

BMJ Open Cross-sectional analysis of UK research studies in 2015: results from a scoping project with the UK Health Research Authority

Tim Clark,¹ Richard H Wicentowski,² Matthew R Sydes³

To cite: Clark T, Wicentowski RH, Sydes MR. Cross-sectional analysis of UK research studies in 2015: results from a scoping project with the UK Health Research Authority. *BMJ Open* 2018;**8**:e022340. doi:10.1136/bmjopen-2018-022340

► Prepublication history for this paper is available online. To view these files, please visit the journal online (<http://dx.doi.org/10.1136/bmjopen-2018-022340>).

Received 14 February 2018
Revised 22 May 2018
Accepted 23 August 2018



© Author(s) (or their employer(s)) 2018. Re-use permitted under CC BY. Published by BMJ.

¹Faculty of Medicine, Institut für Medizinische Informationsverarbeitung, Biometrie und Epidemiologie (IBE), Ludwig-Maximilians University, Munich, Germany
²Computer Science Department, Swarthmore College, Swarthmore, Pennsylvania, USA
³MRC Clinical Trials Unit at UCL, Institute of Clinical Trials and Methodology, University College London, London, UK

Correspondence to
Mr Matthew R Sydes;
m.sydes@ucl.ac.uk

ABSTRACT

Objectives To determine whether data on research studies held by the UK Health Research Authority (HRA) could be summarised automatically with minimal manual intervention. There are numerous initiatives to reduce research waste by improving the design, conduct, analysis and reporting of clinical studies. However, quantitative data on the characteristics of clinical studies and the impact of the various initiatives are limited.

Design Feasibility study, using 1 year of data.

Setting We worked with the HRA on a pilot study using research applications submitted for UK-wide ethical review. We extracted into a single dataset, information held in anonymised XML files by the Integrated Research Application System (IRAS) and the HRA Assessment Review Portal (HARP). Research applications from 2014 to 2016 were provided. We used standard text extraction methods to assess information held in free-text fields. We use simple, descriptive methods to summarise the research activities that we extracted.

Participants Not applicable—records-based study

Interventions Not applicable.

Primary and secondary outcome measures Feasibility of extraction and processing.

Results We successfully imported 1775 non-duplicate research applications from the XML files into a single database. Of these, 963 were randomised controlled trials and 812 were other studies. Most studies received a favourable opinion. There was limited patient and public involvement in the studies. Most, but not all, studies were planned for publication of results. Novel study designs (eg, adaptive and Bayesian designs) were infrequently reported.

Conclusions We have demonstrated that the data submitted from IRAS to the HRA and its HARP system are accessible and can be queried for information. We strongly encourage the development of fully resourced collaborative projects to further this work. This would aid understanding of how study characteristics change over time and across therapeutic areas, as well as the progress of initiatives to improve the quality and relevance of research studies.

INTRODUCTION

The need to improve the quality of clinical research is increasingly understood.¹ With

Strengths and limitations of this study

- First study to draw information from Health Research Authority (HRA's) Assessment Review Portal HARP system into one searchable dataset.
- Anonymised data from the HRA's HARP system can be interrogated with minimal manual intervention.
- Feasibility study, so limited to data from 2015 only and excluding phase I healthy volunteer studies.
- Aligning the questions from the XML document was not straightforward.
- The search terms chosen for the free-text fields were not exhaustive and because it was unfeasible in a pilot to review all applications, the sensitivity and specificity of the text-mining methodology could not be calculated.

a particular focus on clinical studies, the 'gold standard' for evidence-based medicine, there has been a noticeable push towards: improving study protocols¹; developing and implementing newer methodologies, such as adaptive designs²; involving patients in the design, conduct and management of studies³; and ensuring that study results are quickly and accurately reported.⁴

In order to properly evaluate the current state of clinical research and changes over time, it is clearly necessary to have unfettered access to the research protocols and study results.^{5 6} There are a number of limitations in accessing such information. Published evaluations of the state of clinical research are mainly based on the clinical study publication due to the difficulty in obtaining access to unpublished research protocols.⁷ Research registers are incomplete for many jurisdictions and/or are, like study publications, limited in detail compared with the research protocol. For example, clinicaltrials.gov⁸ has good coverage of clinical studies in North America, where registration of clinical trials is mandatory. The WHO International

