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Incidence, nature and causes of avoidable significant harm in primary care in England

Citation for published version:

Avery, AJ, Sheehan, C, Bell, B, Armstrong, S, Ashcroft, DM, Boyd, MJ, Chuter, A, Cooper, A, Donnelly, A, Edwards, A, Evans, HP, Hellard, S, Lymn, J, Mehta, R, Rodgers, S, Sheikh, A, Smith, P, Williams, H, Campbell, SM & Carson-Stevens, A 2020, 'Incidence, nature and causes of avoidable significant harm in primary care in England: retrospective case note review', BMJ Quality & Safety. https://doi.org/10.1136/bmjqs-2020-011405

Digital Object Identifier (DOI):

10.1136/bmjqs-2020-011405

Link:

Link to publication record in Edinburgh Research Explorer

Document Version:

Peer reviewed version

Published In:

BMJ Quality & Safety

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Download date: 03 Dec. 2020

BMJ Quality & Safety

Incidence, nature and causes of avoidable significant harm in primary care in England: retrospective case note review.

Journal:	BMJ Quality & Safety
Manuscript ID	bmjqs-2020-011405.R1
Article Type:	Original research
Keywords:	Primary care, General practice, Patient safety

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Incidence, nature and causes of avoidable significant harm in primary (rospec. care in England:

Abstract

Objectives: To estimate the incidence of avoidable significant harm in primary care in England; describe and classify the associated patient safety incidents, and generate suggestions to mitigate risks of ameliorable factors contributing to the incidents.

Design: Retrospective case note review. Patients with significant health problems were identified and clinical judgements made on avoidability and severity of harm. Factors contributing to avoidable harm were identified and recorded.

Setting: Primary care.

Participants: Thirteen general practitioners undertook a retrospective case note review of a sample of 14,407 primary care patients registered with 12 randomly selected general practices from three regions in England (total list size: 92,255 patients).

Main Outcome Measures: The incidence of significant harm considered at least 'probably avoidable' and the nature of the safety incidents.

Results: The rate of significant harm considered at least 'probably avoidable' was 35.6 (95%CI: 23.3-48.0) per 100,000 patient-years (57.9 (95% CI: 42.2-73.7) per 100,000 based on a sensitivity analysis). Overall, 74 cases of avoidable harm were detected involving 72 patients. Three types of incident accounted for more than 90% of the problems: problems with diagnosis accounted for 45/74 (60.8%) primary incidents, followed by medication-related problems (n=19; 25.7%) and delayed referrals (n=8; 10.8%). In 59 (79.7%) cases, the significant harm could have been identified sooner (n=48) or prevented (n=11), if the GP had taken actions aligned with evidence-based guidelines.

Conclusions:

There is likely to be a substantial burden of avoidable significant harm attributable to primary care in England with diagnostic error accounting for most harms. Based on the contributory factors we found, improvements could be made through more effective implementation of existing information technology; enhanced team coordination and communication, and greater personal and informational continuity of care.

Word count: 280

Introduction

Healthcare-associated harm is an internationally recognised threat to public health and wellbeing. As many countries, across all income settings, aspire towards universal health coverage, attention has focused on the critical role of primary care-led health care systems to help achieve this goal.^{1,2} In countries like the United Kingdom (UK), over 90% of clinical encounters are delivered in community settings,³ but a clear understanding of avoidable harm is needed to enable health care systems to identify and learn from the most serious incidents and the factors amenable to intervention.

Most patient safety research has focused on hospital-based care settings resulting in a greater awareness of the frequency and causes of health care-associated errors, and the resulting burden to patients.⁴ Patient safety research in primary care has been slower^{2,5} although the profile of patient safety in primary care was provided a platform by the World Health Organization's (WHO) Safer Primary Care Expert Group (2012), and catalysed by more recently by the US National Patient Safety Foundation's call to look 'beyond hospitals to the full care continuum' and the OECD's assessment of the economic burden of unsafe primary and ambulatory care.⁶⁻⁸ The WHO's Technical Series for Safer Primary Care, where world experts have explored the existing evidence base for primary care safety, highlighted that major evidence gaps exist and robust high-quality epidemiological studies are needed to definitively establish the burden of unsafe primary care.⁹ Whilst harm from hospital-based care may be more visible, given the volume of patient consultations that occur in primary care, the aggregate burden of harm cannot be ignored.⁷

Our WHO-commissioned systematic review investigating the frequency and burden of harm in general practice concluded 2–3% of primary care encounters involved a patient safety incident, and around one in 25 of those resulted in a significant harm outcome that has a substantial impact on a patient's well-being.⁸ Included studies were notably heterogeneous in study design and definitions of outcome measures. None of the primary care studies in this, or our subsequent systematic review,¹⁰ reported the incidence of avoidable harm^{11,12} based on independent review of medical records, and few distinguished between minor and more significant harms.¹³ Also, we are aware of only one previous study that was large enough to identify substantial numbers of significant harms, but it did not report on these in detail.¹³ This means that based on the literature it has not been possible to reliably quantify the overall burden of avoidable significant patient harm in primary care.

We have addressed this issue in the current study by undertaking a large retrospective case note review study, using independent clinical reviewers, to: 1) estimate the incidence of

avoidable significant harm in primary care in England; 2) quantify, describe and classify the patient safety incidents that result in avoidable significant harm (thus showing the top categories of avoidable harm) and 3) generate suggestions to mitigate risks of ameliorable factors that contributed to the incidents. Our study is different to other primary care studies because of its specific focus on identifying and understanding significant harm, and because have estimated the incidence (rather than prevalence) of harm on the basis that this provides policy makers with a better idea of the potential burden of the problem. We have used a definition of avoidable harm based on a consensus study panel with general practitioners, 14 using real cases of unsafe general practice from our earlier national-level analysis of patient safety incident reports. 15

Methods

Our study protocol describes the methods we employed in detail,¹⁶ and an expanded version of our methods is in the supplementary materials. Box 1 provides the definitions used in the study. The study had NHS research ethics committee approval (15/EM/0411).

Participants

We used a stratified random sampling approach to invite general practices to participate from three different areas of England. We undertook a retrospective case note review of an open cohort of all primary care patients registered with participating general practices (between 1 April 2015 and 31 March 2016) to identify cases of avoidable significant harm.

Recruitment and training of data collectors

General practitioners with at least five years' experience in general practice were recruited to collect data from the participating practices, and were provided with training.¹⁶

Sampling of patient records

We sampled patient records in three stages. In Stage 1, we identified the total patient population of the practices at the start of the retrospective cohort (1 April 2015). In Stage 2, we used electronic registry queries to identify patients at increased risk of significant health problems and/or avoidable significant harm (the 'enhanced sample'). Drawing on suggestions made by the research commissioners, the literature on avoidable harm in primary care⁸ and our own experience of analysing reports of harm associated with primary care, we included patients who had: died¹⁷ been admitted to secondary care¹⁸ were resident in a care home; had multimorbidity¹⁵ or polypharmacy, 20,21 had undergone an

invasive procedure in general practice²² or had been certified unfit for work long-term. In Stage 3, one of the GP data collectors screened the electronic health record of each patient in the 'enhanced sample' to identify any new significant health problems experienced by patients over the 12 months of the study (1 April 2015 – 31 March 2016). The GPs then undertook detailed retrospective reviews of the records of this final sample of patients to identify the extent to which errors in primary healthcare provision contributed to these problems.

For the purposes of sensitivity analysis (recognising that cases might have been missed by our sampling approach), the GP data collectors also undertook a detailed records review for the following:

- 2.5% random sample of the Stage 1 population, not including patients identified for the
 Stage 2 enhanced sample; each record was examined by a single GP reviewer.
- 10% random sample of the Stage 2 enhanced sample; each record was examined by a second GP reviewer.

