Impact of Pharmacy Care upon Adherence to Cardiovascular Medicines: A Feasibility Pilot Controlled Trial

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ClinicalTrials.gov Identifier: NCT01920009
Abstract

Objective
To investigate the feasibility and potential impact of a pharmacy care intervention, involving motivational interviews amongst patients with acute coronary syndrome, on adherence to medication and on health outcomes.

Methods
This article reports a prospective, interventional, controlled feasibility/pilot study. Seventy one patients discharged from a London Heart Attack Centre following acute treatment for a coronary event were enrolled and followed up for six months. Thirty two pharmacies from 6 London Boroughs were allocated into intervention or control sites. The intervention was delivered by community pharmacists face-to-face in the pharmacy, or by telephone. Consultations were delivered as part of the New Medicine Service or a Medication Usage Review. They involved a 15-20 minute motivational interview aimed at improving protective cardiovascular medicine taking.

Results
At 3 months there was a statistically significant difference in adherence between the intervention group (M= 7.7, SD=0.56) and the control group (M= 7.0, SD=1.85), (P= 0.026). At 6 months the equivalent figures were for the intervention group M=7.5, SD=1.47 and for the controls M= 6.1, SD=2.09 (P=0.004). In addition, there was a statistically significant relationship between the level of adherence at 3 months and beliefs regarding medicines (P=0.028). Patients who reported better adherence expressed positive beliefs regarding the necessity of taking their medicines. However, given the small sample size, no statistically significant outcome difference in terms of recorded blood pressure and LDL-C was observed over the six months of the study.

Conclusion
The feasibility, acceptability and potentially positive clinical outcome of the intervention was demonstrated, long with a high level of patient acceptability. It had a significant impact on cardiovascular medicine taking adherence. But these findings must be interpreted with caution. The intervention should be tested in a larger trial to ascertain its full clinical utility.
**Key Messages**

**What is already known on this subject**

- Pharmacist interventions have been shown to be successful in enhancing adherence to cardiovascular medication and improving outcomes of cardiovascular diseases.
- Improved adherence to secondary prevention medication for coronary heart disease would promote better clinical outcomes.
- Motivational interviewing can be an effective approach to improve health behaviour in people with coronary risk factors.

**What this study adds**

- This pilot study suggests that a behavioural intervention, incorporating motivational interviewing and delivered in a community pharmacy setting, can improve adherence to secondary prevention cardiovascular medication, and corresponding clinical outcomes for patients following a myocardial infarction.
Background

Despite progress since the 1950s, cardiovascular disease (CVD) remains a significant cause of mortality and morbidity in the UK. There are currently an estimated 2.3 million people living with CHD who are in need of secondary prevention medication [1]. Yet long-term adherence to secondary prevention therapies is poor. Reported adherence to medication regimens post Myocardial Infarction (MI) ranges from 13-60 per cent [2].

Research indicates that approximately a quarter to a third of CVD patients discontinue their medication [3][4]. This problem is associated with drug wastage and, more importantly a loss of clinical benefit and potentially serious health consequences [5].

There is robust evidence that consistent use of secondary prevention medication after a coronary event is associated with lower adjusted mortality as rates compared with those amongst subjects who are not consistent medicine takers [6]. For example, patients discontinuing clopidogrel within a month after hospital discharge following acute myocardial infarction and drug eluting stent placement are significantly more likely to have an adverse outcome in the subsequent 11 months [7].

Strategies to tackle non-adherence can involve community pharmacy service providers. In England, Medicine Use Reviews (MURs) were first instituted in 2005[8]. They are intended to help identify and address problems that patients experience in relation to taking medicines. More recently, the New Medicines Service (NMS) [9] was introduced in order to promote adherence in patients taking medicines for the first time for a range of long-term conditions. Both these services are NHS (The UK National Health Service) remunerated services of community pharmacists.

