To the Cochrane Scientific Committee

Interim guidance on how to decide whether to include clinical study reports and other regulatory documents into Cochrane reviews

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To the Cochrane Scientific Committee

Summary
The documented presence of reporting bias in biomedical literature of clinical trials is a major threat to the validity and credibility of Cochrane reviews. This interim guidance report outlines the rationale for accessing clinical study reports and other regulatory documents (regulatory data) as a means of addressing reporting bias, and identifies factors that may aid in the decision of whether (or not) to include regulatory data into Cochrane reviews. The guidance includes the origins and current state of regulatory data access, a survey of current systematic reviewers’ practices in considering regulatory data for inclusion in reviews, and a glossary (with hyperlinks and screenshots) of terms and nomenclature used in regulatory documents. The glossary will be of considerable practical use to Cochrane authors, given that most lack familiarity.

This guidance does not address how to access, assess and extract regulatory data.

Cochrane should consider making regulatory data a preferred source, primarily when the intervention in question is of potential high value and when there is evidence of reporting bias, or both. Cochrane should invest in the infrastructure to make this possible.

Scope of this document
Recognising the need to widen the scope of data sources for Cochrane intervention reviews beyond journal publications, conference abstracts or trial registry reports, the Cochrane Methods Innovation Fund (MIF) funded a project to produce interim guidance on the circumstances under which clinical study reports and other regulatory documents should be considered for inclusion in Cochrane reviews, either in addition to or instead of data from more traditional sources.

It is important to note that the scope of this project is limited to consideration of ‘whether’ (or not) to incorporate regulatory data into a Cochrane review and when it might be most important. The project did not intend to address the question of how to use these data sources. The rationale for considering the question of ‘whether’ was an assumption that not all reviewers would have the resources to incorporate clinical study reports and other regulatory documents into their reviews, and therefore some guidance for prioritization would be helpful. We refer to ‘interim’ guidance on how to decide whether to incorporate clinical study reports and other regulatory documents into Cochrane reviews, as we have been unable to identify any research evidence in this area.

For those who elect to include clinical study reports and other regulatory documents in their review, the next issue is ‘how’ to incorporate such data. This project, however, was not funded to address the ‘how’ question.

Background
Systematic reviews of randomised controlled trials play an important role in decision-making. If properly designed and conducted, they identify, evaluate and summarise complex trial-derived information and provide more reliable and precise estimates of intervention effects than individual studies. Up to now, most systematic reviews have used data extracted from journal publications. In a survey of 348 systematic reviews published in 2014, around three-quarters relied solely on data provided in peer reviewed journals. ¹

Of those that accessed other sources, data from trials registries (such as ClinicalTrials.gov), conference proceedings or contacting authors were the most used. No reviews reported using or attempting to obtain clinical study reports even though the majority of the reviews evaluated drug interventions. ¹ A survey of 2184 Cochrane authors also found that contacting ‘trialists/investigators,’ was one of the most common methods for accessing unpublished data and that data from manufacturers or from regulatory agencies
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were rarely obtained.2

“Clinical study reports” (CSRs) are documents prepared and submitted to regulators to obtain a license, and represent the most complete synthesis of the planning, execution, and results of a clinical trial. CSRs contain some of the same information as journal papers (i.e. rationale, objectives, methodology, results, discussion/conclusion), but are substantially more detailed with numerous large tables and figures, and datasets not constrained by page limits. A CSR for a single trial may be hundreds, thousands, or even tens of thousands of pages in length and are easy to navigate throughout (when all components are in a single file). CSRs generally contain, as appendices, important study documents including the study protocol and amendments, statistical analysis plan and amendments, case report forms (CRFs), patient information sheet, certificates of analysis, informed consent forms, and individual patient listings among others. CSRs therefore provide more detailed information and complete data than are usually available in journal articles.

“Regulatory documents” is a term we use to refer to any document produced by, or held by, a regulatory agency. Notable types of regulatory documents are CSRs (which are submitted by sponsors) and Medical Officer Reviews (produced by US Food and Drug Administration medical officers) or European Public Assessment Reports (produced by the European Medicines Agency).

(See definitions of the above terms, and others, in the Glossary).

In late 2010, the European Medicines Agency (EMA) began releasing CSRs of drugs and biologics on request under its Policy 0043.33,45 In October 2016, the EMA began to release CSRs prospectively under its Policy 0070.3,4 Policy 0070 applies only to marketing authorisation applications received since 1 January 2015. Documents available from the EMA under this policy normally include the clinical overview, clinical summary, and CSRs of individual trials. Each CSR describes the trial design, conduct and results of analyses including three selected appendices of each CSR: the study protocol, statistical analysis plan, and sample case report forms.3 In 2017, Health Canada published a report announcing an initiative to publicly release clinical information concerning drugs and devices under an eventual EMA Policy 0070-like mechanism.6 And in January 2018, the FDA announced that it will publicly release CSRs in a pilot program.7

There is an increasing potential for CSRs and other regulatory documents to be considered for inclusion in Cochrane reviews and to alter the way that systematic reviews are conducted in future due to their increasing public availability from a variety of sources.

Not all interventions have regulatory data
At the outset of this project in 2014, we decided to focus on clinical study reports and other regulatory documents relating to pharmaceuticals and biologics for which these documents generally exist. We acknowledge, however, that non-pharmaceutical interventions (such as implantable devices, surgery, rehabilitation, behavioural interventions and diagnostics) are responsible for a large part of healthcare expenditure and that regulatory activity and transparency have been recently increasing in this area, at a slower pace, in particular in the field of devices.

It is important to note that publicly funded trials, even of drugs and biologics, do not usually produce internationally standardised documentation similar to a CSR and are not the focus of this document.

