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# Multidisciplinary intervention reducing readmissions in medical inpatients: a prospective, non-randomized study

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**Background:** The purpose of this study was to examine whether a multidisciplinary intervention targeting drug-related problems, cognitive impairment, and discharge miscommunication could reduce readmissions in a general hospital population.

**Methods:** This prospective, non-randomized intervention study was carried out at the department of general internal medicine at a tertiary university hospital. Two hundred medical inpatients living in the community and aged over 60 years were included. Ninety-nine patients received interventions and 101 received standard care. Control/intervention allocation was determined by geographic selection. Interventions consisted of a comprehensive medication review, improved discharge planning, post-discharge telephone follow-up, and liaison with the patient's general practitioner. The main outcome measures recorded were readmissions and hospital nights 12 months after discharge. Separate analyses were made for 12-month survivors and from an intention-to-treat perspective. Comparative analyses were made between groups as well as within groups over time.

**Results:** After 12 months, survivors in the control group had 125 readmissions in total, compared with 58 in the intervention group (Mann–Whitney *U* test,  $P = 0.02$ ). For hospital nights, the numbers were 1,228 and 492, respectively ( $P = 0.009$ ). Yearly admissions had increased from the previous year in the control group from 77 to 125 (Wilcoxon signed-rank test,  $P = 0.002$ ) and decreased from 75 to 58 in the intervention group ( $P = 0.25$ ). From the intention-to-treat perspective, the same general pattern was observed but was not significant (1,827 versus 1,008 hospital nights, Mann–Whitney test,  $P = 0.054$ ).

**Conclusion:** A multidisciplinary approach, targeting several different areas, could substantially lower readmissions and hospital costs in a non-terminal general hospital population.

**Keywords:** medical inpatients, hospital readmissions, intervention, drug-related problems, cognitive impairment, hospital discharge

## Introduction

Hospital readmissions are common in older inpatients, with one fifth being readmitted within 30 days and 60% within a year, and unplanned readmissions accounting for 90%.<sup>1,2</sup> This issue is becoming increasingly important because many countries have aging populations and are reducing hospital bed capacity.<sup>3,4</sup>

Previous studies propose that a proportion of these readmissions are avoidable.<sup>5–9</sup> Numerous interventions have succeeded by targeting a specific condition, primarily heart failure, which is a known risk factor for readmissions.<sup>1,2,10</sup> Another condition, highly prevalent in medical inpatients, but with fewer successful interventions, is cognitive impairment.<sup>11–15</sup> The methods applied in these studies include improved discharge practices and hospital-based follow-up after discharge.<sup>16–19</sup> However, a high-

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quality intervention study that reduced the incidence of delirium demonstrated disappointing results regarding readmissions.<sup>20,21</sup> Apart from specific conditions, more general approaches have also been applied. Adverse drug reactions have been the focus of interventions succeeding in reducing hospital readmissions.<sup>22,23</sup> Another area suggested for intervention is communication gaps in the transition between caregivers.<sup>24,25</sup>

We designed a multidisciplinary intervention applying methods previously shown to be effective, such as discharge improvement and medication overviews, combined with approaches targeting cognitive impairment and miscommunication. Our aim was to apply this intervention in a broad general hospital population and evaluate it regarding readmissions through a prospective approach.

## Materials and methods

### Setting

The city of Malmö in southern Sweden (population 300,000) consists of ten boroughs. The demographics of the boroughs differ and a majority of the city's elderly population lives in two boroughs, from here on called borough A and B. Every borough has a social services department, managing community care services, and 2–3 primary health care centers. Inpatient care in Malmö is provided by the Skåne University Hospital, a 700-bed tertiary hospital. The department of general internal medicine at this hospital contains four wards with 100 beds in total. The four wards have a similar general medical orientation, treating primarily elderly patients with multiple disorders.

Before discharge, hospital staff initiate the coordination of post-discharge services, including community care and primary care follow-up. First, hospital staff obtain information from community care, patients, and relatives to determine if a significant loss of function has occurred before or during hospitalization. If additional support at home is considered desirable or necessary, a multidisciplinary conference is held at the ward, assessing specific needs. The handover of medical responsibility to the general practitioner is managed through a discharge summary (containing main diagnosis, current medication list, follow-up arrangements) sent to the general practitioner on the day of discharge.