Other strategies for supporting enhanced medicines usage involve motivational interviewing. This can be defined as a client-centred, directive, form of counselling intended to foster behavioural change by increasing awareness of ambiguities and internal dissonance [10]. Motivational interviewing has been employed in many clinical settings and with multiple patient groups. [11],[12],[13]

Pharmacists are increasingly employing patient-centred approaches to support patients taking medicines for long-term conditions. Yet there is currently no adequate evidence base regarding the feasibility and effectiveness of motivational interviewing to promote medication adherence in the pharmacy setting. This study was carried out to address this shortcoming and to evaluate the potential effectiveness of a community pharmacy intervention for patients discharged following a myocardial infarction with secondary prevention medication. It also explored issues relating to improving communication and collaboration between hospital and community pharmacists.
Objectives

To investigate the potential impact on outcomes of a pharmacy care intervention involving hospital pharmacy referral to community pharmacy services and motivational interviewing on adherence to secondary prevention medication amongst recently discharged coronary heart disease patients.

Methods

Design

The study was designed as a prospective, feasibility/pilot, controlled trial. The primary outcome was adherence to cardiovascular medication (Figure 1).

Study setting and Study population

The study was undertaken in collaboration with community pharmacists in East London and the North East London Pharmaceutical Committee (NELLPC) and with practitioners and patients from a London Heart Attack Centre. The study gained research ethical approval from the National Research Ethics Service Committee (North West – Preston), from the R &D Joint Research Management Office, Queen Mary Innovation Centre, and from the R&D Office, University College London. The study population included coronary heart disease patients with a discharge diagnosis of Acute Coronary Syndrome (ACS).

Recruitment

There were two stages of recruitment; recruitment of pharmacies and recruitment of patients. Community pharmacists/pharmacies were recruited through NELLPC and assigned as below to either the intervention or control group. The inclusion criteria were: (1) Pharmacists willing to counsel patients and interested in attending further training; (2) have a consultation area and have access to a telephone (land line or mobile); (3) the pharmacists were knowledgeable about the NMS and MUR, and had contacts with or were willing to contact general practitioners and also willing to contact patients to invite them for a consultation.

Allocation to intervention and control groups

Whilst simple randomisation of the entire sample was not possible, procedures were adopted to ensure comparability of the intervention and control groups for this study. Pharmacy recruitment was all done through NELLPC. Pharmacists informed of study by two different routes. Firstly, by email 22 pharmacies responded that they wished to take part. These were randomised to intervention and control by an independent statistician at UCL School of Pharmacy. This
process was concealed from the researcher and the research team and was performed at pharmacy level to avoid contamination of controls.

To achieve sufficient numbers a second group were invited to participate during a professional meeting and 10 pharmacies met the inclusion criteria. As the dates of motivational interview training had had to be set in advance, pharmacists wishing to take part and able to attend the pre-determined dates were allocated to the intervention group. The control group was a matched sample drawn from remaining pharmacists who expressed a wish to take part. See Figure 2. Eligible patients were prior to discharge given introductory information about the study by the researcher and supplied with further details as requested. They were then asked if they would like to participate. The full eligibility and exclusion criteria are described in Supplementary Table 1. After recruitment, patients were assigned into groups according to the primary care pharmacy at which they usually refill their prescriptions. Patients who normally refill their prescription from the intervention pharmacies were assigned to the intervention group and patients who regularly refilled their prescription in the pharmacies that were control sites were assigned to the control group.

**Blinding**

The research pharmacist responsible for the data analysis was blind to the above group allocations. The General Practitioners/Practices from which data regarding blood pressures and LDL-C levels were collected were also blind, unless referral of a patient by a community pharmacist took place. However, due to the nature of the intervention it was not possible to blind the hospital and community pharmacists delivering the intervention or the patients receiving it.

**Sample size**

Power calculations were based on the findings of previous studies in which the primary outcome was adherence. For instance, a similar study\(^\text{[14]}\) reported a 33 per cent increase in adherence with a margin error of 5 per cent and confidence interval 95 per cent. Given these and allied data the enrolment target was set at 200 patients.