Rationale for the consideration of regulatory documents (including clinical study reports) as sources of data for inclusion in Cochrane reviews
There has been a gradual realisation that sources of evidence historically considered to be reliable (such
as peer-reviewed literature) are affected by reporting bias. Reporting bias generally refers to selective reporting of research depending on the nature and direction of research results. Reporting bias includes publication bias\textsuperscript{8-12} and outcome reporting bias,\textsuperscript{8-14} among many others.\textsuperscript{15}

Studies published in the last decade or so have shed light on reporting biases present in publicly available reports of trials of pharmaceuticals and have highlighted the general lack of transparency that surrounds trial data. Combined, these present a major obstacle in assessing bias in studies included in Cochrane reviews.

Reporting biases can generally only be detected when two or more reports of the same trial are compared: for example, peer-reviewed publications compared with CSRs. In addition to reporting bias, lack of transparency and lack of detail in journal publications may prevent or hinder detailed analyses of data which could be relevant to specific subpopulations potentially benefiting from or being harmed by the intervention.\textsuperscript{16} This situation is likely to be the consequence of compressing thousands of pages of text and tables into the historically restricted confines of a printed journal article.\textsuperscript{17}

There are indications that CSRs may be incomplete and in some cases may be internally inconsistent between different components of the same CSR.\textsuperscript{18} However, a consistent picture emerges when comparing different data sources for the same trial: CSRs provide the greatest breadth and depth of information compared to journal articles, register data and grey literature. Aggregate data on subpopulations are often found in CSRs and can provide a source of further analysis. Such a wealth of information gives a fuller and more reliable picture of trial strengths and weaknesses, as well as a more reliable assessment of the benefits and harms of the studied interventions.

Table 1 contains a selected and illustrative list of studies that have compared different sources of data for the same trial, such as publication vs. CSR or trial register entries vs. publications. Although this is not an exhaustive list of all such studies, it covers more than 50 different interventions and offers glimpses of the ways in which reporting bias affects the biomedical literature.

The studies in Table 1 strongly suggest that discrepancies in the reporting of trials across different sources of data is common. There are limitations to be aware of when interpreting discrepancies. First, different types of trial documents may have very different objectives. CSRs, for example, inform regulators and, by law, provide a comprehensive record of a study. Trials registers, in contrast, are primarily a visible collection of trials and their reporting format is heterogeneous. For example ClinicalTrials.gov does not have a methods section and results can be either absent or incomplete. Under some circumstances (such as for specific funding sources), reporting of trials within trials registers, including the submission of results, is compulsory (but not always adhered to nor adequately policed).\textsuperscript{19} Within the United States, there are requirements for National Institutes of Health (NIH) funded research,\textsuperscript{20} but in many cases worldwide trial registration is voluntary. The EMA’s European Public Assessment Reports (EPARs) and the FDA’s Drug Approval Packages are records of regulators’ work in assessing a medicine for potential registration and are not primarily meant to provide summaries of trials for reviewing purposes (see Glossary of terms and definitions for taxonomy of regulatory documents). Journal articles are the main means of communicating clinical trial results providing short, accessible summaries of trial findings; but there is increasing evidence that articles may be incomplete or biased. Journal trial articles are readily available and provide relatively short, usually readable, summaries. These qualities are offset by the bias they may introduce and the considerable time and effort expended (sometimes in vain) to clear up discrepancies, contradictions and missing information.

The generalisability of each finding of the studies in Table 1 to the larger population of trials or topic areas that exist is debatable, and it is unclear whether reporting biases are lessening over time. Some journals
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have taken steps to limit the bias introduced by the current format of trial reporting, by requiring adherence to CONSORT, by publishing the trial protocol or supplementary online data as an appendix or by requiring data sharing as a condition of publication.\textsuperscript{21–23} As it is impossible to squeeze thousands of pages’ worth of information into a 10-page publication and the resulting information selection is based on unknown criteria, an alternative solution may be that authors can, where these exist, provide links to the relevant CSR and other summary data (e.g. FDA Drug Approval Packages).

We are aware of three examples of four systematic reviews (a Cochrane review of neuraminidase inhibitors, twin reviews of rhBMP-2, and a review of reboxetine) allowing assessment of contributions of regulatory data compared to the same trial data from published journal articles.\textsuperscript{24–27}

In the case of recombinant human bone morphogenetic protein 2 (rhBMP-2), both CSRs and individual participant data were included in the twin reviews,\textsuperscript{25,26} while the Cochrane review of neuraminidase inhibitors and the review of reboxetine were based on CSRs.\textsuperscript{24} In all cases the conclusions of important aspects of the reviews were changed with access to the more complete data available in the CSRs. Access to the CSRs also provided a deeper understanding of the strengths and limitations of the trial evidence. In the case of the review of reboxetine, the inclusion of CSR data changed the conclusions of the review, and allowed quantification of the exaggeration in favour of the effects of reboxetine compared to placebo (99-115%) and other SSRIs (19-23%).\textsuperscript{27} The Cochrane review of neuraminidase inhibitors for influenza also found FDA medical officer reviews to be an important source of data and detail.

As Cochrane reviews are considered to be a gold standard of reliable research synthesis, we need to pay attention to the issue of reporting bias and to address whether, and how to decide when, accessing regulatory data, including CSRs, might offer a solution. The approach, however, is new and unfamiliar to most Cochrane reviewers and at the time of writing, regulatory data are not always immediately available. When available, using such documents involves reviewing very large quantities of information, which is time-consuming and resource intensive. Thus, a framework to help identify where using data from regulatory documents is likely to matter most, and prioritising those reviews which should adopt such an approach, will be helpful for Cochrane groups grappling with how to respond to the increasing availability of these new sources of information.

Current Cochrane practice
To raise awareness of the above issues and to inform our work, we surveyed Cochrane and non-Cochrane authors to gauge how many had considered using regulatory data and how many had actually included them in their reviews. The survey was announced in the Cochrane Digest, and in an email to all Cochrane authors on 10th June 2016. The release intended for authors of non-Cochrane reviews was first advertised on the University of York Centre for Reviews and Dissemination website on the 25th June 2016 and then on the Systematic Reviews journal website. Links to this were also shared via social media. Both surveys were closed on the 19th September 2016 and then the results were combined.

