Patients’ online access to their electronic health records and linked online services: a systematic review in primary care

Abstract

Background
Online access to medical records by patients can potentially enhance provision of patient-centred care and improve satisfaction. However, online access and services may also prove to be an additional burden for the healthcare provider.

Aim
To assess the impact of providing patients with access to their general practice electronic health records (EHR) and other EHR-linked online services on the provision, quality, and safety of health care.

Design and setting
A systematic review was conducted that focused on all studies about online record access and transactional services in primary care.

Method
Data sources included MEDLINE, Embase, CINAHL, Cochrane Library, EPOC, DARE, King’s Fund, Nuffield Health, PsycINFO, OpenGrey (1999–2012). The literature was independently screened against detailed inclusion and exclusion criteria; independent dual data extraction was conducted, the risk of bias (RoB) assessed, and a narrative synthesis of the evidence conducted.

Results
A total of 176 studies were identified, 17 of which were randomised controlled trials, cohort, or cluster studies. Patients reported improved satisfaction with online access and services compared with standard provision, improved self-care, and better communication and engagement with clinicians. Safety improvements were patient-led through identifying medication errors and facilitating more use of preventive services. Provision of online record access and services resulted in a moderate increase of e-mail, no change on telephone contact, but there were variable effects on face-to-face contact. However, other tasks were necessary to sustain these services, which impacted on clinician time. There were no reports of harm or breaches in privacy.

Conclusion
While the RoB scores suggest many of the studies were of low quality, patients using online services reported increased convenience and satisfaction. These services positively impacted on patient safety, although there were variations of record access and use by specific ethnic and socioeconomic groups. Professional concerns about privacy were unrealised and those about workload were only partly so.

Keywords
electronic health records; online access; patient records; primary care; systematic review.

INTRODUCTION
Online access to medical records by patients has the potential to enhance provision of patient-centred care and may improve patient satisfaction. However, it may also prove to be an additional burden for the healthcare provider.1,2 Proposals to link personal records from multiple health providers into a single online record have also not been widely implemented.3,4 Online services may also offer patients greater convenience and satisfaction, although concerns remain about privacy and confidentiality.

While significant progress has been made in other jurisdictions, most notably in some sectors of the US health system,5,6 there has been more limited progress in the UK, at least in part reflecting professional concerns about security, privacy,7–11 legal constraints,12 and the potential pitfalls of allowing access to children’s records.13 Pilot studies demonstrate the feasibility of providing online access.14,15 ‘Online access’ is generally subordinated into access to the medical record itself; provision of e-mail or an equivalent messaging service through a portal; and ‘transactional services’ such as requesting a repeat prescription or booking an appointment. In the US, Kaiser Permanente has had success in securing two-thirds of its 3.4 million members to sign up for online appointment booking, viewing test results, and e-mail.16 Similarly, the Veterans Health Administration has also registered large numbers online, with over 600 000 users making over 20 million ‘visits’ over the internet by 2008; with the most popular service being online repeat prescription requests;17 but uptake on this scale has yet to be replicated in the UK.

The policy context for this review is one where the English NHS (Department of Health/NHS England) intends patients to have free access to their health record by 2015.18 Guidance developed by early innovators of online access in primary care was not widely adopted,19 and has recently been superseded.20 The aim of this review was to assess the
How this fits in

Online services have been successfully piloted by large American organisations (such as Kaiser Permanente) but this success has not been replicated in the UK. This study found that patient satisfaction improved through enabling better self-care. Clinicians had concerns about the additional burden and workload from online access but evidence found their fears were only partly realised. There is a lack of good quality research in these areas in the UK; especially research that measures impact (such as health outcomes).

Impact of providing patients with access to their general practice electronic health records (EHR) and other EHR-linked online services on the provision, quality, and safety of health care.

METHOD

This systematic review was conducted following Cochrane guidelines, and complies with the Preferred Reporting Items for Systematic reviews and Meta-Analyses (PRISMA) framework. The protocol for this review has already been published, including details of the key research questions, and inclusion and exclusion criteria. The study aims were structured in a systematic way, using the elements of a clinical question, including population, intervention, comparator, and outcome (Box 1).