### Patients

#### Control versus intervention

Patients living in borough A and B formed the intervention group, with patients from the eight other boroughs as controls,

using convenience sampling. Data collection commenced in November 2009, starting with the control group. When 101 controls were included, data collection began in the intervention phase in February 2010 (Figure 1).

#### Ineligibility and exclusion criteria

Four ineligibility criteria were applied, ie, age under 60 years, living outside Malmö, living in a nursing home, and prior enrollment (a patient could only be included once).

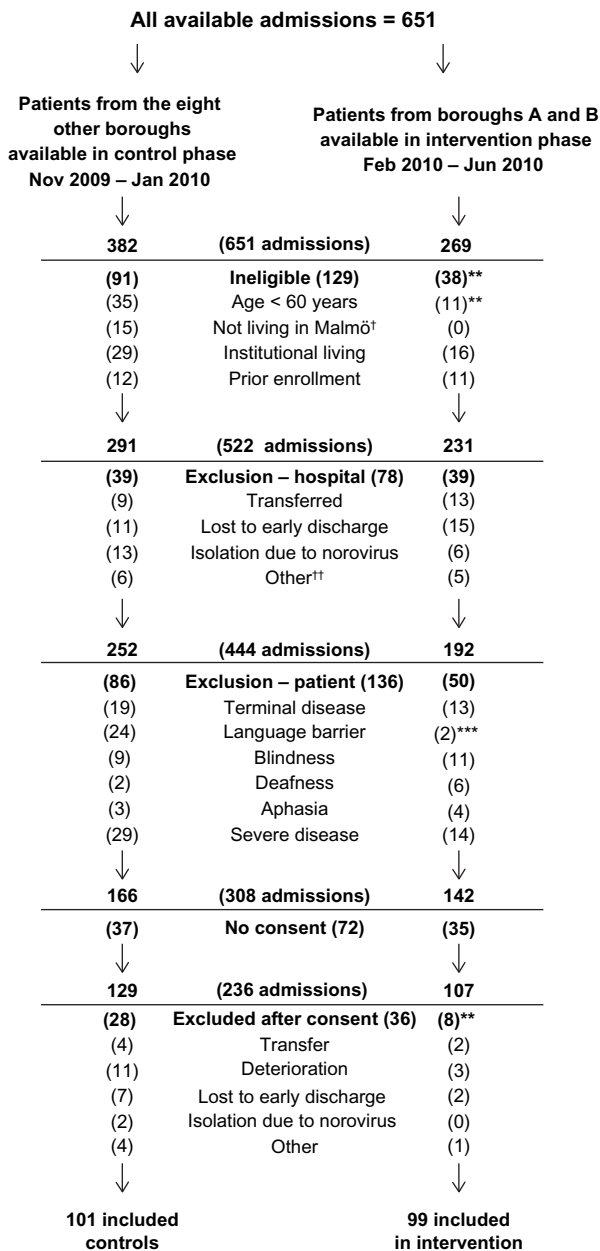


Figure 1 Patient flow and exclusion criteria.

Notes: †Not applicable since address was a criteria for availability in the intervention group; ††five patients were missed due to Easter holidays and six due to Christmas holidays; \*\*P ≤ 0.01, \*\*\*P ≤ 0.001.

Exclusion criteria included aspects related to time restraints and the hospital (transfer to another department, lost to early discharge, isolation due to communicable disease). Patient-related factors that obstructed cognitive tests or made these inappropriate (terminal disease, language barrier, blindness/deafness/aphasia, or severe disease with inability to communicate, eg, because of altered consciousness) also resulted in exclusion.

## Ethics statement

The study was performed according to the Declaration of Helsinki. All included patients gave their written informed consent and the study protocol was approved by the regional ethics committee of Lund University (2009/662).

## Baseline measurements

All baseline measurements were conducted by study staff in a private environment at the wards during office hours. The study staff consisted of one project manager/physician (GT) and three experienced research assistants (one registered nurse and two occupational therapists).