Clinical Trials Registry Portal⁹ provides access to 15 different regional or national registries, but both have markedly less detail than in the study's research protocol. Published evaluations of the literature are likely affected by publication bias and there are often discrepancies between research publications and their underpinning protocols.¹⁰

The ability to analyse a large database of application forms, each of which contain very detailed information taken directly from the research protocol, would enable researchers to perform a detailed examination of the characteristics of clinical studies, particularly those necessary for generating reliable evidence. This would aid understanding of how study characteristics change over time and across therapeutic areas, as well as the progress of initiatives to improve the quality and relevance of clinical studies.^{11–15} Therefore, we approached the UK Health Research Authority (HRA), which oversees national ethical review of health research conducted within the National Health Service (NHS). With the support of the HRA and having signed the appropriate confidentiality declarations, we developed a pilot project to determine the feasibility of interrogating with minimal manual intervention the data contained in HARP (HRA Assessment Review Portal). HARP is a web-based management information system used by the HRA for all research ethics applications that require NHS REC (National Health Service Research Ethics Committee) review and/or HRA approval. We negotiated access to data on clinical studies submitted in 2015. The extract had personal data and organisational identifiers removed by HRA and did not include phase I healthy volunteer studies.

Our pilot focused on automatically summarising the characteristics of clinical studies, including therapeutic area, blinding, randomisation, use of Independent Data Monitoring Committee (IDMC), patient and public involvement (PPI) in the research, dissemination of the study results and use of new methodologies such as adaptive designs.

METHODS

The HRA collects applications for REC 'approval' (a favourable opinion) through the Integrated Research Application System (IRAS)¹⁶ and stores them in HARP. IRAS is a web-based system used to capture the information required by review bodies in the UK, including the Medicines and Healthcare Products Regulatory Agency and HRA. When the forms are completed, IRAS saves the results as an XML document. XML is a commonly used file format that stores structured data in plain text. The XML document can be used to generate PDF copies of the required forms and to repopulate the online web tool so users can later edit their applications.

Once the appropriate permissions were agreed, the HRA provided us with anonymised applications for clinical studies submitted for review from 2014 to 2016, excluding phase I healthy volunteer studies. For this

pilot, we wished to focus on the extraction of clinical study characteristics of interest from the PDF versions of the HRA forms underlying the IRAS application system. We were provided with an incomplete mapping between the questions on the PDF and the data stored in the XML document, such that many questions relevant to our study were not immediately labelled; nor was it apparent which questions were effectively switched on and off for applicants in completing the system's initial filter questions for their application. Therefore, we reverse-engineered the structure of the data in IRAS to determine which XML tags were responsible for creating the content we wanted to extract from the XML files to study.

The IRAS tool has been updated over time; not all of applications were created using the same version. Although the majority (91%) of the XML documents were created with one version (the latest version of the tool in the dataset), eight different versions were present in the dataset we received. We were not provided with the documentation detailing the ways in which the versions differed; we had access only to the latest version. Our inspection of the data indicated that the differences relevant to this pilot study were minor and we corrected for these when they were detected; however, there may be some undetected differences where the distinction between versions was important for our study.

For our pilot project, we chose to collect information on: the number of randomised controlled trials (RCT); the trial phase; the therapeutic area; the number of clinical trials of investigational medicinal products (CTIMPs — trials that require registration with regulatory body); the research method used (eg, RCT, feasibility/pilot study, blinding); the use of blinding; the use of systematic or formal literature reviews in planning the study; the use of adaptive or Bayesian designs; the use of a data monitoring committee; the plans for dissemination of findings; and the role of PPI in aspects of the research. Success would be measured in terms of our ability to access the information and generate descriptive summaries with minimal need for manual data reviews.

The data for our target areas were either held in multiple choice questions that we could identify and summarise or in free-text fields from which information needed to be determined and extracted. For those questions, we preprocessed the text using Spacy (<https://spacy.io/>), a natural language processing (NLP) package for Python. This involved identifying sentence boundaries and performing morphological analysis to convert words into their canonical dictionary form. We then used regular expressions to match phrases we were interested in, ignoring differences relating to spacing, hyphenation and capitalisation; Roman numerals were standardised to Latin numbers. For example, when looking for evidence that a systematic review was either performed or an existing review used during the study planning process, we searched for phrases such as 'systematic review', 'reviewed systematically', 'literature review', 'evidence review' and 'evidence-based review' in the answers provided to

