Integrating clinical pharmacists within general practice: protocol for a pilot cluster randomised controlled trial

Aisling Croke,¹ Frank Moriarty,¹,² Fiona Boland ²,³, Laura McCullagh ⁴,⁵ Karen Cardwell,¹ Susan M Smith ¹, Barbara Clyne ¹

ABSTRACT

Introduction Managing patients with multiple conditions (multimorbidity) is a major challenge for healthcare systems internationally, particularly in older patients. Multimorbidity and subsequent polypharmacy increase treatment burden and the risk of potentially inappropriate prescribing, and both are complex to manage in primary care. Limited evidence suggests integration of pharmacists into general practice teams could improve medication management for patients with multimorbidity and polypharmacy. Building on findings from a non-randomised, uncontrolled General Practice Pharmacist (GPP) feasibility study conducted in Irish primary care, the aim of this study is to conduct a pilot cluster randomised controlled trial (cRCT) of the GPP study, to assess feasibility, intervention impact, costs and appropriateness of continuing to a definitive cRCT.

Methods and analysis This pilot cRCT will involve 8 general practitioner (GP) practices and 120 patients. Practices will identify and recruit patients aged ≥65 years, who are taking ≥10 regular medications. Practices will be allocated to intervention or control after baseline data collection. Intervention practices will have a pharmacist integrated in their service, working with GPs, patients and practice staff to optimise prescribing and other medication-related activities. Control practices will provide standard GP care. The primary feasibility outcomes will include recruitment rate, uptake of medication reviews and study retention. For the primary clinical outcome, the number of potentially inappropriate prescribing incidences per patient will be collected. Secondary outcomes will include medication-related outcomes, patient-reported outcome measures, and data pertaining to the role and impact of the pharmacist on prescribing. In addition, economic and process evaluations will be conducted.

Ethics and dissemination This trial has been approved by the Irish College of General Practitioners Research Ethics Committee and will be performed in accordance with the Declaration of Helsinki. The results will be reported in peer-reviewed journals and be presented at national and international conferences.

Trial registration number ISRCTN Registry (https://doi.org/10.1186/SRCTN18752158).

INTRODUCTION

Managing patients with multiple conditions (multimorbidity) and associated polypharmacy is recognised as a major challenge for healthcare systems. A recent systematic review of 70 studies reported a 33% prevalence of multimorbidity (defined as two or more chronic conditions) in community settings, with a large proportion (more than 50% in many cases) of individuals aged 65 years and over having multimorbidity.¹ The volume of medications being prescribed to older patients has increased over the past number of years also. A study analysing prescribing trends over a 15-year period found that polypharmacy (commonly defined as patients taking five or more regular medications) affects 60.4% of the Irish population aged 65 years or older.² Complex polypharmacy, defined as patients taking 10 or more regular medications, affects 21.9% of Irish patients.² Polypharmacy is often associated with multimorbidity as best practice and clinical guidelines are typically derived...
from populations which do not reflect these patients and tend to be single disease focused. Overall management of these multiple medications, commenced by various specialists, is usually provided by a patient’s general practitioner (GP). There is an increased medication burden in this patient cohort and this leads to the risk of potentially inappropriate prescribing (PIP). PIP can be described as suboptimal prescribing, and may lead to adverse drug reactions (ADRs), hospitalisations and subsequent costs. Older people are most at risk of negative consequences of reactions (ADRs), hospitalisations and subsequent costs. as suboptimal prescribing, and may lead to adverse drug reactions (ADRs), hospitalisations and subsequent costs. Older people are most at risk of negative consequences of suboptimal prescribing, and may lead to adverse drug reactions (ADRs). PIP can be described as suboptimal prescribing, and may lead to adverse drug reactions (ADRs).

