REVIEW

Electronic case report forms and electronic data capture within clinical trials and pharmacoepidemiology

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AIMS

Researchers in clinical and pharmacoepidemiology fields have adopted information technology (IT) and electronic data capture, but these remain underused despite the benefits. This review discusses electronic case report forms and electronic data capture, specifically within pharmacoepidemiology and clinical research.

METHODS

The review used PubMed and the Institute of Electrical and Electronic Engineers library. Search terms used were agreed by the authors and documented. PubMed is medical and health based, whereas Institute of Electrical and Electronic Engineers is technology based. The review focuses on electronic case report forms and electronic data capture, but briefly considers other relevant topics; consent, ethics and security.

RESULTS

There were 1126 papers found using the search terms. Manual filtering and reviewing of abstracts further condensed this number to 136 relevant manuscripts. The papers were further categorized: 17 contained study data; 40 observational data; 27 anecdotal data; 47 covering methodology or design of systems; one case study; one literature review; two feasibility studies; and one cost analysis.

CONCLUSION

Electronic case report forms, electronic data capture and IT in general are viewed with enthusiasm and are seen as a cost-effective means of improving research efficiency, educating participants and improving trial recruitment, provided concerns about how data will be protected from misuse can be addressed. Clear operational guidelines and best practises are key for healthcare providers, and researchers adopting IT, and further work is needed on improving integration of new technologies with current systems. A robust method of evaluation for technical innovation is required.

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WHAT IS ALREADY KNOWN ABOUT THIS SUBJECT
- Information technology (IT) has tangible benefits to assisting high quality research.
- Investment in IT is underfunded in healthcare and research.

WHAT THIS STUDY ADDS
- A governance support framework is necessary to assist healthcare providers and researchers to maximize the benefits of IT.
- Further work is required in improving interoperability between IT systems for research and pharmacoepidemiology.
- An unambiguous legislative framework is needed to ensure high quality research can continue successfully whilst continuing to adhere to good clinical practice, data protection and ethics.
- Generic and adaptable solutions are required to meet the software needs of researchers and healthcare providers.

Introduction

Information technology (IT) provides a fast and efficient way to collect scientific and clinical data and has become the most effective way to collaboratively share data. The benefits have underpinned the incremental introduction of electronic patient records in healthcare organizations which has been suggested as the principal reason for the increasing allocation of healthcare industry funding to IT; from 2% of total revenue, in the 1990s, to 5–7% in recent years [1]. This in turn has contributed to investment in the use of IT and electronic case report forms (eCRFs) in clinical research. Whilst these systems are designed and used differently, they share a common goal of storing, and communicating in a safe and confidential way private clinical data in a structured format [2]. Pharmacoepidemiology and clinical research have undoubtedly benefitted from IT; however, developments in these areas have continued to lag behind the healthcare sector, with investment limited due to various concerns. Reasons cited for not further using IT in research include: technical issues in setting up infrastructure, financing and maintaining the newest technology, and ethical fears [3]. Additionally, different funding streams and personnel involved in development of electronic patient records used for healthcare purposes, and those used for data capture for research, make it difficult to integrate solutions that would satisfy both aims. The objectives of both types of system are often different, which can also lead to conflicts.

Different regulatory processes govern systems used in routine healthcare and research. However, clinical research relying on IT and electronic data capture (EDC) often depends on interfacing with healthcare IT systems, which generally comprise numerous dissimilar software systems and storage formats for storing patient data. Clinical research also often operates over large geographical areas, incorporating several different healthcare providers, further compounding challenges when interfacing with diverse local systems. Although there is a drive towards IT unification in the National Health Service primary care practises and hospital trusts in the UK are under no obligation to use collaborative IT systems or storage formats, nor are they required to make these data available for research purposes. While the need to exploit healthcare data for research to cost effectively drive healthcare improvements has never been greater, it is largely for these reasons that the task of collecting, storing and amalgamating health service data is likely to become increasingly difficult in the future.

Objective

The objective of this review is to assess the advantages and disadvantages of eCRF and EDC technologies in pharmacoepidemiology and clinical research, and to explore where further research should be best directed. For the purpose of this paper the term eCRF will refer to a system used to capture clinical data for research and EDC will refer to the generic process of data capture.

Methods

A literature review was conducted to identify articles pertaining to pharmacoepidemiology (drug epidemiology) and clinical research, and their use of eCRFs and EDC. Whilst the use of IT in routine healthcare is increasingly commonplace, the emphasis of this review was on the use of EDC and eCRFs in the conduct of clinical research. Common themes relating to these topics emerged covering a broad range of issues including technical and practical matters, consent, ethics, and security. PubMed and the Institute of Electrical and Electronic Engineers (IEEE) libraries were searched using to cast a wide net over the subject area; electronic case report form, eCRF, electronic data capture, and electronic data collection. Filters were applied to search terms to condense results to relevant articles (see Appendix). The search was conducted between 2014 and 2015 with a final analysis of the literature completed in August 2016. PubMed is a clinical library while IEEE is technology based.