Search strategy

Searches were developed and run across 10 bibliographic databases (1999 to September 2012): MEDLINE®, Embase, Cumulative Index to Nursing and Allied Health Literature (CINAHL), Cochrane Database, Cochrane Effective Practice and Organisation of Care Group (EPOC), Database of Abstracts of Reviews of Effects (DARE), King’s Fund, Nuffield Health, and PsycINFO. A search for unpublished material was conducted using the database OpenGrey. An online suggestion box was also created on the project team’s website for other articles to be considered for review. Search strings were tailored to each database according to each source using Medical Subject Heading (MeSH), index terms, and key words. The total number of papers identified was 9877, and of these, 3971 were duplicate articles, therefore after this initial filter 6191 papers remained. Examples of the search string are available from the authors on request.

Box 1. PICO criteria

Population
Defined as users or non-users of online record access and remote contact services, including both patients (including carers and children) and clinicians/support staff in primary care

Intervention
Any intervention related to online access and services used in primary care. The types of intervention will be classified by whether they are principally designed to impact on patients or clinicians, or are technical in nature. Also whether online access is primarily synchronous or asynchronous

Comparator
Any comparison including usual care

Outcomes
Attributes of the user, use of access and services on quality and safety, and impact of use on patient and professional groups. This includes professional–patient relationships, access, satisfaction, and workload/workflow, and any possible impact on health outcomes. No limits were placed on how these outcomes were measured within specific intervention timescales

RESULTS

Study selection, screening, data extraction, and risk of bias assessment

Papers were screened by three authors using a detailed inclusion and exclusion guide and selection flowchart to identify relevant papers. Results from these searches, and all decisions regarding inclusion and exclusion were stored using Endnote (v.7.2.1.4) [Table 1]. Disagreement was resolved by reviewing and discussing full text versions. The included papers were then sourced and full text papers were divided across the review team for review. The review team were volunteers who had expressed interest in joining Working Group 7 (and evaluation of the evidence) of a larger Royal College of General Practitioners’ exercise to define a Road Map for providing patients online access to their medical records. A purposeful sample of academics, practitioners, and patient representatives with relevant expertise were recruited. Each reviewer was provided with a data extraction form to ensure consistency. Two exercises were conducted to test and refine the electronic and web-based data extraction forms. All reviews were independently checked by a third reviewer.

The data extraction form also included a risk of bias (RoB) form (available from the authors on request) for each paper, to capture study design limitations and offer insight into future study design. The RoB assessment was grouped into six domains: sequence generation, allocation concealment, blinding, incomplete outcome data, selective reporting, and other bias. It was intended to eventually use the Grading of Recommendations Assessment, Development and Evaluation (GRADE) instrument to assess the strength of evidence across particular study outcomes.

Analysis considerations

Most formal testing of hypotheses within studies used t-tests and χ² tests. Because of the variation in study designs, no single analysis method was undertaken across all the studies. Where necessary, all relevant statistical information is provided for each paper. A meta-analysis could not be undertaken because trials were not sufficiently homogeneous in terms of primary outcome measures to provide a meaningful summary. Further analysis, such as sensitivity or subgroup analysis, could not be undertaken.

Included studies and risk of bias

From the 176 papers sourced, only 17
papers were experimental in design, or cohort studies. Nine studies were randomised controlled trials\(^{29-36}\) (including one randomised control pilot study with qualitative element),\(^{37}\) and four were cohort studies\(^{38-41}\) (including one retrospective cohort study,\(^{41}\) one cross-sectional cohort study,\(^{41}\) one retrospective cohort and matched controlled study).\(^{41}\) Three were cluster randomised control trials.\(^{42-44}\) One study was defined as using a quasi-experimental non-randomised design.\(^{45}\) Most of these studies originated from the US (\(n = 13\)), with three studies being undertaken in Norway, and only one from the UK. Further study characteristics are summarised in Table 1. Included studies were based on the Oxford Centre for Evidence-based Medicine Levels of Evidence,\(^{46}\) and represented the high-quality evidence from the wider systematic review. Supplementary key findings are