Identification of avoidable significant harm, and factors associated with this

For those patients with significant health problems, the GP data collectors recorded whether they found any evidence of avoidable harm. If so, the GPs provided a detailed written account of the principal problem in the patient's primary care that led to the significant health problem, a narrative describing the manner in which the significant health problem could have potentially been prevented within primary care, and a judgement on the avoidability of the significant health problem using a validated six-point scale (see Box 2). 17,23 All cases were considered in detail by the study team, and the GP data collectors were asked to provide additional information if any clarification were needed. To ensure consistency the study team made the final judgement, through consensus, in terms of the classification of avoidable significant harm.

Data collection and coding

Each of the participating general practices was visited by an informatician from the study team who collected baseline data on the practice population and ran a computer search to identify patients for the enhanced sample and for the sensitivity analyses. Using encrypted tablet computers and a Virtual Private Network (VPN) connection, the GP data collectors entered anonymised data directly into a database on a secure server at Cardiff University. The nature of the avoidable harm was recorded by the GP data collectors using the

comprehensive patient safety classification system developed in the Primary Care Patient Safety Classification (PISA) study.²⁴

Analysis

We estimated the incidence of significant harm that was considered at least probably avoidable (our primary outcome – avoidability score 4 or more) and at least possibly avoidable (avoidability score 3 or more) accompanied by 95% confidence intervals (95% CI). We assessed inter-rater reliability of judgements made using the Cohen's Kappa statistic (with 95% CI).

Members of the study team then undertook a detailed analysis of the information provided on each case of potentially avoidable significant harm and included cases with at least 'slight to modest' (score 2 or more) evidence of avoidability, as we judged that even in these cases there were important insights. We analysed the data recorded on the cases and examined the relationships between different types of incident and the factors that contributed to these incidents. As a result, we identified the most important factors contributing to avoidable significant harm.

Results

Twelve practices were recruited (as shown in Figure 1) and Table 1 shows their characteristics compared with national averages for England. The practices were similar to the English average in terms of list size, Index of Multiple Deprivation, and age and gender of patients, but had a higher percentage of non-White patients. Also, all the participating practices were rated overall as 'good' or 'outstanding' by the CQC, whereas almost 12% of the practices in England received 'inadequate' or 'requires improvement' scores.

The total list size for the 12 general practices at the start date of the study cohort (1 April 2015) was 92,255 (Stage 1). The total number of patient-years of clinical data available for the 92,255 patients over the year of the cohort (1 April 2015 - 31 March 2016) was 89,779.

The flow of patient records through the study is shown in Figure 2. The computer searches identified 12,080 patients (13.1%) for the enhanced sample (Stage 2). Their records were all examined by at least one GP data collector (first GP data collector in Figure 2), and 1,271 (10.5% random sample) were examined independently by a second GP data collector. From

the Stage 1 population of 92,255, a random sample of 2,327 (2.5%) patients (but not included in the enhanced sample) was examined by one of the GP data collectors.

Based on the assessment of the GP data collector doing the first assessment on the enhanced sample, there were 2,131 new significant health problems for 2,116 patients between 1 April 2015 and 31 March 2016 (Stage 3 – see Figure 2). For 2,054 (96.4%) of the significant health problems, the GP data collector judged that the patient had received an adequate standard of care and therefore classified these cases as having 'virtually no evidence of avoidability'.

For the remaining 77 (3.6%) cases, the first GP reviewer formally assessed avoidability and the distribution of avoidability scores, following moderation by the study team, is shown in Table 2. A further 10 cases had 'virtually no evidence of avoidability', meaning that in total 2,064 (96.9%) of the 2,131 significant health problems were considered unavoidable in primary care.

There were 32 cases (1.5%) of significant harm considered to be at least probably avoidable and 51 (2.4%) considered at least possibly avoidable. This translates into a rate of 35.6 per 100,000 patient-years (95% CI: 23.3-48.0) for significant harm considered at least probably avoidable and 56.8 per 100,000 patient-years (95% CI: 41.2-72.4) at least possibly avoidable.

Sensitivity analysis

The examination of the 2.5% sample of the patient population did not identify any additional cases of significant harm considered at least possibly avoidable. The examination of the 10% sample of the enhanced sample by a second GP reviewer identified two further cases of significant harm considered to be at least probably avoidable and four cases considered at least possibly avoidable, based on the final judgement of the study team. This means that had all the patient records in the enhanced sample been assessed independently by two GPs, there could have been an additional 20 cases considered at least probably avoidable, and 40 cases of significant harm considered at least possibly avoidable. In the sensitivity analysis, this translated into rates of 57.9 (95% CI: 42.2-73.7) per 100,000 patient-years for significant harm considered at least probably avoidable, and 101.4 (95% CI: 80.5-122.2) per 100,000 patient-years for significant harm considered at least possibly avoidable.

Interrater reliability

Where an assessment of avoidability was done, there was 77.0% agreement between GP data collectors and the study team about whether the case was considered at least possibly avoidable (Cohen's Kappa: 0.49 (95% CI: 0.29-0.69). For the 10% sample of the enhanced sample, there was 71.5% agreement between the first and second GP reviewer that a patient had at least one significant health problem (Kappa: 0.33 (95% CI: 0.27-0.38), and where an avoidability assessment was done independently by two GP data collectors, there was a 67.6% agreement about whether the significant harm was considered at least possibly avoidable (Kappa: 0.34 (95% CI: 0.02-0.66)).

Analysis of the nature and causes of avoidable significant harm

For the detailed analysis of the nature and causes of avoidable significant harm, we included
74 cases involving 72 patients. The distribution of avoidability scores for these cases is
shown in Table 3.

The distribution of different types of primary incidents for the 74 cases is shown in Table 4, with problems with diagnosis accounting for 60.8%; medication-related problems for 25.7% and delayed referrals for 10.8% (the latter relating to situations where a clinician had decided that a referral was needed, but there was such a delay in the referral being made that the patient may have been harmed as a result). Examples of these incidents are shown in Box 3 and Box 4. In relation to the 74 primary incidents, 114 underlying 'contributory factors' were identified, and these are shown in Table 5. Patient factors accounted for 71.9%, with co- or multi-morbidities the most important categories (24.6% of all contributory factors), whilst 17.5% of factors included issues such as not taking medicines as prescribed, problems with eliciting relevant information from patients or caregivers, not following medical advice, and presenting with multiple issues in a single consultation. Factors such as multimorbidity and frailty contributed either through offering alternative explanations for symptoms or by presenting clinicians with multiple competing demands. Organisational factors accounted for 21.1% of contributory factors whilst staff factors such as inadequate knowledge, skills or mistakes by healthcare professionals accounted for 7.0%. In 59 (79.7%) of the 74 cases, the significant harm could have been identified sooner (48 cases), or prevented (11 cases), if the GP had taken actions aligned with evidence-based guidelines (see examples in Box 3).

These 74 cases involved 115 healthcare professionals (81 (70.4%) GPs and 10 (8.7%) practice nurses), and only four of these (3.5%) were clearly identifiable as being from outside the participating general practices (community nurse, community optometrist, community physiotherapist, community psychiatric nurse).

Discussion

Principal findings

The estimated incidence of significant harm in English primary care considered at least 'probably' avoidable is between 35.6 and 57.9 per 100,000 patient-years (the latter figure being based on sensitivity analysis). Extrapolating our findings to the English population of 55.6 million (mid-year 2017), there are likely to be between 19,800 and 32,200 cases of 'probably avoidable' significant harm to patients each year.

The three major sources of significant avoidable harm in general practice were diagnostic error (60.8% of the avoidable incidents), medication incidents (25.7%) and delayed referrals (10.8%). In 79.7% of cases, the significant harm could have been identified sooner, or prevented, if the GP had taken actions aligned with evidence-based guidelines. The study identified a mix of organisational, clinician and patient contributory factors associated with the avoidable incidents. The majority of these were patient factors (71.9% of the total contributory factors identified) including multimorbidity, old age and complexity arising from pathophysiological factors such as frailty. Most of these factors are not ameliorable, but highlight the challenges that healthcare professionals face when trying to avoid patients coming to harm. Of the organisational factors, problems relating to continuity and coordination of care (between providers and within primary care) were most important (14.1% of the total). For example: patient did not experience a 'seamless service' due to failures in coordination and sharing of information between different providers across the health and social care system; disconnect between multiple members of the primary care in the same practice; and lack of care coordination as a patient transitions from secondary back to primary care. Mitigating risk for future patients could be achieved through targeting the organisational structures and processes underpinning the most frequent contributing factors.

