RESEARCH PAPER

The nurse-coordinated cardiac care bridge transitional care programme: a randomised clinical trial

PATRICIA JEPMA†,1,2, LOTTE VERWEIJ†,1,2, BIANCA M. BUURMAN†,2,3, MICHEL S. TERBRAAK2, SARA DALIRI4, CORINE H.M. LATOUR2, GERBEN TER RIET1,2, FATMA KARAPINAR - ÇARKIT4, JILL DEKKER5, JOSE L. KLUNDE6, SU-SAN LIEM7, ARNO H.M. MOONS6, RON J.G. PETERS1, WILMA J.M. SCHOLTE OP REIMER1,8

1Amsterdam UMC, Department of Cardiology, Amsterdam, the Netherlands
2Amsterdam University of Applied Sciences, Faculty of Health, Centre of Expertise Urban Vitality, Amsterdam University of Applied Sciences, Amsterdam, the Netherlands
3Amsterdam UMC, Department of Internal Medicine, Section of Geriatric Medicine, Amsterdam, the Netherlands
4OLVG Hospital, Department of Clinical Pharmacy, Amsterdam, the Netherlands
5Bovenij Medical Centre, Department of Cardiology, Amsterdam, the Netherlands
6OLVG Hospital, Department of Cardiology, Amsterdam, the Netherlands
7Amstellaard Hospital, Department of Cardiology, Amstelveen, the Netherlands
8HU University of Applied Sciences Utrecht, Research Group Chronic Diseases, Utrecht, the Netherlands

Address correspondence to: Patricia Jepma, Amsterdam University Medical Centre, department of Cardiology, Centre of Expertise Urban Vitality, Amsterdam University of Applied Sciences, Tafelbergweg 51 Amsterdam, 1105 BD the Netherlands. Email: p.jepma@amsterdamumc.nl

†Authors Patricia Jepma and Lotte Verweij equally contributed to this manuscript.

Abstract

Background: after hospitalisation for cardiac disease, older patients are at high risk of readmission and death.
Objective: the cardiac care bridge (CCB) transitional care programme evaluated the impact of combining case management, disease management and home-based cardiac rehabilitation (CR) on hospital readmission and mortality.
Design: single-blind, randomised clinical trial.
Setting: the trial was conducted in six hospitals in the Netherlands between June 2017 and March 2020. Community-based nurses and physical therapists continued care post-discharge.
Subjects: cardiac patients ≥ 70 years were eligible if they were at high risk of functional loss or if they had had an unplanned hospital admission in the previous 6 months.
Methods: the intervention group received a comprehensive geriatric assessment-based integrated care plan, a face-to-face handover with the community nurse before discharge and follow-up home visits. The community nurse collaborated with a pharmacist and participants received home-based CR from a physical therapist. The primary composite outcome was first all-cause unplanned readmission or mortality at 6 months.
Results: in total, 306 participants were included. Mean age was 82.4 (standard deviation 6.3), 58% had heart failure and 92% were acutely hospitalised. 67% of the intervention key-elements were delivered. The composite outcome incidence was 54.2% (83/153) in the intervention group and 47.7% (73/153) in the control group (risk differences 6.5% [95% confidence intervals, CI −4.7 to 18%], risk ratios 1.14 [95% CI 0.91–1.42], P = 0.253). The study was discontinued prematurely due to implementation activities in usual care.
Conclusion: in high-risk older cardiac patients, the CCB programme did not reduce hospital readmission or mortality within 6 months.
Trial registration: Netherlands Trial Register 6,316, https://www.trialregister.nl/trial/6169
hospitalised cardiac patients.

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Introduction

The incidence of readmission and mortality in older patients with cardiovascular disease is rising [1, 2]. Hospital treat-
ment of older cardiac patients is commonly disease-oriented with interventions based on disease-specific guidelines. However, geriatric conditions such as functional impair-
ment, fall risk and malnutrition [3] often go unreconised although they increase the risk of adverse events [4, 5].