### Table 1. Study characteristics

<table>
<thead>
<tr>
<th>Study type</th>
<th>Country</th>
<th>(n)</th>
<th>Reference</th>
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</thead>
<tbody>
<tr>
<td>Randomised control trials (including one randomised control pilot study)</td>
<td>US</td>
<td>5</td>
<td>29, 31, 35, 36, 37</td>
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<tr>
<td></td>
<td>Norway</td>
<td>3</td>
<td>30, 33, 34</td>
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<tr>
<td></td>
<td>UK</td>
<td>1</td>
<td>32</td>
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<tr>
<td>Cohort studies (including one retrospective cohort study, one cross-sectional cohort study, one retrospective cohort, and matched-control study)</td>
<td>US</td>
<td>4</td>
<td>38, 39, 40, 41</td>
</tr>
<tr>
<td>Cluster randomised control trials</td>
<td>US</td>
<td>3</td>
<td>42, 43, 44</td>
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<tr>
<td>Quasi-experimental control trial</td>
<td>US</td>
<td>1</td>
<td>45</td>
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<table>
<thead>
<tr>
<th>Finding</th>
<th>Study type</th>
<th>Country</th>
<th>Reference</th>
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<tbody>
<tr>
<td>Users of these services were more likely to be female (OR 1.15), middle aged or older, with much greater morbidity than non-users (OR 5.6)</td>
<td>Cohort with match controls, cross-sectional cohort study</td>
<td>US</td>
<td>39, 40</td>
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<tr>
<td>People with Medicaid cover were less likely to use these services (OR 0.8) than people with commercial insurance</td>
<td>Cross-sectional cohort study</td>
<td>US</td>
<td>40</td>
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<tr>
<td>African-American, other non-white ethnicities, and females with lower socioeconomic classification were less likely to use the services</td>
<td>Randomised control trial, randomised control pilot study</td>
<td>US</td>
<td>29, 37</td>
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<tr>
<td>70% of clinicians felt that online access strengthened relationships, enhanced trust, and improved decision making</td>
<td>Quasi-experimental non-randomised design</td>
<td>US</td>
<td>45</td>
</tr>
<tr>
<td>Patients who used online access to test results were more satisfied than those who did not</td>
<td>Cluster randomised control trial</td>
<td>US</td>
<td>44</td>
</tr>
<tr>
<td>Use of a secure messaging tool improved patient satisfaction with care from 48% to 59% ((P &lt; 0.04))</td>
<td>Randomised control trial</td>
<td>US</td>
<td>35</td>
</tr>
<tr>
<td>Patients felt themselves better able to express their concerns in writing (34%), to communicate about difficult topics (38%), and to send psychosocial information when using online tools versus telephone call</td>
<td>Randomised control trials (and interviews)</td>
<td>US, US</td>
<td>34, 35</td>
</tr>
<tr>
<td>Online systems saved patients time, telephone calls, and clinic visits</td>
<td>Randomised control trials (and interviews)</td>
<td>Norway, US</td>
<td>34, 35</td>
</tr>
<tr>
<td>About half of US patients were willing to pay for online access, and 21% were unwilling. Of those who were willing, the median value assigned was $2 per message</td>
<td>Randomised control trials, randomised control pilot study</td>
<td>Norway, US, US</td>
<td>33, 35, 37</td>
</tr>
<tr>
<td>Patients given online access to their records identified more than twice as many medication list discrepancies with potential for severe harm than those who did not</td>
<td>Cluster-randomised trial</td>
<td>US</td>
<td>43</td>
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<tr>
<td>Users of online services received 84% of preventive care services compared with 68% of controls, including influenza vaccine (22% versus 14%) and mammography (49% versus 30%); children whose parents used online services received 96% of immunisations versus 67% in controls. Online services also facilitated health maintenance/appointment reminders</td>
<td>Cluster randomised control trial</td>
<td>US</td>
<td>42</td>
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<td></td>
<td>Randomised control trials</td>
<td>UK, US</td>
<td>32, 36</td>
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<tr>
<td>Only 1-8% of patients say that viewing their record online caused confusion, worry, or offence</td>
<td>Quasi-experimental non-randomised design</td>
<td>US</td>
<td>45</td>
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<tr>
<td>The impact of online record access on patient use of face-to-face services and telephone calls was unclear</td>
<td>Randomised control trial (and interviews), cohort with match controls, quasi-experimental non-randomised design</td>
<td>Norway, US, US</td>
<td>34, 39, 45</td>
</tr>
<tr>
<td></td>
<td>Randomised control trial, randomised control trial (and interviews), cohort and matched-control study, cluster randomised control trial</td>
<td>Norway, Norway, US, US</td>
<td>30, 34, 41, 42</td>
</tr>
<tr>
<td>Clinicians had concerns about the additional burden and workload from online access but found their fears only partly realised</td>
<td>Randomised control trial (and patient survey), three-part cohort study, quasi-experimental non-randomised design</td>
<td>US, US, US</td>
<td>31, 38, 45</td>
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\(\text{OR} = \text{odds ratio.}\)
Records identified through 10 database searches (n = 137844 (not including DH database))