Table 1 Overview of research activities submitted for REC approval

Characteristic	RCT		Other		Total	
	N	%	N	%	N	%
N	963		812		1775	
Therapeutic area*						
Cancer	168	17	192	24	360	20
Cardiovascular	94	10	94	12	188	11
Musculoskeletal	97	10	30	4	162	9
Respiratory	97	10	30	4	159	9
Paediatrics	56	6	30	4	148	8
Neurological	73	8	30	4	145	8
Mental health	66	7	30	4	117	7
Inflammatory and immune system	72	7	30	4	113	6
Oral and gastrointestinal	64	7	30	4	113	6
Blood	44	5	67	8	111	6
Diabetes	58	6	38	5	96	5
Infection	55	6	30	4	92	5
Renal and urogenital	39	4	30	4	82	5
Generic health relevance	46	5	30	4	76	4
Metabolic and endocrine	32	3	30	4	70	4
Dementias and neurodegenerative	27	3	36	4	63	4
Skin	36	4	30	4	57	3
Reproductive health and childbirth	30	3	30	4	56	3
Stroke	27	3	30	4	53	3
Eye	24	2	28	3	52	3
Injuries and accidents	22	2	30	4	36	2
Congenital disorders	10	1	20	2	30	2
Ear	6	1	7	1	13	1
Count of therapeutic areas claimed						
None	52	5	49	6	101	6
1	660	69	495	61	1155	65
2	201	21	200	25	401	23
3	29	3	49	6	78	4
≥4	21	2	19	2	40	2
Research methods used*						
RCT	963	100	2	0	965	54
Feasibility/pilot study	177	18	300	37	477	27
Questionnaire	98	10	139	17	237	13
Cohort observation	23	2	161	20	184	10
Qualitative research	90	9	76	9	166	9
Controlled trial, no randomisation	3	0	157	19	160	9
Laboratory study	30	3	47	6	77	4
Case series/case note review	7	1	45	6	52	3
Cross-sectional study	4	0	31	4	35	2
Case-control study	5	1	25	3	30	2
Database analysis	8	1	17	2	25	1
Epidemiology	7	1	5	1	12	1

Continued

Table 1 Continued

Characteristic	RCT		Other		Total	
	N	%	N	%	N	%
Meta-analysis	0	0	0	0	0	0
Other	68	7	269	33	337	19
Count of research methods used						
1	641	67%	522	64	1163	66
2	192	20	160	20	352	20
3	73	8	97	12	170	10
4	48	5	25	3	73	4
5	7	1	7	1	14	1
6	2	0	1	0	3	0
CTIMP						
Unlicensed	315	33	189	23	504	28
New use	146	15	70	9	216	12
Within SmPC	66	7	26	3	92	5
Other	10	1	5	1	15	1
<i>Any of these</i>	515	53	284	35		
Involves ionising radiation						
Yes	666	69	582	72	1248	71
No	297	31	222	28	519	29
Missing	0	NA	8	NA	8	NA
REC opinion						
Favourable	942	98	798	98	1740	98
Favourable	200	21	217	27	417	23
Favourable (extra info)	742	77	581	72	1323	75
Unfavourable	21	2	14	2	35	2

*Not mutually exclusive.

CTIMP, clinical trials of investigational medicinal product; RCT, randomised controlled trial; REC, Research Ethics Committee; SmPC, summary of product characteristics.

the relevant questions on the form, or instances where these words were used near each other, where 'near' was defined as being within three words. For adaptive designs, we employed the US Food and Drug Administration (FDA) definition, namely 'a study that includes a prospectively planned opportunity for modification'.¹⁷ We extracted the sentence containing the target phrase as well as one sentence before and after into a separate document. These textural extracts were then reviewed by the authors to identify true and false positives.

The data extracted from the separate XML files were collated into one dataset and descriptive analyses of this dataset were done with Stata.

RESULTS

We received (30 June 2016) the XML files for 1814 application records submitted for ethical review during the period specified, and extracted by HRA from the IRAS system. Of these, 1659 (92%) were from 2015, 154 (9%) from 2014 and 1 was from 2016. Three records were

corrupted and could not be processed. In discussion with HRA, we discarded some records to remove duplicates. We kept one of four entries for the WHEAT trial which had been sent to multiple committees as part of research on consistency of REC opinions.¹⁸ We discarded one of two entries for a further study, which was initially given a favourable opinion and then submitted additionally to a specialist REC in Scotland (again, favourably). Thirty-two further records were discarded that initially had an unfavourable opinion before re-submission as a (near) identical, separate application for review. We discarded the first applications and kept the re-submissions, regardless of the subsequent review's outcome. Therefore, our dataset included 1775 studies.