The transitional phase, when patients transfer from hos-
torial hospital, is a high-risk period for adverse events [6]. Medication-related problems are common [7] and symp-
toms of physical deterioration often stay unreconised [8]. Furthermore, participation in cardiac rehabilitation (CR) programmes is low [9]. As CR is effective in older patients [9], non-participation could increase the risk of recurrent cardiovascular events and mortality [10].

Transitional care has been shown effective in reducing hospital readmission and mortality [11–13]. However, results are inconclusive in older cardiac patients [14–17]. Most transitional care interventions are provided from a case management perspective, delivering interventions with a broad focus on patients’ needs [6, 17]. The integration of disease management and tailored home-based CR into transitional care interventions may be necessary.

The purpose of this study was to evaluate the effects on unplanned hospital readmission and mortality of the nurse-coordinated 'cardiac care bridge (CCB) transitional care programme' which combines case management, disease management and home-based CR in high-risk older hospitalised cardiac patients.

Methods

Study design and setting

We tested the CCB programme in a parallel single-blind multicentre randomised trial, performed between 5 June 2017 and 31 March 2020 in six hospitals surrounding Amsterdam, the Netherlands. Community nurses (CNs) and community-based physical therapists (PT) continued care post-discharge. The trial design has been published [18]. The study was approved by the Medical Ethics Committee of the Amsterdam University Medical Centre (Protocol ID: MEC2016_024) and registered in the Dutch Trial Register (NTR6316, April 6, 2017). The results are reported according to the Consort (Consolidated Standards of Reporting Trials) statement [19].

Study population

Cardiac patients of ≥70 years, admitted to the departments of cardiology or cardiothoracic surgery and admitted ≥48 h were eligible if they were at high risk of functional loss according to the screening instrument for frail older people of the Dutch Safety Management System (DSMS) [20]. Four geriatric conditions (limitation in activities of daily living [ADL], falls, malnutrition and delirium) are part of this frailty tool, and the DSMS-score ranges between 0 and 4 (Appendix 1, Supplementary data are available in Age and Ageing online). Patients were considered at high risk with a DSMS-score ≥ 2 in patients aged 70–79 years or DSMS-score ≥ 1 in patients aged ≥80 years [21]. Regardless of the DSMS-score, we also included patients with an unplanned hospital admission in the prior 6 months as this is associated with increased risk for adverse events [22].

Exclusion criteria were (i) inability to provide consent and follow instructions due to severe cognitive impairment (mini-mental state examination, MMSE < 15) or delirium as confirmed by the treating physician, (ii) congenital heart disease, (iii) life expectancy of ≤3 months as estimated by the treating physician, (iv) transfer from or planned discharge to a nursing home, (v) planned discharge to another depart-
ment or hospital not participating in this study and (vi) inability to communicate.

Randomisation

The consent procedure and randomisation were performed ≤72 h after admission. According to the postponed informed consent procedure of Boter et al. [23], study participants were blinded to the specific study aims to prevent a potential Hawthorne effect [24]. At the end of the study, participants were fully informed about the intervention and treatment allocation. Stratified block randomisation to the intervention or control group, allocation ratio 1:1, was used with pre-stratification by study site and cognitive status (MMSE 15–23 vs ≥24). Allocation concealment was ensured by a web-based data management programme (Research Manager, https://my-researchmanager.com/en/) and random permuted blocks of two, four and six were used.

Usual care

All patients received a comprehensive geriatric assessment (CGA) at baseline by a cardiac research nurse. The control group continued with usual care including consultation by other disciplines during hospitalisation, outpatient visits to
Transitional care in older cardiac patients

Figure 1. Overview of the CCB programme.

the cardiologist and cardiac nurse specialist and centre-based CR if indicated. In addition, standard care was provided by the family physician. The Dutch healthcare system is described in Appendix 2 (Supplementary data are available in Age and Ageing online).