Duplicates removed (n = 3971)

Records (n = 9877) (Endnote)/ (n = 235 (other sources))

Stage 1. Records excluded at title screen (n = 3921)

Stage 2. Records excluded at abstract screen (n = 5959)

Full-text articles assessed for eligibility (n = 232)

Stage 3. Full-text articles excluded, with reasons (n = 44) and unobtainable (n = 10)

Stage 4. Reviewers request to remove items with reasons (n = 33)

Stage 5. Exclude (n = 159) observational/qualitative

Studies included in wider review (n = 174)

Final number of trial studies included in review (n = 17)

Patient Satisfaction & Value
44. Matheney et al 2007, US
Willingness to Pay 33. Bergmo & Wangberg 2007, Norway
Patient Safety, Prevention and Privacy Concerns
42. Nagykaldi et al 2012, US
43. Schnipper et al 2012, US
Prevention 32. Fairhurst & Sheikh, 2008, UK
36. Wright et al 2011, US
Workload Concerns and Change
30. Bergmo et al 2005, Norway
34. Kummerfeldt et al 2004, Norway
40. Ralston et al 2009, US

Given in Table 2. Details of the searches are presented in the Preferred reporting items for systematic reviews and meta-analyses (PRISMA) flowchart (Figure 1).

Outcomes of these studies varied, with most focusing specifically on online record access, or online service utilisation, including impact on workload and attendance rates. Several focused on patient or provider satisfaction, two on patient safety, medication accuracy, and preventive reminders, and one study on willingness to pay for electronic communication with their GP. These studies were subject to RoB assessment. These studies illustrated a wide variation in their RoB and most showed a high risk of bias in more than one assessment area. Figures 2 and 3 show risk of bias tables and summary graphs.

User characteristics
Online record access and service users tended to be slightly older (t-test, P < 0.001), or middle aged (50–65 years), and were more likely to be female (χ²; P = 0.002), female (OR 1.15; 95% confidence interval [CI] = 1.10 to 1.19). Users also tended to have much greater overall morbidity (OR 5.64, 95% CI = 5.07 to 6.28). Differences were also observed between insurance types, with users of messaging being less likely to be funded by Medicaid insurance than commercial insurance (OR 0.81, 95% CI = 0.68 to 0.96). African-American and other non-white ethnicities (42.9% and 50.3%, respectively, P < 0.01) were less likely than white and Asian patients to use online access (73.6% and 93.4%, respectively), and so were females of lower socioeconomic status. Further information on each of the included studies is available from the authors on request.

Patient satisfaction and value
Self-care: Patient satisfaction improved through enabling better self-care. The introduction of a system which enabled patients to view their test results, manage their medication list, and have secure messaging with their practice, resulted in 80% of responders (n = 448/560) saying the system facilitated their participation in their own care. Another post-intervention survey found that 77–87% of 5391 patients across three sites who had online access to visit notes felt more in control of their care. Of the clinician responses, 73 out of 104 (70%) felt that enabling patients to read their own notes online strengthened relationships, enhanced trust, and improved decision making.