The filter questions switched off, for some applications, questions in which we were interested. For example, trial phase was only recorded for CTIMPs and the use of data monitoring committees was infrequently recorded. Furthermore, the use of blinding was not sufficiently well captured in the system to allow us to present reliable data.

A total of 963/1775 (54%) of the applications were stated as being RCTs. Table 1 describes the disease setting, research method, CTIMP and REC opinion by whether they were an RCT. The most common research area, using the categorisation of the IRAS system, was cancer, followed by cardiovascular, musculoskeletal and respiratory diseases; around one-third of records specified more than one area. This is broadly consistent with previous reviews.^{19 20} Around one-fifth of RCTs were pilot studies; one-third of records were employing more than one method. The REC gave a favourable opinion to the vast majority of applications (1740/1775, 98%), but required additional information for most of these 1323/1740 (75%).

Over half of the RCTs (515/963, 53%) were CTIMPs, of which 315/515 (61%) were CTIMPs of unlicensed products, 146/515 (28%) licensed CTIMPs in a new setting and 66/515 (13%) CTIMPs used according to their

summary of product characteristics (SmPC); the categories were not mutually exclusive. Over a third (284/812, 35%) of the other studies (not RCTs) were also CTIMPs.

Table 2 describes the plans in design and dissemination. Most RCTs (895/963, 93%) were not recorded as being preceded by a formal systematic review of the literature. Only 15 (2%) RCTs were detectably designed under a Bayesian framework; 20 (2%) were detectable as having an adaptive design. There were 34 cluster randomised and 5 stepped wedge trials, overall. Six hundred fourteen (63%) RCTs were requesting new biological samples; 120 (12%) RCTs were seeking access to previous biological samples.

The large majority of research (1667/1775, 94%) was planned for dissemination in the peer-reviewed literature and most (1585/1775, 89%) were planned for conference presentations. A total of 475/963 (49%) RCTs and 352/812 (43%) other studies were planned for submission

Table 2 Issues in design and dissemination in all entries

Characteristic	RCT		Other		Total	
	N	%	N	%	N	%
N	963		812		1775	
Review of data as part of development						
Neither	895	93	759	93	1654	93
Systematic review only	38	4	32	4	70	4
Meta-analysis only	18	2	13	2	31	2
Both	12	1	8	1	20	1
Design characteristics						
Neither	926	96	778	96	1704	1
Adaptive design	20	2	9	1	29	2
Bayes design	15	2	24	3	39	2
Both	2	0	1	0	3	0
Cluster randomised	26	3	8	1	34	2
Stepped wedge	3	0	2	0	5	0
Sample collection*						
Taking new samples	614	64	422	52	1036	58
Accessing stored samples	120	12	110	14	230	13
Plans for dissemination†						
Peer-reviewed scientific journal	912	95	755	93	1667	94
Conference presentation	869	90	716	8	1585	89
Internal report	641	67	526	65	1167	66
Publication on website	536	56	367	45	903	51
Submission to regulatory authority	475	49	352	43	827	47
Other publication	239	25	178	22	417	23
Access to raw data	191	20	133	16	114	6
Other	0	0	3	0	324	18
No plans to report or disseminate					3	0

*Not asked for non-regulatory RCTs.

†Not mutually exclusive.

RCT, randomised controlled trial.

Table 3 Reported PPI

Characteristic	RCT		Other		Total	
	N	%	N	%	N	%
N	963		812		1775	
Areas of PPI activity*						
Design of the research	403	42	323	40	726	41
Dissemination of findings	375	39	284	35	659	37
Undertaking the research	242	25	219	27	461	26
Management of the research	225	23	129	16	354	20
Analysis of results	72	7	75	9	147	8
Number of areas of PPI activity						
None	389	40	302	37	691	39
1	205	21	209	26	414	23
2	139	14	117	14	256	14
3	125	13	93	11	218	12
4	66	7	37	5	103	6
All	39	4	32	4	71	4

*Not mutually exclusive.

PPI, patient and public involvement.

to regulatory authorities. A small number, 114/1775 (6%), were planning to offer raw data to external applicants as a key form of dissemination.