Intervention

The CCB programme was performed in three phases (Figure 1): the clinical, discharge and post-clinical phase. The intervention consisted of three care components: (i) case management, (ii) disease management and (iii) home-based CR. The intervention key-elements are described below and in Appendix 3 (Supplementary data are available in Age and Ageing online). All involved healthcare professionals received a post-Bachelor-level training in case management, disease management and CR (Appendix 4, Supplementary data are available in Age and Ageing online). Informal caregivers were involved in the intervention if they were present.

In the clinical phase, health issues identified by the CGA were discussed and prioritised by the cardiac research nurse and the participant. An integrated care plan based on patients’ goals was formulated, which was leading during the intervention. A geriatrician and other disciplines (e.g. dietician) were consulted based on CGA findings.

The discharge phase started when the discharge date was set. The cardiac research nurse contacted the CN and PT to arrange the post-clinical phase. In hospital, the CN visited the participant and the cardiac research nurse for a handover of the integrated care plan, and information about participants’ medical condition and treatments. In addition, the medical discharge letter was sent to all post-discharge CCB healthcare professionals.

In the post-discharge phase the CN planned home visits within 3 days, and 1, 3 and 6 weeks after discharge and an additional home visit within 12 weeks if necessary. During home visits, the CN reviewed the integrated care plan, participants’ health status, medication and potential drug-related problems (DRPs) including side-effects and inappropriate use. Together with the CCB pharmacist, medication reconciliation was performed during the first home visit. DRPs were signalled by the CN using the Red Flag instrument [25]. Issues were discussed with the pharmacist who proposed adjustments. For questions regarding participants’ health status, the CN contacted e.g. the general practitioner or cardiologist based on indication.
Furthermore, the PT provided one or two home-based CR sessions per week, with a maximum of nine sessions during the first 6 weeks post-discharge according to the Dutch CR guideline [26]. The first home visit by the PT was a joint intake with the CN and the participant to discuss goals and desired activities, which led to a rehabilitation plan. Depending on participants’ functional status a stepwise graded exercise approach was followed, including improving functional activities (e.g. rising from chair, walking and climbing stairs) and increasing muscle strength.

### Primary and secondary outcomes

The primary outcome was a composite of first all-cause unplanned readmission or mortality within 6 months after randomisation. We defined an unplanned readmission as a non-elective admission ≥1 night. Secondary outcomes included the composite outcome at three and 12 months after randomisation and the incidence of the first all-cause unplanned hospital readmission and mortality separate at 3, 6 and 12 months. Mortality data were collected from medical files and the Dutch National Personal Records Database [27]. Data on readmissions were collected from medical files in the participating hospitals and supplemented with participants’ self-reported readmissions to other hospitals. Data collection were performed by research nurses who were blinded to the treatment allocation.

### Sample size calculation

The sample size calculation was based on a comparable study of 101/674 hospitalised cardiac patients ≥65 years at high risk of functional loss [13]. Based on a 6 month incidence of 44% (readmission and mortality combined) in the usual care group and a minimal important difference of 12.5% in absolute risk reduction (from 44 to 31.5%) in participants in the intervention arm (chi-square test and a 2-sided alpha of 0.05; power of 80%), a sample size of 235 participants per group was required. To compensate for an assumed 5% loss of 0.05; power of 80%), a samplesize of 235 participants per group was required. To compensate for an assumed 5% loss to follow-up, the total intended sample size per group was 250.

### Statistical analyses

Analyses were performed according to a predefined statistical analyses plan based on the intention-to-treat principle (Appendix 5, Supplementary data are available in Age and Ageing online).

We reported univariable outcomes and presented the multivariable models in the appendices as both analyses revealed comparable results. The treatment effect of the primary and secondary outcomes was expressed as risk differences (RD) and risk ratios (RR) and the corresponding 95% confidence intervals (CIs) based on a chi-square test.

Multivariable logistic and Cox regression analyses were performed and resulting adjusted odds ratios (OR) were transformed into RRs [28]. We adjusted for frailty status, study site, age, sex, any admissions in the previous 6 months, Charlson comorbidity score, MMSE, cardiovascular diagnosis and living arrangement. In addition, we checked for treatment interaction with the following predefined subgroup analyses: age, frailty status, any unplanned hospital admission in the previous 6 months, cognitive impairment and diagnosis at index admission. Correction for (semi-)competing risk was performed by a unidirectional transition multistate model (illness-death model; [29, 30]).