All returned abstracts were read and articles deemed irrelevant to eCRFs and EDC, or articles that did not involve pharmacoepidemiology or clinical research, were excluded. Unlike clinical studies, IT has no universally accepted quality scoring system for academic papers. Therefore, it was decided that any published and peer reviewed article that was returned from the IEEE or PubMed search would be included. Exceptions to this were where there was an overt conflict of interest or the journal was not available in English. Figure 1 depicts a flow diagram of the review process. The authors endeavoured to adhere fully to the PRISMA checklist [4] in structuring this review; however, the nonstandard output of
Results

A total of 1126 papers were returned from all search topics. After review and consideration, 136 manuscripts were deemed relevant to the review. Each topic was further separated into manuscript types. There were 17 papers documenting a study or clinical trial that used EDC where the system was the primary focus of the manuscript; 40 papers discussed observational studies comparing or evaluating EDC; 27 papers contained anecdotal evidence or opinion regarding EDC; 47 papers detailed EDC models or designs. There was one literature review, one cost benefit analysis, two feasibility studies, and one case study comparing the use of EDC in five studies (Table 1). During this review, papers were further discarded that were found to be of poor overall quality or adding little to the topic. For a list of all included publications see Table 2.

Research has been conducted into ways to maximize data accuracy and efficiency using IT. Trials have taken data from patient’s electronic medical records (EMRs) and transferred these directly into eCRFs. The cost savings, quality improvements, and reduction of data entry errors, were significant [5–7]. Whilst not all required data ARE available from the patient EMR, studies have found varying results with as much as 69% of data required being found and used to prepopulate trial eCRFs [8]. Discussions around the design and theoretical modelling of EMRs, eCRFs and ECD were prevalent within the included papers [9–17]. The electronic systems reported vary in quality, with some being used in mock environments and others being purely theoretical. Commercial software packages are available, but are generally not cost effective and in some circumstances it is unclear who owns the data entered into them [18–20]. Observational studies have compared paper based systems against EDC or canvassed opinion on the use of EDC systems [21–24]. These papers were overwhelmingly in favour of EDC as long as security could be maintained.

Obtaining patient consent is an ethical necessity, and up until recently, has almost always required a physical signature. Varnhagen et al. [25] considered obtaining informed consent online and questioned whether it is ethical to obtain consent electronically. Recently, electronic consent has been accepted by the National Health Service as a viable alternative to a written signature [26]. This review found one trial where consent had successfully been captured online [27]. Collecting participant consent electronically is a novel, and largely unexplored, method that invites further innovation. There are ethical implications of conducting research entirely online. IT is advancing faster than ethical review panels can address and there is a need for greater ethical consideration of conducting research online and how we share data between IT systems and within organizations [28–35]. Government attempts to legislate – the Health Insurance and Portability Act [36] in the USA, and the Data Protection Act [37] in the UK, have had little impact on alleviating public scepticism [38]. Patient privacy is critical and despite the well-intentioned zeal for the mass adoption of IT within healthcare, serious security concerns remain [39, 40]. However, patients are open to technology being used to store their medical data if trust and privacy concerns can be addressed [41].

Table 1
Characteristics of journal papers

| Report characteristics                        | n  | %   |
|----------------------------------------------|----|-----|
| Report included one or more benefit/ disadvantage of EDC | 136|     |
| Main objective(s) of report:                 |    |     |
| Studies using EDC                            | 17 | 12.5|
| Observational studies evaluating EDC use     | 40 | 29.4|
| Opinion/discussion piece                     | 27 | 19.9|
| Description of model EDC system              | 47 | 34.6|
| Feasibility studies                          | 2  | 1.5 |
| Literature review                            | 1  | 0.7 |
| Cost–benefit analysis                        | 1  | 0.7 |
| Case study                                   | 1  | 0.7 |