## Demographics and comorbidity

Patients were interviewed about living arrangements, current community care utilization, and educational level. All comorbidities recorded in the electronic medical record during the current and three preceding admissions were noted. Cumulative comorbidity was determined using the Charlson comorbidity index, assigning different weights from 1 to 6 for different disorders, eg, coronary heart disease is weighted 1 and tumor with metastasis 6.<sup>26</sup>

## Cognitive tests and activities of daily living

We applied the Mini-Mental State Examination, scored from 0 to 30, and the Clock-Drawing Test, scored from 0 to 5.<sup>27,28</sup> In both tests, low scores indicate cognitive impairment. Ability to perform activities of daily living (ADL) was quantified using the ADL subset of the GBS (Gottfries-Bråne-Steen) scale.<sup>29</sup> The GBS-ADL scale contains six items (dressing, eating, physical activity, spontaneous activity, personal hygiene, continence), each scored from 0 to 6 for a total score of 0 to 36, with higher scores signifying ADL impairment.

## Preceding health care utilization

For each patient, emergency department visits, hospital admissions, and hospital nights for the preceding 12 months were extracted from the hospital's electronic medical record.

The number of general practitioner visits in the preceding 6 months was determined using the regional health care registry.

## Interventions

### Pharmacist intervention

A clinical pharmacist performed a medication review, using a method called the Lund Integrated Medicines Management model.<sup>30</sup> First, the patient's most accurate list of medications was established from structured interviews, records from primary care, community care, and the National Pharmacy Register.<sup>31</sup> The list was compared with the current list at the hospital. Unintentional discrepancies, known as medication errors, were noted and classified into five groups, ie, omission of drug, erroneous addition of drug, dose too high, dose too low, and wrong dosage form (eg, sustained release).<sup>32</sup>

Moreover, throughout the hospitalization, the pharmacist identified and monitored drug-related problems using interviews, hospital records, laboratory values, and physiologic data.<sup>33</sup> Drug-related problems were classified as unknown indication for treatment, dose not adapted to renal/liver function, inappropriate drug in the elderly (according to hospital policy, based on the recommendations of the National Board of Health and Welfare), adverse drug reaction, untreated or not optimally treated indication, transferring error at discharge, non-adherence, and drug handling (eg, problems with swallowing or crushing).<sup>34</sup> Based on medication errors and drug-related problems, a recommendation was developed and delivered to the ward physician, who made all decisions regarding medications.

### Discharge conference

If a discharge conference was required, the social services were informed beforehand by study staff regarding cognitive test results. Thus, everyone was aware of any cognitive deficits in advance and could prepare accordingly for the conference. Study staff attended all conferences, conveying cognitive and ADL impairment in a standardized way, based on the Mini-Mental State Examination, Clock-Drawing Test, and GBS-ADL scales. Numbers of discharge conferences, discharge destinations, and length of stay were recorded for evaluation.

### Telephone follow-up

The registered nurse was assigned the role of communication nurse. The communication nurse met with all patients and relatives at the hospital, providing them with a booklet containing contact information, and encouraged them to call in case of any worries after discharge. The communication nurse called all

discharged patients in their homes within one week, asking the set of questions found in Table S1. If a problem had occurred, the communication nurse could provide counseling, book an appointment at the community health center, or initiate a home visit from social services, usually on the same day. Numbers of calls and minutes on the phone were noted.

### General practitioner liaison

The ordinary discharge summary sent to general practitioners was accompanied by a separate document containing cognitive test results and a recommendation on how to proceed with investigations. The recommendation was based on an algorithm using Mini-Mental State Examination, Clock-Drawing Test, and age (see Table S2). The general practitioners had the opportunity to discuss the results and recommendations with the study physician. After 12 months, the number of patients who had obtained a registered diagnosis of dementia were recorded.

### Standard care in control group

In the control group, all baseline measurements were performed but none of the interventions. Regular staff were informed of cognitive test results verbally and through the electronic medical record. Apart from this, the control group received standard care.

### Health care utilization after 12 months

Emergency department visits, readmissions, and hospital nights were recorded after 12 months, using the hospitals electronic medical record. All overnight readmissions were recorded, regardless of department and presenting complaint, except for hospice admissions. General practitioner visits were noted from the PASIS regional electronic registry. All data regarding health care utilization was analyzed twice, in a random sequence by two persons blinded to group allocation.