There were 160 respondents who completed the Cochrane (n=153) and non-Cochrane (n=7) surveys combined (Table 2). However, it is not clear how many authors received, opened the digest, or read the invitation to participate. 20/160 (13%) of the respondents had previously requested or used CSRs and other regulatory documents, 7/160 (4%) had considered it, and 133/160 (83%) had never considered it. Data sought by survey respondents were mainly from the EMA and/or the FDA (19 (40%) of the 47 requests made by those previously requesting CSRs in total) and/or directly from pharmaceutical companies (18/47 (38%).) 5/47 (11%) of the requests included non-regulatory data requests to authors of published trials. Amongst the 20 respondents that requested regulatory data, 12 (60%) involved CSRs, five obtained medical and statistical reviews from the FDA and two European public assessment reports (EPARs). The main reasons for accessing CSRs were concerns about reporting biases 11/20 (55%),
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outcome reporting bias and publication bias (5/20 - 25%).

Fourteen (70%) of the authors that had used or requested regulatory data, and 6 (86%) authors who only considered regulatory data had faced barriers to access. These were identified mainly as the restricted and limited sharing of trials data, and the time constraints involved in searching and requesting the data.

The survey results in brief show a lack of familiarity with regulatory sources of data, barriers to access and lack of resources to do so. The main rationale for authors seeking regulatory data, however, was minimisation of bias.

The circumstances under which clinical study reports and / or other regulatory documents should be considered for inclusion in Cochrane reviews

Because of unfamiliarity and the additional investment of resources required, we do not consider that Cochrane reviews can be converted immediately to include routine searching for, or inclusion of, regulatory data without a period of preparation and consideration of the consequences of such an action. It may not be necessary for all reviews to adopt such an in-depth approach. It is also important to take stock of current practice. Selection of likely candidate reviews is therefore required.

We were unable to identify any research on the topic of how to decide whether to incorporate clinical study reports and other regulatory documents into systematic reviews, i.e. a rule for determining which reviews would most benefit from the inclusion of such data.

We therefore created an initial list of reasons (or triggers) for seeking and using such data through discussion amongst our group. Our list was a product of our opinion and experience. We then carried out a follow-up targeted survey in which we asked respondents to rate the importance of each criterion in our list. This survey was sent to the 21 (of 27) systematic review authors who had used, requested, or considered using regulatory data in their review and had agreed to participate in a follow-up survey. Fourteen of 21 (66%) provided a response. A description of the criteria are in Table 3 and the results are presented in Figure 1 (where criteria are listed in order of importance).

When authors were asked which criteria were considered most important when considering access to regulatory data (Figure 1), omission and underreporting of trial outcomes and results were the most frequently cited because of likelihood of reporting bias (criteria 10-14, 17 and 18). However, the other criteria listed in Figure 1 and Table 3 were also considered important by most authors.

The variables are self-explanatory, reflecting either known or suspected bias in published results or the potential for greatest impact in terms of public health - for example, what are the human costs of acting on biased estimates of effectiveness or harm?

There is no proposed scoring or algorithm for combining criteria to identify priority topics or topic area. The relative importance of criteria listed in Table 3 will depend very much on context, and prioritisation is inevitably a somewhat subjective process.

The authors would be interested in receiving suggestions or reports of experience regarding accessing regulatory data and including such data in systematic reviews. However, as mentioned elsewhere, that this is beyond the scope of this current project.

Discussion

Regulatory documents are a complex and underutilised source of highly detailed data for Cochrane reviews. Although the methodological steps for their inclusion, extraction and analysis are broadly the
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same as those with other sources of data, the resource implications of their use are not. For example, resource use for constructing an index of all pharmaceutical comparative studies from multiple sources and reading and extracting data from drug approval packages are 6 months for one whole time equivalent for two drugs of the same class and one vaccine. The results of our surveys and our own experience indicate that the use of regulatory documents should be considered, especially when the intervention in question is of potential high value and when there is evidence of reporting bias, or both.

We think that Cochrane should consider making regulatory data a preferred source of data in such circumstances and should invest in the infrastructure to make this possible. This ranges from supporting a regulatory data option in reviews of pharmaceuticals, to training aids on the content and use of such data, to investing in a research programme to identify priorities and limitations of the use of regulatory data.

Acknowledgments (peer reviewers)

We are grateful for helpful comments from the following; Chris Champion, Senior Programme Manager, CEO’s Office, Cochrane, Mark Helfand, Professor of Medicine and Professor of Medical Informatics and Clinical Epidemiology at the Oregon Health & Science University, Rene Allard, Sarah Nevitt, Jean-Mac Ferran, Sherry Meeh and Kim Musgrave of Pharmaceutical Users Software Exchange (PhUSE), Erick Turner, Senior Scholar, Oregon Health & Science University, and Beate Wieseler, Head of the Institute for Quality and Efficiency in Healthcare (IQWiG) Drug Assessment Department.

We are very grateful to those who responded to calls for feedback and we endeavoured to incorporate their comments. Incorporation of feedback comments and/or listing of names above does not infer respondents’ endorsement of our work.

Acknowledgments (support)

Kristina Charlotte Dietz, Centre for Reviews and Dissemination, University of York who helped to organise and summarise the survey results.
Toby Lasserson, Senior Editor, Cochrane Editorial Unit, for providing an approximate number of active Cochrane reviews over the last 2 years to complement our survey results.
Table 1. Examples of studies comparing different sources of data for the same trials.

