in treatment optimization.^{12,13} However, the St Vincent's Screening to Prevent Heart Failure (STOP-HF) trial was the first randomized controlled trial to indicate that an NP-guided screening approach in asymptomatic people, linked to targeted prevention, reduced the progression of asymptomatic LVSD and the development of HF in high-risk groups.¹³

Globally, HF guidelines differ in their recommendations on NP screening to detect LVSD: European guidelines do not currently advocate screening, whereas North American guidelines recommend NP screening in high-risk patients but do not specify optimal NP screening thresholds or make any recommendation for general populations.4,14-17 Moreover, the overall performance of NP as a screening tool for LVSD remains unclear. How NP performs as a screening test across different populations (e.g. high-risk vs. general community populations) and what NP threshold is appropriate for biomarker-based screening is also uncertain. Previous systematic reviews have focused on the accuracy of NP as a diagnostic tool for HF in symptomatic, presenting patients,^{17,18} or analysed accuracy in combined diagnostic and screening studies.^{18,19} Those reviews that have assessed NP performance in a screening context have often combined primary and secondary care studies, rather than focussing specifically on screening in the community.^{18,20,21}

The aim of this study was to provide an up-to-date summary of the accuracy of NP screening for LVSD, considering both brain natriuretic peptide (BNP) and N-terminal prohormone BNP (NT-proBNP) compared with echocardiography and cardiac MRI in community populations, and to determine the optimal NP screening threshold.

Methods

This systematic review and meta-analysis protocol has been prospectively registered in PROSPERO (registration number: CRD42018087498) and separately published.²² This review has been produced in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-analyses (PRISMA) guidelines and recommendations from the Cochrane Collaboration.²³ The full search strategy is available in *Appendix S1*.

This study was discussed with our patient and public involvement group, who agreed that improving the detection of HF was a research priority. The group was familiar with the consequences of delayed diagnosis, some with personal experience, and welcomed more research on screening.

Search methodology

We searched Ovid Medline, Embase, Cochrane Database of Systematic Reviews and Cochrane Central Register of Controlled Trials, Cochrane CENTRAL, DARE, and Science Citation Index from inception on 17 April 2019 and updated the search on 13 August 2020. We did not employ a study design filter or add any language restriction. We searched reference lists to identify more publications. Screening was carried out by two reviewers independently (CG and either CP, NJ, or OJ). Disagreements were resolved by discussion or referral to a third reviewer (CT).

Screening methodology

We included studies of over 100 adult participants in a community setting to limit bias from small studies. Included studies compared the performance of NP testing with either echocardiography or cardiac MRI for the detection of LVSD. We included studies that recruited screened community populations. Community screening may identify patients with preclinical HF, such as asymptomatic LVSD, also termed stage B HF. Screening may also detect patients with clinical HF who had not been appropriately diagnosed previously. Given this, we took a pragmatic approach and included all patients who had participated in screening studies.

We excluded studies of patients recruited through secondary care, such as cardiology clinics, and studies presenting insufficient data to construct 2×2 tables. We also excluded duplicate datasets, selecting the papers that most closely aligned with inclusion criteria or were most recently published. The inclusion and exclusion criteria for participants in included studies are found in *eTable 1*.

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Target condition

Consistent with the overall aim of community screening, LVSD was defined broadly by evidence of reduced ejection fraction (EF) including either quantitative or narrative descriptions of reduced systolic function and/or other echocardiographic parameters. An EF cut-off of 40% (or nearest) was selected in studies reporting results for more than one EF threshold.

Data extraction and quality assessment

Data extraction was performed independently by two reviewers (CG and either CP, NJ, or OJ).²² A risk of bias template incorporating QUADAS-2 criteria was used to assess methodological quality.²⁴ Disagreements were discussed or referred to a third reviewer (CT). Diagnostic accuracy data were extracted from all studies,²⁵ and 2 × 2 tables were constructed in accordance with current reporting guidelines, at all NP thresholds reported.²⁶

Statistical analyses

Meta-analysis was performed where there were at least four studies with available data. Subgroup analyses were performed for both types of NP (to include BNP and NT-proBNP) as well as high-risk and general populations. High-risk populations were defined as having at least one cardiovascular (CV) risk factor or ischaemic heart disease (IHD) or were selected non-general populations, such as cohorts of nursing home residents or patients with chronic obstructive pulmonary disease (COPD), as the overlap between COPD and HF is known to be high.²⁷