All statistical tests were 2-sided. P-values < 0.05 were considered statistically significant. Analyses were performed with SPSS 25.0 (SPSS Inc., Chicago, IL, USA) and Stata Statistical Software: Release 13 (College Station, TX: StataCorp LP).

### Intervention fidelity

Fidelity to key-elements of the intervention was registered by CCB healthcare professionals and evaluated by quality indicators [31]. For each participant, the denominator of the intervention key-elements was set to the number of feasible key-elements. Key-elements missed due to e.g. hospital readmission, death or disabilities that precluded participants from taking part in any key-element, were not deemed feasible and not counted in the denominator. The mean fidelity rate was calculated per intervention key-element and in addition for each participant, we calculated the mean fidelity percentage across all key-elements that a participant was entitled to. The overall adherence percentage across all 153 participants was calculated as an unweighted average of the participant-specific percentages.

### Results

We screened 6,857 patients for enrolment, 623 patients (9%) were eligible for participation (Figure 2). Most exclusions were due to low DSMS-scores (59%). In total, 306 eligible patients provided informed consent (49%) and were randomised (153/153). Inclusion was discontinued prematurely on 31 March 2019 because of contrast problems between intervention and control due to an increase in implementation activities in usual care (e.g. home-based follow-up and the Red Flag instrument) [25]. Outcome data were complete for all included participants (follow-up until 31 March 2020).

On average, participants were 82.4 years old (SD 6.3) and 51% were male. Participants were mostly admitted for HF (58%) and 45% had had an unplanned hospital admission in the previous 6 months. In total, 56% were at risk of delirium, 47% had fallen in the 6 months prior to admission, 39% had ADL-limitations and 33% had malnutrition (Table 1).

### Primary outcome

The incidence of the 6-month composite outcome of first all-cause readmission or mortality was 54.2% (83/153) in the intervention group and 47.7% (73/153) in the control group.
Table 1. Baseline characteristics