Table 3 notes the reported patients and public involvement (PPI) using the categories specified by IRAS. A total of 726/1775 (41%) of studies claim PPI in the design and 659/1775 (37%) claim planned PPI in dissemination of the findings. PPI engagement in undertaking and management of the research was less common, and few studies (147/1775, 8%) involved planned PPI in analysis. Around one-third of studies involve PPI in two or more of these IRAS-defined areas.

Table 4 shows within CTIMPs the similarities and differences across the phase of research. Trial phase was asked as a series of separate yes/no questions for phase I, II, III and IV. These were not mutually exclusive. For the purpose of summarising, we selected the highest level if more than one option was selected. Early phase trials were more likely to be designed within a Bayesian framework; phase III trials most often reported the use of a placebo and the use of an Independent Data Monitoring Committee; phase IV trials were least likely to be submitted to a regulatory authority or put on a website, but most often reported PPI in all five broad areas.

Table 5 shows the outcome of the manual review of the extracted free-text fields. Accuracy, defined as number of true positives divided by overall total ranged from 43% for adaptive designs to 100% for stepped wedge. The false positives related to studies described as 'phase 1/2' or 'phase 2/3', but with no evidence of an adaptive step, for example, phase 1/2 was often used to describe a classical pharmacokinetic study in patients rather than healthy volunteers; the database did not contain healthy

volunteer studies. For 'systematic review' and 'meta-analysis', the search often picked up references to previous studies and not the planned study.

DISCUSSION

We achieved our primary feasibility aim of negotiating access to the centralised UK approvals system and devising a way of extracting information from a series of separate XML files. We anticipate that the programmes we developed could extract further pertinent points of information with minimal manual involvement. We were also able to achieve our primary descriptive aim of systematically reporting on the state of clinical research in the UK over around 1 year, focused on 2015.

We found that nearly 1000 RCTs were submitted for approval in the UK during that period and >800 other studies. A key message is the volume of research activity in the UK, with the country demonstrably research active. The majority of applications received a favourable opinion but most required further information first, suggesting that applications could be better completed, saving time and effort for all parties.

It was notable that the reported use of adaptive designs was low and was not markedly different from earlier estimates.^{19 20} This is disappointing, as many members of this design family have scope to reduce time to answer and/or reduce the average cost per answer, particularly by moving away early from treatment approaches that are not likely to improve outcomes sufficiently for patients.²¹ However, Bayesian approaches have penetrated 10% of early phase CTIMPs. The reported use of IDMCs for phase III CTIMPs was, perhaps, low at 58%, but this

Table 4 Characteristics of the CTIMPs by trial phase

Trial phase*	Ph I		Ph II		Ph III		Ph IV		Total	
Characteristic	N	%	N	%	N	%	N	%	N	%
All entries, N	99		287		324		98		808	
Design characteristics										
Neither	86	89	261	93	303	94	90	94	740	93
Adaptive	1	1	7	2	13	4	3	3	24	3
Bayes	10	10	12	4	5	2	2	2	29	4
Both	0	0	2	1	0	0	1	1	3	0
Cluster randomised	0	0	0	0	0	0	0	0	0	0
Stepped wedge	0	0	0	0	0	0	0	0	0	0
Involves placebo	21	22	123	44	163	51	25	26	332	42
Independent data monitoring committee										
No	40	82	84	59	61	42	46	78	231	58
Yes	9	18	59	41	84	58	13	22	165	42
Missing	8		139		176		37		400	
Methods of dissemination										
Peer-reviewed scientific journal	86	89	265	94	290	90	93	97	86	89
Submission to regulatory authority	85	88	215	76	272	85	48	50	85	88
Conference presentation	82	85	253	90	271	84	89	93	82	85
Internal report	71	73	212	75	259	81	60	63	71	73
Publication on website	56	58	161	57	203	63	45	47	56	58
Other publication	26	27	76	27	92	29	17	18	26	27
Access to raw data	8	8	12	4	25	8	8	8	8	8
Other	12	12	47	17	48	15	18	19	12	12
No plans to report or dissemination	0	0	0	0	2	1	0	0	0	0
Areas of PPI activity†										
Design of the research	26	27	82	29	39	12	50	52	26	27
Management of the research	10	10	34	12	28	9	31	32	10	10
Undertaking the research	14	14	47	17	36	11	29	30	14	14
Analysis of results	1	1	15	5	7	2	12	13	1	1
Dissemination of findings	20	21	74	26	64	20	44	46	20	21
None of the above	53	55	149	53	223	69	35	36	460	58
REC opinion										
Favourable	97	98	277	97	318	98	93	95	785	97
Favourable	14	14	51	18	40	12	15	15	120	15
Favourable (after extra info)	83	84	226	79	278	86	78	80	665	82
Unfavourable	0	0	5	2	3	1	3	3	11	1

*Trial phase calculated.