To visually explore the variation in diagnostic accuracy, sensitivity and specificity forest plots with 95% confidence intervals (CIs) for both types of NP, ordered by each threshold reported, were produced in RevMan 5.3.28 Where data included multiple thresholds. R version 3.5.3 (diaameta package)²⁹ was used to produce SROC curves in relation to the positivity threshold with estimation of the single threshold that maximized the sum of sensitivity and specificity.³⁰ A logistic distribution for threshold within the diseased and non-diseased groups was assumed, and NP levels were log-transformed due to skewness. If there were insufficient data to generate SROCs with multiple thresholds, hierarchical SROC curves with 95% CI and prediction regions were drawn (Stata version 15.0, metandi command) using the lowest threshold for each study.³¹ Some studies did not report data for the combined group of men and women together with the threshold used to define positivity. Therefore, for our primary analysis, we decided to include all the available

data for men and totals (i.e. men and women combined if reported, and otherwise men only). Sensitivity analysis was carried out to analyse the available data on women and totals (i.e. men and women combined if reported, and otherwise women only) to explore whether there were any differences that resulted from this analysis decision. Sensitivity analyses were also carried out to compare studies that excluded participants with a previous diagnosis of LVSD with studies that did not and to examine whether there was any difference in accuracy in studies that described participants as entirely asymptomatic compared with other included studies.

Results

From 3131 records (*Figure 1*), 24 studies presented accuracy data for NP screening to detect LVSD, involving 26 565 participants^{32–55}; all included studies were cross-sectional. The included studies, with population characteristics and the lowest threshold reported, are summarized in *Table 1*. Full data for all thresholds including prevalence are available in *eTable 2*. Details of inclusion and exclusion criteria are listed in *eTable 1*; included studies were published over 16 years between 1998 and 2014.

Characteristics of included studies

Eight studies included data on only high-risk populations,^{32–39} 12 included only general populations,^{37,40–50} and four included data on high-risk and general populations.^{51–54} Nine studies assessed NT-proBNP only,^{32–34,40,43,46,47,51,54} and nine studies assessed BNP only; one of these used a point-of-care BNP test, Biosite[®],³⁵ whereas all other tests were processed in a laboratory. Six studies evaluated performance in both biomarkers.^{37,39,42,43,48,52} All studies used echocardiography as the comparator except one that used cardiac MRI.⁴⁵ Subgroup sample sizes corresponding to each meta-analysis were smaller for high-risk populations with NT-proBNP (n = 4193) and BNP (n = 2940), with larger groups seen in the general populations, with NT-proBNP (n = 13 416) and BNP (n = 12970).

Studies differed in the inclusion criteria for the ages of participants they recruited. General populations included participants of younger ages, whereas high-risk populations were older (as summarized in *Table 1*).

Most included studies classified LVSD as reduced EF of either <40% or <50%. In two studies, the outcomes of diastolic dysfunction and LVSD were combined, 32,40 and three studies used narrative descriptions of LVSD rather than quantitative measurements of EF. 32,41,42





The reported prevalence for studies presenting accuracy data for LVSD in total populations ranged from 0.4% to 12.9% (*eTable 2*).

There was some variability in how individual studies defined high-risk populations. Some studies categorised any participant who had more than one CV risk factor as high risk.^{37,38} One only included patients who had IHD in a high-risk group,⁵³ whereas another included post-myocardial infarction (MI) or IHD patients in combination with other risk factors.⁴⁷ There were no included studies that included cohorts of COPD patients; one high-risk study recruited from nursing homes.³⁹

Only seven studies described the population as completely asymptomatic.^{32,33,36–38,42,54} One study described the population as 'mostly asymptomatic'.⁵⁵ McDonagh et al. described the population as 50% symptomatic and 50% asymptomatic.⁵³ Of the remaining studies, the majority recruited randomly selected community populations and did not provide details of whether the participants reported any symptoms. Some described their population cohorts as 'healthy' without any data to explain this.⁴⁵ Two studies described participants as having a high proportion of symptomatic participants^{35,39} and one study recording only 22.2% as symptomatic.⁴³