Figure 1
Flow diagram of review
### Table 2
Publication review list

| Authors | Title | Journal | Year | Paper Type |
|---------|-------|---------|------|------------|
| Aiello EJ, Taplin S, Reid R, Hobbs M, Seger D, Kamel H, et al. | In a randomized controlled trial, patients preferred electronic data collection of breast cancer risk-factor information in a mammography setting [58] | J Clin Epidemiol | 2006 | Observational |
| Alexander I. | The impact of future trends in electronic data collection on musculoskeletal research and evidence-based orthopaedic care [71]. | Arthroscopy | 2003 | Anecdotal |
| Ariza AJ, Binns HJ, Christoffel KK, Paediatric Practice Research Group. | Evaluating computer capabilities in a primary care practice-based research network. | Ann Fam Med | 2004 | Observational |
| Ashar R, Lewis S, Blazes DL, Chretien JP. | Applying information and communications technologies to collect health data from remote settings: A systematic assessment of current technologies | J Biomed Inform | 2010 | Anecdotal |
| Ashley L, Jones H, Thomas J, Newsham A, Downing A, Morris E, et al. | Integrating patient reported outcomes with clinical cancer registry data: a feasibility study of the electronic Patient-Reported Outcomes From Cancer Survivors (ePOCS) system. | J Med Internet Res | 2013 | Observational |
| Atreja A, Achkar JP, Jain AK, Harris CM, Lashner BA. | Using technology to promote gastrointestinal outcomes research: a case for electronic health records. | J Gastroenterol | 2008 | Anecdotal |
| Ayatollahi H, Mirani N, Haghani H. | Electronic health records: what are the most important barriers? [3] | Perspect Health Inf Manag | 2014 | Observational |
| Azad T, Kalani M, Wolf T, Kearney A, Lee Y, Flannery L, et al. | Building an electronic health record integrated quality of life outcomes registry for spine surgery. | J Neurosurg Spine | 2016 | Observational |
| Bellamy N, Wilson C, Hendrikz J, Patel B, Dennison S. | Electronic data capture (EDC) using cellular technology: implications for clinical trials and practice, and preliminary experience with the m-Womac® Index in hip and knee OA patients. | Inflammopharmacology | 2009 | Model |
| Bellary S, Krishnankutty B, Latha MS. | Basics of case report form designing in clinical research [9]. | Perspect Clin Res | 2014 | Model |
| Bock M, Moore D, Hwang J, Shumay D, Lawson L, Hamolsky D, et al. | The impact of an electronic health questionnaire on symptom management and behaviour reporting for breast cancer survivors. | Breast Cancer Res Treat | 2012 | Observational |
| Borlawsky TB, Lele O, Jensen D, Hood NE, Wewers ME. | Enabling distributed electronic research data collection for a rural Appalachian tobacco cessation study. | J Am Med Inform Assoc | 2011 | Study |
| Brandt CA, Cohen DB, Shifman MA, Miller PL, Nadkarni PM, Frawley SJ. | Approaches and informatics tools to assist in the integration of similar clinical research questionnaires. | Methods Inf Med | 2014 | Model |
| Brewster W, Gibbs T, Lacroix K, Murray A, Tydeman M, Almenoff J. | Evolving paradigms in pharmacovigilance. | Curr Drug Saf | 2006 | Anecdotal |
| Bruland P, Forster C, Breil B, Ständer S, Dugas M, Fritz F. | Does single-source create an added value? Evaluating the impact of introducing x4T into the clinical routine on workflow modifications, data quality and cost-benefit. | Int J Med Inform | 2014 | Model |
| Burnstead B, Furlan G. | Unifying drug safety and clinical databases [72]. | Curr Drug Saf | 2013 | Anecdotal |
| Bushnell DM, Martin ML, Parasuraman B. | Electronic versus paper questionnaires: a further comparison in persons with asthma [23]. | J Asthma | 2003 | Observational |
| Bushnell DM, Reilly MC, Galani C, Martin ML, Ricci JF, Patrick DL, et al. | Validation of electronic data capture of the Irritable Bowel Syndrome – Quality of Life Measure, the Work Productivity and Activity Impairment Questionnaire for Irritable Bowel Syndrome and the EuroQol. | Value Health | 2003 | Observational |

(continues)
| Authors                          | Title                                                                 | Journal                  | Year | Paper Type |
|---------------------------------|----------------------------------------------------------------------|--------------------------|------|------------|
| Carvalho JC, Bottenberg P, Declerck D, van Nieuwenhuysen JP, Vanobbergen J, Nyssen M. | Validity of an information and communication technology system for data capture in epidemiological studies [66]. | Caries Res | 2011 | Observational |
| Cleland J, Caldow J, Ryan D.    | A qualitative study of the attitudes of patients and staff to the use of mobile phone technology for recording and gathering asthma data. | J Telemed Telecare | 2007 | Study |
| Collins M, Ross E, Meropol NJ, Lazev AB. | Using metadata to generate web-based Electronic Data Capture Forms. | AMIA Annu Symp Proc | 2006 | Model |
| Coons SJ, Gwaltney CJ, Hays RD, Lundy JJ, Sloan JA, Revicki DA, Lenderking WR, Cella D BEI ePRO TF. | Recommendations on evidence needed to support measurement equivalence between electronic and paper-based patient-reported outcome (PRO) measures: ISPOR ePRO Good Research Practises Task Force report. | Value Health | 2009 | Model |
| Courtney KL, Craven CK.         | Factors to weigh when considering electronic data collection.         | Can J Nurs Res | 2005 | Anecdotal |
| Crichton C, Davies J, Gibbons J, Harris S, Tsui A, Brenton J, et al. | Metadata-driven software for clinical trials [10]. | SEHC | 2009 | Model |
| Curcin V, Soljak M, Majeed A.   | Managing and exploiting routinely collected NHS data for research [6]. | Inform Prim Care | 2012 | Anecdotal |
| Curcin V, Woodcock T, Poots AJ, Majeed A, Bell D. | Model-driven approach to data collection and reporting for quality improvement [17]. | J Biomed Inform | 2014 | Model |
| Dale EL, Mueller MA, Wang L, Fogerty MD, Guy JS, Nthumba PM. | Epidemiology of operative burns at Kijabe Hospital from 2006 to 2010: pilot study of a web-based tool for creation of the Kenya Burn Repository. | Burns | 2013 | Model |
| Dillon DG, Pirie F, Rice S, Pomilla C, Sandhu MS, Metala AA, et al. | Open-source electronic data capture system offered increased accuracy and cost-effectiveness compared with paper methods in Africa [52]. | Clin Epidemiol | 2014 | Observational |
| Dugas M, Dugas-Breit S, Getz K, Hearn J, Sullivan R, Stewart D, et al. | Integrated data management for clinical studies: automatic transformation of data models with semantic annotations for principal investigators, data managers and statisticians. | PLoS One | 2014 | Model |
| Dunsmauir DT, Payne BA, Cloete G, Petersen CL, Gorges M, Lim J, et al. | Development of mHealth applications for pre-eclampsia triage. | IEEE J Biomed Heal Informatics | 2014 | Model |
| Dupont A, Wheeler J, Herndon JE, Coan A, Zafar SY, Hood L, et al. | Use of tablet personal computers for sensitive patient-reported information [64]. | J Support Oncol | 2009 | Observational |
| Dy CJ, Schmicker T, Tran Q, Chadwick B, Daluisi A, Hudak PL, et al. | The use of a tablet computer to complete the DASH questionnaire [24]. | J Hand Surg Am | 2012 | Observational |
| Eisenstein EL, Collins R, Cracknell BS, Podesta O, Reid ED, Sandercop P, et al. | Sensible approaches for reducing clinical trial costs. | Clin Trials | 2008 | Study |
| El Emam K, Jonker E, Sampson M, Krljez-Jeric K, Neisa A. | The use of electronic data capture tools in clinical trials: web-survey of 259 Canadian trials. | J Med Internet Res | 2009 | Observational |
| El Fadly A, Lucas N, Rance B, Verplancke P, Lastic P-Y, Daniel C. | The REUSE project: EHR as single datasource for biomedical research. | Stud Health Technol Inform | 2010 | Model |
| El Fadly A, Rance B, Lucas N, Mead C, Chatellier G, Lastic P-Y, et al. | Integrating clinical research with the Healthcare Enterprise: from the RE-USE project to the EHR4CR platform [57]. | J Biomed Inform | 2011 | Model |
| Ene-Iordache B, Carminati S, Antiga L, Rubis N, Ruggenenti P, Remuzzi G, et al. | Developing regulatory-compliant electronic case report forms for clinical trials: experience with the demand trial [53]. | J Am Med Inform Assoc | 2009 | Model |
Table 2
(Continued)