†Not mutually exclusive.

CTIMP, clinical trials of investigational medicinal product; PPI, patient and public involvement; REC, Research Ethics Committee.

is a substantial increase over the prevalence in trials published in high impact-factor journals in 1990 and 2000.²² Involvement of patients and the public in various aspects of research is increasingly seen as important, but reported rates of engagement were quite low: more than half of studies had no PPI involvement.²³ Cursory

review of the free-text fields associated with these PPI categories suggests that some applicants may have been a little generous in choosing to select a particular category, suggesting that the numbers we report may be overestimates. Other comments suggest that PPI is not needed, perhaps reflecting that the value of PPI is yet to

Table 5 Review of possible search terms

Design element Search term	Performance of search term*					
	At sentence level			At study level		
	True positive	False positive	Accuracy	True positive	False positive	Accuracy
Systematic review						
Review (near) systematically	2	0	100	2	0	100
Evidence (near) review	4	1	80	4	1	80
Systematic (near) review	98	39	72	58	23	72
Literature (near) review	29	13	69	26	12	68
<i>Evidence-based review</i>	0	0	NA	0	0	NA
Summary*†	133	53	72	90	36	71
Meta-analysis						
Pooled analysis	2	0	100	2	0	100
Meta-analysis	77	31	71	49	19	72
<i>Integrated analysis</i>	0	0	NA	0	0	NA
Summary*†	79	31	72	51	19	73
Adaptive design						
Adaptive design	19	0	100	13	0	100
Adaptive randomisation	8	0	100	5	0	100
Continual reassessment	2	0	100	1	0	100
Sample size re-estimation	15	0	100	9	0	100
Seamless design	1	0	100	1	0	100
Phase I/II	4	43	9	3	37	8
Phase II/III	2	25	7	2	23	8
<i>CRM design</i>	0	0	NA	0	0	NA
<i>Drop-the-loser</i>	0	0	NA	0	0	NA
<i>MAMS design</i>	0	0	NA	0	0	NA
<i>Multiarmed multistage</i>	0	0	NA	0	0	NA
<i>Pick-the-winner</i>	0	0	NA	0	0	NA
Summary*†	51	68	43	31	60	34
Bayes						
Bayes	83	2	98	42	1	98
Summary*†	83	2	98	42	1	98
Cluster RCT						
Cluster RCT	26	0	100	11	0	100
Control cluster	6	0	100	4	0	100
Randomised cluster	2	0	100	1	0	100
Cluster random	46	1	98	30	1	97
Summary*†	80	1	99	34	1	97
Stepped wedge						
Step (near) wedge	1	0	100	1	0	100
Stepped (near) wedge	16	0	100	5	0	100
Summary*†	17	0	100	5	0	100

NB The table reports on whether each search term accurately identified a specific design element of the study in selected free-text fields. All variations of the spelling, capitalisation, hyphenation and so on were covered in the search terms used.

*The left side of the table separately considers each sentence which matched a search term accurately identified the design property. There were 613 such sentences. The Summary row in each subsection is the sum of the constituent search terms.

†The right side of the table separately considers whether any sentence in a study matching a search term accurately identified the design property. Some studies matched a search term repeatedly, therefore the numbers are smaller. The Summary row in each subsection reflects a search across all of its constituent search terms. Note that some studies matched more than one search term in a subsection, so the Summary row may not be the sum of the rows above for example, a study that uses both 'cluster random' and 'control cluster' would appear only once in the Summary row, and would appear separately in the 'cluster random' and 'control cluster' rows.

CRM, continual reassessment method; MAMS, multi-arm multi-stage; RCT, randomised controlled trial.

be recognised by some researchers.³ Further research in this area is required.