### Statistical analysis

Sample size estimation was based on a comparable intervention study with a mean difference of 2.6 hospital days between control and intervention groups.<sup>18</sup> The standard deviation for readmission was 6.6 days. To detect a similar difference with a statistical power of 0.8 and  $\alpha$  of 0.05,  $2 \times (2.8 \times 6.6/2.6)^2 = 101$  patients were needed in each group. For baseline measurements, *t*-tests, Mann–Whitney *U* tests, and Chi-square tests were used where appropriate.

The primary outcome measure was health care utilization after 12 months. This was analyzed following the intention-

to-treat principle and for 12-month survivors, separately. To estimate hospital costs, we applied the costs of the Swedish Association of Local Authorities and Regions, equivalent to 281 € per emergency department visit and 651 € per hospital night, using an exchange rate of 8.614 Swedish crowns to 1 €, as of February 4, 2013.<sup>35</sup> Health care utilization was compared between the groups using the Mann–Whitney *U* test.

Due to the non-randomized design, health care utilization after 12 months was compared with that in the 12 months preceding the index hospitalization. This analysis was performed using the matched-pairs Wilcoxon signed-rank test for the control and intervention groups separately. Thus, it was possible to decide whether health care utilization had increased, decreased, or remained constant over time. To obtain equivalent time periods only, 12-month survivors were included in this analysis.

In all statistical tests, a two-sided *P*-value of  $<0.05$  was considered to be statistically significant. The statistical procedures were performed using Statistical Package for the Social Sciences version 19.0 software (SPSS Inc, Chicago, IL, USA).

## Results

### Patients

In total, 594 unique patients were admitted 651 times, with 382 admissions in the control phase and 269 in the intervention phase (Figure 1). The patient was ineligible in 129 admissions, excluded for hospital-related reasons in 78, excluded for patient-related factors in 136, lack of consent in 72, and was excluded due to events occurring between giving consent and starting the baseline measurements in 36 cases (Figure 1). There were no differences in age between these categories by analysis of variance [ $F(5, 599) = 1.07$ ,  $P = 0.38$ ]. The age group  $<60$  years was not included in this analysis. There were no differences in gender between the groups [ $\chi^2(5, n = 651) = 7.38$ ,  $P = 0.19$ ].

### Baseline measurements

Age and education level were higher in the intervention group. Combined comorbidity and cognitive tests did not differ between groups. On the GBS-ADL, there were no differences in the separate items (data not shown) or in total score. Previous health care utilization did not differ between the groups regarding hospitalizations, but there was a trend toward the control group having had more emergency department visits and fewer general practitioner visits (Table 1).

The patients were distributed across the four wards in the department as follows: ward A (control/intervention)

**Table 1** Baseline measurements

	Intervention	Controls	P-value
Age, years	84.6 (7.3)	82.3 (8.7)	0.04*
Female gender	64%	66%	0.77
Living alone	62%	72%	0.13
Home care	57%	58%	0.89
Education >9 years	56%	37%	0.01*
Neurocognitive disorder	11%	4%	0.06
Coronary heart disease	34%	33%	0.88
Arrhythmia	38%	32%	0.37
Heart failure	31%	25%	0.35
Hypertension	50%	48%	0.89
COPD	16%	23%	0.29
Gastrointestinal disease	19%	15%	0.46
Stroke	23%	17%	0.29
Diabetes	15%	31%	0.01*
Cancer, non-skin	30%	26%	0.53
CCI	2.4 (1.4)	2.1 (1.6)	0.27
MMSE	23.0 (4.7)	22.7 (3.7)	0.66
CDT	3.3 (1.3)	3.5 (1.2)	0.33
GBS-ADL	6.6 (5.3)	6.9 (6.0)	0.71
GP visits	1.9 (1.7)	1.6 (2.3)	0.08
Emergency department visits	1.3 (1.6)	2.4 (3.7)	0.08
Admissions	1.1 (1.5)	1.4 (2.0)	0.61
Hospital nights	8.0 (12.9)	11.3 (17.0)	0.31

**Notes:** All data are mean (standard deviation) or percentages. General practitioner visits were recorded for the preceding 6 months, emergency department visits, admissions and hospital nights for 12 months. \* $P \leq 0.05$ .