| Authors                               | Title                                                                 | Journal                          | Year   | Paper Type |
|----------------------------------------|-----------------------------------------------------------------------|----------------------------------|--------|------------|
| Farnell DJJ, Routledge J, Hannon R, Logue JP, Cowan RA, Wylie JP, et al. | Efficacy of data capture for patient-reported toxicity following radiotherapy for prostate or cervical cancer. | Eur J Cancer                      | 2010   | Observational |
| Faulds MC, Bauchmuller K, Miller D, Rosser JH, Shuker K, Wrench I, et al. | The feasibility of using “bring your own device” (BYOD) technology for electronic data capture in multicentre medical audit and research. | Anaesthesia                      | 2016   | Feasibility Study |
| Fontaine P, Mendenhall TJ, Peterson K, Speedie SM. | The “Measuring Outcomes of Clinical Connectivity” (MOCC) trial: investigating data entry errors in the Electronic Primary Care Research Network (ePCRN). | J Am Board Fam Med                | 2007   | Observational |
| Fraccaro P, Dentone C, Fenoglio D, Glacomin M. | Multicentre clinical trials’ data management: a hybrid solution to exploit the strengths of electronic data capture and electronic health records systems. | Informatics Heal Soc Care         | 2013   | Model |
| Franklin JD, Guidry A, Brinkley JF. | A partnership approach for Electronic Data Capture in small-scale clinical trials [18]. | J Biomed Inform                   | 2011   | Anecdotal |
| Fritz F, Tilahun B, Dugas M. | Success criteria for electronic medical record implementations in low-resource settings: a systematic review [68]. | J Am Med Inform Assoc             | 2015   | Review |
| Fu L, Ding S, Chen T. | Clinical Data Management System. | 2010 International Conference on Biomedical Engineering and Computer Science | 2010   | Model |
| Gallagher SA, Smith AB, Matthews JE, Potter CW, Woods ME, Raynor M, et al. | Roadmap for the development of the University of North Carolina at Chapel Hill Genitourinary Oncology Database – UNC GOLD. | Urol Oncol                        | 2014   | Anecdotal |
| Galliher JM, Stewart T V, Pathak PK, Werner JJ, Dickinson LM, Hickner JM. | Data collection outcomes comparing paper forms with PDA forms in an office-based patient survey. | Ann Fam Med                      | 2008   | Observational |
| Gibbons C, Caudwell P, Finlayson G, King N, Blundell J. | Validation of a new hand-held electronic data capture method for continuous monitoring of subjective appetite sensations. | Int J Behav Nutr Phys Act         | 2011   | Study |
| Gioli-Pereira L, Bernardez-Pereira S, Goulart Marcondes-Braga F, Rocha Spina JM, Muniz Miranda da Silva R, Evangelista Ferreira N, et al. | Genetic and Electronic medical records to predict outcomes in Heart Failure patients (GENIUS-HF) – design and rationale. | BMC Cardiovasc Disord             | 2014   | Study |
| Goodman K, Krueger J, Crowley J. | The automatic clinical trial: leveraging the electronic medical record in multisite cancer clinical trials [5]. | Curr Oncol Rep                    | 2012   | Model |
| Gupta SK. | Paperless clinical trials: myth or reality? [2] | Indian J Pharmacol                | 2015   | Anecdotal |
| Haak D, Samsel C, Gehlen J, Jonas S, Deserno TM. | Simplifying electronic data capture in clinical trials: workflow embedded image and biosignal file integration and analysis via web services [11]. | J Digit Imaging                   | 2014   | Model |
| Haller G, Hailer DM, Courvoisier DS, Louis C. | Handheld vs. laptop computers for electronic data collection in clinical research: a crossover randomized trial [67]. | J Am Med Inform Assoc             | 2009   | Observational |
| Hammond WE, Bailey C, Boucher P, Spohr M, Whitaker P. | Connecting information to improve health. | Health Aff                        | 2010   | Anecdotal |
| Harding JP, HAMM LR, EHSANULLAH RSB, HEATH AT, SORRELLS SC, HAW J, et al. | Use of a novel electronic data collection system in multicentre studies of irritable bowel syndrome. | Aliment Pharmacol Ther            | 1997   | Study |
| Harris PA, Taylor R, Thielke R, Payne J, Gonzalez N, Conde JG. | Research electronic data capture (REDCap) – a metadata-driven methodology and workflow process for providing translational research informatics support [44]. | J Biomed Inform                   | 2009   | Model |