Most but not all studies were planned for publication; there is still some way to go on this transparency aim. That raw data would be made available for some studies is encouraging and we hope that this will increase with time for appropriate projects from qualified researchers.²⁴

Extracting the information from the HRA system was more effortful than expected. Moreover, important characteristics of clinical trials like blinding could not be analysed and certain search terms clearly had limitations. The issue of blinding is particularly important, as it is one of the desirable characteristics in generating reliable evidence from clinical trials.¹¹ However, now that we have completed this feasibility step, it is possible for future research projects to access an expanded dataset and work with the HRA to organise the database in a way that will facilitate its analysis. Access to several years of data would allow researchers, as well as the HRA itself, to examine the progress of initiatives to improve the quality of clinical research, such as those designed to encourage PPI and publication of clinical trial results.^{13 15} We would also encourage the HRA to adopt standard definitions to describe the characteristics of clinical trials, as done in the USA.²⁵ Confusion over the definition of terms greatly complicates efforts to automatically analyse clinical research.

Our pilot work has some limitations. We had no straightforward way of aligning the questions in the PDF document with the information stored in the XML document. Perhaps if we had negotiated access to the backend software used by IRAS to generate these PDF forms, we may have been able to do this automatically; instead, we needed to reverse-engineer the IRAS system in order to figure out which XML tags were responsible for creating the content we wanted to study. Not all of the applications had been created using the same version of the IRAS tool.¹⁶ Although a large majority of the documents were created using the latest version of the tool, eight different versions were present in the dataset we received. We did not receive any guidance on the ways in which one version of the tool differed from another version, and we only had access to the latest version. We made efforts to determine the specific situations in which questions were not asked rather than asked but not answered, in order to make this distinction ourselves, though ideally this could be indicated directly in the XML dataset. The labelling of missing answers and inactivated questions was not consistent. There is scope for applicants to misunderstand the questions and misrepresent the data. For example, three trials were noted as using RCT methodology but answered 'no' to the question: 'Will participants be allocated to groups at random?' We checked the data and found these were not actually RCTs; we included them with the 'other studies'. Likely this error by the applicants activated or de-activated some questions inappropriately. In this example, the RCT method question actually asked about randomisation of individuals and these three

applicants who submitted cluster randomised trials had answered this question inconsistently. Finally, the search terms chosen for the free-text fields were not exhaustive and because it was unfeasible in a pilot to review all applications, the sensitivity and specificity of the free-text extraction could not be calculated.

Some interesting information was held only in the free-text fields. We used standard methods to extract this information using automation supplemented by manual checking. There is a wealth of free-text information, which through the application of text-mining techniques could provide incredibly valuable insights into the characteristics of health research. For this pilot study, we did not have the resources to review all of the questions and we particularly could not spend time reading free-text fields. Therefore, we felt obliged not to correct categorical questions if we noticed they were contrasted by free-text; however, a fully-resourced study could do this. For example, we noticed at least one entry, which ticked boxes for PPI engagement and expanded on this by claiming that, as doctors, they knew what patients wanted and did not need to trouble patients for their time on these aspects.

In conclusion, we have demonstrated that anonymised data from the HRA's research system are accessible and can be queried for information. We strongly encourage the development of fully resourced collaborative projects to delve more deeply into this data. We believe that it is imperative that the characteristics of clinical research in the UK are understood, as these underpin clinical guidance. Finally, there are many ongoing initiatives to improve the quality of clinical research, but only by fully understanding the state of, and changes to, the research profile of the UK, can we appreciate their impact.

Acknowledgements The authors would like to thank Hazel Gage, Amanda Hunn, Janet Messer—UK Health Research Authority for providing data, engaging in discussion and commenting on the draft manuscript and Professor Dr Ulrich Mansmann, IBE, Munich, Germany for commenting on the draft manuscript.

Contributors TPC and MRS: conceptualised the study, designed the study, collected the data, analysed and interpreted the data, wrote critical sections of manuscript, and reviewed and approved final manuscript. RHW: designed the study, collected the data, analysed and interpreted the data, wrote critical sections of manuscript, and reviewed and approved final manuscript.

Funding This study was funded by Medical Research Council (grant no: MC_UU_12023/24).

Competing interests TPC: worked as a consultant for the clinical research organisation ICON in the previous 3 years.

Patient consent Not required.

Provenance and peer review Not commissioned; externally peer reviewed.

Data sharing statement The data used in this analysis were obtained from the Health Research Authority under a confidentiality agreement and is not under the researcher's gift to share. Applications for these data or expanded data must involve the HRA. Access to the processing code is available through the MRC Clinical Trials Unit at UCL's usual Data Release Request approach.

