A systematic review of heart failure dyadic self-care interventions focusing on intervention components, contexts, and outcomes

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TITLE: A Systematic Review of Heart Failure Dyadic Self-care Interventions Focusing on Intervention Components, Contexts, and Outcomes

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ABSTRACT

Background: Having support from an informal carer is important for heart failure patients. Carers have the potential to improve patient self-care. At the same time, it should be acknowledged that caregiving could affect the carer negatively and cause emotional reactions of burden and stress. Dyadic (patient and informal carer) heart failure self-care interventions seek to improve patient self-care such as adherence to medical treatment, exercise training, symptom monitoring and symptom management when needed. Currently, no systematic assessment of dyadic interventions has been conducted with a focus on describing components, examining physical and delivery contexts, or determining the effect on patient and/or carer outcomes.

Objective: To examine the components, context, and outcomes of dyadic self-care interventions.

Design: A systematic review registered in PROSPERO, following PRISMA guidelines with a narrative analysis and realist synthesis. Data Sources: PubMed, EMBASE, Web of Science, PsycINFO, and Cochrane Central Register of Controlled Trials were searched using MeSH, EMTREE terms, keywords, and keyword phrases for the following concepts: dyadic, carers, heart failure and intervention. Eligible studies were original research, written in English, on dyadic self-care interventions in adult samples.

Review methods: We used a two-tiered analytic approach including both completed studies with power to determine outcomes and ongoing studies including abstracts, small pilot studies and protocols to forecast future directions.

Results: Eighteen papers – 12 unique, completed intervention studies (two quasi- and ten experimental trials) from 2000 to 2016 were reviewed. Intervention components fell into three groups – education, support, and guidance. Interventions were implemented in 5 countries, across multiple settings of care, and involved 3 delivery modes – face to face, telephone or technology based. Dyadic intervention effects on cognitive, behavioral, affective and health services utilization outcomes were found within studies. However, findings across studies were inconclusive as some studies reported positive and some non-sustaining outcomes on the same variables. All the included papers had methodological limitations including insufficient sample size, mixed intervention effects and counter-intuitive outcomes.

Conclusions: We found that the evidence from dyadic interventions to promote heart failure self-care, while growing, is still very limited. Future research needs to involve advanced sample size justification, innovative solutions to increase and sustain behavior change, and use of mixed methods for capturing a more holistic picture of effects in clinical practice.
Contribution of the paper:

What is already known about the topic?

- Having support from an informal carer is important for heart failure (HF) patients
- Dyadic (patient and informal carer) HF self-care interventions seek to improve patient self-care such as adherence to medical treatment, exercise training, symptom monitoring and symptom management
- No systematic assessment of dyadic interventions has been conducted with a focus on describing components, examining physical and delivery contexts, or determining the effect on patient and/or carer outcomes

What this paper adds

- The body of evidence for dyadic interventions in HF is small
- All the papers in the current review had some methodological limitations, mixed intervention effects and counter-intuitive outcomes.
- It is time to change the design, development, and implementation of dyadic HF interventions
- What is needed are co-design methods, judicious use of technology, careful development of theoretical frameworks, with clear hypotheses and descriptions of mechanisms and relevant outcome measures.
Introduction

Heart failure is a common condition worldwide, with a prevalence of 1-2% in the population; rising to ≥10% among persons above 70 years of age. (Ponikowski et al., 2016; Yancy et al., 2013) Heart failure self-care interventions which improve heart failure patients’ necessary knowledge and management skills for this chronic and progressive condition are advocated for in clinical guidelines and widely implemented in heart failure care. (Ponikowski et al., 2016; Yancy et al., 2013) A recent meta-analysis of 20 studies (n=5624 patients) found that self-care interventions which targeted patients alone without involving informal carers were effective in reducing hospitalization or all-cause death, delaying time to hospitalization, and improving quality of life during the study. (Jonkman et al., 2016) However, a critical question facing clinicians and researchers is why heart failure self-care interventions have often resulted in non-sustained effects on patient outcomes when examined over time. (Liljeroos, Agren, Jaarsma, Arestedt, & Stromberg, 2015) A series of recent systematic reviews (Alexander M Clark et al., 2014; Currie et al., 2015; Harkness, Spaling, Currie, Strachan, & Clark, 2014; Spaling, Currie, Strachan, Harkness, & Clark, 2015; Strachan, Currie, Harkness, Spaling, & Clark, 2014) suggest that one explanation for this could be that these past self-care interventions which have targeted only the patient have missed a critical component – the informal carer. In actual practice, patients rarely engage in heart failure self-care in isolation. (H. G. Buck, Harkness, et al., 2015)

Many heart failure patients live within a family system as part of a patient/informal carer dyad. A dyad is typically defined as two individuals maintaining a sociologically significant relationship. ("Merriam-Webster," n.d.) Members of heart failure patient/carer dyads have been found to influence each other’s behaviour, physical and mental well-being. (H. G. Buck, Mogle, Riegel, McMillan, & Bakitas, 2015; Kitko, Hupcey, Pinto, & Palese, 2014; Vellone et al., 2014) For
example, higher self-care maintenance in patients has been associated with better mental well-being in carers. (Vellone et al., 2014) In a recent review of 45 qualitative studies, two factors were identified: 1) the importance of carers and social support in patients’ self-care and 2) the carers’ relative under-representation in current self-care programs. (Strachan et al., 2014) In another systematic literature review of heart failure self-care determinants, the researchers found significant linkage between increased involvement by others in patient self-care and greater intervention effectiveness. (Alexander M Clark et al., 2014) Yet, self-care interventions which include patient and informal carer dyads remain a relatively recent, but growing area of exploration.

Self-care interventions target daily heart failure patient self-care behavior, such as adherence to medical treatment, exercise training, routine self-monitoring and symptom management when needed. (Riegel, Dickson, & Faulkner, 2016) Dyadic self-care interventions are delivered to both a patient and his/her informal carer with expectations that both dyad members will be actively engaged in the patient’s heart failure self-care. Dyadic self-care interventions generally mirror current patient interventions by including educational materials as well as some form of support and guidance. Currently, no systematic assessment of heart failure self-care dyadic interventions has been conducted with a focus on describing components, examining contexts, or determining the effect on patient and/or carer outcomes. Therefore, the purpose of this systematic literature review was to examine in Aim 1) the components; in Aim 2) the contexts; and in Aim 3) the outcomes of dyadic self-care interventions. This study’s findings will enable researchers to chart future directions in the science of dyadic interventions.