**Pharmacist training**

Pharmacists in the intervention delivery group participated in a two day training session on motivational interviewing, followed by a subsequent booster session, given by an expert psychologist (KF), all the training sessions on motivational interviewing including the booster session were completed before inclusion of patients. An additional training session on the use of secondary prevention medicines after a myocardial infarction was given by a consultant pharmacist (SA).

**Liaison with General Practitioners**

The GPs were asked for their written consent to providing the results of blood pressure measurements and LDL-C levels during the duration of the study with patient consent.
The intervention

The intervention was developed on the basis of a previous systematic review \[15\]. A ‘consultation chart’ (a pro forma guide the motivational interview process) was developed by referring to a previous randomised controlled trial involving hypertensive patients \[16\] which generated statistically significant impacts on adherence. In this instance trained research assistants rather than pharmacists used motivational interviewing techniques. The intervention was designed to include elements of motivational interviews and to be integrated into the existing NMS and MUR pharmacy services so that the participating community pharmacists would be able to claim funding for their work.

On discharge patients receiving the intervention were initially given usual care from a hospital pharmacist. This consisted of a review of medications use, counselling on secondary prevention and any other additional prescribed medication usage, an antiplatelet medication leaflet and referral to cardiac rehabilitation. Patients were subsequently contacted by a pharmacist to arrange a community pharmacy consultation.

The first community pharmacy consultation typically took place at around 2 weeks after hospital discharge on either a face to face basis or by telephone as recent evidence shows that motivational interviewing can be effectively delivered by telephone \[17\] and lasted for 15-20 minutes. The substance of these sessions is detailed in Supplementary Box 1, also comparison of motivational interviewing with traditional counselling can be found in Supplementary Table 2.

The control group

On discharge control group patients received usual care from a hospital pharmacist. As described above.

Outcome measures and Data collection

The primary outcome measure used was self-reported adherence with the coronary artery disease medication regimen prescribed, assessed via the Morisky Medication Adherence Scale MMAS8\[18\]. The Beliefs about Medicines Questionnaire-Specific BMQ-S \[19\] was also used at 3 months after discharge; to evaluate the effect of the intervention on patients beliefs regarding their medication and to examine the relationship between patients’ beliefs regarding their medicines and adherence, this study did not evaluate changes in patients’ beliefs over time.

Secondary outcome measures included blood pressure and LDL-C. Baseline data collected from the hospital included gender, age, diagnosis, blood pressure, LDL-C, ethnicity, post code and GP practice, all patients enrolled in the study were discharged on four classes of medication (antiplatelets, beta blockers, ACE inhibitors or ARBs and statins) as recommended for secondary care of patients following a myocardial infarction \[20\]. Data collection took place at two weeks after hospital discharge and at 3 and 6 months (figure 1).
Analysis

Data was analysed by using the Statistical Package for the Social Sciences (SPSS) version 22 for Windows. An independent T-test was used to compare the differences in the intervention group and control group adherence means and also to compare the differences between the blood pressures and LDL-C levels (Significance was set at the 5 percent level). A chi-square test was performed to examine the relationship between beliefs about medication and adherence to the cardiac medication at 3 months. The scores from the BMQ-S were handled according to standard procedures for analysis of the questionnaire [19].

Results

In the 4 months available for recruitment for this study 71 patients were enrolled consecutively. Recruitment is commonly one of the biggest challenges for any study. In this instance it was undertaken by a single researcher. On average it was possible to recruit 2–3 patients per day, excluding those occasions on which no eligible patients presented. Out of a total of 233 patients assessed for eligibility only 14 individuals refused to participate. Others were excluded because they did not meet the inclusion criteria as illustrated in the consort diagram—see Figure 3.

The NHS users recruited were predominantly male (76%) and as shown in Supplementary Table 3, most were in their sixties and seventies. It was found that 51 of the patients involved had had an ST-Elevation Myocardial Infarction (STEMI). The remaining 20 had suffered a Non ST-Elevation Myocardial Infarction (NSTEMI).

As a feasibility/pilot study, this was not powered to measure clinical outcomes and was designed only to provide an indication of potential effectiveness. Hence the findings presented here should be interpreted with caution.