<table>
<thead>
<tr>
<th>Reference</th>
<th>Type of study</th>
<th>Intervention comparisons</th>
<th>Source comparison</th>
<th>Take home message</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chan 2004</td>
<td>Cohort study of 102 randomized trials registered with scientific-ethical committees in Denmark, 1994-1995</td>
<td>75% drug trials, 12% counseling/lifestyle trials, 11% surgery/procedure, 2% equipment</td>
<td>Protocols vs publications</td>
<td>“62% of trials had at least 1 primary outcome that was changed, introduced, or omitted.” In 40 of 82 trials, pre-specified primary outcomes were not presented as such in the journal publication. In 11 trials, outcomes not pre-specified were reported as the “primary outcome” in the publication. “The reporting of trial outcomes is not only frequently incomplete but also biased and inconsistent with protocols.”</td>
</tr>
<tr>
<td>Turner 2008</td>
<td>Review of 74 RCTs for 12 antidepressants reviewed by the Food and Drug Administration, and their corresponding publication (or lack thereof) in the literature</td>
<td>12 antidepressants vs. placebo</td>
<td>Medical officer reviews vs publications</td>
<td>Non-publication and selective reporting occurred frequently, and can change the apparent risk-benefit assessment of drugs. Publicly available medical officer reports are a valuable source of unbiased information about clinical trial design and results.</td>
</tr>
<tr>
<td>Eyding 2010</td>
<td>Systematic review of 13 trials. 76% of patient data unpublished: 86% (1946 of 2256 patients) for reboxetine vs placebo and 67% (1760 of 2641 patients) for reboxetine vs SSRI</td>
<td>Reboxetin for depression vs placebo or vs other SSRIs included in IQWIG HTA report</td>
<td>CSRs vs publications</td>
<td>The addition of unpublished data changed the direction and conclusions of the efficacy and harms analyses. Published data vs full dataset overestimate benefits by 99-115% vs placebo and 19-23% vs other SSRIs.</td>
</tr>
<tr>
<td>Jefferson 2012</td>
<td>Cochrane review of 25 trials (15 oseltamivir, 60% unpublished, those published had been ghostwritten and corresponding “authors” had no access to study data)</td>
<td>Neuraminidase inhibitors for influenza vs placebo</td>
<td>CSRs vs publications</td>
<td>Lack of detail in publication and unexplained discrepancies when compared to CSRs led the authors to change methods compared to previous version of the review and include only regulatory data, significantly changing the conclusions of the review.</td>
</tr>
<tr>
<td>Coyne 2012</td>
<td>Review of the Normal Hematocrit Trial (NHT) run in the 1990s on 1265 hemodialysis patients with cardiac disease</td>
<td>Epoetin lower (9–11 g/dl) vs higher (13–15 g/dl) doses to increase hematocrit to reduce mortality and improve survival and QoL</td>
<td>CSR vs publication</td>
<td>“Disclosure of these [CSR] results in the 1998 publication or access to the FDA filed report on the NHT in the late 1990s would likely have led to earlier concerns about epoetin safety and greater doubts about its benefits.”</td>
</tr>
<tr>
<td>Wieseler 2012</td>
<td>Systematic review of 29 studies included in 16 HTA reports prepared by IQWIG during 2006-2011</td>
<td>16 different pharmaceuticals mainly for depression and type I and II diabetes</td>
<td>CSRs vs publications vs register entries</td>
<td>CSR consistently reported more information than registers or journal publications.</td>
</tr>
</tbody>
</table>
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<table>
<thead>
<tr>
<th>Author(s)</th>
<th>Type of Review</th>
<th>Details</th>
<th>Findings/Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wieseler 2013</td>
<td>Systematic</td>
<td>Systematic review of 101 trials with full CSR available included in 16 HTA reports prepared by IQWIG. The study population is the same as Wieseler 2012 but in this study the authors quantified information gain for patient-relevant outcomes graded from 1 to 4. 16 different pharmaceuticals mainly for depression, asthma and type I and II diabetes. CSRs vs publications vs register entries (unclear which trials have been registered where. Also some trials were conducted in the late 1980s). CSRs reported complete information on 78%-100% of benefit outcomes vs 20% - 53% in combined publicly available sources. The authors estimated 13% publication bias. CSRs reported complete information on 84% - 92% of harm outcomes vs 27% to 72% of combined publicly available sources. 15% NR by publicly available sources for both general harms and withdrawals due to possible harms.</td>
<td></td>
</tr>
<tr>
<td>Rodgers 2013 &amp; Fu 2013</td>
<td>Systematic</td>
<td>Systematic review of 13 trials and 4 single arms studies (10 and 1 journal published) Recombinant human bone morphogenetic protein 2 (rhBMP-2) for spinal fusion vs iliac crest bone graft. IPD vs CSRs vs journal publications. Wealth of extra detail from CSRs provided by manufacturer. “Early journal publications misrepresented the effectiveness and harms through selective reporting, duplicate publication, and underreporting.” Fu et al conclude that “Early journal publications misrepresented the effectiveness and harms through selective reporting, duplicate publication, and underreporting.”</td>
<td></td>
</tr>
<tr>
<td>Doshi 2013</td>
<td>Descriptive</td>
<td>Descriptive review of 78 CSRs. 14 different pharmaceuticals and biologics. CSRs vs publications (comparison in size). The ratio of CSR pages to publication pages for available full CSRs with a corresponding publication (“compression factor”) ranged from 379 to 8805.</td>
<td></td>
</tr>
<tr>
<td>Vedula 2013</td>
<td>Review of</td>
<td>Review of transparency and accuracy of reporting of the numbers of participants, description of types of analyses, and criteria for including participants in the analysis in 11 published trials. Gabapentin vs placebo for four off-label uses (migraine prophylaxis, treatment of bipolar disorders, neuropathic pain, and nociceptive pain). CSRs accessed from litigation with their published counterparts (21 trials identified, 11 assessed, 8 trials excluded because unpublished, 1 not randomised, 1 no CSR available). Probably biggest discrepancies occurred between protocol and publication. Authors conclude “we found that the trial publication was not a transparent, or accurate (presuming that the research report truly describes the facts), record for the numbers of participants randomized and analyzed for efficacy”.</td>
<td></td>
</tr>
<tr>
<td>Maund 2014</td>
<td>Review of</td>
<td>Review of nine trials in 1999-2001 (7 journal published). Duloxetine vs placebo. CSR vs publications vs register entries. 1/9 R1 and 9/9 R2. 7 S published 2 NS unpublished. 1 NS published as S after post hoc analysis not mentioned in the paper. Harms 50% and 25% participant reporting inaccuracy in 2 trials, 1 death in active arm in unpublished trial; lack of clarity on phase of deaths Suicide NR &lt; 2% in register reports. SAE 3 articles failed to report, register entries unclear.</td>
<td></td>
</tr>
<tr>
<td>Le Noury 2015</td>
<td>RIAT publication, restoring GSK’s trial 329 run in the 1990s and journal published in 2001. Paroxetine vs placebo &amp; imipramine vs placebo. IPD with CRFs for 34% (93/275) participants and CSR vs publication. Paroxetine was reported as safe and effective in company sponsored ghost written publications. Access to CSR data led the authors to conclude that the drug was no more effective than placebo and was toxic in adolescents. The authors identified 4 outcomes cited in the protocol but not reported in the CSR and publication.</td>
<td></td>
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</tr>
<tr>
<td>Author</td>
<td>Year</td>
<td>Study Description</td>
<td>Comparative Groups</td>
</tr>
<tr>
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</tr>
<tr>
<td>Köhler</td>
<td>2015</td>
<td>Systematic review of 15 dossier assessments by AMNOG submitted to IQWIG between 2011 and 2015. The authors assessed completeness of reporting in each document category.</td>
<td>15 different drugs including anti HIV and oncology</td>
</tr>
<tr>
<td>Beaumier</td>
<td>2015</td>
<td>Cochrane review update of 4 CSR (3 journal-published in 4 publications)</td>
<td>Olanzapine vs placebo</td>
</tr>
<tr>
<td>Cosgrove</td>
<td>2016</td>
<td>Review of data considered by regulators for registration vs other data available to them vs publications and comparison of regulatory vs SR process</td>
<td>Vortioxetine vs placebo (4 RCTs) or active comparator (6 studies) for depression</td>
</tr>
<tr>
<td>Hodkinson</td>
<td>2016</td>
<td>Exploratory review to assess the reporting of harms in Orlistat trials</td>
<td>Orlistat vs placebo</td>
</tr>
<tr>
<td>Jureidini</td>
<td>2016</td>
<td>Litigation documents vs publication</td>
<td>Citalopram vs placebo</td>
</tr>
</tbody>
</table>
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<table>
<thead>
<tr>
<th>Schroll 2016 (^{40})</th>
<th>Descriptive review of 7 RCTs to assess the reporting of AEs</th>
<th>Orlistat vs placebo</th>
<th>7 CSRs from Roche vs. Protocols vs. Journal publications</th>
<th>“Study identified important disparities in the reporting of adverse events between protocols, clinical study reports, and published papers. Reports of the trials systematically understated adverse events. Based on the study findings, systematic reviews of drugs might be improved by including protocols and CSRs in addition to published articles.”</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mayo-Wilson 2017 (^{41})</td>
<td>Impact assessment to determine whether disagreements among multiple data sources of the same trials affected meta-analytic effect estimates, statistical significance and interpretation</td>
<td>Gabapentin and quetiapine</td>
<td>21 gabapentin RCTs (74 reports, six IPD) and seven quetiapine RCTs (50 reports, one IPD)</td>
<td>“Disagreements across data sources affect the effect size, statistical significance, and interpretation of trials and meta-analyses.”</td>
</tr>
</tbody>
</table>