Methods

Protocol and registration
We conducted a systematic review with a narrative analysis and realist synthesis to answer our questions. Meta-analysis was deemed not appropriate given the small number of studies and heterogeneity of the interventions. The systematic review team consisted of six international nurse researchers with significant experience in conducting studies to support heart failure self-care, one clinical psychologist with dyadic self-care intervention expertise, and two medical librarians. The team developed the study protocol based on PRISMA statement criteria (Liberati et al., 2009) and methods developed in previous systematic reviews (Harleah G. Buck, Akbar, Zhang, & Bettger, 2013; H. G. Buck, Harkness, et al., 2015; H. G. Buck et al., 2012). The protocol was then registered with PROSPERO (CRD42016050214).

Eligibility criteria

The heart failure dyadic intervention body of literature is limited. Therefore, we were as inclusive as possible to capture the extant literature across disciplines. A decision was made a priori to hold 1) meta-analyses or reviews for reference list hand-searches rather than analysis; and 2) abstracts (without full papers), feasibility/small pilot studies, or protocol papers for a separate analysis to forecast future directions in this line of inquiry. This design decision resulted in a two-tiered analytic approach. Specifically, we analyzed 1) completed studies with sufficient information and power to determine patient/carer outcomes, and 2) ongoing studies including abstracts, small pilot studies, and protocols to explore future directions in dyadic intervention research. Inclusion criteria for all papers were: adult samples (18 years of age or greater); dyad consisting of a patient with heart failure and at least one informal carer; both dyad members must be the target of the intervention, present at the intervention and outcomes (primary or secondary) from both OR either one of the participants measured; English language papers; intervention description provided (with hand-searches if intervention described in a second paper); and with an experimental or quasi-experimental design. Meta-analyses and
systematic reviews (except as noted above); duplicative papers; case reports, opinion pieces, editorials and letters to the editor were excluded.

Information sources and search

The two medical librarians developed and performed searches in PubMed (1946 to present), EMBASE (Elsevier, 1947 to present), Web of Science (Core Collection, 1900 to present), PsycINFO (EBSCO, 1887 to present), and Cochrane Central Register of Controlled Trials (CENTRAL, through August 2016). MeSH terms, EMTREE terms, keywords, and keyword phrases were used to search for the following concepts: dyadic, carers, heart failure, and intervention. Search terms were shared with the team to elicit feedback. Searches were limited to English language. Appendix A shows the detailed search strategies.

The combined searches yielded 1,217 papers as of September 29, 2016. All papers were exported into EndNote and 327 duplicates were removed mechanically. The remaining papers (890) were imported into an MS EXCEL worksheet for purposes of documenting the inclusion/exclusion analysis performed by the team.

Study selection

A two-step screening approach was used to select the final papers to include in the review. In the first screen (title/abstract screen), the 890 papers were divided amongst five team members with a planned overlap of 5% of the papers given the large sample size. Approximately 200 papers were reviewed per team member. The focus was on retaining as many papers as possible, therefore team members were instructed to retain any ambiguous papers for the next level of review. For example, if a title appeared to meet the inclusion criteria but there was no abstract the paper was retained until the criteria could be applied to the full paper. Team members were also instructed to retain papers that met the criteria for the hand search and separate analysis (ongoing studies). One team member was held out of this
screening step and analyzed the overlapping papers to assess inter-rater reliability. Only four papers from the 5% overlap (n= 45 papers) resulted in disagreement (91% inter-rater agreement). These four papers were then reviewed and discussed by two other team members and a decision to keep or discard resulted. The title/abstract screen resulted in 54 papers for full-text review. In the second screen, the full-text of the 54 papers were reviewed once again using our criteria. If there were multiple papers from one author or research team, they were reviewed by the same reviewer to assure that duplicate papers would be identified and discarded. The papers were divided amongst the team and went through a 100% overlap review. Two team members had to agree on inclusion and exclusion determination of the article resulting in 100% inter-rater agreement.

Data collection process and data items

Data extraction elements were selected a priori to address our aims. These included: intervention components, contextual details and main study outcomes. We also collected: year of publication, journal, study design, sample size, participant age and relationships in the dyad (spouse/partner, adult child, relative, friend, etc.). The data extraction form was adapted from previous studies and team discussion. Team members also received an EndNote library of all the papers with the full-text PDFs attached. Each paper was abstracted by one team member and then confirmed by a second team member.

Risk of bias analysis

Critical Appraisal Skills Programme (CASP) criteria("Critical Appraisal Skills Programme:

Box.1 Quality assurance steps

- Use of a standardized protocol
- Frequent, regular team communication
- Eligibility confirmation at each step
- Inter-rater assessment at each step
- Standardized process for resolving reviewer disagreement
- Standardized data extraction process
- CASP (Critical Appraisal Skills Programme) criteria used to asses for risk of bias in all studies
- Final approval of full team on all papers for inclusion
Making Sense of Evidence," 2017) were used to assess for risk of bias in all the papers; CASP provides standardized, study design specific, criteria that allowed us to analyze multiple study designs using a single program. Each team member conducted the CASP appraisal while extracting study data. For example, randomized control trials (RCTs) were analyzed for blinding, similarity between groups on baseline data, equal treatment besides experimental condition and allocation, while quasi-experimental papers were analyzed for representativeness, exposure measurement, identification of confounders and assessment of follow-up time. The quality assurance steps are summarized in Box 1.

**Synthesis of results**

Realist synthesis techniques (Pawson, Greenhalgh, Harvey, & Walshe, 2005; Wong, Greenhalgh, Westhorp, Buckingham, & Pawson, 2013) as in previous heart failure systematic reviews(H. G. Buck, Harkness, et al., 2015; A. M. Clark et al., 2016) were used to interpret the data elements across papers, analyze the nature and relationships between the data elements and draw conclusions about the higher order abstractions. In the first aim specific techniques included: data elements (actual intervention components) were identified in individual papers, aggregated across papers and then categorized. Conclusions related to the categories were then drawn. In the second aim, implementation contexts were the data elements and patterns across papers were identified and conclusions drawn. In the third aim, all reported outcomes were identified individually, then categorized across papers and finally analyzed to determine patterns in responses. To forecast the direction of future dyadic intervention studies, we analyzed ongoing studies that our search uncovered, and to the degree possible, evaluated them using the same criteria used in our analyses of the completed studies.