Within this context it is important to develop interventions for patients with multimorbidity and polypharmacy which reduce PIP as a public health measure. Interventions to improve appropriate prescribing include those aimed at prescribers (eg, computerised decision support), patient education and changes to care delivery arrangements, such as staffing models or skills-mix. One such intervention is the integration of clinical pharmacists into general practices to address PIP, deprescribing, medication reviews and general practice workload. Evidence suggests that pharmacists in general practice can have positive impacts on clinical outcomes with one systematic review reporting a significant reduction in glycated haemoglobin between pharmacist intervention groups and control (mean difference −0.88%, 95% CI −1.15% to −0.62%, p<0.001) and prescribing safely. In particular, the PINCER trial in the UK demonstrated that a pharmacist-led information technology intervention was an effective method for reducing a range of medication errors in general practice. Integrating pharmacists into primary care may also reduce GP workload (particularly medication-related administration), emergency department attendance and medication-related hospitalisations. The current evidence base is however varied with some high-quality studies such as the PINCER trial demonstrating effectiveness, and some smaller studies of mixed quality with mixed results. Further high-quality research is needed to assess the impact of this new role on patient outcomes, GP workload, stakeholder experiences and cost-effectiveness.

Rationale for study
Unlike countries such as the UK and Canada, pharmacists in Ireland are not formally integrated into general practice, nor do they have prescribing rights. Primary care in Ireland is delivered in a mixed public and private healthcare system, described in more detail in online supplemental appendix 1. The General Practice Pharmacist (GPP) feasibility study demonstrated that an intervention involving pharmacists working within general practices in Ireland is feasible to implement and has potential to improve prescribing quality in older patient populations.

The aim of this study is to conduct a pilot cluster randomised controlled trial (cRCT) of the GPP study, to assess feasibility, potential intervention impact and costs, and assess whether it is appropriate to continue to a definitive cRCT.

Development of the GPP Medicines Optimisation Study
This pilot cRCT has been informed by the non-randomised, uncontrolled feasibility study as outlined in table 1. All but two of the continuation criteria from the uncontrolled feasibility study reached the threshold of ‘Proceed with RCT’. Based on these findings and the process evaluation, areas highlighted for change to improve the pilot cRCT included (table 1): Patient recruitment. 2. Evaluating the role and impact of the pharmacist when integrated within the GP practice. 3. Pharmacist isolation in GP practice. 4. Issues surrounding space in GP practices and individual case discussion time allocation with GPs. 5. Immediacy of communication with GPs. 6. Standard framework for medication review.

METHODS/DESIGN

Objectives of study
This study will involve the conduct and evaluation of a pilot cRCT of an intervention of clinical pharmacists based in general practice, to determine if general practice-based pharmacists are feasible and can potentially improve the management of, and outcomes for, patients with complex polypharmacy, in comparison with usual GP care.

The pilot cRCT will also inform the conduct (using formal continuation criteria) and sample size of a definitive trial.

Study design
A cluster design was chosen to address potential contamination of GPs as a result of exposure to working with a pharmacist during the intervention. GP practices are the units of randomisation (the clusters), and individual patients with polypharmacy are the units of analysis (the participants) with adjustments for clustering. We will report the trial according to the Consolidated Standards of Reporting Trials guidelines adapted for pilot studies.

The pilot cRCT design was informed by the Medical Research Council (MRC) Framework for the design and evaluation of complex interventions. Informed consent will be obtained from participants (practices and patients) prior to data collection.

Study setting
This study will take place in the Irish General Practice setting. Both single-handed and group practices from the Health Research Board (HRB) Primary Care Clinical Trials Network Ireland (PC CTNI) (http://primarycare-trials.ie) will be included.
The feasibility study highlighted a need to examine the role the pharmacist assumed in the practice in greater detail. The pharmacists in the feasibility study took part in a wide range of activities involving quality of practice, administration, medication review and education.

**Adaptations for proposed pilot cRCT:**

This proposed study will also evaluate the role and impact of a pharmacist on care provision within the general practice when integrated into the practice team in more detail than previously. As per the previous study, qualitative interviews will be conducted with participants. Additionally, a description of the activities that the pharmacist undertakes and the length of time undertaken to complete those activities will be collected. For the pilot cRCT, a pharmacist activity log will be used to capture pharmacist activities. This activity log will be updated weekly by the research pharmacists, and will not include any identifiable patient data. The purpose of the activity log is to document pharmacist tasks and actions, which will vary depending on practice requirements. This is key information in terms of implementation of this type of intervention.