| Abhayaratna 2006§Canberra, AustraliaNT-proBNPBetti 2009§Unclear, ItalyNT-proBNPBetti 2003§Valencia, SpainNT-proBNPCostello-BoerrigterOlmsted County,BNP2006Minnesota, USANT-proBNP2006Dallas, Texas, USANT-proBNPBasko 2009Dallas, Texas, USANT-proBNPGalasko 2005London, UKNT-proBNPGalasko 2005London, UKNT-proBNPGavazzi 2014Lombardy, ItalyNT-proBNPGoetze 2006Copenhagen, DenmarkNT-proBNPGroenning 2004Louisiana, USABNPHebert 2010Louisiana, USABNPHeberg 2004England, UKBNPHobbs 2004England, UKBNPHobbs 2004England, UKBNPHobbs 2004England, UKBNP | SNP 1229 SNP 1012 SNP 203 SNP 203 SNP 2429 SNP 734 SNP 734 523 SNP 672 SNP 672 145 145 133 | $60-74 \\ > = 67 \\ = 67 \\ 49-81 \\ <65 \\ <65 \\ <65 \\ -864 \\ 18-64 \\ 18-64 \\ -8-$ | General population High-risk General population General population General population General population High-risk High-risk General population General population | 151 125 12 12 32 32 7.1 28.9 28.9 44 144 144 | EF ≤ 40% and advanced DD-NEF Pooled systolic and moderate-severe DDf LVEF ≤50% LVEF ≤40% /LVEF ≤50% LVSD EF < 55 or LVH†† |
|---|--|---|---|---|--|
| Cosin Aguilar 2003Valencia, SpainNT-proBNPCostello-BoerrigterOlmsted County,BNP2006Minnesota, USANT-proBNP2006Minnesota, USANT-proBNPDe Lemos 2009Dallas, Texas, USANT-proBNPGalasko 2005London, UKNT-proBNPGavazzi 2014Lombardy, ItalyNT-proBNPGoetze 2006Copenhagen, DenmarkNT-proBNPGroenning 2004Louisiana, USABNPHebert 2010Louisiana, USABNPHobbs 2004England, UKBNPHobbs 2004England, UKBNPHobbs 2004England, UKBNP | SNP 203 SNP 203 SNP 1869 SNP 2429 SNP 734 761 672 SNP 672 145 145 133 | 49–81 <65 <65 18–64 18–64 18–64 18–64 55–80 55–80 >55–80 >55–80 >55–80 25–80 25–80 25–80 25–80 25–80 25–80 25–80 25–80 25–80 25–80 25–80 25–80 25–80 25–80 25–80 25–80 26–80 27–80 27–80 27–80 27–80 27–80 27–80 27–80 27–80 27–80 27–80 27–70 2 | General population General population General population General population High-risk High-risk General population General population | 215 32 7.1 28.9 44 44 45 | LVEF <50% LVEF <40% /LVEF <50% LVSD EF < 55 or LVH†† |
| De Lemos 2009 Dallas, Texas, USA NT-proBNP De Lemos 2009 Dallas, Texas, USA NT-proBNP Galasko 2005 London, UK NT-proBNP Gavazzi 2014 Lombardy, Italy NT-proBNP Goetze 2006 Copenhagen, Denmark NT-proBNP Groenning 2004 Louisiana, USA BNP Hebert 2010 Louisiana, USA BNP Hobbs 2004 England, UK BNP | SNP 1869 SNP 2429 SNP 2429 SNP 761 761 761 761 761 761 761 761 761 761 | <65 18-64 18-64 > = 45 > = 45 55-80 > 55-80 > = 50 40-75 75 | General population General population General population High-risk High-risk General population General population | 32 7.1 4 + + 4 55 | LVSD EF $<$ 55 or LVH†† |
| Galasko 2005 London, UK NT-proBNP Gavazzi 2014 Lombardy, Italy NT-proBNP Goetze 2006 Copenhagen, Denmark NT-proBNP Groenning 2004 Copenhagen, Denmark NT-proBNP Hebert 2010 Louisiana, USA BNP Hobbs 2004 England, UK BNP MT Dobbs 2004 England, UK BNP | 3NP 2429 3NP 734 3NP 761 3NP 623 3NP 672 3NP 672 407 133 | 18-64 > = 45 55-80 55-80 > = 45 55-80 > = 45 > = 45 > = 45 > = 50 40-75 75 | General population General population High-risk General population General population | 28.9 ++ 14 551 28.9 | |