**Results**
See Figure 1 for numbers of papers screened, assessed for inclusion, and included in the review.
Figure 1. PRISMA Flow diagram

Records identified through database searching
- PubMed: 204
- Embase: 546
- Web of Science (Core Collection): 207
- PsycINFO: 88
- Cochrane Central Register of Controlled Trials: 172
  (n = 1217)

Records after duplicates removed
(n = 890)

Records screened
(n = 890)  →  Records excluded
(n = 836)

Full-text papers assessed for eligibility
(n = 54)  →  Full-text papers excluded
(n = 28)

Papers included in synthesis as complete
(n = 18 representing 12 unique studies)

Papers included in separate synthesis as ongoing
(n = 8)
Eighteen papers involving 12 unique, complete intervention studies, from 2000 to 2016, met the criteria of complete studies. Of the 12 complete studies there were two quasi-experimental studies (Bull, Hansen, & Gross, 2000; Piette et al., 2008) and 10 RCTs. See Table 1 for the characteristics of the main outcome papers from the 12 complete studies. Six papers (Agren, L, Davidson, & Stromberg, 2013; Agren, Stromberg, Jaarsma, & Luttik, 2015; Dunbar et al., 2016; Liljeroos et al., 2015; Liljeroos, Agren, Jaarsma, Arestedt, & Stromberg, 2016; Stamp et al., 2016) reported on secondary outcomes or primary outcome variables at a later time from the complete studies. These secondary outcome papers were included in the systematic analysis to examine additional intervention effect on outcomes. Eight papers met the criteria of ongoing studies. These included six abstracts, (Bakitas, Dionne-Odom, Kvale, Kono, & Pamboukian, 2016; Chung, Lennie, & Moser, 2014; Chung, Moser, & Lennie, 2014; Deek et al., 2015; Demers et al., 2014; McIlvennan, Thompson, Matlock, Cleveland, & Allen, 2015) one feasibility study, (Dionne-Odom et al., 2014) and one protocol paper. (Taylor et al., 2015) Methodological analysis using CASP criteria found some limitations related to blinding, allocation, and effect sizes in all studies.

A total of 1459 dyads (sample sizes range 20 (Piamjariyakul et al., 2015)-372 (Piette, striplin, Marinec, Chen, & Aikens, 2015)) from the complete intervention studies were involved with an average patient age across studies of 59-78 years and an average carer age of 29-67 years. The studies which clearly identified the types of dyads reported spousal (n = 417), adult child (n = 344), or friend/relative (n=209) dyadic partners. Two of the 12 studies included only spouse/partner dyads, (Ågren, Berg, Svedjeholm, & Strömberg, 2015; Agren, Evangelista, Hjelm, & Stromberg, 2012) two studies did not identify the dyad relationships, (Hasanpour-Dehkordi, Khaledi-Far, Khaledi-Far, & Salehi-Tali, 2016; Kenealy et al., 2015) the other eight studies included mixed dyads, primarily spouse or adult child dyads.
<table>
<thead>
<tr>
<th>Authors</th>
<th>Publication Year</th>
<th>Country</th>
<th>Study Design</th>
<th>Type of Dyads</th>
<th>Gender (% male)</th>
<th>Sample Size</th>
<th>Mean Age</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bull et al. (Bull et al., 2000)</td>
<td>2000</td>
<td>US</td>
<td>Pre/post nonequivalent control group</td>
<td>Spouse 50%; Adult child 38%; Family 7%; Friend 5%</td>
<td>Patient ND Carers 27%; Carers 27%</td>
<td>180 dyads</td>
<td>Patient 73.7 ± 8.8 Carer 58.5 ± 14.9</td>
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<tr>
<td>Dunbar et al. (Dunbar et al., 2005)</td>
<td>2005</td>
<td>US</td>
<td>RCT</td>
<td>Spouse 67%; Adult child 22%; Other 11%</td>
<td>Patients 54%; Carers 23%</td>
<td>61 dyads</td>
<td>Patient 61.0 ± 12 Carer 54.0 ± 17</td>
</tr>
<tr>
<td>Piette et al. (Piette et al., 2008)</td>
<td>2008</td>
<td>US</td>
<td>Cohort study</td>
<td>Adult child 75%; Family 15%; Friend 10%</td>
<td>Patients 89%; Carer 42%</td>
<td>52 dyads</td>
<td>Patient 65.9 ± 11.0 Carer 42.3 ± 10.0</td>
</tr>
<tr>
<td>Schwarz et al. (Schwarz, Mion, Hudock, &amp; Litman, 2008)</td>
<td>2008</td>
<td>US</td>
<td>RCT</td>
<td>Intervention group Spouse 64%; Adult child 26%; Other 10%</td>
<td>Patient 48%; Carer ND</td>
<td>102 dyads</td>
<td>Patient 78.1 ± 7.1 Carer 63.4 ± 16.1</td>
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<tr>
<td>Authors</td>
<td>Publication Year</td>
<td>Country</td>
<td>Study Design</td>
<td>Type of Dyads</td>
<td>Gender (% male)</td>
<td>Sample Size</td>
<td>Mean Age</td>
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<tr>
<td>Ågren et al. (Ågren et al., 2012)</td>
<td>2012</td>
<td>Sweden</td>
<td>RCT</td>
<td>Partner 100%</td>
<td>Patient intervention group 69%</td>
<td>155 dyads</td>
<td>Interventio group 67.0 ± 13.0 Control group 70.0 ± 10.0</td>
</tr>
<tr>
<td>Dunbar et al. (Dunbar et al., 2013)</td>
<td>2013</td>
<td>US</td>
<td>RCT</td>
<td>Spouse 53%</td>
<td>Patient 63%</td>
<td>117 dyads</td>
<td>Patient 55.9 ± 10.5 Carer 52.3 ± 33.3</td>
</tr>
<tr>
<td>Ågren et al. (Ågren et al., 2015)</td>
<td>2015</td>
<td>Sweden</td>
<td>RCT</td>
<td>Partner 100%</td>
<td>Patient intervention</td>
<td>42 dyads</td>
<td>Patient intervention</td>
</tr>
<tr>
<td>Authors</td>
<td>Publication Year</td>
<td>Country</td>
<td>Study Design</td>
<td>Type of Dyads</td>
<td>Gender (% male)</td>
<td>Sample Size</td>
<td>Mean Age</td>
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<tr>
<td>Srisuk et al. (Srisuk, Cameron, Ski, &amp; Thompson, 2015)</td>
<td>2015</td>
<td>Thailand</td>
<td>RCT</td>
<td>Patient group</td>
<td>100 dyads</td>
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<tr>
<td>Piette et al. (Piette et al., 2015)</td>
<td>2015</td>
<td>US</td>
<td>RCT</td>
<td>Family 34%</td>
<td>Patient intervention group 61% Family 27% Friend 12%</td>
<td>369 dyads</td>
<td>Patient intervention group 67.8 ± 10.2 Control group 68.0 ± 10.2 Carer intervention group 46.6 ± 12.1 Control group 47.6 ± 14.2</td>
</tr>
<tr>
<td>Piamjariyakul et al. (Piamjariyakul et al., 2015)</td>
<td>2015</td>
<td>US</td>
<td>RCT</td>
<td>Spouse 65% Family 35%</td>
<td>Patient 60% Carer 15%</td>
<td>20 dyads</td>
<td>Patient 62.3 ± 13.5 Carers 61.4 ± 10.0</td>
</tr>
<tr>
<td>Kenealy et al. (Kenealy et al., 2015)</td>
<td>2015</td>
<td>New Zealand</td>
<td>RCT – 3 sites with different</td>
<td>ND</td>
<td>Patient 59% Carer ND</td>
<td>171 dyads</td>
<td>Patient 65.3 Carer ND</td>
</tr>
<tr>
<td>Authors</td>
<td>Publication Year</td>
<td>Country</td>
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<tr>
<td>Hasanpour-Dehkordi et al. (Hasanpour-Dehkordi et al., 2016)</td>
<td>2016</td>
<td>Iran</td>
<td>protocols RCT</td>
<td>ND</td>
<td>Patients intervention group 60% Control group 62% Carers ND</td>
<td>90 dyads</td>
<td>Patient intervention group 60.8 Control group 59.1 Carers ND</td>
</tr>
</tbody>
</table>