| Sociodemographics | Measurement | Intervention (n = 153) | Control (n = 153) |
|-------------------|-------------|------------------------|------------------|
| **Age**           |             |                        |                  |
| 70–79 years       | 40 (26.1%)  | 51 (33.3%)             |                  |
| ≥ 80 years        | 113 (73.9%)| 102 (66.7%)            |                  |
| **Sex**           |             |                        |                  |
| Male              | 70 (45.8%)  | 86 (56.2%)             |                  |
| **Country of origin** |         |                        |                  |
| Netherlands       | 135 (88.2%)| 138 (90.2%)            |                  |
| **Level of education** |       |                        |                  |
| Primary education | 66 (43.1%)  | 61 (39.9%)             |                  |
| Secondary education | 52 (34.0%)| 44 (28.8%)             |                  |
| Higher education  | 35 (22.9%)  | 47 (30.7%)             |                  |
| **Cohabiting**    |             |                        |                  |
| 66 (43.1%)        |             | 68 (44.4%)             |                  |
| **Socioeconomic status** |    |                        |                  |
| Low (< 1 SD)      | 25 (16.3%)  | 27 (17.6%)             |                  |
| Intermediate      | 83 (54.2%)  | 81 (52.9%)             |                  |
| High (> 1 SD)     | 45 (29.4%)  | 45 (29.4%)             |                  |
| **Index hospitalisation** |     |                        |                  |
| Acute hospitalisation | 139 (90.8%) | 141 (92.2%)           |                  |
| Length of stay    | 7 [4–10] | 7 [4–10] |                  |
| **Diagnosis on admission** |           |                        |                  |
| Heart failure     | 86 (56.2%)  | 91 (59.5%)             |                  |
| **Actce coronary syndrome** | 19 (12.4%) | 24 (15.7%) |                  |
| Valve deficits    | 14 (9.2%)   | 12 (7.8%)              |                  |
| Other             | 7 (4.6%)    | 6 (3.9%)               |                  |
| **Treatment during admission** |      |                        |                  |
| Medical treatment only | 115 (75.2%) | 116 (75.8%)         |                  |
| PCI               | 13 (8.5%)   | 15 (9.8%)              |                  |
| TAVR              | 15 (9.8%)   | 11 (7.2%)              |                  |
| Device implantation | 12 (7.8%)  | 10 (6.5%) |                  |
| Other             | 1 (0.7%)    | 4 (2.6%)               |                  |
| **Inclusion criteria** |           |                        |                  |
| Previous hospital admission | ≤ 6 months prior to index event | 66 (43.1%) | 73 (47.7%) |                  |
| Delirium          | 94 (61.4%) | 77 (50.3%)             |                  |
| Activities of daily living | 65 (42.5%) | 54 (35.3%)          |                  |
| Median (KATZ-6)   | 1 [0–3] | 0 [0–2] |                  |
| ADL-functioning   | 72 [38–84] | 76 [63–86] |                  |
| Malnutrition      | 57 (37.3%) | 43 (28.1%)             |                  |
| Fall risk         | 67 (43.8%) | 78 (51.0%)             |                  |
| Fear of falling   | 63 (41.2%) | 66 (43.1%)             |                  |
| DSM5 scorec       | 13 (8.5%)  | 13 (8.5%)              |                  |
| DSM5 1            | 49 (32.0%) | 59 (38.6%)             |                  |
| DSM5 2            | 50 (32.7%) | 57 (37.3%)             |                  |
| DSM5 3            | 33 (21.6%) | 19 (12.4%)             |                  |
| DSM5 4            | 8 (5.2%)   | 5 (3.3%)               |                  |
| **Medical history** |           |                        |                  |
| Heart failure     | 105 (68.6%)| 110 (71.9%)            |                  |
| Hypertension      | 95 (62.1%) | 94 (61.4%)             |                  |
| Acute coronary syndrome | 57 (37.3%) | 53 (34.6%) |                  |
| Atrial fibrillation | 54 (35.3%) | 59 (38.6%) |                  |
| Diabetes mellitus | 52 (34.0%) | 47 (30.7%)             |                  |
| Renal failure     | 51 (33.3%) | 59 (38.6%)             |                  |
| Chronic obstructive pulmonary disease | 29 (19.0%) | 24 (15.7%) |                  |
| Peripheral vascular disease | 29 (19.0%) | 21 (13.7%) |                  |
| Cerebrovascular accident | 23 (15.0%) | 27 (17.6%) |                  |
| **Lifestyle factors** |           |                        |                  |
| Current smoker    | 16 (10.5%) | 14 (9.2%)              |                  |
| Body mass index   | 26.8 (5.9) | 25.8 (4.6)             |                  |
| **Geriatric conditions** |        |                        |                  |
| Cognitive impairment | MMSE 15–23 | 47 (30.7%) | 48 (31.4%) |                  |
| Comorbidities     | Charlson Comorbidity Score | 3 [1–4] | 3 [1–4] |                  |
| Depressive symptoms | GDS ≥ 6   | 22 (14.6%) | 18 (11.8%) |                  |
| Anxiety           | HADS-A ≥ 8 | 18 (11.9%) | 24 (15.7%) |                  |
| Dyspnoea          | Self-reported | 125 (81.7%) | 123 (80.4%) |                  |
| Fatigue           | NRS ≥ 4   | 114 (74.5%) | 114 (74.5%) |                  |

Continued
**Table 1. Continued**

| Sociodemographics | Measurement                             | Intervention (n = 153) | Control (n = 153) | Risk difference (%) | Risk ratio (95% CI) | P-value risk ratio |
|-------------------|-----------------------------------------|------------------------|-------------------|---------------------|---------------------|-------------------|
| Dizziness         | Self-reported                           | 65                     | 42.5%             |                     |                     |                   |
| Urine incontinence| Self-reported                           | 42                     | 27.5%             |                     |                     |                   |
| Polypharmacy      | ≥ 5 (from medication overview)          | 141                    | 92.2%             |                     |                     |                   |
| Medication side effects | Self-reported           | 34                     | 22.2%             |                     |                     |                   |
| Functional status | SPPB                                    | 4                      | [2–6]             | 5                   | [3–7]              |                   |
| Handgrip strength | Male (norm > 30 kg)                     | 26.4                   | 9.2               | 27.0                | (7.8)              |                   |
|                   | Female (norm > 18 kg)                   | 16.1                   | (5.8)             | 15.3                | (4.7)              |                   |