| Gavazzi 2014 Lombardy, Italy NT-proBNP Gavazzi 2014 Lombardy, Italy NT-proBNP Groenning 2004 Copenhagen, Denmark NT-proBNP Groenning 2004 Copenhagen, Denmark NT-proBNP Hebert 2010 Louisiana, USA BNP Hedberg 2004 England, UK BNP Hobbs 2004 England, UK BNP | 3NP 761 3NP 761 3NP 761 8197 672 407 133 | <pre>> = 45 55-80 55-80 > 55-80 > = 50 40-75 75</pre> | High-risk High-risk General population General population | -++ 6 K 4 R | 11/FE /50% |
| Goetze 2006 Copenhagen, Denmark NT-proBNP Groenning 2004 Copenhagen, Denmark NT-proBNP Hebert 2010 Louisiana, USA BNP Hedberg 2004 England, UK BNP Mobbs 2004 England, UK BNP | anp 3497 3NP 3497 672 145 145 133 133 | >= 55 >= 50 40-75 75 | General population General population Hinh risk | - 1 271 | LVEF < 30% LVEF < 40% |
| Goetze 2006 Copenhagen, Denmark NT-proBNP Groenning 2004 Copenhagen, Denmark NT-proBNP Hebert 2010 Louisiana, USA BNP Hedberg 2004 England, UK BNP MT 2004 England, UK BNP | 3NP 3497 3NP 672 145 407 133 | >55 > = 50 > = 40-75 75 | General population General population Hich risk | 144 351 | LVEF <45% LVEF <45% LVFF <50% |
| Hebert 2010 Louisiana, USA BNP Hedberg 2004 Louisiana, USA BNP Hobbs 2004 England, UK BNP | 145 407 133 133 | 40-75 75 | Hinh rick | | LVEF <40% LVEF <40% |
| Hebert 2010 Louisiana, USA BNP Hedberg 2004 Västerås, Sweden BNP Hobbs 2004 England, UK BNP | 145 407 133 | 40–75 75 | Hinh risk | - | LVSD LVEF <40% LVSD LVEF <45% |
| Hobbs 2004 Västerås, Sweden BNP BNP Hobbs 2004 England, UK BNP NT 2004 | 407 133 207 | 75 | | θŪ | LVSD LVEF <50% |
| Hobbs 2004 England, UK BNP BNP | 133 | | General population | 28 | LVSD based on wall motion index |
| | | > = 45 | Hiah risk | 103.8 | corresponding to LVEF <40% LVSD<40% |
| | 100 | > = 45 | General population | 103.8 | LVSD<40% |
| | 3NP 133 | > = 45 | High risk | 338 | LVSD <40% |
| I choc Boisrano 2012 Madrid Snain BND BND | 3NP 307 | < 45 | General population | 338 71 | LVSD <40% |
| Luchner 2000 Augsburg, Germany BNP | 672 | 50-67 | General population | 34 | LVD (defined as fractional |
| Lines 2010 Goodtingon Gormany BND | C 7 J | 07/ | Lizh rick | 105 | shortening <28%)/general population |
| Lukowicz 2005 Augsburg, Germany BNP | 1678 | 25-75 | General population | 27 | LVSD (EF < 40%) |
| Mason 2013 North-East England, UK BNP | 392 | VI 7 | High risk | 145 | LVSD $EF < 50\%$ |
| McDonach 1008 Classon IIV BND | 3NP 393 1757 | ל0 | High risk General population | 1000 | 2000 - 2000 |
| | 1252 | 25-74 25-74 | derrerar population High risk | 17.9 | |
| Mureddu 2013 Lazio region, Italy NT-proBNP | 3NP 435 | 65–84 67 04 | High risk | 278 | LVSD EF < 50 |
| MI-ProBNP Mi | 2017 JUS | 65-84 | General population اتعلہ ہندار | 8/7 | |
| | 014 | >40 | ліуп-пуп | 70 | LVSD < 40% |
| Ng 2003 Leicestershire, UK BNP | 1331 | 45-80 | General population | 66 | Definite systolic $EF \le 35\%$ |
| | 5NP 1331 | 45-80 | | α18 7.13 | |
| Smith 2000 Poole, Dorset, UK BNP | CC 1 | //< | General population | 64.7 | Reduced left ventricular ejection fraction/LVSD confirmed by echo |
| Vasan 2002 Framingham, BNP Massachusetts, USA | 3177 | >40 | General population | 21 | |