Legend: RCT- randomized control trial; ND – no data
<table>
<thead>
<tr>
<th>Authors</th>
<th>Intervention Components</th>
<th>Intervention Physical Contexts</th>
<th>Intervention Outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bull et al. (2000)</td>
<td>1 time: Staff training included: Patient and Carer assessment, videotape structured communication guide for use with clinicians Medication form to fill out Brochure on community services</td>
<td>Hospitals</td>
<td>Significantly higher Patient preparedness, continuity of care, health perception, and vitality at 2 weeks post-hospital discharge</td>
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<td></td>
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<td>Significantly better continuity of care at 2 months post-discharge</td>
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<td></td>
<td>No statistically significant differences in scores on satisfaction or difficulties in managing care at 2 weeks or 2 months post-discharge</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Significantly higher Carer scores on continuity of information about condition, services available, general health perceptions, vitality, mental health; less negative reaction to caregiving 2 weeks post-discharge</td>
</tr>
<tr>
<td></td>
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<td>Significantly higher Carer scores on care continuity scales at 2 months post-discharge</td>
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<tr>
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<td></td>
<td>No significant differences on hospital readmission and emergency room use at 2 weeks post-discharge</td>
</tr>
<tr>
<td>Authors</td>
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<td>Intervention Physical Contexts</td>
<td>Intervention Outcomes</td>
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</tbody>
</table>
| Dunbar et al. (Dunbar et al., 2005) | 1 face-to-face education and counseling session  
Video  
Follow up phone call with patient data  
Newsletter  
Same components as 1st group  
Plus 2 additional sessions on family support and patient choice  
Case scenarios  
Group discussion  
Role-play | General Clinical Research Center | Both groups decreased dietary NA+ at 3 months  
2nd group showed greater decrease in NA+ and greater percentage of patients with decreased NA+  
Significant increases in Patient and family heart failure knowledge from pre- to post- education sessions in both groups but did not differ in degree of knowledge change  
Both groups declined in knowledge by 3 months  
Significant Group X Time interaction when accounting for time-varying measures of body mass index  
No significant changes in autonomy support in either group |
| Piette et al. (Piette et al., 2008) | Weekly interactive voice response calls to Patient for 6-15 weeks  
Automated report | Academic medical center and VA hospital | 92% completion rate and reported problems that might otherwise have gone unidentified. 75% made changes in self-care as a result of the intervention                                                                                   |
<table>
<thead>
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<th>Authors</th>
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<th>Intervention Physical Contexts</th>
<th>Intervention Outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Schwarz et al. (2008)</td>
<td>1 home visit to train on equipment which automatically uploaded daily weight</td>
<td>Academic medical center</td>
<td>No significant differences for any outcomes</td>
</tr>
<tr>
<td>Ågren et al. (2012)</td>
<td>3 face-to-face sessions</td>
<td>Outpatient clinic in one university and, one county hospital or patient’s home</td>
<td>Significantly increased patient-perceived control at 3 months</td>
</tr>
<tr>
<td></td>
<td>Computer-based educational program</td>
<td></td>
<td>No effect on dyads quality of life and depression</td>
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<tr>
<td></td>
<td>Written materials</td>
<td></td>
<td>No effect on Carer burden</td>
</tr>
<tr>
<td>Authors</td>
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<td>Intervention Outcomes</td>
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<tr>
<td>Dunbar et al. (Dunbar et al., 2013)</td>
<td>1 face-to-face education session 1 group reinforcement session Written materials DVD Written individual feedback 1 telephone booster session Newsletters</td>
<td>Outpatient clinic</td>
<td>No significant changes in medication adherence, autonomy support in either group</td>
</tr>
<tr>
<td></td>
<td>Same components as 1st group Plus 2 small group sessions with breakout sessions with dyad together and separate</td>
<td></td>
<td>Increased heart failure knowledge immediately after intervention but not sustained</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Significant changes in dietary NA+ in both groups</td>
</tr>
<tr>
<td>Ågren et al. (Ågren et al., 2015)</td>
<td>1 face-to-face session 2 telephone sessions Psychoeducational support</td>
<td>Outpatient clinic</td>
<td>No significant differences on SF-36 after 3 and 12 months</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Significantly improved SF-36 dimensions over time</td>
</tr>
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<td></td>
<td></td>
<td></td>
<td>No difference in depression</td>
</tr>
<tr>
<td>Srisuk et al. (Srisuk et al., 2015)</td>
<td>1 face-to-face session</td>
<td>Outpatient</td>
<td>Significantly increased dyad knowledge at</td>
</tr>
<tr>
<td>Authors</td>
<td>Intervention Components</td>
<td>Intervention Physical Contexts</td>
<td>Intervention Outcomes</td>
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<tr>
<td></td>
<td>session 9 telephone sessions Heart failure manual DVD</td>
<td>clinic</td>
<td>3 and 6 months</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Significantly increased Patient self-care maintenance, confidence, and health-related quality of life at 3 and 6 months, and self-care management at 6 months</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Significantly increased Carer perceived control at 3 months.</td>
</tr>
<tr>
<td>Piette et al. (Piette et al., 2015)</td>
<td>Weekly interactive voice response calls to Patient for 12 months, Carer mailed information on heart failure self-care</td>
<td>Veteran’s Administration outpatient clinics</td>
<td>Significantly less Carer strain at 6 and 12 months</td>
</tr>
<tr>
<td></td>
<td>Weekly interactive voice response calls to Patient for 12 months Automated report to Carer Links to additional online resources Communication printed guidelines</td>
<td></td>
<td>Interaction effect between arm and baseline on Carer strain/depression at both endpoints</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Significantly less time spent in high time commitment Carers; greater participation in clinic visits and medication adherence</td>
</tr>
<tr>
<td>Authors</td>
<td>Intervention Components</td>
<td>Intervention Physical Contexts</td>
<td>Intervention Outcomes</td>
</tr>
<tr>
<td>----------------------------------------------</td>
<td>----------------------------------------------------------------------------------------</td>
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<tr>
<td>Piamjariyakul et al. (Piamjariyakul et al., 2015)</td>
<td>Logbook Laminate reminder and tips cards 4 weekly telephone sessions Printed materials (caregiving guide, list of local support organizations) Pill organizer Referral to Social Worker if needed</td>
<td>Outpatient clinic</td>
<td>Significantly fewer heart failure rehospitalizations at 6 months</td>
</tr>
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<td></td>
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<td></td>
<td>Significantly higher Carer confidence and social support scores and significantly lower Carer depression</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>No significant difference in preparedness or burden</td>
</tr>
<tr>
<td>Kenealy et al. (Kenealy et al., 2015)</td>
<td>1 home visit to train on equipment which provided instruction, asked pre-programmed questions, gave short message Patient entered self-care data manually Monitored by clinicians</td>
<td>Hospital, primary care site</td>
<td>No significant changes in quality of life, self-efficacy and disease-specific measures</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Significant changes in anxiety and depression</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>No significant changes in hospital admissions, days in hospital, emergency department visits, outpatient visits and costs</td>
</tr>
<tr>
<td>Hasanpour-Dehkordi et al. (Hasanpour-Dehkordi et al., 2016)</td>
<td>3 face-to-face sessions</td>
<td>Hospital</td>
<td>Significant difference in physical activity limitation following physical problems,</td>
</tr>
<tr>
<td>Authors</td>
<td>Intervention Components</td>
<td>Intervention Physical Contexts</td>
<td>Intervention Outcomes</td>
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</tr>
<tr>
<td></td>
<td>Education</td>
<td></td>
<td>energy and fatigue, social performance, physical pain and general health 6 months Significant difference in quality of life at 6 months Significant differences in hospital readmissions and referring to physicians Significant difference in mean health care cost</td>
</tr>
</tbody>
</table>