Note: (SD), [25–75 percentile]. Abbreviations: ALDS, Amsterdam Linear Disability Scale; CABG, Coronary Artery Bypass Grafting; DSMS, Dutch Safety and Management System; GDS, Geriatric Depression Scale; HADS-A, Hospital Anxiety and Depression Scale-Axiety; MMSE, Mini-Mental State Examination; NRS, numeric rating scale; PCI, Percutaneous Coronary Intervention; SNAQ, Short Nutritional Assessment Questionnaire; SPPB, Short Physical Performance Battery; TAVR, Transcatheter Aortic Valve Replacement. *Primary education: elementary or primary school. Secondary education: pre-vocational, senior general or pre-university. Higher education: higher professional or university. †Socioeconomic status was calculated from the postal code of patients’ residence by the Netherlands Institute for Social Research (SCP) and based on income, employment and educational level. ‡Dutch Safety Management System [20]: the score between 0 and 4 points, based on four domains of frailty (malnutrition, risk of impairments in daily functioning, risk on delirium and fall risk). A higher score on the DSMS indicates a higher risk of functional loss. Dominant hand highest value.

**Table 2. Primary and secondary outcomes in the CCB study**

| Secondary outcome | Intervention n = 153 (%) | Control n = 153 (%) | Risk difference (%) (95% CI) | Risk ratio (95% CI) | P-value risk ratio |
|-------------------|--------------------------|---------------------|-------------------------------|---------------------|-------------------|
| Composite outcome |                          |                     |                               |                     |                   |
| 3 months          | 63 (41.2)                | 59 (38.6)           | 2.6 (−8.4–13.6)              | 1.07 (0.81–1.41)    | 0.641             |
| 6 months          | 83 (54.2)                | 73 (47.7)           | 6.3 (−4.7–18.0)              | 1.14 (0.91–1.42)    | 0.253             |
| 12 months         | 101 (66.0)               | 88 (57.5)           | 8.5 (−2.4–19.3)              | 1.13 (0.96–1.37)    | 0.126             |
| Unplanned readmission† |                 |                     |                               |                     |                   |
| 3 months          | 45 (29.4)                | 48 (31.4)           | −1.9 (−12.2–8.3)             | 0.94 (0.67–1.32)    | 0.709             |
| 6 months          | 60 (39.2)                | 59 (38.6)           | 0.7 (−10.3–11.6)             | 1.02 (0.77–1.35)    | 0.907             |
| 12 months         | 73 (47.7)                | 70 (45.8)           | 1.9 (−0.2–13.1)              | 1.04 (0.82–1.32)    | 0.731             |
| Mortality         |                          |                     |                               |                     |                   |
| 3 months          | 26 (17.0)                | 20 (13.1)           | 3.9 (−4.1–12.0)              | 1.30 (0.76–2.23)    | 0.337             |
| 6 months          | 36 (23.5)                | 28 (18.3)           | 5.2 (−3.9–14.3)              | 1.29 (0.83–2.00)    | 0.261             |
| 12 months         | 59 (38.6)                | 41 (26.8)           | 11.8 (1.3–22.2)              | 1.44 (1.04–2.00)    | 0.028             |

*Results are not corrected for (semi-)competing risk. Appendix 8 of supplementary data presents the for (semi-)competing risk corrected outcomes in a multi-state (illness-death) model.

The analysis showed similar results (Appendix 6, Supplementary data are available in Age and Ageing online). Uni- and multivariable Cox regression analyses are presented in Appendix 6 (Supplementary data are available in Age and Ageing online). Appendix 8 of supplementary data shows the results of the multi-state illness-death models up to 12 months.