(continues)
Table 2
(Continued)

| Authors | Title | Journal | Year | Paper Type |
|---------|-------|---------|------|------------|
| Hasekew J, Keny V, William R, Puri A, Mostafa Y, et al. | Use of Mobile Information Technology during Planning, Implementation and Evaluation of a Polio Campaign in South Sudan. | PLoS One | 2015 | Observational |
| Hensel DJ, Fortenberry JD, Harezlak J, Craig D. | The feasibility of cell phone based electronic diaries for STI/HIV research [63]. | BMC Med Res Methodol | 2012 | Study |
| Hetland ML. | DANBIO – powerful research database and electronic patient record [54]. | Rheumatology (Oxford) | 2011 | Model |
| Holzer B, Giesinger JM, Pinggera J, Zugal S, Schöpf F, Oberguggenberger AS, et al. | The Computer-based Health Evaluation Software (CHES): a software for electronic patient-reported outcome monitoring [12]. | BMC Med Inform Decis Mak | 2012 | Model |
| Huffstutter J, David Craig W, Schimizzi G, Harshbarger J, Lisse J, Kasle S, et al. | A multicentre, randomized, open study to evaluate the impact of an electronic data capture system on the care of patients with rheumatoid arthritis. | Curr Med Res Opin | 2007 | Study |
| Hye RJ, Inui TS, Anthony FF, Kiley M-L, Chang RW, Rehring TF, et al. | A multiregional registry experience using an electronic medical record to optimize data capture for longitudinal outcomes in endovascular abdominal aortic aneurysm repair. | J Vasc Surg | 2015 | Study |
| Installé AJ, Van den Bosch T, De Moor B, Timmerman D. | Clinical data miner: an electronic case report form system with integrated data preprocessing and machine-learning libraries supporting clinical diagnostic model research. | JMIR Med informatics | 2014 | Model |
| Jamison RN, Raymond SA, Levine JC, Slawsbay EA, Nedeljkovic SS, Katz NP. | Electronic diaries for monitoring chronic pain: 1-year validation study [69]. | Pain | 2001 | Study |
| Jamison RN, Raymond SA, Slawsbay EA, McHugo GJ, Baird JC. | Pain assessment in patients with low back pain: comparison of weekly recall and momentary electronic data [22]. | J Pain | 2006 | Observational |
| Jansen ME, Kollbaum PS, McKay FD, Rickert ME. | Factors influencing the electronic capture of patient-reported contact lens performance data. | Cont Lens Anterior Eye | 2013 | Observational |
| Jensen RE, Rothrock NE, DeWitt EM, Spiegel B, Tucker CA, Crane HM, et al. | The role of technical advances in the adoption and integration of patient-reported outcomes in clinical care. | Med Care | 2015 | Case Study |
| Katzan I, Speck M, Dopler C, Urchek J, Bielawski K, Dunphy C, et al. | The Knowledge Program: an innovative, comprehensive electronic data capture system and warehouse. | AMIA Annu Symp Proc | 2011 | Model |
| Kessel KA, Bohn C, Engelmann U, Oetzel D, Bougatf N, Bendr R, et al. | Five-year experience with setup and implementation of an integrated database system for clinical documentation and research. | Comput Methods Programs Biomed | 2014 | Model |
| Kho A, Zafar A, Tierney W. | Information technology in PBRNs: the Indiana University Medical Group Research Network (IUMG ResNet) experience. | J Am Board Fam Med | 2007 | Anecdotal |
| King C, Hall J, Banda M, Beard J, Bird J, Kazembe P, et al. | Electronic data capture in a rural African setting: evaluating experiences with different systems in Malawi. | Glob Health Action | 2014 | Model |
| King JD, Buolamwini J, Cromwell EA, Panfel A, Teferi T, Zerihun M, et al. | A novel electronic data collection system for large-scale surveys of neglected tropical diseases. | PLoS One | 2013 | Observational |
| Kinnula S, Renko M, Tapiainen T, Pokka T, Uhari M. | Post-discharge follow-up of hospital-associated infections in paediatric patients with conventional questionnaires and electronic surveillance [51]. | J Hosp Infect | 2012 | Observational |
| Kohl CD, Garde S, Knaup P. | Facilitating secondary use of medical data by using openEHR archetypes. | Stud Health Technol Inform | 2010 | Model |