Impact on adherence

As indicated in Figure 4 there was at baseline no difference in self-reported adherence rates between the intervention group (M=7.45, SD=0.79) and the control group (M=7.5, SD=0.93) (P=0.85). However, at 3 months there was a statistically significant difference in adherence between the intervention group (M=7.7, SD=0.56) and the control group (M=7.0, SD=1.85), (P=0.026). There was also a statistically significant difference at 6 months between the intervention group (M=7.5 (93.75%), SD=1.47) and the controls (M=6.1 (76.25%), SD=2.09) (P=0.004). Note: (M=Mean, SD=Standard Deviation)
Beliefs about Medicines

There was a statistically significant relationship between the level of adherence at 3 months and the beliefs regarding medicines as evaluated by the BMQ-S \((P=0.028)\). Patients with greater levels of self-reported adherence showed more positive beliefs regarding the necessity of their medicines.

Results on clinical outcomes: blood pressure and LDL-C

It was disappointing that for both BP and LDL-C around two-thirds of patients in both groups did not have a follow-up evaluation from their GPs. This may help explain why at 3 months there was no statistically significant difference between the intervention group \((M=127, SD=20)\) and the control group in systolic blood pressure \((M= 121, SD=20), P=0.3\).

Similarly at 6 months there was no statistically significant result between the intervention group \((M=132, SD= 11)\) and the control group \((M=129, SD= 12), P=0.6\) (Figure 4). Nevertheless, systolic blood pressure in the intervention group at 3 months decreased by 5 mmHg and at 6 months returned to the same as baseline. By contrast, Figure 4 also shows, in the control group systolic blood pressure had decreased by 3 mmHg at 3 months but increased by 5mmHg at 6 months.

Likewise, there was no significant difference in diastolic blood pressure between the intervention group at baseline \((M= 74, SD=7.2)\) and the control group \((M=73, SD= 11), P=0.8\). At 3 months there was again no statistically significant difference between the intervention group \((M=73, SD= 11.5)\) and the controls \((M=72, SD= 9.9), P=0.84\). At 6 months there was similarly no statistically significant result in the intervention group the figures were \((M=68, SD= 11.7)\) and in the controls they were \((M=75, SD=4.8), P=0.2\). Nevertheless, at six months mean diastolic blood pressure in the intervention group had decreased by 6mmHg from baseline. In the control group diastolic blood pressure had by then increased by 2 mmHg from baseline.

With regard to the LDL cholesterol levels reported, there was no statistically significant difference between the intervention group’s LDL-C at baseline \((M=2.75, SD= 1.05)\) and the control group figures \((M=2.79, SD=1.4), P=0.9\). At six months there was a 0.79 mmol/l difference in LDL-C between the intervention group and control group (Figure 4). However, although suggestive of a material difference this result was once again non-significant, possibly because of the small numbers of subjects for whom data were available.

Discussion

This study reports positive findings regarding the potential outcomes of the community pharmacy intervention investigated. Numerous studies have examined patients’ views on services provided by community pharmacists. It has been commonly found that patient awareness of the pharmacist’s role outside that of dispensing and non-prescription drug supply is generally low. This could to date have led to an under-utilisation of pharmacist provided clinical
services \textsuperscript{[21][22]}. Initiatives like the study reported here may over time enhance awareness of the value of pharmacy services in ‘serious’ contexts like post-hospital discharge following a cardiac event. Such initiatives might also contribute to the uptake and utility of existing services (ie the NMS and MURs), and promote improved hospital and community pharmacy communication.