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Quality of included studies

The majority of studies had low or unclear risk of bias with <20% of ratings in the high-risk category (*eFigure 1*, *eFigure 2*). Studies recorded with high risk for patient selection generally excluded patients who had a previous MI even though this patient group may benefit from screening for LVSD.^{32,42,45} In reporting the index test, some studies presented incomplete data.^{44,49,55} Not all studies performed the reference test blinded to the index test,⁵⁵ and some failed to give information on how echocardiography was conducted.⁵³

Accuracy of NP to screen for LVSD in high-risk populations

Most studies of high-risk populations demonstrated a trade-off between sensitivity and specificity as shown in the forest plots for NT-proBNP and BNP (*Figure 2A*). For NT-proBNP (*Figure 3A*), the pooled sensitivity was 0.87 (95% CI 0.73–0.94) and specificity 0.84 (95% CI 0.55–0.96) for detecting LVSD in screened high-risk populations. For BNP in high-risk populations (*Figures 2B and 3B*) the pooled sensitivity was 0.75 (95% CI 0.65–0.83) and specificity 0.78 (95% CI 0.72–0.84).

Accuracy of NP to screen for LVSD in general populations

There was extensive variability in sensitivity and specificity for studies of general populations (*eFigure 3*), some of which was due to differences in reported threshold. For NT-proBNP in general populations (*eFigure 4*), the pooled sensitivity was 0.72 (95% CI 0.42–0.90) with specificity 0.82 (95% CI 0.60–0.93), and the optimal threshold was 274 pg/mL. For BNP in general populations, the pooled sensitivity was 0.62 (95% CI 0.32–0.85) with specificity 0.83 (95% CI 0.61–0.94) and optimal threshold 46 pg/mL (*eFigure 1*).

Optimal screening thresholds

It was not possible to calculate optimal thresholds for NP screening for LVSD in only high-risk populations as there were not enough studies of this population that provided data at multiple thresholds. However, the pooled accuracy of NT-proBNP in high-risk and general community populations combined (*Figure 4*) gave an optimal cut-off of 311 pg/mL with sensitivity of 0.74 (95% CI 0.53–0.88) and specificity 0.85 (95% CI 0.68–0.93). The pooled accuracy data for BNP (*eFigure 6*) yielded an optimal screening threshold for the detection of LVSD at 49 pg/mL with a sensitivity of 0.68 (95% CI 0.45–0.85) and a specificity of 0.81 (0.67–0.90).

Figure 2 (A) Paired sensitivity and specificity plot for NT-proBNP compared with echocardiography for detecting LVSD in screened high-risk populations. Thresholds are measured in pg/mL, and studies are ordered by NP threshold. Threshold were age specific for Gavazzi et al., based on 95th percentiles (data not provided) and for Galasko et al., based on 97.25th percentiles (100 pg/mL males aged 45–59; 164 pg/mL females aged 45–59; 172 pg/mL males aged > = 60; 225 pg/mL females age> = 60). (B) Paired sensitivity and specificity plot for BNP compared with echocardiography for detecting LVSD in screened high-risk populations. Thresholds are measured in pg/mL, and studies are ordered by NP threshold.

| Study | TP | FP | FN | TN | Age gro | up Thre | shold Ser | sitivity (95% CI) | Specificity (95% CI) | Sensitivity (95% CI) | Specificity (95% CI) |
|------------------|------|-----|------|------|---------|---------|-----------|-------------------|------------------------|--------------------------|----------------------|
| Gavazzi 2014 | 6 | 0 | 3 | 610 | 55- | 80 | + | 0.67 [0.30, 0.93] | 1.00 [0.99, 1.00] | | |
| Galasko 2005 | 36 | 223 | 5 | 497 | 2 | 45 | * | 0.88 [0.74, 0.96] | 0.69 [0.66, 0.72] | | |
| Luers 2010 | 15 | 62 | 8 | 457 | > | 40 | 105.0 | 0.65 [0.43, 0.84] | 0.88 [0.85, 0.91] | | |
| Betti 2009 | 51 | 192 | 1 | 768 | 2 | 87 | 125.0 | 0.98 [0.90, 1.00] | 0.80 [0.77, 0.82] | - | |
| Goode 2007 | 27 | 219 | 5 | 176 | > | 50 | 150.0 | 0.84 [0.67, 0.95] | 0.45 [0.40, 0.50] | | • |
| Mureddu 2013 | 6 | 82 | 0 | 347 | 65- | 84 | 278.0 | 1.00 [0.54, 1.00] | 0.81 [0.77, 0.84] | | • |
| Hobbs 2004 | 10 | 66 | 0 | 57 | 2 | 45 | 338.0 | 1.00 [0.69, 1.00] | 0.46 [0.37, 0.56] | | |
| Mason 2013 | 25 | 86 | 9 | 273 | 2 | 65 1 | 0.000 | 0.74 [0.56, 0.87] | 0.76 [0.71, 0.80] | | |
| | | | | | | | | | | 0 0.2 0.4 0.6 0.8 1 | 0 0.2 0.4 0.6 0.8 1 |
| | | | | | | | | (-) | | | |
| | | | | | | | | (A) | | | |
| | | | | | | | | | | | |
| Study | | TP | F | P FN | TN A | e group | Threshold | Sensitivity (95% | CI) Specificity (95% (| CI) Sensitivity (95% CI) | Specificity (95% CI) |
| McDonagh 1998 | | 118 | 5 26 | 7 22 | 847 | 25-74 | 17.9 | 0.84 [0.77, 0.9 | 0.76 [0.73, 0.7 | 9] | |
| Murtagh 2012a | | 8 | 18 | 2 3 | 618 | >40 | 50.0 | 0.73 [0.39, 0.9 | 0.77 [0.74, 0.8 | oj ——— | |
| Hebert 2010 | | 11 | 1 | 95 | 110 | 40-75 | 60.0 | 0.69 [0.41, 0.8 | 0.85 [0.78, 0.9 | 1] | |
| Lobos Bejarano 2 | 2012 | 13 | 3 5 | 2 4 | 123 | >63 | 71.0 | 0.76 (0.50, 0.9 | 93] 0.70 [0.63, 0.7 | 7] | - |
| Hobbs 2004 | | 5 | 5 4 | 1 5 | 82 | ≥45 | 103.0 | 0.50 (0.19, 0.8 | 31] 0.67 (0.58, 0.7 | 5] | |
| Luers 2010 | | 14 | 5 | 2 9 | 467 | >40 | 105.0 | 0.61 (0.39, 0.8 | 0.90 [0.87, 0.9 | 2] | |
| Mason 2013 | | 28 | 5 9 | 0 8 | 268 | ≥65 | 145.0 | 0.76 (0.59, 0.8 | 39] 0.75 (0.70, 0.7 | | |
| | | | | | | | | | | 0 0.2 0.4 0.6 0.8 1 | 0 0.2 0.4 0.6 0.8 1 |
| | | | | | | | | | | | |