Online communication. Several trials showed that online access to records and services improved communication. Patients were more satisfied with automated communication of test results (OR 2.35; 95% CI = 1.05 to 5.25; P = 0.03) and with online information about their treatment or condition (OR 3.45; 95% CI = 1.30 to 9.17; P = 0.02), compared with those who accessed this information in person or by telephone. One trial measuring the frequency of portal use and satisfaction with secure messaging found that patients’ communication with their clinic improved; with 44% (n = 77/174) in the intervention arm compared with 12% (n = 18/146; χ² = 38.8, P < 0.001) in the control group. Use of the messaging portal also resulted in higher satisfaction with overall care 59% (n = 103/174) in the intervention group compared with 48% (n = 78/162; χ² = 4.1, P = 0.04).
The contents, tone, and timeliness of online messaging to clinicians also improved communication and satisfaction. Some patients (34%, n = 68/200) felt better able to express their concerns in writing; 36% (n = 72/200) felt it easier to communicate about difficult topics, and felt that they would be more likely to send psychosocial information via online systems than by telephone. Online services, such as online communication, also saved patients time; however, time in these trials was not quantified, and was found not to necessarily help reduce the number of telephone calls and clinic visits. In addition, patients’ satisfaction with their primary care providers’ general communication skills may not necessarily improve with the adoption of online access and services. Research into the communication experience for patients was largely at low risk of bias in several areas, including allocation concealment, addressing all outcome data, and being free from reporting bias (Figure 2).

How patients valued online access and services. Many patients would not be willing to pay for, or only placed a low value on, online services, such as online communication with clinicians. In one US study, only 51% of study participants (n = 77/151) expressed a willingness to pay for electronic contact with their clinician, and 21% (n = 31/151) expressed unwillingness. Those who had communicated with their clinician electronically for at least a year had a lower willingness to pay than those who did not have access (P = 0.0028). Two further studies reported similar findings, with less than half of participants (42%, n = 111/266) reporting an unwillingness to pay a fee to have e-mail or online messaging with their clinician. The value assigned to this service was low: a median of $2 (mean $4.10) per message, and another stating either €0, €6, or €12. Future research into willingness to pay, needs to be better designed, as research presented here is classified as either unclear or at high risk of bias (Figure 2).

Patient safety, prevention, and privacy concerns

Patient safety was largely improved through patients identifying errors in their medication list. Patients given access to their medication list online corrected more than twice as many medication discrepancies with potential for severe harm (0.03 versus 0.08 per patient, adjusted risk ratio 0.31, 95% CI = 0.10 to 0.92; P = 0.04). Online access to consultation notes was also found to increase reported medication

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**Figure 2.** Patient issues risk of bias table and summary graph.
adherence as 60–78% (n = 5391) of patients taking medications reported increased adherence.45 Online services facilitated uptake of preventive care services by providing prevention or health maintenance reminders.36,42 Nearly all (84.4%) patients in the intervention group received all recommended preventive services, compared with 68% of controls (n = 384, P<0.0001). Children in the intervention group received 95.5% of immunisations compared with 87.2% in the control arm (n = 105 parents, P = 0.044). Patients provided with online prompts were significantly more likely to receive influenza vaccines (22.0%, n = 50/227 versus 14.0% control; P = 0.018); and undergo mammography than control groups (48.6%, n = 51/105 versus 29.5%; P = 0.006). No significant improvement was noted in uptake rates of other screening tests.36

Only 1–8% (n = 5391) of patients stated that viewing consultation notes caused confusion, worry, or felt offended with note comments.45 Similarly, clinicians acknowledged their fears about potentially offending or worrying patients; however, these concerns did not materialise in practice.45 The studies about medication safety (inevitably) lacked allocation concealment and these papers reported blinding was not applicable to their study methods.42,43,45 The prevention outcomes studies had low risk of bias (Figure 3).32,36