**Abbreviations:** COPD, chronic obstructive pulmonary disease; CCI, Charlson comorbidity index; MMSE, Mini-Mental State Examination, scored from 0 (worst) to 30 (best); CDT, Clock-Drawing Test, scored from 0 (worst) to 5 (best); GBS-ADL, activities of daily living subset of the GBS scale, scored from 0 (best) to 36 (worst); GBS, Gottfries-Bråne-Steen.

20/22, ward B 42/36, ward C 34/28, and ward D 4/14, with a significant difference for the latter [ $\chi^2$  (1,  $n = 200$ ) = 6.63,  $P = 0.01$ ]. The entire ward D was put in isolation for a substantial time due to a Norovirus epidemic.

## Interventions

The distributions of medication errors and drug-related problems are shown in Table 2. The pharmacist gave recommendations to the ward physician for 73 of 99 patients, which were followed by the physician completely in 53, partially in 16, and not at all in four.

There was no difference between groups in number of discharge conferences [ $\chi^2$  (1,  $n = 200$ ) = 0.002,  $P = 1.0$ ]. Neither were there any differences regarding length of stay [ $U(200) = 4883$ ,  $z = 0.16$ ,  $P = 0.87$ ] or discharge destination [ $\chi^2$  (1,  $n = 196$ ) = 0.060,  $P = 0.56$ , see Table 2].

The communication nurse reached 65 of 81 patients discharged to their homes, of whom 38 had experienced problems after discharge. For 31 patients, an action was taken by the communication nurse (Table 2). Only four of 38 patients with a problem actively contacted the communication nurse. Ten patients

**Table 2** Details of interventions

	Intervention	Controls
<b>Medication overview</b>		
Any medication error	52%	–
Median (range)	1 (0–6)	–
Medication errors, total	111	–
Omission of drug	70	–
Addition of drug	15	–
Dose too high	11	–
Dose too low	9	–
Wrong dosage form	6	–
Any drug-related problem	65%	–
Median (range)	1 (0–7)	–
Drug-related problems, total number	145	–
Unclear indication	43	–
Dose not adapted to renal/liver function	29	–
Inappropriate in elderly	23	–
Adverse reaction	21	–
Untreated indication	19	–
Transferring error	8	–
Non-adherence	5	–
Drug handling	3	–
<b>Discharge conference</b>		
Discharge conference held	69%	70%
Discharged to nursing home	18%	14%
Length of stay, days, median (range)	9 (3–41)	9 (2–29)
<b>Telephone follow-up</b>		
Action taken, total	31	–
Medication counseling	5	–
Other counseling	6	–
Appointment made	14	–
Complementing prescriptions	6	–
<b>GP liaison</b>		
Recommendation sent	100%	–
No follow-up needed	14%	–
New cognitive tests promptly	38%	–
New tests in 6 months	30%	–
New tests in 12 months	18%	–

**Note:** Data are presented as percentages, numbers, or median (range).

**Abbreviation:** GP, general practitioner.

were called more than once for a total of 78 calls. The time spent on the phone by the communication nurse was 604 minutes with a median of 5 (range 0–80) minutes per call.

The recommendations based on the algorithm in Appendix 2 and sent to general practitioners on the day of discharge are shown in Table 2. After 12 months, 23 patients in the intervention group had obtained a diagnosis of dementia compared with 12 in the control group [ $\chi^2$  (1,  $n = 200$ ) = 4.46,  $P = 0.04$ ].

## Health care utilization after 12 months

After 12 months, 63 patients were deceased (31 in the control group and 32 in the intervention group). The median (interquartile range) survival was 96 (32–222) days.

In intention-to-treat analysis, the control group had more emergency department visits, readmissions, and hospital nights as well as higher hospital costs than the intervention group but the difference was not statistically significant, with a trend for more hospital nights in the control group ( $P = 0.054$ ). For the 12-month survivors ( $n = 137$ ), the differences were statistically significant regarding readmissions, hospital nights, and hospital costs (Table 3).