(B)



Figure 3 (A) SROC curves of NT-proBNP compared with echocardiography for detecting LVSD in screened high-risk populations. (B) SROC curves of BNP compared with echocardiography for detecting LVSD in screened high-risk populations.

Sensitivity analysis

Sensitivity analysis demonstrated that overall NP performance was similar when studies that excluded participants with a previous diagnosis of LVSD were compared with studies that did not (eFigures 7-10). Seven of the included LVSD studies described the screened population as entirely asymptomatic and sensitivity analysis examined whether there were any differences when results of these studies were compared to the other included studies; performance of NP screening was comparable across both groups. Sensitivity analysis was also performed to exclude studies identified as having high risk of bias (eFigures 13-16), and no major differences in performance were noted. We analysed the available data on women and totals (i.e. men and women combined if reported, and otherwise women only) in a further sensitivity analysis, as we had decided to base our primary analysis on all the available data for men

and totals (i.e. men and women combined if reported, and otherwise men only) because not all studies reported data for the combined group of men and women together (*eFigures 17* and *18*). We felt this analysis was particularly relevant given sex-specific differences in the manifestation of CV diseases including HF.⁵⁶ There was a drop in the sensitivity of NT-proBNP from 0.72 (95% CI 0.42–0.90) to 0.68 (95% CI 0.36–0.89) and for BNP from 0.62 (95% 0.32 to 0.85) to 0.59 (95% CI 0.30–0.83), whereas specificity was similar [NT-proBNP 0.82 (95% CI 0.60–0.93) vs. 0.83 (95% CI 0.63–0.94)] and BNP 0.83 (95% CI 0.61–0.94) vs. 0.82 (95% CI 0.64–0.90). The differences in sensitivity were small, however, and in the context of wide CIs, they may not be clinically meaningful.

One of the included studies, Mason et al.³⁹ used only one NT-proBNP threshold set at 1000 pg/mL (which was higher than most other included studies), and no other results at lower thresholds were included. A sensitivity analysis was

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Figure 4 SROC curve of NT-proBNP at multiple thresholds compared with echocardiography for detecting LVSD in screened general and high-risk populations combined.

therefore performed to examine whether excluding this study affected the pooled accuracy results (*eFigure 19*); there was no significant change in the pooled sensitivity and specificity for detecting LVSD in screened high-risk populations with Mason et al. excluded.