Strengths and limitations of the study

This is one of the most comprehensive studies of avoidable harm in primary care, 8,10 and one of only two records review studies we are aware of that is large enough to report on substantial numbers of significant harms. 13 It is the only study of which we are aware that has reported the incidence of avoidable harm based on independent review of primary care clinical records. In terms of other potential methods of investigation, independent retrospective case note review has significant advantages over incident reports, which are more at risk of selection bias and are not well suited to accurately estimating the incidence of avoidable harm. It also has advantages over database studies, because detailed examination is required of the healthcare records (including hospital correspondence) that is not possible through clinical databases. One major limitation of case note review is the onerous task of searching for and identifying important information to build a narrative, based on what is explicitly stated or from what is absent based on the clinician's knowledge of the relevant evidence-based guidelines.

We used a stratified random sampling approach to recruit 12 general practices from three geographically different regions of England, and the independent GPs involved in data collection were experienced and were given thorough training. We used a validated method for judging the avoidability of harm, 14 and a comprehensive validated system for classifying the underlying causes of patient harm. 15 Our methodological approach used the Recursive Model of Incident Analysis and permitted us to capture the series of 'contributing incidents' that led to the final 'principal incident' prior to the patient experiencing a harmful outcome. 25 This meant we could provide the most robust and comprehensive assessment of the patient safety incidents implicated in significant avoidable harm outcomes, as well as consider the apparent underlying events when formulating our recommendations to mitigate future risk to patients.

Only a quarter of the general practices in the stratified random sample agreed to participate and this is a limitation from an epidemiological perspective. The most common reason cited for not participating was lack of time. Although the characteristics of the practices recruited were similar to those in England in most respects, none of the study practices received an overall CQC rating of 'inadequate' or 'requires improvement', whereas 2.6% and 9.1% (respectively) of all English practices received these ratings. If CQC ratings are associated with patient safety, then our study may underestimate the overall incidence of avoidable significant harm in English general practices. Our inter-rater reliability assessments showed that there was moderate agreement between the GPs in their identification of patients with significant health problems, and their judgements as to whether a patient had experienced

avoidable significant harm. Even with our sensitivity analysis, the upper limit of our estimates of the incidence of avoidable significant harm may be an underestimate. This highlights the uncertainties of estimates of frequency of harm originating from case note reviews that rely on clinical judgement. It suggests that our study could have missed some cases of avoidable significant harm but could also have included cases that others might not consider to be 'significant' or 'avoidable'.

Through our 'enhanced sample' we successfully identified patients most likely to have avoidable significant harm, but the criteria we used might be difficult to replicate in other countries. We did however manage to identify these patients through electronic medical records, and so a similar approach should be possible in countries with comprehensive primary care electronic records. Our study was not designed to detect near misses.

Comparison with other studies

We recognise from our previous systematic reviews,^{8,10} that comparing studies of avoidable harm is difficult because of different study designs and different ways of applying definitions of avoidable harm. One key difference between our study and almost all previous studies is that we report the incidence of avoidable harm rather than the prevalence (per consultation). Our approach allows for a clearer estimate of the public health burden of avoidable harm, while also recognising that some harms, especially in a primary care setting, may occur over several consultations (e.g. delayed diagnosis). Also, we have focused specifically on 'significant' harm (such as a clinically important delay in cancer diagnosis) to ensure that our findings reflect a health burden that is unquestionably of importance to patients, the public, clinicians and policymakers.

The only study we are aware of that was of a similar large size to ours, while also reporting on severity, was from a convenience sample of 48 health centres in Spain with health professionals reporting any incidents causing harm. From 96,047 consultations, 773 harms were detected with 46 of these considered 'severe'. Of all the harms, 64.3% were considered preventable, and applying this percentage to the severe harms suggests a prevalence of 30.8 severe harms per 100,000 consultations. It is not possible to directly compare this with the incidence figures from our study (where the same harm may have been apparent across several consultations over the course of the 12 months), but the overall rates of significant harm are probably not widely dissimilar.

There is considerable variation in studies reporting categories of avoidable harm as well as contributory factors. Nevertheless, our findings are in keeping with a systematic review that found that diagnostic errors were among the most important causes of avoidable harm, 10 and a review of the global burden of diagnostic errors in primary care, 26 while the systematic review¹⁰ (and other studies) have highlighted the importance of prescribing errors.¹³ In relation to diagnostic delay in cancer, a recent study has highlighted that in almost half of cases this is attributable to primary care.²⁷ with problems with clinical appraisal of the patient and referral being particularly important. Our study has specifically highlighted the importance of delays in making a referral, 15,23 and this has been highlighted as an important problem by the Institute for Healthcare Improvement.²⁸ In comparison with other studies. ours is unusual in reporting such a high level of patient factors contributing to patient harm. While many of these cannot be considered the reason for the harm being avoidable, the findings suggest that factors such as multimorbidity, frailty and complex presentations may make it more difficult for clinicians to make timely and accurate diagnosis and avoid medication errors. In relation to contributory factors that are avoidable, our findings are in keeping with other studies that have highlighted organisational issues and communication problems.8,12,29

Implications for clinicians and policy makers

This study has estimated the incidence of significant harm in English primary care considered at least 'probably' avoidable, which translates to 3-4 cases per year for an average general practice of 8,000 patients. Efforts to make improvements should focus on addressing the structures and processes underpinning the identified patient, clinician and organisational contributing factors. For example, better organisation of key systems (e.g. referrals, test result management, identifying non-adherence) and related administration could have prevented most incidents. Some of the earliest patient safety studies carried out in primary care over two decades ago pointed to administrative failures, such as the mismanagement of test results, as the root cause of the commonest incidents concerning diagnosis and medication,³⁰⁻³² and an Australian study concluded about 70% of incidents were related to processes of providing healthcare, rather than gaps in the knowledge and skills of health professionals.¹¹ The current study provides considerable insights into the ameliorable contributory factors associated with avoidable significant harm in primary care, which in turn have generated the following suggestions for improvement.

More effective implementation of existing information technology solutions could ensure that planned action such as referrals take place in a timely way.²⁶ Enhanced team coordination

and communication could ensure that patients are seen (or have necessary investigations), or that they are recalled for follow-up investigations or assessment, when needed. Currently it is largely down to individual primary care teams, and individual healthcare practitioners, to develop their own strategies. Without stifling innovation, however, it might be helpful to model 'what best practice looks like' in relation to preventing patients from coming to harm. Interventions like the 'QRISK®3-2018' algorithm to calculate a person's risk of developing a 'heart attack' or stroke are now commonly integrated into electronic health record systems and used by clinicians to explain and manage risk and support patient decision-making.³³ In a similar way, the factors implicated in patient safety incidents that we have identified might be considered as signals for future algorithms for development and validation, either to flag up patients for timely clinical review to mitigate current risk levels or to proactively detect risks of future unsafe care.

Our study suggests that lack of continuity of care may contribute to avoidable significant harm in some cases.³⁴ Recent systematic reviews suggest that low continuity of care is associated with a higher risk of mortality across different healthcare settings, 35 and specifically in general practice.³⁶ In some cases in our study, follow up by the same primary healthcare practitioner could have been helpful to enable earlier recognition of the progression of a serious health problem. In other cases, better 'informational continuity'37 could have helped to ensure that the assessment and suggested follow-up plans from a previous consultation better informed the next consultation. This should be facilitated by electronic health records, but we found several examples where recommendations from one consultation were not acted upon in a subsequent consultation involving a different healthcare practitioner. Nevertheless, high levels of personal continuity may not always be best for patients. A recent study qualitative showed a mixed picture in terms of patients' perceptions of whether personal continuity improved safety, or not,38 while a cross-sectional ecological study found that general practices that appeared to have high levels of personal continuity, did fewer urgent referrals;³⁹ this does not necessary mean they were less safe, but it is a potential cause for concern.