Over time, the yearly admissions in the control group increased from 77 to 125 (Wilcoxon signed-rank test, 36 positive differences, 13 negative, 21 ties,  $z = 3.16$ ,  $P = 0.002$ ). In the intervention group, the yearly admissions decreased insignificantly from 75 to 58 (Wilcoxon signed-rank test, 18 positive differences, 25 negative, 24 ties,  $z = 1.14$ ,  $P = 0.25$ ). A similar pattern was seen for hospital nights (Figure 2).

## Discussion

In this prospective study in medical inpatients, an intervention group had substantially fewer hospital readmissions than a group receiving standard care. After 1 year, survivors of the intervention group had spent 492 nights in hospital compared with 1228 in the control group. In the year preceding the intervention, the same groups had had 501 and 549 hospital nights, respectively. However, the difference

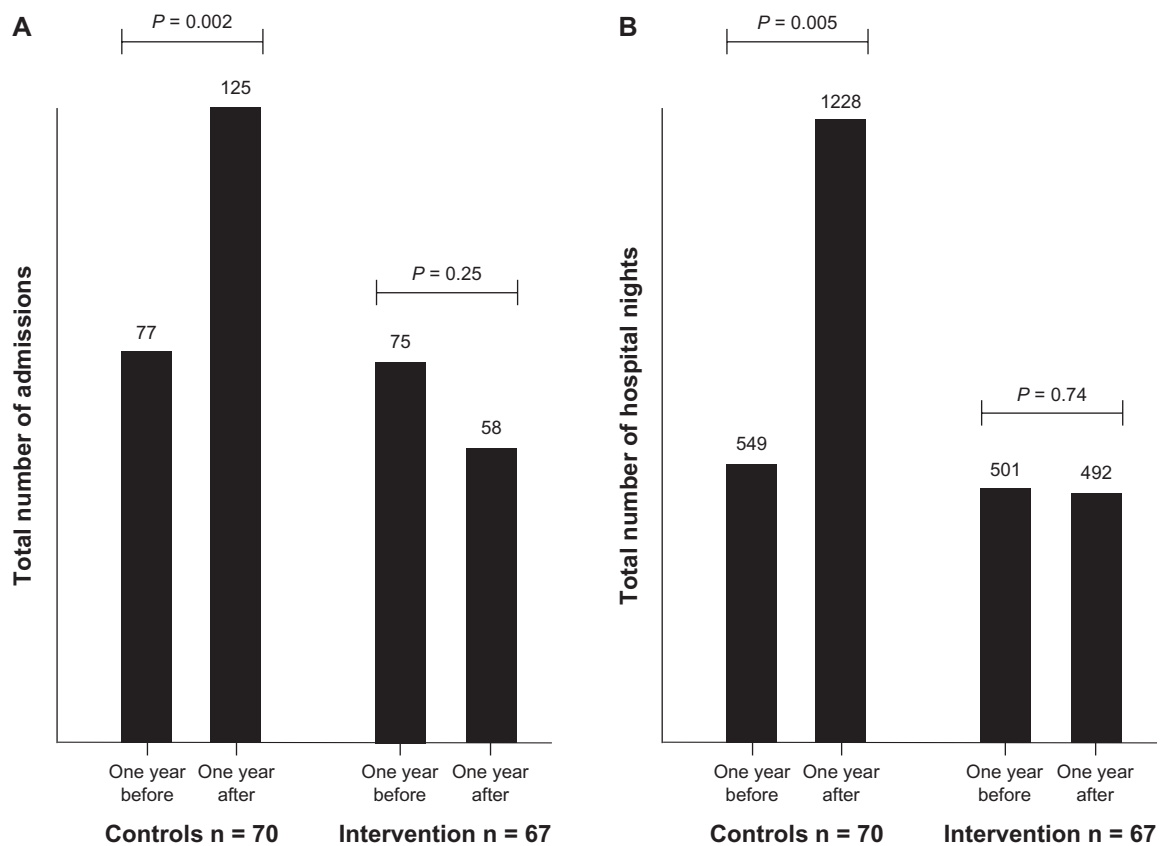
between groups was statistically significant for 12-month survivors only. From an intention-to-treat perspective, there was a substantial arithmetic difference, albeit with a lower significance level ( $P = 0.054$ ). One reason for this might be that some patients in the intervention group did not have time to benefit fully from the interventions, eg, those who passed away shortly after discharge. Another possible explanation is that health care utilization increases dramatically during the last months of life and interventions in the last year are seldom effective.<sup>36,37</sup>