Discussion

We found that in studies recruiting high-risk community populations, screening with NP testing accurately detected LVSD: NT-proBNP testing demonstrated a high pooled sensitivity of 0.87 (95% CI 0.73–0.94) and specificity of 0.84 (95% CI 0.55– 0.96). In studies of general community populations, the pooled sensitivity of NT-proBNP to detect LVSD was lower at 0.72 (95% CI 0.42–0.90) with specificity 0.82 (95% CI 0.60–0.93). The pooled accuracy of NT-proBNP in high-risk and general community populations combined gave an optimal cut-off of 311 pg/mL with sensitivity of 0.74 (95% CI 0.53–0.88) and specificity 0.85 (95% CI 0.68–0.93). The pooled accuracy data for BNP yielded an optimal screening threshold for the detection of LVSD at 49 pg/mL with a sensitivity of 0.68 (95% CI 0.45–0.85) and a specificity of 0.81 (0.67–0.90).

Comparison with previous studies

We compared our results with similar studies exploring the use of NP testing in screening non-presenting patients for LVSD. Although previous systematic reviews have included NP screening, studies from secondary and community populations have been combined.^{20,57} or community settings were analysed but specifically in nursing homes only.⁵⁸ Ewald et al. calculated a pooled diagnostic odds ratio (DOR) of 9.3 (95% CI 4.7–17.4) for NT-proBNP screening for severe LVSD.²⁰ Based on the pooled estimate of sensitivity and specificity, our DOR for NT-proBNP to detect LVSD is higher at 42, but there is more heterogeneity within our studies. A recent individual patient data study has also developed and validated a prediction model that identified older patients with HF.⁵⁹ The clinical model improved with the addition of NT-proBNP and modelling in combination with NP screening may provide the most accurate screening strategy.

Our results for optimal screening thresholds are consistent with previous research in this area. For general and high-risk populations combined, the optimal threshold for BNP was 49 pg/mL in our study. This aligns closely with the STOP-HF study that used BNP \geq 50 pg/mL as a cut-off for intervention, and it is the results from this study that underpin the current North American guideline recommendations.^{13,14} Moreover, findings from another meta-analysis of screening studies that

combined primary and secondary care studies found that the optimal sensitivity was achieved when BNP was below the cut point of 50 pg/mL.⁵⁷ To put this in context, this level of BNP threshold is similar to 2021 European Society of Cardiology (ESC) guidelines for the diagnosis of chronic HF for patients who present with symptoms in the non-acute settings in which the upper limit of normal for BNP is 35 pg/ mL (125 pg/mL for NT-proBNP) although these do not refer to screened community populations.⁴ However, the optimal threshold for NT-proBNP in our analysis of 311 pg/mL was notably higher than the current recommended ESC guideline cut-off of 125 pg/mL. NT-proBNP is increasingly used in favour of BNP in some countries, so agreement on an optimal threshold is needed.⁶⁰ A recent German study found that there was an age-related incremental increase in NT-proBNP levels in asymptomatic older adults, with significant sex differences also observed.⁶¹ The Heart Failure Association of the ESC has published practical guidance on the use and interpretation of NP tests, including in the context of screening.62

Overall, there is a paucity of HF diagnostic accuracy studies in screened community-based populations. More research has been conducted in acute HF with more published diagnostic accuracy data available. A meta-analysis by Roberts et al., which informed the current National Institute for Health and Care Excellence acute HF guideline, pooled data from 37 acute HF diagnostic accuracy studies, including 15 263 NP test results, to evaluate NP test performance.⁶³ They found the NP thresholds defined in the ESC HF guideline 2012 performed well, particularly at the lower threshold (BNP < 100 pg/mL, NT-proBNP<300 pg/mL) for ruling out acute HF.⁶³ A similar increase in the number and quality of community-based screening studies is needed to further evaluate the role of NP testing to detect or rule out LVSD.⁴

Implications for policy and practice

There is an urgent need for further prospective studies as well as an individual participant data meta-analysis to more precisely evaluate diagnostic accuracy and identify optimal screening thresholds in specifically defined sub-populations within the community, including comparing performance by EF, in populations who are aged under or over 65 years, and this should include comparisons of accuracy in men and women. Biological variables such as BMI and renal function also need to be better understood in a screening context.

Current policy on the use of NP screening to detect LVSD varies globally. European guidelines do not recommend NP screening, whereas North American guidelines advocate screening without specifying where the positivity threshold should be set.¹⁴ Our results provide evidence to support NP screening in high-risk populations to detect LVSD and contribute to the evidence base on screening thresholds.