Some of the cases of avoidable significant harm in our study were associated with GPs having too many problems to deal with adequately in a single consultation, with significant health problems not detected early enough because of lack of effective and timely clinical history taking, examination or investigation. Some of the contributory factors associated with patient behaviours may have resulted from their concerns being unrecognised or unresolved, as highlighted in a study of missed opportunities in cancer diagnosis.⁴⁰ When such incidents occur in general practice, it is essential that practice teams know how to

generate learning from the incident, including how to identify vulnerabilities in their existing structures and processes, and feel confident to plan and test changes that could achieve improved outcomes for future patients.⁴¹ The introduction of quality improvement domains into 2019/20 Quality and Outcomes Framework by NHS England represents a promising commitment for supporting practices to learn about and develop their approach to systems improvement⁴² as does the 2019 NHS Patient Safety Strategy.⁴³

Conclusion

There is likely to be a substantial burden of avoidable significant harm attributable to primary care in England with diagnostic error accounting for most harms, followed by medication error and delays in making a referral once a referral decision had been made. Based on the contributory factors we found, improvements could be made through more effective implementation of existing information technology; enhanced team coordination and communication, and greater personal and informational continuity of care.

Word count for the manuscript following editing of text in the methods section (full version of methods moved to supplementary materials): 4500 (excluding the abstract and summary box):

Acknowledgements

We thank Dr Christine Johnson and others who helped us to recruit the GPs for the retrospective case note review. We also thank all of the practices that willingly took part in the retrospective case note review study; they were all very welcoming to the GPs that undertook the data collection, and provided space, access to computers, and support to allow them to undertake their work. We are particularly indebted to the GPs who did the data collection as they were highly committed to the study, and many of them went the extra mile in terms of fitting in extra data collection sessions to enable us to complete the study within the timeframe agreed. In particular, we thank Dr Richard Thomas for providing most of the training for the GPs. We thank Ed Longridge for running most of the baseline computer searches in the practices, and for collecting data to allow us to calculate 'patient-years' (the denominator for our study). We thank Dr Sukhmeet Panesar for advice at the development stage of the project, particularly concerning literature on avoidable harm. We are extremely grateful to the members of our external advisory group (chaired by Professor Charles Vincent, and including Professor Susan Dovey, Dr Frances Healey (funder representative) and Professor Gordon Schiff) who gave very helpful advice throughout the study, and were

particularly helpful in terms of the interpretation of our findings. We also thank members of the East Midlands Academic Health Science Network PPI Senate and the Greater Manchester Patient Safety Translational Research Centre and Health Innovation Manchester Patient Experience Group for reviewing the findings and providing helpful comments from the perspective of members of the public.

Contributorship statement

All authors made a substantial contribution to the conception or design of the work; or the acquisition, analysis, or interpretation of data for the work, Specifically, AJA, ACh, ACS, AD, AE, AS, DA, HE, MB, SC, SJA, SR conceived the study; AJA, ACh, ACS, AD, AE, AS, DA, HE, JL, MB, PS, RM, SC, SJA, SR designed the study; AJA, ACS, CS, DA, SC, SR recruited the GP reviewers and practices; AJA, ACS, AE, CS, HE trained the GP reviewers; AJA, ACh, ACo, ACS, AD, AE, CS, DA, HE, JL, MB, SC, SH, SR designed the data collection process; AJA, ACo, ACS, AE, BB, HW, SH processed the data; AJA, ACo, ACS, AE, HW, SH contributed to team judgements of avoidability of harm; AJA, ACo, ACS, AE, BB, HW analysed the data, and all authors were involved in interpretation of the data; AND AJA, ACS, AE, AS, BB, CS, DA, SC drafted the paper and all authors revised it critically for important intellectual content; AND all authors gave final approval of the version to be published; AND all authors gave agreement to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. AA is guarantor for the paper and accepts full responsibility for the work and/or the conduct of the study, had access to the data, and controlled the decision to publish.

Data sharing statement

Anonymised summary data extracted from patient records regarding the avoidable harms detected in this study will be available upon reasonable request.

Dissemination declaration

We plan to disseminate the results to the general practices that participated in the study.

Ethical and regulatory approvals

The study was granted a favourable opinion by the East Midlands Nottingham 2 Research Ethics Committee on 15 January 2016 (reference: 15/EM/0411) and Confidentiality Advisory Group (CAG) approval for access to medical records without consent under section 251 of the NHS Act 2006 on 11 April 2016 (reference: 15/CAG/0182). Research & Development (R&D) approvals were obtained for the Clinical Commissioning Groups (CCGs) where the study occurred.

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Funding acknowledgement and disclaimer

This paper is based on independent research commissioned and funded by the NIHR Policy Research Programme ('Understanding the Nature and Frequency of Avoidable Harm in Primary Care', Ref: PR-R11-0914-11001). GP time was funded by the NIHR Greater Manchester Patient Safety Translational Research Centre (NIHR Greater Manchester PSTRC). The views expressed in the publication are those of the authors and not necessarily those of the NIHR or the Department of Health and Social Care.

Patient and public involvement and engagement statement

This project had a dedicated lay co-applicant [initials removed for triple blind review] and lay patient [initials removed for triple blind review] who contributed to the study application and the early study development meetings to provide their views on the operationalisation of definitions of avoidable significant harm in the study.

Our lay co-applicant and lay patient supported and checked the patient-facing documents, including the study information sheets and helped develop the care home information documents about the study. Both were members of the project management group, which convened monthly. They actively contributed to discussions about the conduct of the study and had a particularly active role in the interpretation of the findings; attending case analysis meetings dedicated to examining the narratives of each of the cases and discussing the patient-related contributory factors. Both supported the contacting of general practice and CCG Patient and Participation Groups (PPGs) to inform patients about the study and attended dissemination events for members of the public.

Transparency declaration

As the lead author, [name removed for triple blind review] affirms that this manuscript is an honest, accurate, and transparent account of the study being reported; that no important aspects of the study have been omitted; and that any discrepancies from the study as planned (and, if relevant, registered) have been explained.

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Box 1: Definitions used in our study

Significant harm

Our definition of significant harm was informed by the international classification of patient safety definitions of moderate harm, severe harm and death outcomes.⁴⁴ The definition used was as follows:

'A patient outcome is symptomatic, which required more intensive intervention than might otherwise have been required (eg, additional operative procedure) and resulted in an escalation of care (eg, hospital admission), or death. This caused a loss of function of at least one bodily organ, which may have been a temporary or permanent loss of its function'

Avoidability

Our definition of avoidability was informed by our RAND / UCLA appropriateness methods study¹⁴ to contextualise our definition of significant harm. The definition used was as follows:

'a patient safety incident could have probably, or totally been avoided by the timely intervention of a health care professional in family practice (e.g. investigations, treatment, safety netting) and / or an administrative process (e.g. referrals, alerts in electronic health records, procedures for following up results) in accordance with accepted standards of evidence-based practice and / or clinical governance and / or the Bolam test.'44



Box 2: Six-Point Avoidability Scale 17,23

Rating	Category	Description
1	Totally unavoidable	Virtually no evidence of avoidability
2	Unavoidable	Slight to modest evidence of avoidability
3	Possibly avoidable	Possibly avoidable, less than 50-50, but close call
4	Probably avoidable	Probably avoidable, more than 50-50, but close call
5	Probably avoidable	Strong evidence of avoidability
6	Totally avoidable	Virtually certain evidence of avoidability



Table 1: Characteristics and summary statistics of the 12 participating general practices compared with English averages