The pharmacists will join the practice in this study as healthcare professionals, and are bound legally and ethically by the ‘Code of Conduct’ as presented by the Pharmaceutical Society of Ireland. They would be considered a member of the practice team for the duration of the intervention and will enter into a confidentiality agreement with the individual practices. As with any healthcare professional contributing to care, these factors would ensure confidentiality of individual patients’ information.
generator in Excel to identify a random selection of 15 eligible patients. The practice will use a random number by the research team, will run this search to identify 10 or more regular medications. Practice staff, supported electronic record finder tool that can identify patients on will identify a list of potentially eligible patients using an practices will be excluded if they have <500 older patients. Exclusion criteria

<table>
<thead>
<tr>
<th>Study domain</th>
<th>Lessons learnt from uncontrolled feasibility study in 4 practices and adaptations for proposed pilot cRCT</th>
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<tbody>
<tr>
<td>Pharmacist isolation in GP practice</td>
<td>Pharmacist isolation was a theme identified during the feasibility study.</td>
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<td>Adaptations for proposed pilot cRCT:</td>
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<td></td>
<td>For the pilot cRCT, integrative workshops will be conducted with the practices to ensure the pharmacist is familiar with the practice, staff and operating systems. These workshops will provide an opportunity to educate practice staff as to the role of the pharmacist in the practice and identify the pharmacist as a resource for all members of the team, as in the feasibility study some staff stated they were not aware of this. The content of the workshops will be designed to introduce the pharmacist to the team and to meet the needs of the practice. All activities related to these workshops will be recorded using the pharmacist activity log. The impact of education sessions such as this integrative workshop will be assessed in the post-intervention interviews.</td>
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<tr>
<td>Issues surrounding space in GP practices and time with GPs</td>
<td>This will be addressed by forward planning with the practices.</td>
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<td>Adaptations for proposed pilot cRCT:</td>
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<td>The research team will engage with practice staff to examine which practice staff will be present or absent during the intervention period and also if any movement of premises is planned. This will coordinate GP and pharmacist time to avoid any potential additional work burden to either parties.</td>
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<tr>
<td>Immediacy of communication with GPs</td>
<td>Immediacy of query resolution was a factor for GPs during the feasibility study.</td>
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<td></td>
<td>Adaptations for proposed pilot cRCT:</td>
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<td>It is suggested for the pilot cRCT that practices can arrange with the pharmacist how best to contact them when they are not in the practice. In clinical practice, it is not always possible for clinicians to make immediate contact with each other but this issue will be raised as important during the pharmacist and practice training for the intervention.</td>
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**Trial status**

The trial is ongoing; practice and patient recruitment to commence in March 2021. Anticipated intervention completion is in November 2022.

**Population and recruitment**

We will enrol a total of eight practices and each practice will recruit 15 patients (a total of 120 patients) with complex polypharmacy. All practices from the PC CTNI will be invited to participate via an email (or letter where email address is unavailable). Among eligible practices that express an interest in participating, we will recruit a purposive sample reflecting different practice sizes and locations (urban and rural).

**Clusters (GP practices)**

Practices will be eligible to participate if they have at least 500 older patients (aged ≥65 years) on their patient panel to ensure adequate numbers of eligible patients and can use a GP software finder tool to identify patients with complex polypharmacy.

**Exclusion criteria**

Exclusion criteria

- Under the age of 65 years.
- Terminally ill leading to high likelihood of death or major disability during study follow-up period, as judged by the patient’s GP.
- Severe cognitive impairment, or psychiatric/psychological morbidity sufficient to impair informed consent, as judged by the patient’s GP.
- Resident in nursing home.