Key:

AMNOG = Arzneimittelmarktneuordnungsgesetz (Germany’s Act on reform of the market for medicinal products);
CSR = clinical study reports;
CV = cardiovascular;
IQWIG = Institute for Quality and Efficiency in Health Care, Germany;
NA = Not applicable;
NK = Not known;
NR = Not reported (by the authors);
NS = statistically not significantly different;
QoL = quality of life.
R1 (Registration 1) = in public register;
R2 (Registration 2) = in manufacturer register;
S = statistically significantly different;
SAE = serious adverse events
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Table 2: Characteristics of respondents and their experiences with regulatory data

<table>
<thead>
<tr>
<th>Question</th>
<th>Requested regulatory data</th>
<th>Considered regulatory data</th>
<th>Not considered regulatory data</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n=20</td>
<td>n=7</td>
<td>n=133</td>
</tr>
<tr>
<td>Should regulatory data be used in Cochrane reviews?</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>15 (75)</td>
<td>3 (43)</td>
<td>43 (32)</td>
</tr>
<tr>
<td>In some cases</td>
<td>5 (25)</td>
<td>3 (43)</td>
<td>66 (50)</td>
</tr>
<tr>
<td>No</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>17 (13)*</td>
</tr>
<tr>
<td>Rationale for using regulatory data?</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Under reporting of harms</td>
<td>3</td>
<td>2</td>
<td>N/A</td>
</tr>
<tr>
<td>ORB</td>
<td>11</td>
<td>3</td>
<td>N/A</td>
</tr>
<tr>
<td>Publication bias</td>
<td>5</td>
<td>0</td>
<td>N/A</td>
</tr>
<tr>
<td>Missing data</td>
<td>2</td>
<td>1</td>
<td>N/A</td>
</tr>
<tr>
<td>Other</td>
<td>2*</td>
<td>2**</td>
<td>N/A</td>
</tr>
<tr>
<td>Familiarity with the regulatory process for pharmaceutical and biologics?</td>
<td>N/A</td>
<td>N/A</td>
<td>n=133</td>
</tr>
<tr>
<td>Yes - detailed understanding</td>
<td>N/A</td>
<td>N/A</td>
<td>8 (6)</td>
</tr>
<tr>
<td>Yes - basic understanding</td>
<td>N/A</td>
<td>N/A</td>
<td>83 (62)</td>
</tr>
<tr>
<td>No</td>
<td>N/A</td>
<td>N/A</td>
<td>42 (32)*</td>
</tr>
<tr>
<td>Awareness of debate for improved access to clinical trial data?</td>
<td>N/A</td>
<td>N/A</td>
<td>n=133</td>
</tr>
</tbody>
</table>
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<table>
<thead>
<tr>
<th></th>
<th>Yes</th>
<th>N/A</th>
<th>N/A</th>
<th>113 (85)***</th>
</tr>
</thead>
<tbody>
<tr>
<td>No</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>20 (15)</td>
</tr>
</tbody>
</table>