General practice	List size*	Mean age in years†	Age ≥ 65 years n (%) [‡]	Gender: Male n (%) Female n (%) [†]	Ethnicity: non-White ethnic groups n (%)§	Index of Multiple Deprivation (Decile)	Rural/ urban [¶]	CQC safety rating**	CQC overall rating**
Α	23687	42.59	4937 (21.0)	11497 (48.9) 12014 (51.1)	611 (2.6)	10.1 (9)	Rural	Good	Good
В	6780	37.03	1021 (15.3)	3123 (46.8) 3551 (53.2)	1041 (15.6)	45.0 (1)	Urban	Good	Good
С	4128	39.18	535 (13.2)	2113 (52.1) 1942 (47.9)	965 (23.8)	26.9 (4)	Urban	Good	Outstanding
D	9533	41.24	1724 (17.8)	4756 (49.1) 4931 (50.9)	436 (4.5)	18.3 (7)	Urban	Good	Good
F	8044	34.76	735 (8.7)	4070 (48.2) 4373 (51.8)	4120 (48.8)	28.2 (4)	Urban	Requires Improvement	Good
G	7311	31.45	541 (7.5)	3592 (49.8) 3621 (50.2)	2936 (40.7)	55.5 (1)	Urban	Requires Improvement	Good
Н	3841	34.90	218 (5.1)	2205 (51.5) 2077 (48.5)	1494 (34.9)	23.3 (5)	Urban	Good	Good
I	6636	37.18	814 (13.2)	3181 (51.6) 2983 (48.4)	1467 (23.8)	26.9 (2)	Urban	Good	Good
J	3447	47.96	980 (30.1)	1560 (47.9) 1696 (52.1)	94 (2.9)	7.1 (10)	Rural	Good	Good
К	9310	41.35	1697 (18.8)	4478 (49.6) 4551 (50.4)	153 (1.7)	21.8 (6)	Urban	Good	Good
L	5202	37.36	744 (13.4)	2676 (48.2) 2875 (51.8)	983 (17.7)	22.1 (5)	Urban	Good	Good
M	4336	33.46	326 (6.5)	2651 (52.9) 2360 (47.1)	3357 (67.0)	23.2 (5)	Urban	Good	Good

	Mean list size (SD)*	Mean age in years [†] (SD)	Mean % aged ≥ 65 years (SD)‡	% Male (SD) % Female (SD)†	% non-White ethnic groups [§] (SD)	Mean Index of Multiple Deprivation [®] (SD)	Number rural/ urban [¶]	CQC safety rating**(%)	CQC overall rating** percentage
All study practices++	7688 (5453)	38.87 (4.03)	15.4 (6.0)	49.4 (1.54) 50.6 (1.54)	19.0 (19.6)	23.5 (13.2)	10 urban 2 rural	Good (83.3%) Requires improvement (16.7%)	Good (91.7%) Outstanding (8.3%)
	Mean list size [‡]	Mean age in years [†]	% aged ≥ 65 years‡	% Male % Female [†]	% non-White ethnic groups (%)‡‡	Mean Index of Multiple Deprivation [®]			Overall verage**
English average	7586	39.85	17.2	49.83 50.17	14.0	21.8		For all English Overall rating v Outstanding (4 (84.2%); Requi Improvement (9 Inadequate (2.0	vas: .1%); Good ires 9.1%);

^{*}Taken from NHS Digital on 01-04-2015 http://digital.nhs.uk/catalogue/PUB17356

†Taken from NHS Digital April 2017. https://digital.nhs.uk/catalogue/PUB23475

‡For 2016 Accessed from Public Health England National General Practice Profiles. http://fingertips.phe.org.uk/profile/general-practice/data

§Taken from 2011 Census. Accessed from Public Health England National General Practice Profiles. http://fingertips.phe.org.uk/profile/general-practice/data

Illndex of Multiple Deprivation 2015. Accessed from Public Health England National General Practice Profiles. http://fingertips.phe.org.uk/profile/general-practice/data

Taken from the 2011 census figure for the population of the city or town where the practice was located.

‡‡Taken from 2011 Census. https://www.ons.gov.uk/peoplepopulationandcommunity/culturalidentity/ethnicity/articles/ethnicityandnationalidentityinenglandandwales/2012-12-11 §§As of the end of February 2017

^{**}Taken from Care Quality Commission (CQC) on February 2017 http://www.cqc.org.uk/what-we-do/services-we-regulate/doctorsgps

^{††}The practice average and standard deviation use values that are weighted by the practice list size.

Table 2: Avoidability of the 2131 new significant health problems identified by the first **GP** data collector

Avoidability classification	Numbe	r (%)
Totally avoidable: Virtually certain evidence of avoidability	0	(0.0)
Probably avoidable: Strong evidence of avoidability	14	(0.7)
Probably avoidable: Probably avoidable, > 50:50, but close call	18	(8.0)
Possibly avoidable: Possibly avoidable, <50:50, but close call	19	(0.9)
Unavoidable: Slight to modest evidence of avoidability	16	(0.7)
Totally unavoidable: Virtually no evidence of avoidability (based on study team avoidability assessment)	10	(0.5)
Totally unavoidable: Virtually no evidence of avoidability (based on GP assessment that there had been an 'adequate standard of care')	2054	(96.4)
Total	2131	(100)
		27

Table 3: Summary of cases judged by the study team to have significant harm with at least slight to modest evidence of avoidability

0.000	Avoidability rating following moderation of all cases by the study team							
Cases	Slight to modest evidence of avoidability	modest avoidable, avoidable, evidence of <50-50, >50-50,		Strong Virtually evidence of avoidability evidence of avoidability		Total		
Cases from enhanced sample (1st GP data collector)	16	19	18	14	0	67		
Additional cases from 10% sample of enhanced sample (2 nd GP data collector)	2	2	1	1	0	6		
Additional case from 2.5% sample (not from enhanced sample)	1	0	0	0	0	1		
Total	19	21	19	15	0	74		

Table 4: Distribution of different types of primary incidents

Types of primary incident: incidents occurring proximal (chronologically) to the patient outcome		At least slight to modest evidence of avoidability n (%)		possible nce of ability %)	prob evide avoid	At least probable evidence of avoidability n (%)	
Diagnostic errors	45	(60.8)	34	(61.8)	22	(64.7)	
Wrong diagnosis – original diagnosis is found to be incorrect because the true cause is discovered later.	16	(21.6)	13	(23.6)	11	(32.4)	
Delayed diagnosis (non-cancer) – diagnosis could have been made earlier if care was evidence-based.	21	(28.4)	15	(27.3)	10	(29.4)	
Delayed cancer diagnosis	8	(10.8)	6	(10.9)	1	(2.9)	
Medication errors	19	(25.7)	13	(23.6)	6	(17.6)	
No drug treatment given	4	(5.4)	3	(5.4)	2	(5.9)	
Insufficient drug treatment given	4	(5.4)	4	(7.3)	1	(2.9)	
Prescribing errors	6	(8.1)	4	(7.3)	1	(2.9)	
Monitoring errors	2	(2.7)	2	(3.6)	2	(5.9)	
Adverse drug reaction	1	(1.3)	-		-		
Medication not commenced in a timely manner	1	(1.3)	-		-		
Vaccine administration	1	(1.3)	-		-		
Referral errors	8	(10.8)	7	(12.7)	6	(17.6)	
Delayed referral	7	(9.4)	6	(10.9)	6	(17.6)	
Referral not performed when indicated	1	(1.3)	1	(1.8)	-		
Other	2	(2.7)	1	(1.8)	-		
Patient communication not sent from secondary to primary care	1	(1.3)	1	(1.8)	-		
Incorrect test ordered	1	(1.3)	-		-		
Total (%)	74	(100)	55	(100)	34	(100)	

Box 3. Examples of avoidable significant harm

30-week delay in diagnosing throat cancer (avoidability rating: 5; strong evidence of avoidability)

A middle-aged patient attended the GP with a hoarse voice and difficult swallowing. He reported that his mouth felt like the time he had thrush which required a referral to a gastroenterologist for an endoscopy and it eventually settled with an anti-fungal medication. In the medical records the GP described signs of oral thrush in the mouth and wrote, 'if does not settle, consider oral treatment or referral back to gastroenterologist'. Over the next 19 weeks the patient returned, with six visits to the original and different GPs, with on and off 'red flag' symptoms that were either improving or worsening or of varying intensity (hoarseness, swallowing issues, odd breathing pattern, difficulty talking, sensation of a lump in throat) and each time was given a course of anti-fungal medication. From week 12, the medical records describe 'food getting stuck'. At week 15, a referral for endoscopy was made and the patient was seen one week later. At this point, the patient informed the gastroenterologist he was losing weight. No abnormalities were seen on endoscopy and the patient was told to go back to his GP and have his hoarse voice investigated further (with the same advice in a letter to the GP). Four weeks later the patient was seen by the GP and a non-urgent referral to ENT was made. At week 30 the patient was seen by ENT and a rare type of neck cancer was diagnosed requiring chemotherapy.