Participants (patients)

Patients will be eligible if they are 65 years of age or older, have complex polypharmacy (defined as ≥10 repeat medications) and must have an ability to attend a medication review with the pharmacist. PIP is highly prevalent in older patient populations and thus this patient group is most at risk of negative consequences of polypharmacy. The implicit and explicit inappropriate prescribing screening tools used in this study have been designed for older patients and alternative indicators would need to be used in younger populations.

**Table 1 Continued**
Currently participating in a related study.

To ensure transparency, a highly visible poster will be displayed in the practices and an information leaflet will be available outlining the role of the pharmacist, and also steps to take if an individual does not want their clinical information reviewed by the pharmacist.

Randomisation

Once consent and all baseline data collection has been completed, practices will be allocated into control or intervention groups using minimisation (see figure 1). This approach offers the advantage of ensuring balance between the groups in terms of prognostic factors, in this case, practice size (number of whole-time equivalent GPs) and practice location. Sequence generation and practice allocation will be carried out remotely by an independent statistician. Allocation of practices will be done in blocks to enable the pharmacists to deliver the intervention in different practices and stagger data collection.

Intervention

Pharmacists will be recruited via a competitive interview process. All pharmacists who currently operate in a clinical role will be considered eligible for interview, though it will be more desirable for the recruited pharmacists to have experience of working in community care and primary care settings. Two pharmacists will work between four intervention practices (two intervention practices per pharmacist). It is anticipated that the split of GP practices will be on an urban/rural basis and will be flexible to fit with pharmacist requirements. Intervention delivery will be staggered in practices. The pharmacist will integrate into the GP practice as per practice preference regarding time allocation for 10 hours per week for a period of 4 months. Integrative workshops prior to pharmacist commencement will ensure they are familiar with the practice, staff and operating systems. Written training material based on the previous feasibility study will be

Figure 1 Flow of practices through the GPP Medicines Optimisation Programme pilot cRCT. cRCT, cluster randomised controlled trial; GPP, General Practice Pharmacist.
provided to pharmacists also. When in the practices, pharmacists will support prescribing-related activities for consented patients as agreed with the GPs. They will also be available to support other related activities such as practice education and support with practice audits as requested. Table 2 describes in more detail the potential activities the pharmacist may undertake in the GP practice including, but not limited to: liaison at transitions of care, medications information query management, determination of blood samples due and repeat prescriptions management. This list is not finite. The specific focus of the intervention will be the medications optimisation component delivered by targeted patient medication reviews, based on improving safety and addressing national medications management priorities and guidelines. Review of anonymous aggregate data within practices by the pharmacist will also identify specific areas to focus medication optimisation. Medication reviews for consented patients (and practice audit activities) will focus on high-risk prescribing, PIP and deprescribing as per the criteria used in the non-randomised feasibility study.16

Patients in the intervention arm will make an appointment with the pharmacist for medication review, preferably in the first 2 months of the intervention. The intervention will run for 4 months in each practice with follow-up data to be collected at completion. Depending on GP availability, further follow-up may be possible after the 4-month intervention period. Figure 2 provides an overview of the process of the GPP intervention.

**Control**

Control practices will continue to provide care as usual with no input from the pharmacists. As a form of wait list control, patients in control practices will be offered a medication review with the pharmacist at study completion.

**Outcome measures**

The primary outcome of feasibility will include recruitment rate, uptake of medication reviews and study retention. Participant experience will be explored through the process evaluation. We will use continuation criteria (see Table 3) to determine if progressing this pilot cRCT to a definitive pragmatic cRCT is warranted based on data from the pilot cRCT and process evaluation.