N/A: question was not asked in the survey as it was not applicable; ORB: outcome reporting bias

*(1) was a request from a reviewer and, (1) for detailed medical information from manufacturer about the product

**(1) Because of uncertainty in risk of bias assessment domains and (1) based on national Australian guidelines for regulatory approval

γ Reasons why regulatory data should not be considered: (9) interventions non-pharmacological, (5) lack of guidance on how to include the data and (3) too time-consuming

η Reasons why not familiar: (2) Respondents conducted non-pharmacological reviews that do not require familiarity with regulatory data

***(2) respondents mentioned the AllTrials initiative, (2) mentioned the “Tamiflu review”, (1) respondent was involved in the EMA policy 0070 regarding access to clinical trial data in 2014 and (1) Ben Goldacre’s Bad Pharma.
Table 3. Criteria for assessing whether to include regulatory data of a drug or biologic in a Cochrane review (not in order of priority)

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Description of criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Monetary cost of the intervention on the healthcare budget (i.e. considering both the price of a course and the number of people in the population that are being - or will be treated)</td>
</tr>
<tr>
<td>2</td>
<td>Burden of disease of the indication this product is meant to treat/prevent</td>
</tr>
<tr>
<td>3</td>
<td>Number of people using or likely to use the product</td>
</tr>
<tr>
<td>4</td>
<td>Product new to the market</td>
</tr>
<tr>
<td>5</td>
<td>Product from a new drug class or has a new mechanism of action</td>
</tr>
<tr>
<td>6</td>
<td>Has important interactions with other drugs (e.g. drug-drug interactions)</td>
</tr>
<tr>
<td>7</td>
<td>High proportion of RCTs evaluating this product are industry funded</td>
</tr>
<tr>
<td>8</td>
<td>Prominent claims of safety and/or efficacy advantage of this product over currently available treatments</td>
</tr>
<tr>
<td>9</td>
<td>High degree of media attention surrounding this product</td>
</tr>
<tr>
<td>10</td>
<td>High proportion of trials of this product are unpublished</td>
</tr>
<tr>
<td>11</td>
<td>Post-marketing surveillance has identified safety concerns</td>
</tr>
<tr>
<td>12</td>
<td>Important or standard outcome measures (also known as 'endpoints') have not been published</td>
</tr>
<tr>
<td>13</td>
<td>Concerns regarding a lack of published data on potential harms of the product</td>
</tr>
<tr>
<td>14</td>
<td>Marketing authorization based on surrogate outcomes (rather than clinical outcomes)</td>
</tr>
<tr>
<td>15</td>
<td>When protocol(s) are publicly available</td>
</tr>
<tr>
<td>16</td>
<td>When statistical analysis plan(s) publicly available</td>
</tr>
<tr>
<td>17</td>
<td>Known errors or concerns about trial publications of this product</td>
</tr>
<tr>
<td>18</td>
<td>Important discrepancies between the journal publication and the trial registry entry</td>
</tr>
</tbody>
</table>
Figure 1: Criteria for considering using regulatory data by order of importance according to 14 authors who had used, requested, or considered using regulatory data
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Glossary of terms and definitions for taxonomy of regulatory documents
(Also see supplemental file of screenshots)

Acronyms

- **AusPAR.** See Australian Public Assessment Report
- **CRF.** See Case Report Form.
- **CSR.** See Clinical Study Report.
- **CTD.** See Common Technical Document.
- **EMA.** See European Medicines Agency.
- **EPAR.** See European Public Assessment Report.
- **FDA.** See Food and Drug Administration.
- **FOI.** See Freedom of Information.
- **FOIA.** See Freedom of Information Act.
- **ICF.** See Informed Consent Form
- **IMRAD.** See Introduction, Methods, Results, and Discussion.
- **IPD.** See Individual Participant Data.
- **ISE.** See Integrated Summary of Effectiveness
- **ISS.** See Integrated Summary of Safety
- **MAA.** See Marketing Authorization Application.
- **MOR.** See Medical Officer Review
- **NDA.** See New Drug Application.
- **PIL.** See Patient Information Leaflet.
- **PSUR.** See Periodic Safety Update Report.
- **RAP.** See Report and Analysis Plan
- **SAP.** See Statistical Analysis Plan.
- **SmPC.** See Summary of Product Characteristics.