Seven-year delay in diagnosis of prostate cancer (avoidability rating: 4; probably avoidable, more than 50:50, but close call)

An elderly patient with Type 2 diabetes mellitus attended a nurse appointment stating that he was experiencing nocturnal frequency. It was suspected this was due to poorly controlled diabetes and amendments to his medication regime were made. Six weeks later, the patient had a telephone consultation with the GP, since he was concerned about weight loss, a loss of appetite, increased urinary frequency and night cramps. He was booked with the GP for a face-to-face consultation the following day. It was also noted he had a weight loss of 2-3Kg, he was urinating at least 5-6 times per night, and he felt nauseous. The GP felt the signs and symptoms were related to poorly controlled diabetes and arranged for the patient to be reviewed by the practice nurse. Over the next week, blood tests (glycosylated haemoglobin and 'urea and electrolytes' (U&E)) and urine analysis were undertaken, and his antihyperglycaemic medications were amended. The nurse followed the patient up a few days later, where a further drop in weight was noted. The patient reported some improvement in symptoms since he was now getting up at night to pass urine four instead of six times. The nurse advised a follow-up appointment with the GP, which did not occur until four weeks later. At this time, the GP noted the patient had seven years previously had a raised prostate specific antigen (PSA). However, the patient had been unable to tolerate a biopsy for a definitive diagnosis, so six-monthly PSA testing was advised; however, the patient did not have a followup PSA in the subsequent seven year period. On noting this, the GP advised the patient to have a PSA test which was undertaken three weeks later. The PSA measured very high (>100 ng/mL). The patient had a GP appointment one week later when he was informed his PSA was raised. An urgent 'suspected cancer' referral was made. He was seen by a urologist the following week and diagnosed with localised prostate cancer requiring a transurethral resection of the prostate.

16-month delay in diagnosing non-insulin dependent diabetes mellitus (Avoidability rating: 3; possibly avoidable, less than 50-50, but close call)

An impaired fasting glucose was identified in a middle-aged patient and was followed up with a glucose tolerance test (GTT). The patient was seen by GP (A) soon after and was informed the GTT revealed an impaired glucose tolerance and was given dietary and lifestyle advice.

The patient was told to have a repeat test four months later. The patient was seen four months later by a different GP (B); however, the focus of the consultation was on yellow sclerae and liver function tests (LFTs) were ordered. A test to assess diabetes was not requested. The patient presented two months later with weight loss, and GP (B) referred the patient for an endoscopy and a repeat liver function test. A follow-up telephone call one month later occurred to discuss the LFTs with GP (B). Three months later, the patient presented to GP (B) with tiredness and fatigue. Again, blood tests were requested but did not include tests for diabetes. Six months later, a blood glucose was undertaken as part of an annual review and following two fasting blood glucose tests one week apart, poorly controlled non-insulin dependent diabetes mellitus was diagnosed.

Four-month delay in referral for an ischaemic limb (avoidability rating: 5; strong evidence of avoidability)

A patient in his early 60s stubbed his big toe three weeks prior to attending a nurse appointment at the general practice. The patient was known to have cardiovascular disease including hypertension (prescribed two antihypertensives) and raised cholesterol (prescribed a statin). The nurse noted the toe was bruised, painful, red and had a foul odour. The patient was prescribed antibiotics for a presumed infection and a referral was made to podiatry for removal of an associated in-growing toenail. Four weeks later, a podiatrist was unable to detect a dorsalis pedis or posterior tibial pulse in the affected foot and the patient was advised to see a GP urgently. The patient was reviewed by the GP and a referral to a vascular surgeon was discussed, but not made. Instead a further consultation with the same GP in four weeks was agreed, with regular nursing reviews of wound healing in the interim. At the first follow up nurse review one week later, the nurse noted that the toe was healing, but there were no pulses with the Doppler scan, and that the patient informed the nurse he needed to sleep with the foot outside the bed because it was so painful. The GP saw him three days later and decided to make a non-urgent vascular referral, but the letter was not sent to the vascular surgeons for six weeks. The patient was seen in a vascular clinic nearly four months after the podiatrist noted absent pulses and was informed that he had critical leg ischaemia and needed surgery. He underwent a right superficial femoral artery (SFA) and posterior tibial artery (PTA) stent four weeks later. Some nine months later, he required amoutation of his big toe and second digit.

Long-term nephrotoxic medications in older adults

We observed two cases where patients with known reduced kidney function were receiving long-term potentially nephrotoxic drugs. One of the patients was prescribed naproxen 'as required' for gout but received a twice daily monthly supply for >12 months whilst concurrently receiving long-term nitrofurantoin. A hospital admission for acute-on-chronic kidney injury was required (avoidability rating: 5; strong evidence of avoidability).

Another patient was taking lithium and should have had three-monthly U&E blood tests to monitor their kidney function. This did not happen for 15 months and the patient was admitted with acute kidney injury (avoidability rating: 4; probably avoidable, more than 50:50, but close call).

Box 4. Underlying incidents resulting in delayed diagnoses (history taking, examination, investigation, communication and referral)

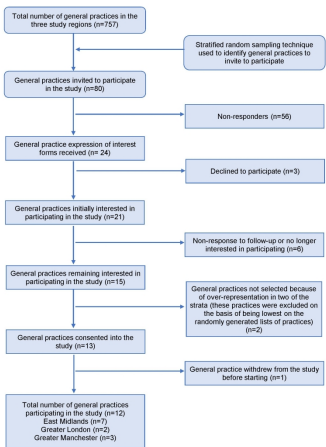
- Incomplete history taking (one case), e.g. not enquiring about red flags and not documenting salient negatives, inaccurate medical records (two cases), and inadequate documentation of care delivered (one case).
- Absent or delayed physical examination (five cases), e.g. advising patient to book another visit to undertake a pelvic examination.
- Not ordering correct investigations (three cases), e.g. no follow up or investigations ordered for an older adult with a three-week history of diarrhoea with blood and mucous; the patient eventually required an emergency admission and Crohn's disease was diagnosed.
- Failing to order a necessary investigation (one case), e.g. not testing for diabetes mellitus when presenting with lower urinary tract symptoms.
- Inappropriate responses to laboratory (three cases) or imaging (one case) investigations, e.g. i) not recognising the cut off for diagnosing Type 2 diabetes; ii) not arranging a follow-up chest x-ray (that had been advised by a radiologist) in a patient with an opacity seen on serial radiographs (this resulted in a delay in referral to respiratory medicine for an eventual diagnosis of lung cancer).
- Transfer of information about the patient which included delays in the communication being sent (two cases) or not sent at all by secondary care (one case), or a communication received but not actioned in primary care (three cases).
- Incorrect advice being given to the patient (one case), e.g. patient with insulin-dependent diabetes not given information about how to prepare for an endoscopy and GP did not inform secondary care the patient was diabetic.
- Delayed referral (seven cases), referral not made (one case) or referral sent to the wrong location.