Legend: NA+ - sodium; VA – Veterans Administration; SF-36 – Short Form (36) Health Survey
Dyadic intervention components

Component groups. The components of the 12 interventions aligned into three groups – those that could be categorized as including a) education; b) support, and c) guidance. Education components were defined as either information related to heart failure, its management or educational strategies. Examples of education informational components included information about heart failure,(Ågren et al., 2015; Agren et al., 2012; Dunbar et al., 2005; Dunbar et al., 2013; Piette et al., 2015; Schwarz et al., 2008; Srisuk et al., 2015) skill-building,(Ågren et al., 2015; Agren et al., 2012; Bull et al., 2000; Dunbar et al., 2005; Dunbar et al., 2013; Piamjariyakul et al., 2015) and relationship information.(Ågren et al., 2015; Agren et al., 2012) Educational strategies included technological (use of computer/telephone(Ågren et al., 2015; Agren et al., 2012; Piamjariyakul et al., 2015; Piette et al., 2015)) or methodological (use of teach-back,(Srisuk et al., 2015) printed materials(Ågren et al., 2012; Bull et al., 2000; Piamjariyakul et al., 2015; Piette et al., 2008; Srisuk et al., 2015) or role play/ group discussion(Dunbar et al., 2005; Dunbar et al., 2013)) strategies. Support components were defined as ancillary support resources or actual support provided during the intervention. Examples of supportive components were equally diverse and included referral to additional resources,(Piamjariyakul et al., 2015; Piette et al., 2008) information on the importance of support(Ågren et al., 2015) or actual supportive calls.(Dunbar et al., 2005; Dunbar et al., 2013) Guidance components were defined as specific verbal or written directions given during the intervention to improve care. All intervention studies included guidance component. See Table 2 for a list of intervention components for each study.

Component numbers. The number of individual components per intervention varied greatly but ranged from two to six components. The 12 interventions included variable numbers of sessions in which the components were delivered; ranging from one home visit to deliver equipment(Kenealy et al., 2015) to weekly telephone calls for 12 months.(Piette et al., 2015) The time spent during a session
ranged from a 15 minute phone call (Srisuk et al., 2015) to a 90 minute phone call (Piamjariyakul et al., 2015) with most face-to-face sessions lasting approximately 60 minutes (or multiples of this).

We attempted to derive a dose (number of sessions) and strength of the dose (comprised of amount of time for each of the session, length of the intervention and length of any follow-up) for each study. However, there was great heterogeneity and incompleteness in reporting both the number of sessions and time spent per session precluding an accurate description of dose and strength of dose for some interventions. As an example of this heterogeneity, Ågren et al. (Agren et al., 2012) reported an intervention comprised of three components (face-to-face counseling, computer-based program and written materials). The intervention involved three sessions of 60 minutes each suggesting 180 minutes of direct contact with an interventionist during the intervention. A second intervention (Bull et al., 2000) included a detailed description of a multi-component intervention but provided no clear information on number of components nor length of time per contact. Therefore, we were unable to summarize an accurate dose of the intervention to make a comparison.