The difference in readmissions corresponds to a substantial economic impact. For 12-month survivors, the hospital costs of the intervention group were 337,000 € compared with 831,000 € in the control group, for a difference of 494,000 €. In comparison, the budget of the project as a whole, including planning and collection of data in both groups, was approximately 150,000 €. Although being indirect ways of assessment, the number of general practitioner visits, discharge planning, and discharge destinations gave no indication toward increased costs elsewhere. The intervention provided other interesting results as well. Fifty-two percent of patients had medication errors on admission and 65% had actual or potential drug-related problems, in line with other research in similar settings.<sup>32</sup> Given the high occurrence of cognitive impairment, the need for structured

**Table 3** Health care utilization after 12 months from (A) intention-to-treat perspective ( $n = 200$ ) and (B) for survivors ( $n = 137$  with 67 in the intervention group and 70 in the control group)

	Intervention	Control	Difference	Test statistic	P-value
<b>(A) Intention-to-treat</b>					
GP visits, total	335	316	-19		
Mean (SD)	3.4 (3.3)	3.4 (3.2)		U(200) = 5440, $z = 1.49$	0.14
ED visits, total	133	219	84		
Mean (SD)	1.4 (1.7)	2.2 (3.8)		U(200) = 4389, $z = 1.20$	0.23
Readmissions, total	104	171	66		
Mean (SD)	1.1 (1.3)	1.7 (2.3)		U(200) = 4183, $z = 1.75$	0.08
Hospital nights, total	1008	1827	819		
Mean (SD)	10.3 (16.0)	18.5 (29.6)		U(200) = 4105, $z = 1.92$	0.054
Hospital costs, ×1000 €	681	1227	546		
Mean (SD)	7.0 (10.6)	15.7 (19.6)		U(200) = 4134, $z = 1.82$	0.07
<b>(B) 12-month survivors</b>					
GP visits, total	271	245	-26		
Mean (SD)	4.0 (3.6)	3.5 (4.6)		U(137) = 2786, $z = 1.92$	0.06
ED visits, total	83	170	87		
Mean (SD)	1.2 (1.7)	2.4 (4.4)		U(137) = 2013, $z = 1.49$	0.14
Readmissions, total	58	125	67		
Mean (SD)	0.9 (1.2)	1.8 (2.5)		U(137) = 1840, $z = 2.41$	0.02
Hospital nights, total	492	1228	736		
Mean (SD)	7.3 (15.4)	17.5 (29.7)		U(137) = 1770, $z = 2.61$	0.009
Hospital costs, ×1000 €	337	831	494		
Mean (SD)	4.9 (10.1)	11.8 (19.6)		U(137) = 1776, $z = 2.52$	0.01

**Abbreviations:** GP, general practitioner; ED, emergency department; SD, standard deviation.



**Figure 2** Pairwise comparisons within the two groups regarding readmissions (A) and hospital nights (B).

**Notes:** All comparisons are made for the year preceding the index hospitalization versus the year after the index hospitalization. P-values represent matched-pairs Wilcoxon signed-rank test.

medication reviews in this population is probably extensive. Although the communication nurse had met with all patients and relatives, supplied them with a direct way of contact, and encouraged them to call, only four of 38 with a post-discharge problem did so. This requires a more proactive approach of reaching out to patients rather than merely instructing them to contact the health care system or local authorities if problems arise.