The performance of NP screening to detect LVSD is comparative in accuracy to some cancer screening approaches such as faecal immunochemical testing (FIT) for haemoglobin to detect colorectal cancer. The FIT test is now integrated into the UK national bowel cancer screening programme.⁶⁴

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Screening in healthcare remains a contentious issue. The costs and benefits of any screening programme would need to be considered on a national level. In many European countries, echocardiography services are already under strain and the potential for overdiagnosis when imaging large numbers of people is considerable. Future research should investigate whether false positive rates could be reduced with refinement of target populations, potentially through improved risk prediction, as well as to explore the acceptability of screening from a patient perspective.⁶⁵ The burden on the patient and healthcare system needs to be outweighed by improvements in mortality and quality of life and reduced costs of caring for patients with advanced HF, which may result from earlier diagnosis and treatment. Future research needs to examine both the impact of NP screening for LVSD on patient outcomes and the resources required for management of screen-detected patients, including up-titration of medications.66

Strengths and limitations

To our knowledge, this is the first systematic review to focus on the diagnostic accuracy of NP testing in screened community populations alone for the detection of LVSD to compare performance in both high-risk and general populations and to analyse NP thresholds used in screening studies. Decisions regarding inclusion criteria were based on a pragmatic approach underpined by clinical experience of community diagnostics. To increase applicability to a broad range of clinical contexts, we expanded on the definition of high-risk populations that were used in STOP-HF.¹³ This review has been produced in accordance with the PRISMA guidelines and the Cochrane Handbook for Systematic Reviews of Diagnostic Test Accuracy.²³ There was significant heterogeneity among the included studies due to variation in the included clinical populations, definition of LVSD used by investigators, and disparities in reporting of diagnostic accuracy parameters at different NP thresholds. Study populations differed in the ages of included participants, presence of symptoms, and the prevalence of LVSD. We were unable to undertake planned subgroup analysis by age due to the wide variation in age ranges recruited across studies (see Table 1, eTable 1, and eTable 2 for age definitions) and the unavailability of individual patient data. Attempts were made to mitigate for this by analysing data separately for high-risk and general populations, an approach not taken by most previous systematic reviews that have focussed on NP screening, and examined the population as a whole.^{20,21} We included all studies from screened community populations but only some studies (n = 7, 29%) described participants as entirely asymptomatic. Many patients with HF describe being unaware of symptoms initially, particularly when these are very mild, and therefore do not present to a healthcare professional.⁵ The presence or absence of symptoms is therefore variable, and the included studies are aligned with the real-world experience of community screening where there might be symptomatic patients still presenting for a screening test, particularly if symptoms are mild.

The definitions of LVSD are listed under target condition in *Table 1, eTable 1*, and *eTable 2*. As the overall aim of community screening is to identify previously undetected LVSD, we chose to include both narrative and quantitative descriptions of reduced left ventricular EF although this may have contributed to the clinical heterogeneity. We planned to analyse test performance by EF (e.g. <40%, <50%), but there were too few studies to enable this subgroup analysis to be performed.

The different NP thresholds reported by included studies are also listed in Table 1, eTable 1, and eTable 2, all forest plots were also ordered by threshold so that different studies that reported performance at similar thresholds were grouped together. We have attempted to provide an estimation of where optimal screening thresholds might lie. The statistical model that was used to pool sensitivity and specificity to provide a summary estimate is different to the model that is required to identify the optimal threshold.³⁰ Fitting this model requires studies to report results at multiple thresholds, and there were not enough such studies in the subgroup of studies in high-risk populations to allow this. The inability to recommend an optimal screening threshold in high-risk populations is a major study limitation. To provide an indication of where the appropriate positivity threshold might lie, we estimated an optimal threshold from the pooled high-risk and general population studies combined. We acknowledge that this also limits the current clinical applicability of the findings although identifying this evidence gap is an important finding and provides a focus for future research that can build on this more general exploration of the accuracy of NP screening for LVSD.

Conclusions

In high-risk community populations, it is likely that NP screening may accurately detect LVSD. Given the huge public health burden of missed HF diagnoses in the community, this finding presents a potentially important opportunity for diagnosis and early intervention. Our study highlights an urgent need for further prospective studies, as well as an individual participant data meta-analysis, to evaluate diagnostic accuracy more precisely, to identify optimal screening thresholds in specifically defined sub-populations within the community, and to further examine the impact of NP screening on both general and high-risk populations.

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Conflict of interest

All authors have completed the ICMJE uniform disclosure form at www.icmje.org/coi_disclosure.pdf and declare: CT reports speaker fees from Vifor and Novartis and non-financial support from Roche outside the submitted work. FDRH reports personal fees and other from Novartis and Boehringer Ingelheim and grants from Pfizer outside the submitted work. CG and the other authors have nothing to disclose.

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Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Appendix S1. Search Strategy.

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