**Dyadic intervention contexts**

**Physical context.** Countries where interventions were tested included the United States, (Bull et al., 2000; Dunbar et al., 2005; Dunbar et al., 2013; Piamjariyakul et al., 2015; Piette et al., 2008; Piette et al., 2015) Sweden, (Ågren et al., 2015; Agren et al., 2012) New Zealand, (Kenealy et al., 2015) Iran, (Hasanpour-Dehkordi et al., 2016) and Thailand. (Srisuk et al., 2015) Settings where the intervention was initiated or where the majority of the interventions were delivered included hospitals, (Agren et al., 2012; Bull et al., 2000; Hasanpour-Dehkordi et al., 2016; Kenealy et al., 2015; Piette et al., 2008; Schwarz et al., 2008) clinics, (Ågren et al., 2015; Agren et al., 2012; Dunbar et al., 2013; Piamjariyakul et al., 2015; Srisuk et al., 2015) home, (Agren et al., 2012) a U.S. Veterans Administration (clinic or hospital), (Piette et al., 2008; Piette et al., 2015) and in a clinical research center. (Dunbar et al., 2005) Two studies (Agren et
al., 2012; Piette et al., 2008) included multiple sites to support the transition from hospital to home or increase recruitment.

**Delivery context.** The 12 interventions aligned into three delivery contexts – those which could be categorized as face-to-face; (Ågren et al., 2015; Agren et al., 2012; Bull et al., 2000; Dunbar et al., 2005; Dunbar et al., 2013; Hasanpour-Dehkordi et al., 2016; Srisuk et al., 2015) telephone; (Piamjariyakul et al., 2015) or telehealth interventions (see Table 2). (Kenealy et al., 2015; Piette et al., 2008; Piette et al., 2015; Schwarz et al., 2008) Most face-to-face interventions also used some form of technology including DVD/video, (Bull et al., 2000; Dunbar et al., 2005; Srisuk et al., 2015) website, (Agren et al., 2012) or phone calls (Ågren et al., 2015; Dunbar et al., 2005; Dunbar et al., 2013; Srisuk et al., 2015) blurring the line between “high touch” and “high tech” modalities. It appears that all the studies were investigator-initiated and led.

**Dyadic intervention outcomes**

**Outcome groups.** This analysis included both primary and secondary outcome papers (n=18). The synthesis resulted into four patient outcome categories comprised of cognitive outcomes (e.g., perceived control, preparedness to care, knowledge); behavioral outcomes (e.g., self-care, carer attending clinic visits); affective outcomes (e.g., depression, social support, strain); and health services utilization outcomes (e.g., hospitalization, quality adjusted life years) (Table 2).

**Effect on outcomes.** See Table 3 for information on positive, null, and mixed outcomes.
Table 3. Categorization and Summary of Dyadic Intervention Outcomes

<table>
<thead>
<tr>
<th>Positive Outcomes</th>
<th>Null Outcomes</th>
<th>Mixed Outcomes</th>
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</thead>
<tbody>
<tr>
<td><strong>Cognitive Outcomes</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patient perceived control at 3 months (Ågren et al., 2015; Agren et al., 2012) and 12 months (Ågren et al., 2015)</td>
<td>Patient perceived control at 12 months (Agren et al., 2012) Carer perceived control at 6 months (Srisuk et al., 2015) Patient/Carer perceived control at 24 months (Liljeroos et al., 2015)</td>
<td>Perceived control for both Patient and Carer</td>
</tr>
<tr>
<td>Carer perceived control at 3 months (Srisuk et al., 2015)</td>
<td>Patient quality of life at 3 months (Schwarz et al., 2008) Patient/Carer quality of life at 3 months (Agren et al., 2012) Patient quality of life at 6 months (Kenealy et al., 2015)</td>
<td>Quality of life for Patient</td>
</tr>
<tr>
<td>Patient quality of life at 3 (Srisuk et al., 2015) and 6 months (Hasanpour-Dehkordi et al., 2016; Srisuk et al., 2015)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patient preparedness to care at 2 weeks (Bull et al., 2000)</td>
<td>Carer preparedness to care at 6 months (Piamjariyakul et al., 2015)</td>
<td>Preparedness to care for both Patient and Carer</td>
</tr>
<tr>
<td>Carer reaction to caregiving (Bull et al., 2000)</td>
<td>Patient/Carer satisfaction with care (Bull et al., 2000)</td>
<td></td>
</tr>
<tr>
<td>Patient knowledge (Dunbar et al., 2013) Patient/Carer knowledge (Dunbar et al., 2005; Srisuk et al., 2015)</td>
<td>Patient (Liljeroos et al., 2015)/Carer (Liljeroos et al., 2016) morbidity</td>
<td></td>
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<tr>
<td>Positive Outcomes</td>
<td>Null Outcomes</td>
<td>Mixed Outcomes</td>
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<tr>
<td><strong>Behavioral Outcomes</strong></td>
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<tr>
<td>Patient self-care at 3 and 6 months (Srisuk et al., 2015)</td>
<td>Patient self-care at 6 months (during intervention) (Kenealy et al., 2015)</td>
<td>Self-care for Patient</td>
</tr>
<tr>
<td>Carers attending visits (Piette et al., 2015)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patient medication adherence (Dunbar et al., 2013)</td>
<td></td>
<td></td>
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<tr>
<td>Carer tasks (Agren et al., 2015)</td>
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<td></td>
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<tr>
<td><strong>Affective Outcomes</strong></td>
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<td></td>
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<tr>
<td>Patient/Carer depression at 6 months (Kenealy et al., 2015; Piamjariyakul et al., 2015; Piette et al., 2015)</td>
<td>Patient depression at 3 months (Dunbar et al., 2005; Schwarz et al., 2008)</td>
<td>Depression for both Patient and Carer</td>
</tr>
<tr>
<td>Patient social support at 6 months (Piamjariyakul et al., 2015)</td>
<td>Patient/Carer depression at 12 months (Ågren et al., 2015; Agren et al., 2012) and 24 months (Liljeroos et al., 2015)</td>
<td>Social support for both Patient and Carer</td>
</tr>
<tr>
<td>Carer confidence (Piamjariyakul et al., 2015)</td>
<td>Carer social support at 3 months (Schwarz et al., 2008)</td>
<td></td>
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<tr>
<td>Patient anxiety (Kenealy et al., 2015)</td>
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<tr>
<td>Carer strain (Piette et al., 2015)</td>
<td>Carer burden (Agren et al., 2012; Agren et al., 2015)</td>
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<tr>
<td><strong>Economic Outcomes</strong></td>
<td>Patient rehospitalization at 6 months (Hasanpour-Dehkordi et al., 2015)</td>
<td>Rehospitalization for Patient</td>
</tr>
<tr>
<td>Patient rehospitalization (Kenealy et al., 2015) at 2 (Bull et al., 2000)</td>
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<tr>
<td>Positive Outcomes</td>
<td>Null Outcomes</td>
<td>Mixed Outcomes</td>
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<tr>
<td>al., 2016; Piamjariyakul et al., 2015</td>
<td>and 3 months(Schwarz et al., 2008)</td>
<td>Patient cost(Kenealy et al., 2015)</td>
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<td></td>
<td></td>
<td>Patient QALY(Agren et al., 2013)</td>
</tr>
</tbody>
</table>