Table 5: Distribution of contributory factors

Types of contributory factor: circumstances, actions or influences which are thought to have played a part in the origin or development, or to increase the risk, of a patient safety incident ⁴⁶	At least slight to modest evidence of avoidability n (%)		At least possible evidence of avoidability n (%)		At least probable evidence of avoidability n (%)	
Patient factors	82	(71.9)	59	(69.4)	32	(68.1)
Multimorbidity: patient has two or more chronic medical conditions	20	(17.5)	15	(17.6)	11	(23.4)
Co-morbidity: the presence of one or more additional diseases	8	(7.0)	5	(5.9)	3	(6.4)
Rare presentation: an uncommon pattern of signs or symptoms	8	(7.0)	5	(5.9)	3	(6.4)
Previous medical / medication history	8	(7.0)	5	(5.9)	3	(6.4)
Patient age	7	(6.1)	6	(7.1)	3	(6.4)
Pathophysiological factors: the patient's physical and medical well-being and health inclusive of frailty	6	(5.3)	5	(5.9)	3	(6.4)
Clinician perception of patient behaviours: the way in which patients or caregivers act towards clinicians	6	(5.3)	5	(5.9)	1	(2.1)
Response to medical advice: patient does not appear to follow the advice or instructions given by the clinician	6	(5.3)	4	(4.7)	-	
Complex agenda: patient presents with multiple issues in a single consultation	4	(3.5)	3	(3.5)	1	(2.1)
Medication taking: patient does not appear to take medication as prescribed	2	(1.8)	2	(2.4)	-	
Clinical history taking: problems with eliciting relevant information	2	(1.8)	1	(1.2)	1	(2.1)
Language: patient unable to communicate in English	2	(1.8)	2	(2.4)	2	(4.3)
Disability: a physical or mental condition that limits a person's movements, senses, or activities	2	(1.8)	1	(1.2)	1	(2.1)
Does not leave the house or home	1	(0.9)	-		-	
Staff factors	8	(7.0)	7	(8.2)	5	(10.6)
Inadequate knowledge/skill set	6	(5.3)	6	(7.1)	4	(8.5)
Mistake	2	(1.8)	1	(1.2)	1	(2.1)

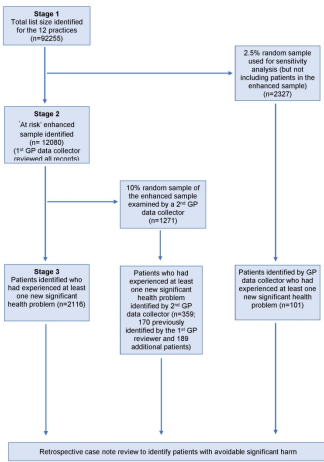
Types of contributory factor: circumstances, actions or influences which are thought to have played a part in the origin or development, or to increase the risk, of a patient safety incident ⁴⁶	At least slight to modest evidence of avoidability n (%)		At least possible evidence of avoidability n (%)		At least probable evidence of avoidability n (%)	
Organisational issues	24	(21.1)	19	(22.4)	10	(21.3)
Continuity of care across system: problem with the delivery of a 'seamless service' through integration, coordination and the sharing of information between different providers	8	(7.0)	8	(9.4)	4	(8.5)
Continuity of care within primary care: seen by multiple members of team within the same practice	6	(5.3)	3	(3.5)	1	(2.1)
Continuity of care between secondary and primary care: lack of coordinated care	2	(1.8)	1	(1.2)	-	
Protocols/ policies/ standards/guidelines inadequate, inefficient, absent or not available (specific problems noted below)	2	(1.8)	1	(1.2)	1	(2.1)
Investigations	2	(1.8)	2	(2.4)	1	(2.1)
Repeat prescribing	1	(0.9)	1	(1.2)	1	(2.1)
Referral	1	(0.9)	1	(1.2)	-	
Locum or agency staff	1	(0.9)	1	(1.2)	1	(2.1)
Waiting lists for 'urgent' referrals	1	(0.9)	1	(1.2)	1	(2.1)
Total (%)	114	(100)	85	(100)	47	(100)

Figure 1. Flow chart showing how practices were recruited



209x297mm (300 x 300 DPI)

Figure 2. Stages of the study and flow of patient records through the study



209x297mm (300 x 300 DPI)

Supplementary materials

Incidence, nature and causes of avoidable significant harm in primary care in England: retrospective case note review

Details of methods used in the study

Our study protocol describes the methods we employed in detail.¹ Box 1 in the main manuscript provides the definitions used in the study. The study had NHS research ethics committee approval (15/EM/0411), Confidentiality Advisory Group (CAG) support under section 251 to process patient identifiable information without consent (15/CAG/0182) and Research and Development (R&D) approvals.

Participants

We undertook a retrospective case note review of an open cohort of all primary care patients registered with participating general practices (between 1 April 2015 and 31 March 2016) to identify cases of avoidable significant harm. The study took place in 12 general practices from three different areas of England: East Midlands (n=7), Greater London (n=2) and Greater Manchester (n=3). The East Midlands and Greater Manchester were chosen for convenience as this is where most of the English members of our team are based. London was selected to provide geographical and demographic balance.

General practices were eligible to participate if they provided written informed consent, delivered NHS services, had electronic health records and used one of the three main computer systems in England (i.e. EMIS Web, TPP SystmOne, or INPS Vision). General practices were excluded from the study if they were involved in a major reorganisation (such as a merger with another practice) between 1 April 2015 and 31 March 2016 since this would have made it difficult to identify the practice list size for the retrospective case note review.

We aimed to sample general practices with characteristics representative of English practices as a whole, with a total population of up to 100,000. This figure was based on a pilot study, which demonstrated that this was the largest sample we could manage within the substantial available funding, whilst also conducting the study to a high standard. We estimated the precision of our study based on different possible rates of avoidable significant harm. For example, for a rate of avoidable significant harm of 40 per 100,000 patients per

year, the precision (based on 95% confidence intervals) was estimated to be between 28 and 52 per 100,000 patients per year.

From the three regions, we identified a total of 757 practices: East Midlands (n=266), Greater London (n=366) and Greater Manchester (n=125). We used stratified random sampling to identify the general practices to approach. Firstly, the practices from each area were stratified by list size into quartiles, with list sizes taken from the NHS Digital website.² Secondly, the practices from each area and each quartile were listed in computer-generated random order. We then selected the 80 practices appearing at the top of the stratified random lists, consisting of 40 practices from the East Midlands (10 practices in each quartile) and 20 practices from each of Greater London and Greater Manchester (five practices in each quartile). Practices were over-recruited from the East Midlands given most of the GPs recruited for data collection were based in this region.

We emailed and/or wrote to general practices (via the practice manager and general practitioners within the practices) inviting participation. We used a range of approaches to encourage participation, including prior publicity about the study, engaging local opinion leaders and providing reassurance about data confidentiality. Of the 80 practices approached, 12 were included in the study (see Figure 1).

Patients in the practices were excluded if they had a computer code in their clinical records indicating that they did not wish to be included in research studies. Patients were also excluded if they completed an opt-out form.

Recruitment and training of data collectors

General practitioners with at least five years' experience in general practice were recruited to collect data from the participating practices. These GPs were recruited from the East Midlands, Greater Manchester and Greater London via the Royal College of General Practitioners and existing contacts. Thirteen general practitioners were recruited and trained to ensure a consistent approach to identifying and classifying patients with avoidable significant harm. Further details are provided in our protocol paper.¹

Sampling of patient records

We sampled patient records in three stages. In Stage 1, we identified the total patient population of the practices at the start of the retrospective cohort. In Stage 2, we identified patients at increased risk of avoidable significant harm (the 'enhanced sample'), and in

Stage 3, we identified those from Stage 2 who had experienced a significant new health problem during the 12-month retrospective review period.