The primary clinical outcome which will be collected will be the number of PIP incidences per patient, measured at baseline and 4 months. However, a range of

<table>
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<tr>
<th>Table 2 Proposed activities for pharmacists in GP practice</th>
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<tbody>
<tr>
<td><strong>Proposed activities</strong></td>
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| ▶ Clinical audit | ▶ Topic requested by GP  
▶ Area determined by pharmacist based on analysis of aggregate data  
▶ As part of chronic disease management  
▶ Practice CPD requirements |
| ▶ Medication review | ▶ Review complex patients with GP and with patients themselves |
| ▶ Transitions of care | ▶ Acting liaison between hospital and primary care settings.  
▶ Forward discharge letters to community pharmacists |
| ▶ Repeat prescriptions management | ▶ Medication review  
▶ Discuss dose adjustments  
▶ Medication initiation or discontinuation  
▶ Medication that require extra monitoring |
| ▶ Management of chronic illness | ▶ Manage medication reviews within chronic disease management programme in line with national guidelines and best practice  
▶ Compile educational resources for patients and practice |
| ▶ Determine when bloods are due | ▶ Medication monitoring requirement  
▶ Chronic disease management  
▶ Regular monitoring |
| ▶ Education sessions | ▶ Therapeutic area of interest  
▶ GP directed  
▶ Pharmacist directed  
▶ Practice staff directed |
| ▶ Query management | ▶ Phone call communication when not in practice to ensure immediacy of query resolution  
▶ GP queries  
▶ Patient queries  
▶ Other healthcare professional queries |
| ▶ Medicines information role | ▶ Evidence based answers to GP queries relating to medications  
▶ Liaising with patients in relation to medication alerts |
medication-related, patient-reported outcome measures (PROMs) and pharmacist impact data will be collected, informed by findings from the non-randomised pilot study, and core outcome sets for multimorbidity and polypharmacy.

**Medication-related outcomes from patient’s medication record**
- Number of PIP per patient, as per criteria used in the non-randomised pilot study and pharmacist clinical judgement.
- Number of repeat medications.
- Proportion of patients with polypharmacy.
- Medication changes:
  - Deprescribing (tapering or stopping) of medications that may cause harm or are no longer providing benefit.
  - Medications started.
  - Number of medications prescribed generically.
- ADRs, defined as an appreciably harmful or unpleasant reaction, resulting from an intervention related to the

**Figure 2** Flow chart for GPP Medicines Optimisation Programme intervention. GP, general practitioner; GPP, General Practice Pharmacist.
use of a medicinal product, which predicts hazard from future administration and warrants prevention or specific treatment, or alteration of the dosage regimen, or withdrawal of the product.\textsuperscript{22}

- Adverse drug withdrawal events, defined as either recurrence of the condition for which the medication was prescribed or a physiological reaction to medication withdrawal.\textsuperscript{23,24}

Patient-reported outcome measures

- Health-related Quality of Life (EQ-5D-5L).\textsuperscript{25}
- Revised Patients’ Attitudes Towards Deprescribing (rPATD).\textsuperscript{26}
- Multimorbidity Treatment Burden Questionnaire.\textsuperscript{27}

Role and impact of pharmacists

- Activities undertaken and total time to complete.
- Condition-specific domains.
- Engagement with other healthcare professionals and practice staff and time spent on each interaction with:
  - GPs.
  - Community pharmacists.
  - Other healthcare professionals/practice staff.

Data collection

Patient data will be collected at baseline, prior to randomisation, and at intervention completion (4 months after baseline). Medication-related data will be collected from patient’s medication records (PMRs). For intervention practices, these records will be reviewed at baseline once the practice has been randomised to the intervention arm, and then at intervention completion. For control practices, records will be reviewed in the PMR at trial completion using previous ‘current prescriptions’ at baseline and all prescription data for the study period to provide contemporaneous medication data. To ensure the correct baseline prescription has been selected, researchers can review prescriptions issued prior to the baseline data, but that were contemporaneous for the patient at that time.

PROMs will be collected from patient questionnaires and will include the following validated measures: EQ-5D-5L,\textsuperscript{25} burden of treatment\textsuperscript{28} and rPATD.\textsuperscript{26} PROMs data will be collected from control practice patients at matched time points. The variability, consistency, response rates and data completeness for each outcome will be determined.