(continues)
| Authors                          | Title                                                                 | Journal                      | Year  | Paper Type   |
|---------------------------------|----------------------------------------------------------------------|------------------------------|-------|--------------|
| Kuchinke W, Ohmann C, Yang Q, Salas N, Lauritsen J, Gueffier F, et al. | Heterogeneity prevails: the state of clinical trial data management in Europe - results of a survey of ECRIN centres. | Trials                        | 2010  | Observational |
| Kush R, Alschuler L, Ruggeri R, Cassells S, Gupta N, Bain L, et al.   | Implementing Single Source: the STARBRITE proof-of-concept study.     | J Am Med Inform Assoc         | 2007  | Feasibility Study |
| Laird-Maddox M, Mitchell SB, Hoffman M.                             | Integrating research data capture into the electronic health record workflow: real-world experience to advance innovation [8]. | Perspect Health Inf Manag     | 2014  | Model        |
| Le Jeannic A, Quelen C, Alberti C, Durand-Zaleski I.                 | Comparison of two data collection processes in clinical studies: electronic and paper case report forms [45]. | BMC Med Res Methodol          | 2014  | Observational |
| Levin E, Levin A.                                                     | Evaluation of spoken dialogue technology for real-time health data collection. | J Med Internet Res            | 2006  | Model        |
| Lin CH, Wu NY, Liou DM.                                               | A multi-technique approach to bridge electronic case report form design and data standard adoption. | J Biomed Inform               | 2014  | Model        |
| Long MD, Kappelman MD, Martin CF, Lewis JD, Mayer L, Kinneer PM, et al.| Development of an internet-based cohort of patients with inflammatory bowel diseases (CCFA Partners): methodology and initial results [13]. | Inflamm Bowel Dis             | 2012  | Model        |
| López-Carrero C, Arriaza E, Bolaños E, Ciudad A, Municio M, Ramos J, et al. | Internet in clinical research based on a pilot experience [55]. | Contemp Clin Trials           | 2005  | Model        |
| Lu M, Rupp LB, Moorman AC, Li J, Zhang T, Lamerato LE, et al.        | Comparative effectiveness research of chronic hepatitis B and C cohort study (CheCS): improving data collection and cohort identification. | Dig Dis Sci                   | 2014  | Observational |
| Lu Z.                                                              | Technical challenges in designing post-marketing eCRFs to address clinical safety and pharmacovigilance needs [14]. | Contemp Clin Trials           | 2010  | Anecdotal    |
| Lu Z.                                                              | Electronic Data-Capturing Technology for Clinical Trials.             | IEEE Engineering in Medicine and Biology Magazine | 2010  | Anecdotal    |
| Mahaffey KW, Wampole JL, Stebbins A, Berdan LG, McAfee D, Rorick TL, et al. | Strategic lessons from the clinical event classification process for the Assessment of Pexelizumab in Acute Myocardial Infarction (APEX-AMI) trial. | Contemp Clin Trials           | 2011  | Model        |
| Mall S, Akmatov MK, Schultz A, Ahrens W, Obi N, Pessler F, et al.    | Web-based questionnaires to capture acute infections in long-term cohorts: findings of a feasibility study. | Bundesgesundheitsblatt Gesundheitsforschung Gesundheitsschutz | 2014  | Observational |
| Maokola W, Willey BA, Shirima K, Chemba M, Armstrong Schellenberg JRM, Mshinda H, et al. | Enhancing the routine health information system in rural southern Tanzania: successes, challenges and lessons learned. | Trop Med Int Heal             | 2011  | Model        |
| Marley JE.                                                          | Safety and efficacy of nifedipine 20 mg tablets in hypertension using electronic data collection in general practice. | J R Soc Med                   | 1989  | Study        |
| Matza LS, Patrick DL, Riley AW, Alexander JJ, Rajmil L, Piel AM, et al. | Paediatric patient-reported outcome instruments for research to support medical product labelling: report of the ISPOR PRO good research practises for the assessment of children and adolescents task force. | Value Heal                    | 2013  | Anecdotal    |
| Meyer J, Fredrich D, Piegsa J, Habes M, van den Berg N, Hoffmann W. | A mobile and asynchronous electronic data capture system for epidemiologic studies [15]. | Comput Methods Programs Biomed | 2013  | Model        |
| Middleton RJ, Gavin AT, Reid JS, O'Reilly D.                        | Accuracy of hospital discharge data for cancer registration and epidemiological research in Northern Ireland. | Cancer Causes Control         | 2000  | Study        |