Terms

- **Advisory Committee documents.** See FDA Advisory Committee meeting materials.
- **Aggregate data.** A colloquial term referring to summary data resulting from combining individual level data (e.g. mean age). **Individual listings** data can be combined to form aggregate data, but this cannot occur in reverse.
- **Amendments.** May refer to Study Protocol amendments or Statistical Analysis Plan. Amendments, documents that list the various versions and changes made to a protocol over time. Amendments can vary in detail. Sometimes they document the original text, the new text, and the reason for the change.
- **Annotated Case Report Form.** An empty Case Report Form (CRF) in which the variable names are noted (annotated) next to fields, indicating how entries were to be recorded in the electronic dataset. Such information can be used to understand how data recorded on CRFs were transformed into an electronic patient level dataset.
  - Example: Zanamivir trial NAI30031, ACRF Contact, PDF p. 6
  - Also see an example in the Screenshots Gallery
- **Appendices.** See Clinical Study Report Appendices.
- **Australian Public Assessment Report (AusPAR).** A public assessment report, authored by the Australian regulator Therapeutic Goods Administration, that summarizes the evaluation and considerations of TGA in deciding to approve or not approve a marketing application for a prescription medicine. Whereas one EMA EPAR is written for each medicine, an AusPAR is created for a single marketing application, and is not updated following publication. Additional AusPARs are published for generic medicines, major variations and extensions of indications.
The first AusPAR was published in Nov 2009. Also see European Public Assessment Report.

**Drug Approval Package**
- Example: AusPAR for Vytorin
- Look up other AusPARs here
- More information here

**Biologic License Application (BLA).** The regulatory vehicle through which sponsors submit a biologic for possible marketing approval to the Food and Drug Administration. The requirements are similar, but not identical, to those of a New Drug Application.

**Blank Case Report Form.** A sample Case Report Form (CRF), of unique pages only, that is, empty forms not yet filled in. One copy of all CRFs used in a trial is typically contained in section 16.1.2 of Clinical Study Reports formatted according to the ICH E3 guidelines.
- Example: Tamiflu (oseltamivir) trial NV18671 PDF page 336-527
- Also see an example in the Screenshots Gallery

**Case Report Form (CRF).** The original paper or electronic forms on which individual participants’ data (demographic, efficacy, safety, etc) are recorded during the clinical trial. The forms are typically the most ‘raw’ form of detailed data available for understanding what happened in a clinical trial, and the data they contain are statistically analysed only after they have been entered into an electronic database of individual patient data. Forms can vary in length, from a few pages to hundreds of pages, and each trial can have multiple forms—for example, for different visits or for the different tests or procedures the participant undergoes.
- Example: Arthronat trial MA-CT-10-002 PDF pp. 3985-4749.
- Also see an example in the Screenshots Gallery

**Centralised procedure.** See European Medicines Agency (EMA).

**Certificate of analysis.** A short report in a CSR describing a chemical analysis and physical appearance of the contents of the medications (including any placebo) used in the clinical trial.
- Example: Tamiflu (oseltamivir) trial WP16263 page 422-3.
- Also see an example in the Screenshots Gallery

**Clinical Overview.** See Module 2.5 (Clinical Overview) & Module 2.7 (Clinical Summary).

**Clinical Study Report (CSR).** An unabridged report of a clinical study written for regulators following the E3 reporting guidelines developed by the regulatory-industry collaborative effort International Conference on Harmonisation of Technical Requirements for Registration of Pharmaceuticals for Human Use (ICH). CSRs represent the most complete synthesis of the planning, execution, and results of a clinical trial. CSRs contain some of the same information as journal articles (i.e rationale, objectives, methodology, results, discussion/conclusion), but are substantially more detailed with numerous large tables and figures, and datasets not constrained by page limits. In addition, CSRs generally contain, as appendices, important study documents including the study protocol and amendments, statistical analysis plan and amendments, case report forms (CRFs), patient information sheet, certificates of analysis, informed consent forms, and individual patient listings among others.
- Example: Tamiflu (oseltamivir) trial M76001
- Also see an example in the Screenshots Gallery

**Clinical Study Report Appendices.** Clinical Study Reports generally contain numerous appendices. The ICH E3 guideline document lists recommended appendices which start in section 16 of the document. These include the study protocol and amendments (section 16.1.1), statistical analysis plan and amendments (section 16.1.9), blank case report form (section 16.1.2), blank informed consent form (section 16.1.3), randomization scheme and codes (section 16.1.7), audit certificates (section 16.1.8), and patient data listings including discontinued patients (section 16.2.1), protocol deviations (section 16.2.2), adverse event listings for each patient (section 16.2.7), case report forms for deaths, other serious adverse events, and withdrawals for adverse events (section 16.3.1), and individual patient data listings (section 16.4).
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- **Clinical Summary.** See Module 2.5 (Clinical Overview) & Module 2.7 (Clinical Summary).
- **Common Technical Document (CTD).** The name adopted by the ICH which refers to the way of structuring quality, safety, and efficacy information in support of a marketing authorization application (called a New Drug Application at the FDA). The CTD format is used by the EMA, FDA and Japanese PMDA. The CTD contains five modules (Modules 1, 2, 3, 4, and 5). Clinical Study Reports are contained in Module 5. The CTD is depicted by the ICH as a pyramid.
  - Also see the pyramid in the Screenshots Gallery
- **Drug Approval Package.** Reviews of clinical study reports and related documents for approved drugs, written by staff from the US Food and Drug Administration (FDA). Drug Approval Packages can be found of the Drugs@FDA website. Drug Approval Packages generally include the approval letter, summary review, medical review, chemistry review, pharmacology review, statistical review, clinical pharmacology biopharmaceutics review and microbiology review. Drug Approval Packages may also include the printed labeling, officer/employee list, office director memo, proprietary name review and administrative documents and correspondence and other reviews. FDA makes similar documents available for biologics under the Center for Biologics Evaluation and Research (CBER) website. Also see: European Public Assessment Report, Australian Public Assessment Report.
  - Example: [Sivextro (Tedizolid Phosphate)](http://www.fda.gov/drugsatfda)
  - Also see an example in the Screenshots Gallery
- **Drug label.** (Also known as prescribing information, product information, labelling, package insert, summary of product characteristics) The content of this document varies by regulator, but generally is an official description of a medical product that includes the indication (for what the medicine is used, and in which population), contraindications, adverse events, instructions for safe use, and technical details. Labels may also include information on clinical pharmacology, toxicology, and clinical trials. This information generally has a primary audience of healthcare professionals, and in the United States can be searched for on DailyMed (NIH), the FDA Online Label Repository, and Drugs@FDA. In the EU, drug labels are referred to as the Summary of Product Characteristics, and can be searched for on EMA’s website (at present, they can be found under the “Product information” tab of the drug’s page on EMA’s website). Information specifically written for patients is found in Medication Guides, Patient Package Inserts, and Patient Information Leaflets, often found attached to the healthcare professional information.
  - Example: [Dalvance (dalbavancin) FDA approved label](http://www.fda.gov/drugsatfda)
  - Example: [Xydalba (dalbavancin) EMA Summary of Product Characteristics](http://www.fda.gov/drugsatfda)
  - Also see an example in the Screenshots Gallery
- **Drugs@FDA.** Searchable database of regulatory data maintained by the FDA. Drugs@FDA offers public access to drug labels, patient information (cf. patient information leaflet), approval letters, medical officer reports, statistical officer reports, and other elements of the drug approval package for drugs approved since 1998. For drugs prior to this date, a Freedom of Information Act request is necessary to obtain these documents. A comparable database does not exist for biologics, but similar information is available by searching the “Vaccines, Blood & Biologics” section of FDA’s website.
  - Drugs@FDA: [http://www.fda.gov/drugsatfda](http://www.fda.gov/drugsatfda)
  - For biologics, look for FDA reviews on the webpage of each individual product. FDA has a webpage that lists of all licensed biologics with supporting documents.
- **EMA Policy 0043.** The European Medicines Agency Policy 0043 governs the agency’s approach to the retrospective release of certain documents when in the agency’s possession. This includes Clinical Study Reports (and other parts of the Common Technical Document including Modules 2.5 and 2.7), Investigator’s Brochures, and Periodic Safety Update Reports. In colloquial terms, it is the agency’s freedom of information policy. The policy is
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dated 30 November 2010, effective from 1 December 2010 and its official title is “European Medicines Agency policy on access to documents (related to medicinal products for human and veterinary use).”