**Intervention fidelity**

In total, the mean participant fidelity percentage across all key-elements that a participant entitled to was 67%. However, the fidelity rates varied widely across the various key-elements (median 60%, IQR [41–69], range [17–100]). Table 3 presents the measures of intervention fidelity per key-element. In total, 75% of all intervention key-elements in the clinical phase were performed, 37% in the discharge phase and 64% in the post-clinical phase.
Discussion

In this study, we found that the CCB programme did not reduce hospital readmission or mortality within 6 months following hospitalisation. Similarly, for the secondary outcomes of unplanned hospital readmission and mortality alone, no statistically significant differences were found. Based on our findings, there is only a limited possibility that the CCB programme would be beneficial.

Systematic reviews on transitional care interventions in patients with HF found that high intensity interventions and (nurse) home visiting programmes reduce the incidence of readmission [11, 14, 15], mortality [11] and the composite endpoint of all-cause readmission and mortality [15]. The discrepancy of these reviews [11, 15] with our findings may be related to a higher mean age (82.4 years versus 70–74 years) and the frailty of the older cardiac population in our trial. In line with our findings, two recent randomised
Figure 3. Kaplan–Meier curve of the composite outcome within 12 months. Dashed line at 90 days marks the end of the intervention period. The curves of the intervention and control group in the primary outcome diverged after the intervention was completed at 90 days follow-up.

Table 3. Intervention fidelity

| Intervention key-elements                                      | N\(^a\) | %  |
|---------------------------------------------------------------|---------|----|
| **Clinical phase**                                            |         |    |
| CGA and CGA-based integrated care plan                        | 153/153 | 100|
| Geriatric consultation based on indication\(^b\)               | 11/66   | 17 |
| **Discharge phase**                                           |         |    |
| Handover                                                      |         |    |
| Face-to-face                                                  | 49/134  | 37 |
| Telephone                                                     | 19/134  | 14 |
| Written                                                       | 66/134  | 49 |
| **Post-clinical phase**                                       |         |    |
| Community nurse home visits\(^c\)                            | 82/133  | 62 |
| First home visit within 72 h after discharge                 | 76/133  | 57 |
| Number of community nurse home visits                        | Median 3| IQR 2–4|
| Medication reconciliation including the Red Flag instrument \(^{[25]}\) | 118/133 | 89 |
| Follow-up of the integrated care plan                         | 71/132  | 54 |
| Lifestyle promotion                                           | 91/132  | 69 |
| Joint home-visit of the physical therapist and community nurse| 33/81   | 41 |
| Home-based CR\(^d\)                                          | 70/116  | 60 |
| Number of home-based rehabilitation sessions                  | Median 4| IQR 2–6|

Mean participant-specific fidelity percentage

|                                                | 153 | 67 |

Abbreviations: CGA, comprehensive geriatric assessment; IQR, interquartile range. \(^a^\)The denominator is set on the number of eligible patients per intervention key-element. \(^b^\)Geriatric team consultation was indicated in case of 1 problem within the psychological domain or 5 geriatric problems in total. \(^c^\)Four home visits, according to the CCB protocol. \(^d^\)Max. nine home-based rehabilitation session, according to the CCB protocol.

Trials in patients with HF \(^{[16]}\) and patients with AMI \(^{[17]}\) reported no significant differences on readmission and mortality.

To our knowledge, our study is the first that combined case management, disease management and home-based CR in frail older patients with a variety of cardiac diagnoses.
However, we did not find that integration of these intervention components improves outcomes. Several factors may have contributed to the results. First, we included a severely frail study population with a high mean age, many disabling comorbidities and geriatric conditions and an extensive medical history. In both groups, mortality rates were high. These factors suggest that the included population may have been beyond the reach of prevention programmes such as the CCB programme. Second, within the high-quality Dutch standard healthcare system many services are being offered to frail older patients which possibly diminished the contrast between the groups (Appendix 2, Supplementary data are available in Age and Ageing online). Third, we observed that real-world circumstances affected the fidelity of this intervention. Our intervention fidelity may have contributed to the lack of effect. A higher fidelity on the intervention key-elements could have resulted in a greater contrast between the intervention and control group. However, we cannot exclude the possibility that full fidelity would have led to even more deleterious effects on mortality due to the detrimental trend in the intervention group, through yet unexplained mechanisms.