Open access This is an open access article distributed in accordance with the Creative Commons Attribution 4.0 Unported (CC BY 4.0) license, which permits others to copy, redistribute, remix, transform and build upon this work for any purpose, provided the original work is properly cited, a link to the licence is given, and indication of whether changes were made. See: <https://creativecommons.org/licenses/by/4.0/>.

REFERENCES

1. Ioannidis JP. Why most clinical research is not useful. *PLoS Med* 2016;13:e1002049.
2. Hatfield I, Allison A, Flight L, et al. Adaptive designs undertaken in clinical research: a review of registered clinical trials. *Trials* 2016;17:150.
3. Elliott J. *HRA strategy for public involvement (v1.1)*. London, UK: Health Research Authority, 2013.
4. AllTrials. All Trials Registered | All Results Reported. alltrials.net (accessed 24 May 2013).
5. English R, Lebovitz Y, Griffin R. *Forum on drug discovery development and translation, institute of medicine. transforming clinical research in the United States: challenges and opportunities: workshop summary*: Institute of Medicine of the National Academies, Naitonal Academies Press, 2010:151.
6. Chalmers I, Glasziou P, Godlee F. All trials must be registered and the results published. *BMJ* 2013;346:f105.
7. Chan AW, Upshur R, Singh JA, et al. Research protocols: waiving confidentiality for the greater good. *BMJ* 2006;332:1086–9.
8. ClinicalTrials.gov. www.clinicaltrials.gov (accessed 22 May 2018).
9. Organisation WH. WHO clinical trials registration portal. <http://www.who.int/ictrp/en/> (accessed 22 May 2018).
10. Chan AW, Hróbjartsson A, Jørgensen KJ, et al. Discrepancies in sample size calculations and data analyses reported in randomised trials: comparison of publications with protocols. *BMJ* 2008;337:a2299.
11. Califf RM, Zarin DA, Kramer JM, Sherman RE, Aberle LH, Tasneem A. Characteristics of clinical trials registered in ClinicalTrials.gov, 2007–2010. *JAMA* 2012;307:1838–47.
12. Grignolo A. The Clinical Trials Transformation Initiative (CTTI). *Ann Ist Super Sanita* 2011;47:14–18.
13. INVOLVE. *Public involvement in research: values and principles framework*. Eastleigh, 2015.
14. Chan A-W, Tetzlaff JM, Altman DG, et al. SPIRIT 2013: new guidance for content of clinical trial protocols. *The Lancet* 2013;381:91–2.
15. Moorthy VS, Karam G, Vannice KS, et al. Rationale for WHO's new position calling for prompt reporting and public disclosure of interventional clinical trial results. *PLoS Med* 2015;12:e1001819.
16. Health Research Authority. IRAS - Integrated Research Application System. <https://www.myresearchproject.org.uk/> (accessed 22 May 2018).
17. FDA. *Draft guidance for industry adaptive design clinical trials for drugs and biologics*, 2010.
18. Gale C, Hyde MJ, Modi N. WHEAT trial development group. Research ethics committee decision-making in relation to an efficient neonatal trial. *Arch Dis Child Fetal Neonatal Ed* 2017;102:F291–8.
19. Clark T. Sample size calculations in clinical trials of investigational medicinal products: an analysis of ongoing practice. *Der Andere Verlag* 2013.
20. Clark T, Berger U, Mansmann U. Sample size determinations in original research protocols for randomised clinical trials submitted to UK research ethics committees: review. *BMJ* 2013;346:f1135.
21. Parmar MK, Barthel FM, Sydes M, et al. Speeding up the evaluation of new agents in cancer. *J Natl Cancer Inst* 2008;100:1204–14.
22. Sydes MR, Altman DG, Babiker AB, et al. Reported use of data monitoring committees in the main published reports of randomized controlled trials: a cross-sectional study. *Clin Trials* 2004;1:48–59.
23. Sacristán JA, Aguarón A, Avendaño-Solá C, et al. Patient involvement in clinical research: why, when, and how. *Patient Prefer Adherence* 2016;10:631–40.
24. Taichman DB, Backus J, Baethge C, et al. Sharing clinical trial data: a proposal from the International Committee of Medical Journal Editors. *The Lancet* 2016;387:e9–e11.
25. ClinicalTrials.gov. ClinicalTrials.gov Protocol Registration Data Element Definitions for Interventional and Observational Studies. 2017 <https://prsinfo.clinicaltrials.gov/definitions.html> (accessed 2018-05-22).