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### Table 2
(Continued)

| Authors | Title | Journal | Year | Paper Type |
|---------|-------|---------|------|------------|
| Mitchell JT, Kim YJ, Choi J, Park G, Cappi S, Horn D, et al. | Evaluation of data entry errors and data changes to an electronic data capture clinical trial database [46]. | Drug Inf J | 2013 | Observational |
| Nahm ML, Pieper CF, Cunningham MM. | Quantifying data quality for clinical trials using electronic data capture [63]. | PLoS One | 2008 | Anecdotal |
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|---------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------|------|------------|
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| Schmier JK, Kane DW, Halpern MT. | Practical applications of usability theory to electronic data collection for clinical trials.                                                                                                                                                                        | Contemp Clin Trials                | 2005 | Anecdotal |
| Schreier G, Messmer J, Rauehger G, Modre-Osprian R, Ladenstein R. | A Web-based platform for interdisciplinary biomedical research [42].                                                                                                                                                                                                  | Front Biosci (Landmark Ed)         | 2009 | Model |
| Schrimpf D, Haag M, Pilz LR. | Possible combinations of electronic data capture and randomization systems. Principles and the realization with RANDI2 and OpenClinica.                                                                                                                             | Methods Inf Med                    | 2014 | Anecdotal |
| Shah J, Rajgor D, Pradhan S, McCready M, Zaveri A, Pietrobon R. | Electronic data capture for registries and clinical trials in orthopaedic surgery: open source versus commercial systems.                                                                                                                                              | Clin Orthop Relat Res              | 2010 | Anecdotal |
| Taylor MJ, Stables R, Matata B, Lisboa PJG, Laws A, Almond P. | Website design: technical, social and medical issues for self-reporting by elderly patients [16].                                                                                                                                                                    | Health Informatics J               | 2014 | Observational |
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| Trachtenberg FL, Martin M, Green S, Oliveros O, Carson S, Gerstenberger E, et al. | Use of electronic data collection to assess pain in thalassaemia: a feasibility study.                                                                                                                                                                                | Int J Palliat Nurs                 | 2012 | Observational |
| Vahabzadeh M, Mezghani M, Guzman AE, Schmittner J, Preston K. | An adaptable assessment generation system for clinical trials complementing human research information system [43].                                                                                                                                                  | 18th IEEE Symposium on                | 2005 | Model |
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| Wang SJ, Middleton B, Prosser LA, Bardon CG, Spurr CD, Carchidi PJ, et al. | A cost–benefit analysis of electronic medical records in primary care [56].                                                                                                                                                                                          | Ann J Med                           | 2003 | Cost benefit analysis? |
| Weiler K, Christ AM, Woodworth GG, Weiler RL, Weiler JM, Meltzer E, et al. | Quality of patient-reported outcome data captured using paper and interactive voice response diaries in an allergic rhinitis study: is electronic data capture really better?                                                                                           | Ann Allergy, Asthma Immunol         | 2004 | Observational |
Clinical research and pharmacoepidemiology often involve interdisciplinary research. This not only means that various researchers deal with different data sources and formats but also that they have different workflows and organizational structures. As a consequence, there are no off-the-shelf solutions to facilitate this. This often results in individual solutions being developed that, over time, evolve and are ultimately difficult to maintain [42]. Unfortunately, it is often much easier to change time points, interventions, and assessment tools on paper than it is to suddenly change the programming of a computerized system. The reality demands future IT systems be flexible and adaptable with more automation [43].

### Advantages and disadvantages

There are distinct advantages to EDC in research and pharmacoepidemiology. However, there are pragmatic concerns that need to be addressed. The role of clinical research and pharmacoepidemiology is to improve healthcare by generating and providing access to high quality data. Due to the limitations of paper based records this is not possible with the status quo [44]. The objectives of ECD are to improve the quality of clinical trials, halt the development of ineffective or unsafe drugs earlier, reduce unnecessary work, reduce cost, and accelerate time to market of new drugs [45–51]. There are also benefits in relation to data quality, performance, productivity and costs in clinical trial management [52–56]. Observational data suggest that it is now considered a preferred method of data capture in clinical research [57–59]. It is well accepted by users and has been shown to contribute to patient empowerment, allowing them to be more engaged in research and to take direct control of their own data [60–62]. By contrast, paper-based questionnaires can suffer from incomplete forms, questions being answered twice or skipped questions. Paper forms are considered time consuming, require dual checking, and data cleansing [63], whereas EDC can alert people to missing answers before any attempt to proceed, and can be easily incorporated into electronic health records. Remote data collection offers the additional advantage of convenience to patients, particularly those who are incapacitated or live far outside the normal range at the time of entry and not days, weeks or months after. EDC systems have been shown to improve the quality of clinical trials, halt the development of ineffective or unsafe drugs earlier, reduce unnecessary work, reduce cost, and accelerate time to market of new drugs [45–51]. There are also benefits in relation to data quality, performance, productivity and costs in clinical trial management [52–56]. Observational data suggest that it is now considered a preferred method of data capture in clinical research [57–59]. It is well accepted by users and has been shown to contribute to patient empowerment, allowing them to be more engaged in research and to take direct control of their own data [60–62]. By contrast, paper-based questionnaires can suffer from incomplete forms, questions being answered twice or skipped questions. Paper forms are considered time consuming, require dual checking, and data cleansing [63], whereas EDC can alert people to missing answers before any attempt to proceed, and can be easily incorporated into electronic health records. Remote data collection offers the additional advantage of convenience to patients, particularly those who are incapacitated or live far from the nearest clinic [47], and may provide a safer environment for questionnaires than paper-based methods eliciting the answers to potentially sensitive questions [64, 65].