Workload

Workload concerns. Clinicians had concerns about the additional burden and workload from online access but found their fears only partly realised.38,45 Clinicians involved in an initiative promoting e-mail thought e-mail was useful for answering patients’ non-urgent medical questions (81.8%, n = 43 intervention versus 61.4%, n = 46 control; P = 0.06) and helpful in managing patients’ administrative concerns (86.0%, n = 43 intervention versus 58.0%, control n = 46; P = 0.05).29 However, a few intervention clinicians felt that e-mails were too lengthy (14.6%, n = 43; P = 0.04) or were concerned about incomplete information (10%, n = 43; P = 0.11). In one cohort study about the implementation of a video to support the use of a patient’s portal, during office visits 12 of 13 (92%) support staff agreed that it was easy to use, and six (46%) agreed that the technology did not take up more time.38

Effect of online access on face-to-face consultations. Online access has an inconsistent effect on face-to-face consultations, which increased in some studies,34,39,45 and went down in others.30,34,41,42 Findings from four randomised controlled trials indicated the reduction of face-to-face consultations, while findings from other study designs (cohort and quasi-experimental trials) found an increase. Clinicians (n = 99) concerned that access to

![Figure 3. Practice issues risk of bias table and summary graph.](image-url)
open notes would lengthen visits, reported a minor rise in consultation rates across three study sites (0-8%), and spent more time addressing patients’ questions outside of visits (0-8%). Clinicians also reported changes in how they recorded clinical information with between 3% and 36% of clinicians (n = 99) changing record content to allow for online access; and 0-21% reporting taking more time writing notes.45 In a post-intervention survey, a significant difference was found in time spent writing or dictating notes between practice size, with four out of 37 clinicians (10.8%) in smaller practices spending more time writing notes compared with seven out of 24 clinicians (29.2%) in larger practices (P = 0.019).46 Finally, 59-62% of patients (n = 5561) thought they should be able to add comments to their consultation notes, such as approving the notes’ contents.

Four studies found a reduction of office visits by patients using a messaging system.30,34,41,42 In one randomised controlled trial the consulting rate fell from 4.5 to 2.8 per annum in the intervention group compared with 4.3 to 3.6 per annum in the controls (P = 0.034).30 In another, patient portal users had fewer visits over the study period, compared with the control group (average of 2.9 versus 4.3 visits; P<0.0001).42 Similarly, annual visit rates decreased by 9.7% (0.23 visits per member) in a cohort study,41 and in a matched-control study the decrease was 10.3%, or 0.25 visits per member per year (P<0.001). The control group reduction for the matched-control study was 3.7%, or 0.08 visits (P<0.003).41 Across-group differences between the cohort and matched-control group showed a significant difference in annual office visit rates of 6.7% [0.17; P<0.003].41

Face-to-face contact also increased in a before and after study of online test result viewing and secure messaging with clinicians. There were 87 206 users of the system and 71 663 non-users. Office visits rose by 0.7 per member per year (95% CI = 0.6 to 0.7; P<0.001) in the patient group enrolled in the online system. There was also a significant increase in rates of after-hours clinic visits (18.7 per 1000 patients per year 95% CI = 12.8 to 24.3; P<0.001), emergency department encounters (11.2 per 1000 patients per year 95% CI = 7.4 to 19.7; P = 0.01), and hospitalisations (19.9 per 1000 patients per year 95% CI = 14.6 to 25.3; P<0.001) for patients who used the online system compared with non-users.39

Impact of e-mail use on patient contacts. E-mail and web-messaging sometimes increased the volume of patient contact,29,31,41 and in other studies it remained the same, but time had to be put aside to respond to e-mails.30,34,45 The use of a triage-based communication tool led to increased volume of e-mail messages in the intervention group; 46 weekly e-mails per 100 scheduled visits compared with only nine in the control group at the study midpoint (P<0.01).29 However, this increase was not sustained and the e-mail volume diminished after the initial promotion period, leading the authors to question whether e-mail use may not necessarily improve the efficiency of clinical care.29 A modest difference was also seen in the attitudes of intervention patients towards using web-based tools to address administrative and prescription issues, with 61.9% (n = 65) of intervention patients, compared with 54.2% of control patients (n = 67), agreeing that they liked using the web to communicate with clinicians (P = 0.15).31