Data on the role and impact of the pharmacist will be recorded by the practice pharmacists while integrated within the intervention practices using prespecified reporting pharmacist activity log which will record:

1. Activity that the pharmacist undertook.
2. Total time spent on activities, for example, 30 min spent on audit. This will allow for content analysis and breakdown of time spent on each activity.
3. Engagement and time spent with GP.
4. Engagement and time spent with community pharmacists.
5. Engagement and time spent with other practice staff and healthcare professionals.
6. Condition-specific domains for audit, for example, PIP would be ‘safer prescribing’.
7. A ‘free text’ option which will allow the pharmacist to note interesting observations which they may forget during the qualitative interview at intervention completion.

Sample size

While no formal sample size is required for a pilot study, based on data we have collected previously, we have...
estimated a sample size which we feel will be large enough to inform about the practicalities of delivering the intervention, recruitment and retention rates, in addition to exploring potential change in PIP outcomes for going forward to a definitive cRCT.

Based on reducing the mean number of PIP per patient from 0.8 by 20% to 0.64, using an SD of 0.2 (based on data from the OPTI-SCRIPT Study, a previous RCT in a similar population) with 90% power requires a sample size of 68 patients. Based on an intraclass correlation coefficient of 0.025 (OPTI-SCRIPT Study), 12 patients per practice results in 84 patients from seven practices. We will further inflate this to 120 patients from eight practices to account for loss to follow-up for the PROM secondary outcomes and to account for uncertainty given that these are estimates from different but related populations. In total, we will recruit 120 patients from eight GP practices. This sample size also reflects our experience of patient response rates for participation in these studies and that it is feasible to recruit 15 patients per practice on 10 or more medications. This study will help to establish future sample size calculations that can be used to inform a definitive cRCT.

Blinding
The trial statistician will be blinded to practice (and thus patient) allocation. Blinding of participant GP practices, patients and pharmacists will not be possible due to the nature of the intervention.

Data analysis
Descriptive statistics and estimation using CIs will be the main focus of the analysis. Descriptive statistics will be presented for all feasibility outcomes and demographic and clinical characteristics of study participants and practices. For categorical measures, frequencies and percentages will be presented, and for continuous measures, the mean and SD will be reported. For continuous measures which show evidence of some skew, a median and IQR may also be presented or substituted for the mean and SD.

While this study is not powered to explore effectiveness, we will estimate and explore potential differences in all outcome measures, in particular, the difference in the mean number of PIP medications, and the presence/absence of PIP medications, for the intervention group versus the control, with transformation as appropriate after examination of the distribution, and adjusting for baseline PIP and minimisation factors. Random effects linear or logistic regression models will be used as appropriate, including a random practice effect to account for the correlation between patients in practices. Results will be presented as the difference in means or ORs, 95% CIs and p values. Other outcomes will be explored and analysed using similar methods.

The analysis of this study will help inform future sample size calculations.

Process evaluation
To explore participant attitudes towards the intervention and the experience of the intervention delivery, a mixed-methods process evaluation combining both quantitative and qualitative methods will be conducted in line with the MRC Framework for the Process Evaluation of Complex Interventions. This framework includes an assessment of treatment fidelity of the intervention, which will be incorporated into the study design of the process evaluation study. The Bellg treatment fidelity framework will be used to monitor and enhance the validity and reliability of interventions involving behavioural change.

Quantitative data will be compiled from completed validated questionnaires, and activity logs which will be summarised with descriptive statistics.

Qualitative data will be collected using semistructured interviews with key stakeholders in the intervention process: GPs, pharmacists, nurses, practice managers and patient participants. A topic guide will be developed to explore the issues surrounding integration, context, fidelity, implementation and experiences of the intervention. Interviews will take approximately 30–60 min to complete. As it is feasible that pharmacists could increase GP workload, we will explore this issue qualitatively while interviewing GPs. Interviews will take place as per participant preference, either in person or on the phone and they will be recorded to allow for transcription, with participant permission. Data will be transcribed verbatim from the recordings following pseudonymisation of participant identifiers with a unique participant study code. Thematic analysis will be undertaken. Data will be coded and common features will be grouped to develop themes. Overarching themes will then be developed with quotations used as exemplars. Following analysis, triangulation will be employed to combine the results of both quantitative and qualitative components.