Despite the advantages, EDC has not been universally accepted. Perceived disadvantages and concerns regarding EDC include: a lack of available technical support, a lack of investigator motivation, complexity of installation,
maintenance of software, high initial investment cost, and complexity of use [66, 67]. Reliable data handling methods, effective project management, and expert technical architecture and infrastructure are all key factors for successful implementation, and should not be underestimated [68]. There are concerns over patient privacy and the need for computer literacy, which may affect generalisability of any research findings [60]. Study retention is considered to be higher where there is direct patient interaction because of the explicit alignment of patient incentives; the patient learns about the study directly, understands what is required, self-consents to participate, and then self-reports study information [69]. Jamison et al. [70] found better rates of compliance with electronic patient reported outcomes (PROs) than paper based PROs. Despite data suggesting benefits of EDC use, Alexander [71] reports that physicians lack motivation and will only use structured electronic records if the system reduces overhead while at the same time minimizing their work load. In the UK, it has been suggested that development of these technologies suffers from the lack of a clear national direction towards unifying clinical and medical data, with no common format for all data systems. Not only would EDC benefit clinical research, but pharmacovigilance and drug safety regulation could also be improved [72].

Discussion

IT and how it is used in pharmacoepidemiology and clinical research is a relatively new field with no substantial guidelines in place and few recommendations. There is a consensus that EDC has clear benefits for use in research but there are fears over security and data protection which must be addressed. IT offers an opportunity to improve pharmacoepidemiology and clinical research and to facilitate the continual improvement of healthcare. If the use of IT in research is to succeed fully, change is required: specifically, investment in infrastructure and the provision of support for integration of interoperable systems. Further efforts will be essential to alleviate healthcare providers and users legitimate concerns regarding IT. Policy makers will need to find ways to supply adequate financial resources to IT to counter a historical lack of investment within the public sector.

Healthcare providers and researchers require a governance-led support network of technology experts to assist in integrating ever more complex systems and providing guidance on compliance and security. IT security is a challenging and fast moving field and requires careful consideration. There is a need for clearer and more consistent policies and more trained data managers, software architects and semantic web specialists working in medical research groups. These architectures will need ongoing support from a robust legal system protecting patient privacy.

Furthermore, the exchange of information between systems is essential. Data format differences need to be resolved, and a solution found for interoperability between healthcare systems. Motivating software vendors, healthcare providers and researchers to agree on a common path will be difficult but worthwhile endeavour. Future technical development needs to focus on creating adaptable and generic software that can be tailored to specific trial needs without major re-development.

Limitations

This work has several limitations. Firstly, a publication bias is very likely as less successful IT projects are unlikely to be reported in published literature. This review only searched the IEEE library and the PubMed database and we did not include papers in non-English languages. In addition, researchers from low-income countries are known to have lower publication rates. The relative novelty of the field means that evaluation studies, in particular, are missing and rapid developments in the field may not yet have been published at the time of conducting the literature search. There are currently no widely accepted methods to evaluate technical publications in the same way as has been developed for reports of clinical trials, for example. Therefore, subjective interpretation had to be used to decide if a journal was of sufficient quality to be referenced. The authors took steps to avoid selection and objectivity bias by including all peer reviewed and published articles. The only exceptions authors made were where there was an overt conflict of interest, or the journal was not available in English. This review aimed to capture the full range of reported advantages and benefits of IT use. It did not measure the relative frequency or impact of individual factors of the utility of EDC and eCRFs. Despite the limitations detailed above, the authors believe this review to be an unbiased appraisal of publications on EDC and eCRFs in pharmacoepidemiology and clinical research.

Conclusion

It is apparent from the results of this literature review that the following areas would benefit from further development:

- Clearer legislation and operational frameworks governing electronic health records.
- Guidelines and best practises for researchers to follow in the use of IT and EDC.
- Standard methods of reporting and evaluating technical innovation to facilitate comparison.

Regardless of the challenges, it is the imperative that healthcare organizations ensure that patients receive safe medications. Effective clinical research and pharmacoepidemiology are essential to this process.

Competing Interests

There are no competing interests to declare.

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Contributors

D.R. conceived the idea with input from T.M.. The initial draft of the manuscript was created by D.R. T.M., R.F., A.D., K.G., I.M. and A.R. analysed and reviewed the manuscript. All listed authors fulfil the requirements for authorship and agree to submission of the manuscript in its current form.

Appendix

Table A1
Electronic data capture – IEEE

| Search term used                                | Results | Search criteria and filters | Date       |
|-------------------------------------------------|---------|----------------------------|------------|
| electronic case report form                     | 221     | NA                         | 27/07/2016 |
| (“Abstract”: electronic data capture)           | 18      | NA                         | 27/07/2016 |
| ecrf                                            | 14      | NA                         | 27/07/2016 |
| (“Abstract”: electronic data collection)        | 15      | NA                         | 27/07/2016 |

Table A2
Electronic data capture – PubMed

| Search term used                                      | Results | Search criteria and filters         | Date       |
|------------------------------------------------------|---------|------------------------------------|------------|
| (electronic data capture[Title/Abstract])            | 170     | Abstract available, Humans         | 27/07/2016 |
| electronic case report form                          | 546     | Abstract available, Humans         | 27/07/2016 |
| ecrf                                                 | 20      | Abstract available, Humans         | 27/07/2016 |
| (electronic data collection[Title/Abstract])         | 122     | Abstract available, Humans         | 27/07/2016 |

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