The population for Stage 1 comprised those patients registered with the 12 general practices at the start of the retrospective cohort (1 April 2015). To identify patients at increased risk of avoidable significant harm (Stage 2), we drew upon suggestions made by the research commissioners, (the National Institute for Health Research (NIHR) Policy Research Programme), the literature on avoidable harm in primary care³ and our own experience of inductively analysing reports of harm associated with primary care in the National Reporting and Learning System (NRLS).4 We included patients with characteristics considered to be associated with significant health problems and/or increased risk of patient safety incidents. The identification of patients with a higher likelihood of significant health problems allowed us to focus on those cases where any avoidable harm was likely to be significant too. We included those who had: died⁵ or had been admitted to hospital or a mental health facility⁶ as these were likely to have experienced a significant health problem; those that were resident in a care home as they were likely to have significant health problems and increased risk of medication errors; those that had 10 or more repeat medications, 8,9 as they were at greatest risk of harm from medication error; those with four or more major morbidities as our previous study had shown multi-morbidity to be associated with avoidable harm;5 those that had undergone an invasive procedure in general practice, such as a minor operation as safety concerns have been raised about this 10 and those that had been certified unfit for work longterm, as this was suggested by our funder, as it might have resulted from an avoidable harm.

Electronic registry queries at each practice (for 12 months from the start of the retrospective cohort) identified these patients who formed the 'enhanced sample'. Search strategies were developed and tested for the medical record systems of participating practices. This was an iterative process aimed at identifying 10-15% of the population for the enhanced sample and influenced the choice of four or more comorbidities and 10 or more repeat medications (smaller numbers of each would have resulted in an enhanced sample that was too large for the resources available for detailed records review).

The approach we used was different to that used in trigger tool methods¹¹ as were trying to identify a sample for detailed case note review, whereas trigger tool methods are applied to a patient sample that has already been selected. There was overlap in the criterion of

hospital admissions, for example,¹¹ but other 'triggers' (such as repeat medication discontinued) would have identified too many patients.

We also asked the participating general practices to identify any patients they knew who had experienced avoidable harm, e.g. based on significant event analyses;¹² this did not identify any additional patients but some practices did not engage in providing this information to reviewers.

The next stage of sampling (Stage 3) identified patients with significant health problems (irrespective of whether these were avoidable or not). It involved one of the GP data collectors screening the electronic health record of each patient in the 'enhanced sample' to identify any new significant health problems experienced by patients over the 12 months of the study (1 April 2015 – 31 March 2016); this included all deaths. The research team provided the GP data collectors with detailed guidance on the significant health problems we wanted to screen for; this included all new major physical and psychiatric morbidities, and accidents (with examples including acute kidney injury, asthma requiring hospital admission, cancer, diabetes mellitus (including serious complications), deep vein thrombosis, heart failure, myocardial infarction, pulmonary embolism, and stroke). The GPs then undertook detailed retrospective reviews of the records of this final sample of patients to identify the extent to which errors of omission (e.g. failures of prevention) or commission in primary healthcare provision contributed to any of these significant health problems.

For the purposes of sensitivity analysis (recognising that cases might have been missed by our sampling approach), the GP data collectors also undertook a detailed records review for the following:

- 2.5% random sample of the Stage 1 population, but not including patients identified for the Stage 2 enhanced sample; each record was examined by a single GP reviewer.
- 10% random sample of the Stage 2 enhanced sample; each record was examined by a second GP reviewer.

Variables

For participating general practices we obtained data on the following variables: list size (number of patients); age distribution (particularly highlighting the number and percentage of patients aged 65 years and older); number and percentage of males and females; ethnicity (number and percentage of non-White patients); Index of Multiple Deprivation (IMD), the official measure of deprivation in England; whether practices were rural or urban; Care

Quality Commission (CQC) overall rating for the practices, and CQC safety rating for the practices. The CQC is an independent regulator health and adult social care service providers in England and responsible for checking through inspection and ongoing monitoring that care quality and safety standards are being met.¹³ In addition, for each practice we calculated the number of patient-years of data available for the period 1 April 2015 to 31 March 2016 using registration data.

Identification of avoidable significant harm, and factors associated with this For those patients with significant health problems, the GP data collectors reviewed the patient records and recorded whether they considered that the patient had received an adequate standard of care for these problems, or whether there was any evidence of avoidable harm. For the latter cases, the GPs provided a detailed written account of the principal problem in the patient's primary care that led to the significant health problem, a narrative describing the manner in which the significant health problem could have potentially been prevented within primary care, and a judgement on the avoidability of the significant health problem using a validated six-point scale (see Box 2 of the main manuscript).^{14,15} The GP data collectors searched back in patients' records as far as was needed to establish whether the significant health problem was avoidable or not. The evidence recorded by the GP data collectors was typically descriptions of salient signs or symptoms, pertinent past or concurrent medical or psychosocial history detail, and/or the actions or plans recorded by GPs in entries for each clinical encounter. Such descriptions were essentially 'signals' in the case note entries identified by the reviewers informing judgements about avoidability.

All cases were considered in detail by the study team, and the GP data collectors were asked to provide additional information if any clarification were needed. Each case was discussed by the study team and we considered what additional evidence (or signals) we would be seeking in the case notes in order to justify the avoidability score awarded, or to upgrade or downgrade the score. During those discussions, a member of the study team had online access to published guidelines to ensure our study team judgements were compliant with best practice guidelines. If relevant guidelines had been published since the observed study period, we considered the evidence available at that time. Where there was an absence of published guidelines, we considered trial data or systematic reviews (particularly Cochrane reviews). If necessary, we asked the GP data collector to return to the relevant general practice to examine the clinical records again to confirm the presence or absence of the evidence the study team deemed relevant to inform final judgements about avoidability. GPs only recorded what was explicitly stated in the records, or described what

was evidently absent in relation to what would be expected based on relevant guidelines for the condition. To ensure consistency the study team made the final judgement, through consensus, in terms of the classification of avoidable significant harm.

Data collection and coding

Each of the participating general practices was visited by an informatician from the study team who collected baseline data on the practice population and ran a computer search to identify patients for the enhanced sample and for the sensitivity analyses. Using encrypted tablet computers and a Virtual Private Network (VPN) connection, the GP data collectors entered anonymised data directly into a database on a secure server at Cardiff University. The nature of the avoidable harm was recorded by the GP data collectors using the comprehensive patient safety classification system developed in the Primary Care Patient Safety Classification (PISA) study. ¹⁶ The classification system has been empirically derived and aligned to the WHO International Classification for Patient Safety using a constant comparative approach. ¹⁷ The system has been used for analysis of over 72,000 patient safety incident reports from NHS organisations in England and Wales for 26 major studies of patient safety predominantly in primary care. ¹⁸⁻³³

Case narratives were deconstructed using codes from the classification system to describe: incident types (primary and contributory); potential contributory factors which are circumstances, actions of influences that played a part in the origin or development of the incident; incident outcomes; and harm severity. Primary incidents included those proximal (chronologically) to the patient outcome, whereas contributory incidents included those that contributed to the occurrence of another incident. Multiple codes for incident type (e.g. administration, medication), contributory factor (e.g. patient co-morbidity, staff workload), and incident outcome were applied to each case where necessary. The codes were applied systematically and chronologically ^{16,34}.

Analysis

We estimated the incidence of significant harm that was considered at least probably avoidable (our primary outcome – avoidability score 4 or more) and at least possibly avoidable (avoidability score 3 or more) and expressed these as 'per 100,000 patient-years' accompanied by 95% confidence intervals (95% CI). We assessed inter-rater reliability of judgements made by paired GP data collectors using the Cohen's Kappa statistic (with 95% CI).

Members of the study team then undertook a detailed analysis of the information provided on each case of potentially avoidable significant harm and included cases with at least 'slight to modest' (score 2 or more) evidence of avoidability, as we judged that even in these cases there were important insights. This included in-depth case analysis meetings, also involving team members with patient and public involvement background (ACh and AD). We reviewed and discussed the cases with the purpose of identifying commonalities and differences between them. We analysed the data recorded on the cases and examined the relationships between different types of incident and the factors that contributed to these incidents. As a result, we identified the most important factors contributing to avoidable significant harm.

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