Two trials and one quasi-experimental study indicated no change in the volume of e-mail or messaging; although in these studies e-mail was clearly an established part of the doctor–patient relationship. One trial reported clinicians received one message per day for every 250 patients who were health portal users. However, only 27% were sent during clinic hours; 73% of messages were sent during non-work hours, with 52% sent between 5 pm and midnight.30 The volume of electronic messages did not change, in a study providing online access to ‘open notes’ records and was achieved with less impact on clinicians’ working lives than originally anticipated.31 However, 21% of clinicians (n = 39) acknowledged the need to use extra time for writing, editing, or explaining notes to their patients.35

Impact of online access on telephone calls to clinicians. Studies focusing on the impact of online services on telephone use offered mixed messages. Two studies reported a rise in telephone use,39,41 as a consequence of online access; four studies no change,29,31,35 and one study reported a decline.34

Results of a before and after study assessing online access to health records found a significant increase in the per-member rates of telephone encounters, with a small rise of 0.3 per member per year (95% CI = 0.2 to 0.3; P<0.001) in the patient group enrolled in the online system.39 Similar outcomes were found in a matched-control study, when telephone contact rates increased by 16.2% (0.32...
contacts per member) in the secure messaging arm compared with 29.9% (0.52 contacts per member) for the matched-control group. Across-group difference was 13.7% (0.20 contacts) and was significant \((P < 0.01)\). Four studies indicated no change in telephone consultation volume. Two studies indicated no difference in telephone consultations between the intervention and control groups \((P \geq 0.258)\); another reporting how e-mail and web-based tools did not offset either telephone volume; or no change in no-show visit rates.

Only one paper suggested that telephone call volume can decline, with the intervention group seeing a reduction in the annual number of visits and telephone calls by 28% \((\text{intervention } n = 74; \text{OR } 3.19; 95\% \text{ CI } 2.44 \text{ to } 3.94 \text{ versus control } n = 92; \text{OR } 4.45; 95\% \text{ CI } 3.60 \text{ to } 5.29; \text{P } < 0.032)\), and a total reduction in the number of calls to the GP of 10%.

Only two areas were consistently at low RoB across workload outcomes, these being incomplete data outcomes being addressed, and being free from selective reporting in these studies. Several papers reported blinding was not applicable to their study methods, three reporting high RoB, and one had unclear blinding methods.

DISCUSSION

Summary

The findings from this review are important for health systems and professionals. Evidence shows that although online access may be achievable regarding technological implementation and improvements to patient safety, there remain challenges about clinicians’ adoption of systems because of workload and workflow concerns.

Users of online access and services report increased satisfaction in terms of better self-care and communication with clinicians. Online access and services also positively impacted on patient safety, especially when patients are given access to medication lists and are offered prevention or health maintenance reminders. However, there is a disparity in who accesses online records and services, with people of non-white ethnicities and those from lower socioeconomic groups potentially being disadvantaged. There is also a difference between expectations and realities of online services for both patients and clinicians. Patients anticipated empowerment and rapid responses to electronic communication and clinicians feared being overwhelmed by a new workload. However, this fear was largely unwarranted. Many administrative and non-urgent tasks appear to be more readily conducted online. They can also be conducted at times of the day convenient for patients. Although fears have been expressed about access to records being abused or causing patients greater worry, no evidence has been published to indicate what harm or privacy breech has occurred. The inter-relationship between provision of e-mail or messaging, online access, telephone, and face-to-face communication has yet to be systematically and robustly researched.

Strengths and limitations

A strength of this study was the RoB assessment that gave an indication of the likely strength of evidence and how future study design can be improved. The studies reported here largely originate from the US from large health plan-based programmes that incur a fee. There have been few randomised controlled trials examining the effectiveness of online access and services on improving health outcomes, but some work has been undertaken focusing on the impact of new technologies in relation to patient decision making and health outcomes. Like all systematic reviews, evidence was gathered from various resources from a specific time period and, as such, there may be several new studies recently published that have not been included in this review.