An extended process evaluation was performed in parallel to the trial and addresses the barriers and facilitators for intervention fidelity [31]. In brief, low fidelity rates in healthcare professionals were mostly associated with time limits. For example, the short hospital stay and ad hoc discharge planning reduced the opportunity for geriatric consultation or an in-hospital handover of the integrated care plan to the community nurse. For future purpose, geriatric co-management interventions could be considered during hospitalisation in which the responsibility for the treatment is shared between the treating physician and the geriatric team. This kind of intervention intensifies collaboration and has proven to reduce mortality post-discharge [32, 33]. Furthermore, alternative communication routes such as a video call handover between the patient, the hospital and community nurse, may ensure continuity of care while less time-consuming than an in-hospital handover. We explored the unexpectedly high mortality rates in the intervention group. Baseline differences in the population regarding e.g. level of frailty were explored statistically. However, correction in the multivariable analysis yielded essentially the same results. Alternatively, our findings may be due to the play of chance considering the limited statistical power. Previously, Fan et al. [34] performed a comprehensive care programme to reduce hospitalisation in patients with pulmonary disease and found unexplained higher mortality rates among intervention patients.

In this frail older cardiac patients, other interventions with more focus on quality of life may be needed [35]. For example, advance care planning (ACP) may be more suitable as the CCB population seemed unresponsive to high intensity preventive interventions and event rates were high. ACP focuses on patient-centred preferences to increase comfort, quality of life and reduce readmission [36]. Future studies should carefully consider the population eligible for preventive interventions versus those who are eligible for palliative interventions.

Study limitations
The following limitations should be considered. First, only 9% (623/6857) of screened patients were considered eligible for the CCB programme. Most patients were excluded because of low DSMS-scores and having their residence in non-participating residential areas. In total, 49% of eligible patients provided informed consents. Patients more often refuse study participation when they experience mental and physical health problems [37]. Second, we were unable to continue the study until the planned 500 participants due to the quickly (and prematurely) developing regular transitional care for older cardiac patients in our region. This development illustrates that the high rates of readmission and mortality in this high-risk population were being recognised and that professionals seek effective preventive interventions. Unfortunately, as a result of these developments, we were unable to achieve the planned sample size and this clearly impacts on the power of the study.

As it turned out, the overall event rate (51%) was higher than the 44% used in the sample size calculation. This proportion of outcome, which is a much stronger driver of power than sample size, is close to the statistically ideal 50% event rate. The RD point estimate of 6.5% indicates an untoward effect of intervention, leaving only a small tail of the statistical outcome distribution in the range of possible beneficial effects. The trial preserved sufficient statistical precision to render beneficial effects greater than −4.7% in RD (the primary outcome’s lower 95% confidence limit) unlikely. Last, we performed a complex intervention according to a standardised intervention protocol. We invested in an intensive training programme and organised regular follow-up meetings, however, variation in the intervention performance turned out to be inevitable. Our findings reflect the effectiveness and working mechanisms of the intervention under ‘real’ circumstances and the perceived barriers and facilitators showed some important lessons on organizing care for frail older cardiac patients [31].

Conclusion
A randomised trial of nurse-coordinated transitional care compared to usual care in The Netherlands found no reduction of unplanned hospital readmission or mortality within 6 months following hospitalisation in high-risk older cardiac patients. Although the suboptimal intervention fidelity prevented assessment of the effect of the full programme, large beneficial effects are unlikely with our findings. It is also conceivable that the patient population selected may not be responsive to high-intensity preventive strategies.

Supplementary Data: Supplementary data mentioned in the text are available to subscribers in Age and Ageing online.
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