Health economic analysis
The ‘Guidelines for the Economic Evaluation of Health Technologies in Ireland’ will be used to inform the economic evaluation component. The direct costs associated with the intervention will be the focus of the evaluation, and will adopt a payer perspective. The health economic evaluation will focus on calculating direct costs related to the intervention, that is, the costs of recruitment, training and salary of the pharmacist, time/salary of GPs and associated staff, revenue and capital overheads, and travel expenses for pharmacists. In addition, the total costs associated with clinical actions linked to the medication reviews will be calculated including the costs of medications prescribed or deprescribed, cost of laboratory monitoring tests that are recommended, costs saved or incurred. Data to be collected include participants’ healthcare utilisation (from patient healthcare records). Unit costs will be applied to convert data on resource use to resource costs and total cost variables will be calculated.

A cost-effectiveness analysis will be conducted on the basis of the primary and secondary outcomes identified in...
the cRCT, that is, reduction in number of PIPs/high-risk prescriptions, deprescribing and pharmacist time. The primary analysis will evaluate the impact of the placement of a pharmacist in a GP practice for a 4-month period. Costs associated with salary, revenue, capital and travel will be informed directly by our 4-month trial data. Inflation to the 12-month time horizon will allow us to investigate potential longer term sequel avoided due to the PIP intervention made by the pharmacists. It is assumed that the changes made to a patients’ long-term medication (as a result of a pharmacist intervention) will persist beyond the 4-month intervention period. In line with this, costs incurred or saved secondary to the medication reviews will be informed by our 4-month trial data and inflated to a 12-month time horizon. A scenario analysis will evaluate the impact of the placement of a pharmacist in a GP practice for a 12-month period. Costs associated with salary, revenue, capital and travel will be informed directly by our trial data and inflated to a 12-month time horizon. The expected continued rate of pharmacists’ interventions beyond the 4-month time period (available from our data) will be extrapolated to 12 months using data published elsewhere and via expert elicitation. Costs incurred or saved secondary to medication reviews will be informed by these extrapolated data. The incremental cost per PIP intervention will be estimated for both the primary and scenario analyses. The impact on changes made to all uncertain model input parameters will be investigated in both one-way sensitivity analysis and probabilistic sensitivity analysis.

The cost-effectiveness analysis will allow us to consider the potential data requirements for a lifetime horizon in future cost-effectiveness analyses, if a definitive cRCT is carried out. This would allow us to investigate the long-term sequel avoided due to PIP intervention, giving consideration to patient or therapeutic subgroups where there is established evidence between PIP and ADR-related hospital admissions (eg, non-steroidal anti-inflammatory drugs in people with heart failure and admissions for exacerbation). Within the process evaluation, we will employ expert elicitation regarding potential wider effects of the intervention on prescribing practice to inform model assumptions.

**Continuation criteria**

We will use continuation criteria (see table 3) to determine if progressing this pilot cRCT to a definitive pragmatic cRCT is warranted. The criteria for continuation (also referred to as progression criteria) will be based around feasibility and the potential for cost-effectiveness. Quantitative and qualitative process evaluation data will be analysed to consider the following continuation criteria:

- Recruitment levels of GP practices, pharmacists and patients.
- Acceptability of intervention for GPs, pharmacists and patients.
- Feasibility of GPP intervention.

- Potential to be cost-effective.

**Ethics, data management and dissemination**

Ethical approval has been granted by the Irish College of General Practitioners Research Ethics Committee. In the event that a prescribing pattern of concern is identified by the research team, as a safeguard this will be referred to an external, independent clinician who will have no involvement in the study to advise the team on appropriate actions.

Patient healthcare records will be accessed to obtain prescription data and healthcare utilisation data. PROM data will be collected from paper-based completed patient questionnaires. For the process evaluation, data will be collected by the verbatim transcription of audio recordings of semistructured interviews. Transcription of audio recordings will be done by a third party with whom we will have a confidentiality agreement, and patient consent will be obtained for this.