*Time of Measurement in months is provided for mixed outcome findings. Legend: QALY-Quality adjusted life years*
Papers reporting positive outcomes include findings in all four categories (cognitive, behavioral, affective, and health services utilization). In the main outcome papers of the complete studies only one paper reported null findings on all outcomes. (Schwarz et al., 2008) In the five papers which reported secondary outcomes or primary outcome variables at a later time null findings were noted. (Agren et al., 2013; Agren et al., 2015; Liljeroos et al., 2015, 2016; Liljeroos, Agren, Jaarsma, & Stromberg, 2014a) These included outcomes such as quality adjusted life years, (Agren et al., 2013) carer tasks, burden or patient morbidity, (Agren et al., 2015; Liljeroos et al., 2016) or non-sustaining differences at a subsequent measurement time. (Liljeroos et al., 2015; Liljeroos et al., 2014a) Deeper analysis identified perceived control, quality of life, preparedness to care, self-care, depression, social support and rehospitalization as the outcome variables with mixed (positive and null) findings (Table 3). One example highlights this: perceived control, as measured by the Control Attitude Scale, was reported in three papers. (Ågren et al., 2015; Agren et al., 2012; Srisuk et al., 2015) In the first RCT (Agren et al., 2012) patients in the intervention arm showed significant improvements in perceived control at three months which was not sustained at 12 months while perceived control in carers did not change. In the second RCT (Ågren et al., 2015) patients in the intervention arm showed significant improvements in perceived control at both three and 12 months with no changes in perceived control in carers. In the third RCT (Srisuk et al., 2015) perceived control was only measured in carers. Carers in the intervention arm showed significant improvement in perceived control at three months, but as in the first study, this improvement was not sustained at six months. The intervention components were similar (education, support, and guidance) yet heterogeneous enough in terms of interventionist, time allocation, and delivery modes making it difficult to attribute either positive or null outcomes to the intervention.

Future directions/ongoing studies

Eight papers (six abstracts, (Bakitas et al., 2016; Chung, Lennie, et al., 2014; Chung, Moser, et al., 2014; Deek et al., 2015; Demers et al., 2014; McIlvennan et al., 2015) one feasibility study, (Dionne-
Odom et al., 2014) and one protocol paper (Taylor et al., 2015) met the criteria of ongoing studies. We were unable to assess sufficient information on components, contexts, and outcomes to include them in our previous analyses. However, they indicate the direction of future research if these studies advance to full clinical trials. Two of the studies (one abstract, one feasibility study; (Bakitas et al., 2016; Dionne-Odom et al., 2014) both from the same research team), adapted a successful concurrent oncology-palliative care model to heart failure. Two of the studies, (both abstracts; (Chung, Lennie, et al., 2014; Chung, Moser, et al., 2014) again from a single team), were small RCTs which tested, 1) a cognitive educational intervention and 2) a technology intervention. The three remaining studies examined decision aids; (McIlvennan et al., 2015) family education; (Deek et al., 2015) a multi-component trial comprised of a bathroom scale, decision aid, education and discharge summary to the primary care provider. (Demers et al., 2014) Finally, the protocol paper described a health professional facilitated, home-based, rehabilitation intervention (Taylor et al., 2015). Taken together the ongoing studies appear like the complete studies in numbers and types of components, contexts and outcomes measures.

Discussion

Synthesis of the Interventions

The purpose of this paper was to review the existing literature on dyadic self-care interventions targeting patients with heart failure and their informal carers. We focused on intervention components tested to date, the contexts in which they were implemented, and the effect of these interventions on patient or carer outcomes. In summary, the included studies had great heterogeneity attributable to varying trial designs with a wide range of intervention components, follow-up periods, and outcome variables. The 12 complete studies included between two-six components generally delivered in a hospital or clinic, used increased amounts of technology over time, and resulted in mixed patient and/or carer outcomes at two weeks to 24 months. The eight ongoing studies were similarly designed and implemented. All of this taken together makes it challenging to recommend which interventions should
be considered for wide-spread implementation or further development. However, components confirmed as important in qualitative, observational and intervention studies, such as assessing mutuality in patient-carer dyads, receiving joint but individualized education, having long-standing formal and informal social support throughout the illness trajectory should be emphasized. (H. G. Buck, Hupcey, J., Watach, A., 2017; Hooker, Schmiege, Trivedi, Amoyal, & Bekelman, 2017; Liljeroos, Agren, Jaarsma, & Stromberg, 2014b)

All studies (complete and ongoing) were judged to be of low to moderate quality. Each study had some methodological limitations, such as weak linkages to theoretical frameworks, small sample sizes, paucity of reported intervention detail, choice of outcome variables known to have floor and ceiling effects, and mixed intervention effects. In general, the studies should have been described in more detail. This lack of information limited our ability to recommend a particular intervention as a starting point for future development. When compared with the CONSORT guidelines (Moher, Schulz, Altman, & Group, 2001) and the template for intervention description and replication (TIDieR) checklist and guide, (Hoffmann et al., 2014) there were significant amounts of missing information in several of the papers. However, based on the information that was provided, we can conclude that the body of evidence is limited and recent. In the following section, we will review specific findings and make recommendations for future work to advance the field of heart failure care.