An obvious methodologic consideration in our study is the lack of randomization, with patients allocated to control or intervention through geographic selection. The interventions required close cooperation with primary care and local authorities; this was not feasible with ten social service departments and 25 community health centers. With the lack of randomization, the risk of dissimilarities between the groups at baseline cannot be ignored. The rather extensive baseline measurements showed only subtle differences regarding education level (higher in the intervention group), language barrier (more frequent in the control phase), and occurrence of diabetes (higher in the control group). Together, they could suggest socioeconomic differences, which have

previously shown to affect readmissions.<sup>38</sup> Further, there was a trend suggesting more emergency department visits and fewer general practitioner visits in the control group before the study, possibly indicating lower accessibility to primary care in the boroughs of the control group. In addition, social services and primary care in boroughs A and B, with a large share of elderly, could be more conscious of and adapted to cognitive symptoms. However, there were no differences between the groups in hospitalizations preceding baseline. Further, we analyzed both groups internally and separately using the Wilcoxon signed-rank test, which demonstrated that health care utilization increased significantly in the control group while remaining unchanged in the intervention group. This result strongly supports our principal assumption that the intervention did in fact contribute to lower readmission rates.

Strengths of the study include its real-life setting in ordinary clinical practice. This is reflected by the rather complex inclusion procedure. Only a third of available patients were included, but the reasons for exclusion were well documented. Further, baseline measurements

were conducted in a consistent and standardized way and there were very little missing data. All our four interventions were performed in a broad population of general internal medicine patients. General practitioner liaison in particular was targeted specifically to cognitive impairment. However, we chose to perform all interventions in all patients due to the high frequency of cognitive impairment and the fact that two of the interventions (medication review and telephone follow-up) were not targeting cognitive impairment. The material was not considered sufficient for subgroup analyses.

Our results are very promising but further research is needed; the next step should be a larger, randomized study in another location. Such a study could possibly evaluate the interventions separately as well. Unplanned readmissions, drug-related problems, and cognitive impairment are likely to increase in the aging population. With the ongoing reduction of hospital bed capacity, this emphasizes the need for an efficient approach to address these issues. Our results may have several important implications for clinicians and policy-makers because they indicate that managing elderly patients in a multidisciplinary and standardized way could be a cost-efficient method to lower hospital readmissions.

In conclusion, we applied an intervention targeting drug-related problems, cognitive impairment, and discharge routines in an attempt to reduce readmissions among medical inpatients. Drug-related problems and cognitive impairment were found to be very frequent. Our results suggest that an approach targeting these areas could substantially lower hospital readmissions in this population, albeit further research is needed.

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## Disclosure

LS is a member of an expert group writing a report on “acute care of elderly in hospitals” on behalf of the Swedish Council of Health Technology Assessment (a governmental

agency). The other authors declare no conflicts of interest in this work.

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## Supplementary materials

**Table S1**

**The telephone follow-up (to be done within a week after discharge)**

**Interview**

- How have you been since leaving the hospital?
- Have you encountered any problems after discharge?
  - if yes, what kind of problems?
- Do you know who to turn to with these problems/if you should have a problem?
- Do you want me (the communication nurse) to help you with anything?
- Do you have any questions regarding your medications?
- Do you have appropriate knowledge of your medications?
- Are you taking your medications as prescribed?
- Do you feel safe at home?
- Do you wish to be contacted by the communication nurse again?

**After the interview**

Was an action taken by the contact nurse? Yes/no  
 – if yes, what action?

How many times has the patient been called?

Has the patient and/or relative contacted the contact nurse?

Total amount of minutes on the phone with this patient:

**Table S2**

**GP liaison – algorithm for recommendations**

**Clock-Drawing Test abnormal (Shulman 0–3)**

Age (years)	MMSE	Recommendation to GP
60–80	27–30	Renewed evaluation and cognitive tests within 1 year
	24–26	Renewed evaluation and cognitive tests within 6 months
	0–23	Renewed evaluation and cognitive tests promptly
80+	26–30	Renewed evaluation and cognitive tests within 1 year
	23–25	Renewed evaluation and cognitive tests within 6 months
	0–22	Renewed evaluation and cognitive tests promptly

**Clock-Drawing Test normal (Shulman 4–5)**

Age	MMSE	Recommendation to GP
All	27–30	No further action needed at the moment
	25–26	Renewed evaluation and cognitive tests within 1 year
	22–24	Renewed evaluation and cognitive tests within 6 months
	0–21	Renewed evaluation and cognitive tests promptly

**Abbreviations:** GP, general practitioner; MMSE, Mini-Mental State Examination, scored from 0 (worst) to 30 (best).

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