In the latter context, patients’ discharge summaries were forwarded from the participating hospital pharmacy to community pharmacists. Community pharmacy access to patients’ health care records is not as yet usually available in the UK or elsewhere. There is evidence that this restricts the capacity of pharmacists’ interventions to improve adherence and resolve other medication related problems \textsuperscript{[23]}. This study demonstrates the potential importance of record sharing between community and hospital pharmacists in improving patient care. The supply of discharge summaries to community pharmacies was achieved by using secure hospital emails and with patient consent. All the stakeholders involved, including the service users taking part, supported the supply of discharge summaries to community pharmacies. This finding is in line with the approach advocated by the Royal Pharmaceutical Society (RPS) (2014). The RPS has recently launched ‘a hospital referral to community pharmacy innovators’ Toolkit developed in response to the report ‘Now or Never: Shaping Pharmacy for the Future’ \textsuperscript{[24]}. In the Society’s view referrals from hospital to community pharmacies could become routine practice within five years.

After six months self-reported medication adherence amongst those receiving motivational support from community pharmacist was 17 per cent greater than that recorded by control patients. This result can be compared to a recent US study \textsuperscript{[25]} that found that a phone-based motivational interview improved adherence in the case of antiplatelet medicines by 14\% (p < 0.01). It is also similar in magnitude to the reported effect of automated text messaging when used to prompt adherence to cardiovascular preventive treatment \textsuperscript{[26]}. Other research studies have failed to find similar benefits in relation to the treatment of people who have experienced strokes or other forms of vascular disease \textsuperscript{[27][28]}. Nevertheless, there is mounting reason to believe that greater use of well- targeted motivational interventions by community pharmacists could prove to be of substantive value in today’s environment. It is also possible that combinations of different types of approach to enhancing medication taking in high risk patient groups could have even greater effects.

In this study a statistically significant relationship was found between reported adherence and medicine takers’ beliefs regarding the necessity of taking their prescribed treatments. Although there remain uncertainties regarding the causal links underpinning such observations, our findings are consistent with other research undertaken in the UK and elsewhere \textsuperscript{[29][30]}. Investing in pharmacy led interventions to further promote awareness of the value of taking medicines in high risk therapeutic situations like post MI care has the potential to contribute cost effectively to improved health outcomes \textsuperscript{[31-34]}. However, no statistically significant difference in the proportion of patients achieving BP and LDL-C reduction targets was found in this trial, which was not adequately powered to identify such effects. To date, most other similar studies have also failed to demonstrate statistically significant results in relation to such proxy clinical outcomes \textsuperscript{[35-38]}. A relatively recent review
concluded that too few pharmacy based trials are available in this area, and that further larger scale quantitative research involving CVD patients should be conducted.

More qualitative work examining pharmacists’ experiences of using motivational interviews to enhance adherence should also prove useful. In addition, after a life changing event such as a myocardial infarction many patients appear to welcome the additional primary care support that appropriately skilled community pharmacists are capable of providing.

The positive responses of GPs involved in this investigation are also informative. Some previous research has indicated that GPs often tend to have negative attitudes towards extending community pharmacists’ clinical roles. Yet the uptake and outcomes of community pharmacy services such as the intervention evaluated here are likely to improve when they are endorsed by GPs and effectively integrated with other primary care services. The findings of this research indicate that, in addition to recent measures aimed at encouraging the employment of pharmacists in GP surgeries, innovative approaches to developing community pharmacy contributions to the care of patients in need of better overall primary care services are also worth further investigation.

This study’s main limitations relate to the small sample size and that it was focused on improving care in just one area of North East London; also it included a single centre this limits its perceived effectiveness in different locations and patient populations and also limits the confidence with which its findings can be generalised. Other limitations; it was not possible to formally assess the extent to which all elements of motivational interviews were followed in the delivery of the intervention and ideally, measures of adherence that reduce reliance on self-reported data would also have been valuable. The strengths of the study that this article reports, which was designed as a feasibility pilot controlled trial, include that it used well validated instruments such as the Morisky Scale questionnaire and the BMQ, and that effective blinding procedures were put in place.

**Conclusion**

This work indicates how enhanced pharmaceutical care could help further improve adherence to medicines and health outcomes in relation to using medicines for preventive purposes amongst patients recovering from acute coronary events. Moving further towards assuring the optimisation of medicines use in this and other contexts is likely to demand the organisation of a larger multicentre randomised control trial or trials, the design of which should be informed by the findings of this feasibility study.
References


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