**Dyadic intervention components**

Examination of the individual intervention components revealed there was great complexity within interventions and great heterogeneity across interventions. While there was an overarching, common logic that dyads need education, support, and guidance, there was also a sense that investigators were not able to determine what or how much of a specific component would result in better outcomes: the result was the inclusion of multiple components in all the interventions. This uncertainty is a challenge that may benefit from theoretical and methodological solutions. In particular,
there is a need for careful development of theoretical frameworks with clear hypotheses and rationales for why and how an intervention should work. Given the fact that most of the interventions can be labelled as complex interventions, larger sample sizes, as well as mixed methods, should be considered for outcome evaluations.

Dyadic intervention contexts

This review also revealed a shift in delivery context over time. From a dyadic intervention in the early 2000s (Bull et al., 2000) which relied heavily on face-to-face interaction supplemented with paper information sources to several papers published 15 years later (Kenealy et al., 2015; Piamjariyakul et al., 2015; Piette et al., 2015) which primarily employed technology-based interventions, we were able to track the shift into telehealth as a means to provide chronic heart failure management. Notably what did not change over time was the lack of robust improvements in measured outcomes. Other larger technology-based interventions in heart failure patient populations (Chaudhry et al., 2010; Ong et al., 2016) have reported similar null findings to those found in these dyadic papers. In these rigorous patient trials, intervention adherence was a critical failure. Even when operationalized and measured as only 50% adherence to a technology protocol, almost half of participants were non-adherent over time despite adherence boosts such as reminder phone calls. This suggests that technology, itself, will not improve heart failure outcomes. We need to systematically examine the social determinants of self-care and then test, rather than assume, that a technological solution is likely to improve outcomes.

Dyadic intervention outcomes

The lack of robust and sustained intervention outcomes is particularly concerning. A likely contributor to this may be that the papers included in this review had large variations in patient and carer characteristics as well as cultures and health care systems where they were conducted. There may be specific subgroups of patients and carers that might benefit more, or not at all, from dyadic self-care interventions. Developing such knowledge will enable self-care interventions to target dyadic groups
anticipated to benefit most, which may become indispensable in times of decreasing health care resources. A second contributor to the lack of robust outcomes may be the lack (or at least unreported) stakeholder engagement before and during the intervention design phase. Co-design models have considerable potential to address the current limitations, particularly in technology-based interventions. Experience-based co-design is a user-focused approach that facilitates the access of patients’, family members’ and professionals’ experience when designing new innovative interventions in health care. Users and researchers work together as partners throughout the process and stages of change to provide a deeper understanding of the strengths and limitations of an intervention and what needs to be redesigned for the future. (Bate & Robert, 2006) In our current patient-centered care environment, engaging patients early in any practice change is critically important to avoid changes that are unacceptable to patients and families.

Limitations

Every review has certain limitations that should be kept in mind when examining the findings. Search terms used, databases searched, and analytic techniques employed all shaped the results. We attempted to mitigate these limitations as much as possible by engaging two medical librarians with expertise in literature searches, including only team members with dyadic expertise, and careful development and registration of a protocol. However, other dyadic intervention studies, particularly those not published in English, may not have captured.

Conclusion

In conclusion, the body of evidence for dyadic interventions in heart failure is small. All the papers in the current review had some methodological limitations, mixed intervention effects and counter-intuitive outcomes. Clearly, it is time to change the design, development, and implementation of dyadic heart failure interventions. Important steps to improve future interventions involve more advanced sample size estimations, innovative solutions to increase recruitment of dyads and decrease
attrition during follow-up, use of mixed methods for capturing a more holistic picture of effects and finally studying implementation processes are warranted, as this is the last step of a successful intervention.
References


Appendix A: Search Methodology

The medical librarians (AMH, RLP) developed and performed searches in PubMed (1946 to present), EMBASE (Elsevier, 1947 to present), Web of Science (1900 to present), PsycINFO (EBSCO, 1887 to present), and Cochrane Central Register of Controlled Trials (CENTRAL, through August 2016).

MeSH terms, EMTREE terms, keywords, and keyword phrases were used to search for the following concepts: dyadic, caregivers, heart failure, and intervention. Search terms were shared with the subject experts to elicit feedback. The results were cross-referenced with various clinical study types and were not limited to randomized controlled trials. The librarians applied an English-only filter but no publication date limit.

PubMed


EMBASE

('caregiver'/exp OR 'family'/de OR 'adult child'/exp OR 'informal care':ab,ti OR 'spouse'/exp OR spouse*:ab,ti OR husband:ab,ti OR wife:ab,ti OR family:ab,ti OR families:ab,ti OR 'son'/exp OR son:ab,ti OR 'daughter'/exp OR daughter:ab,ti OR 'cohabiting person'/exp OR partner*:ab,ti OR couple*:ab,ti OR carer:ab,ti OR carers:ab,ti OR dyad*:ab,ti) AND ('heart failure'/de OR 'congestive heart failure'/de OR 'diastolic heart failure'/exp OR 'systolic heart failure'/exp OR 'heart failure':ab,ti OR 'cardiac failure':ab,ti OR 'heart decompensation':ab,ti) AND (Intervention*[ab,ti] AND ('cohort analysis'/exp OR 'cohort analysis':ab,ti) OR 'cohort study':ab,ti OR 'cohort studies':ab,ti OR 'follow-up study':ab,ti OR 'follow-up studies':ab,ti OR 'clinical study'/exp OR 'clinical study':ab,ti OR 'clinical studies':ab,ti OR 'longitudinal study':ab,ti OR 'longitudinal studies':ab,ti OR 'prospective study':ab,ti OR 'prospective studies':ab,ti OR 'retrospective study':ab,ti OR 'retrospective studies':ab,ti OR 'randomized controlled trial*':ab,ti OR rct:ab,ti OR 'pre intervention*':ab,ti OR preintervention*:ab,ti
The combined searches yielded 1,271 citations as of September 29, 2016. All citations were exported into EndNote and duplicates were mechanically removed. The remaining citations (890) were imported into an MS EXCEL worksheet for purposes of documenting the inclusion/exclusion analysis performed by the subject matter specialists.