Data will be pseudonymised at practice level. Participants will be known to the research team by study ID number only. One member of the research team (AC) will have access to patient contact details for follow-up data collection purposes. Required contact details will be linked to the study number and stored in a restricted access folder on a secure institutional server. Hard copies of data such as questionnaires will be stored in locked cabinets in the university research offices which are accessible only by swipe card access.

The results will be reported in peer-reviewed journals and be presented at national and international conferences.

**Patient and public involvement**

This study has patient and public (PPI) involvement through a multimorbidity patient advisory group. The patient advisory group are patients with multimorbidity who meet quarterly to discuss issues arising with research projects on multimorbidity funded through the HRB Collaborative Doctoral Award (AC is a PhD student on this programme). This PPI group was asked to contribute to the development and refinement of this intervention, as outlined in table 1.

**DISCUSSION**

Polypharmacy is a contributing factor to the treatment burden associated with multimorbidity. PIP is associated with complex polypharmacy and studies have shown that this leads to medical complications and associated costs. A wide variety of interventions have been developed to address the issue of polypharmacy and PIP in primary care settings but the evidence base for the integration of pharmacists into GP practices is limited as few high-quality RCTs, and modest effect sizes have been reported. A number of large trials have been conducted in this area in other jurisdictions, for example, Canada, the UK, Australia, New Zealand and Sweden. Pharmacists...
have direct prescribing roles in the UK, New Zealand and Canada, but similar to the Irish context, pharmacists in Australia and Sweden do not. Quality of life measures by these trials produced varying results.\textsuperscript{19} Irrespective of prescribing roles, all trials were broadly similar, adopting a model of patient medication review and quality enhancement in practice. Our own trial also has similar interventions of patient medication review and practice-level education and quality enhancement. This suggests that our results will be generalisable to other health jurisdictions. High-quality trials such as PINCER demonstrated a positive effect using a pharmacist-led information technology intervention to address medication errors.\textsuperscript{12} Our trial seeks to build on this experience in a health system with less integration of pharmacists in general practice settings. The outcomes of a non-randomised, uncontrolled feasibility study\textsuperscript{16} have been reported and based on predefined continuation criteria the trial is being progressed to further evaluation with the design of this pilot cRCT. We have reported here the protocol for the pilot cRCT. If the pilot cRCT is feasible to conduct, acceptable to stakeholders, and demonstrates a positive impact and cost-effectiveness, we will aim to proceed to a full, pragmatic cRCT. With the increased medicalisation of managing conditions, the need for interventions to address polypharmacy and PIP is warranted.

Recruitment of GPs may be a potential challenge, however feedback from the feasibility study was positive. We will seek the assistance of the HRB CTNI to support this process. Eligibility criteria for GP practices dictate that there must be \( \geq 500 \) patients aged 65 years or older for the practice to be considered. To allow for feasible delivery of the intervention, allocation of practices will be done in blocks to allow for staggered data collection. Logistical barriers will be addressed with practices to ensure that the pharmacist has access to an office, or space in an office, with access to the practice clinical software to complete their intervention-related activity. It is envisaged that the pharmacist will allocate a single block of time, however, these working practices are flexible to allow for successful integration of the pharmacist into the GP practice. As the study is designed as a pilot cRCT, a power calculation is not required. However, we have calculated that the number of patients recruited will give sufficient power to detect a significant change in PIP. Given the relatively small number of study participants the results will not be generalisable, and the primary purpose of the pilot cRCT is to test feasibility and identify appropriate processes, outcomes and sample size calculation parameters for a definitive cRCT.

In summary, the GPP Medicines Optimisation Programme aims to assess the feasibility and potential impact of general practice-based pharmacists in an Irish setting in relation to addressing PIP and GP workload.

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**Competing interests** None declared.

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**Contributors** AC and BC drafted the manuscript. AC, FM, FB, LM, KC, SMS and BC were involved in the study design and in obtaining ethical approvals. SMS was responsible for study conception. All authors read, provided feedback and approved the final manuscript.

**REFERENCES**