Comparison with existing literature

Compared with other sectors, such as for-profit companies, the adoption and integration of online technologies in UK primary care has been slow, but it is steadily growing with the introduction of new online services such as weight-loss, or counselling programmes; comprehensive health information systems; and e-mail and messaging with provider. Technology has changed the nature of the patient–clinician interaction from one that was traditionally clinician-led to a more patient-centred approach. It has possibly improved trust, patient involvement in the clinical decision-making process, and in the quality of communication experience. Disparities in access to computers and information systems have been termed the ‘digital divide’. However, health literacy is also an important component of being able to use online resources, and technology may enable new forms of clinician–patient interaction.
Previous guidance developed by innovators of online access in primary care was not widely adopted. Possible explanations for limited implementation of systems to date could be due to the sociolegal context of records; security and privacy issues; language and interpretation issues; child protection and family safety issues; and patient-added data and organisational issues [such as workload, risks of litigation, training, and system processes and maintenance].

The potential benefits of online access have yet to be fully realised in research originating from the UK; however, the lessons learned from other studies may help with future implementation.

Implications for research and practice
Online access to records may subtly change the patient–physician relationship. Patients are increasingly signing up for online services that are more convenient. Patients may no longer see the medical record as being under the sole provenance of their physician; rather as something they can view and print. This appears to be empowering. The downside is that ready access may make it easier to always consult rather than self-manage, and moves to online access provision may result in variations in service utilisation by specific population groups. Clinicians find that they have to write more carefully considered records that are more informative for their patients to read, rather than being an aide memoire for the doctor and the clinical team. Medical services may have to adapt to meet this new requirement and managers will have to learn what models of delivery increase overall workload, and what approaches might control it.

Willingness to pay studies show online access to be valued little by US research participants. In the UK’s NHS willingness to pay might be less relevant. However, little work has been undertaken in the UK on accurately assessing the value patients might place on specific online services. Given this, it may be possible to undertake further research as to whether there is a place for fee paying compared with the UK norm of publicly funded health care.

Encouraging patients to access online EHR and services are a promising way to enhance safety. Further research is also needed into how patients use their access to EHR, and how patient-added data are used, and how they may improve care by providing insight into health beliefs and their understanding of their condition.

While there have been a number of trials conducted in the US, such evidence from the UK is scarce. Further research into the feasibility of implementing complete online access (record access and transactional services) across the UK, or in a whole-system demonstrator, would provide valuable insight into how to implement such a service.

Research into online access and services has yet to demonstrate how health outcomes can be improved. Research is also needed for patients with specific long-term conditions, such as diabetes, which may offer greater insights regarding facilitating better health outcomes.

Other areas of further study include prevention and privacy issues involved in providing online access and services; how practices can facilitate patients’ use of technologies that integrate into their existing infrastructure; and to what extent online services can replace or complement existing care. A final area is whether such systems are efficient and cost-effective. The second article from this systematic review focuses on the breadth of knowledge in this area and draws together how online record access and services may impact on the wider business process in primary care.

Initiatives to develop policy and practice are continuing through the NHS England Patient Online programme and the Royal College of General Practitioners. The findings from this review are timely and directly relate to these initiatives, as they highlight patient perspectives, such as satisfaction, safety, prevention, and privacy concerns; and clinicians’ perspectives, such as workload concerns.

Patients using online access to their EHR and services reported increased convenience and satisfaction, and these services positively impacted on patient safety. Although patients report specific benefits of record access, such as enabling better self-care and patient–clinician communication, patients were unwilling to pay for these services. Prior to implementation of online systems clinicians were concerned about any possible workload rise. However, post-intervention work suggests clinicians’ concerns are largely unrealised. The nature of the medical record and the role of the clinician may need to evolve to give greater value to patients and to ensure greater equity in uptake. The business model for primary care may also need to change to enable more effective utilisation of information technology in everyday practice.

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