● EMA Policy 0070. The European Medicines Agency Policy 0070 governs the agency’s approach to the prospective publication of clinical data in the agency's possession. The policy was finalized on October 2, 2014, effective from 1 January 2015 but its implementation is happening in stages, beginning with the publication of Clinical Study Reports (Autumn 2016). A second phase (Phase 2) is planned for the future, to cover third party access to individual participant data. The official title of Policy 0070 is “European Medicines Agency policy on publication of clinical data for medicinal products for human use.”


● EMA Policy on Publication of Clinical Data. Also known as the policy on “Clinical Data Publication.” See EMA Policy 0070.

● European Medicines Agency (EMA). Regulatory agency in the European Union responsible for drugs and biologics approved through a centralised procedure. Medicines can also come to market through other non-centralised procedures, such as in a specific individual country or group of countries.

● European Public Assessment Report (EPAR). Not a single document but a collection of regulatory documents describing the evaluation of all medicines granted or refused marketing authorization by the European Medicines Agency. Documents include a lay summary, labelling, package leaflet, summary of product characteristics, a public assessment report for the initial authorization and subsequent major changes, and an overview of procedural steps taken before and after authorization. Some information is published in all official languages of the EU while other documents are in English only, and some are only available online.

○ Example: Olazax (Olanzapine) EPAR

○ Also see an example in the Screenshots Gallery

● Food and Drug Administration (FDA). Regulatory agency in the United States responsible for food (including dietary supplements), drugs, biologics, medical devices, radiation-emitting electronic products, veterinary products, and tobacco products.

● FDA Advisory Committee meeting materials. The FDA makes use of federal advisory committees in an effort to receive independent advice from outside experts regarding regulatory decision making. Under US law (Federal Advisory Committee Act), meeting materials made available to committee members must be made available to the public at or before the time of the meeting. Meeting materials generally consist of two types: sponsor submitted materials and FDA submitted materials. These materials may contain limited data from clinical trials, but can include data not available elsewhere as well as FDA analyses of data (e.g. pooled analyses or sensitivity analyses). Unlike the Drugs@FDA database, Advisory Committee meeting materials may discuss applications that ultimately are not approved by the FDA, and as such serve as a source of unpublished data. Materials are, however, released to the public subject to the Freedom of Information Act (FOIA), which the FDA interprets as exempting certain types of information from disclosure, and therefore the publicly accessible versions may contain redactions. FDA posts Advisory Committee materials on its website, and generally also posts meeting minutes and a meeting transcript.

○ Advisory Committee meeting materials homepage: http://www.fda.gov/AdvisoryCommittees/CommitteesMeetingMaterials/default.htm
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○ Example: http://www.fda.gov/AdvisoryCommittees/CommitteesMeetingMaterials/Drugs/ReproductiveHealthDrugsAdvisoryCommittee/ucm446101.htm

- **Freedom of Information (FOI).** General term that refers to laws or other governmental mechanisms allowing public access to documents held by governments. For discussion relevant to clinical trial data, for the United States see Freedom of Information Act (FOIA) and for Europe, see EMA Policy 0043 and EMA Policy 0070.

- **Freedom of Information Act (FOIA).** A United States freedom of information law passed in 1967 that gives the public - generally irrespective of citizenship - the right to request records from any US federal agency. Thus far the FDA has generally (but not always) considered clinical trial data to be exempt from release under FOIA.

- **ICH E3 guideline.** A guidance document entitled “Structure and Content of Clinical Study Reports,” developed by the International Conference on Harmonisation of Technical Requirements for Registration of Pharmaceuticals for Human Use (ICH) Expert Working Group. They were formalised in 1995 “to assist sponsors in the development of a report that is complete, free from ambiguity, well organised and easy [for regulators] to review.” Most clinical study reports follow the general structure laid out in the E3 guidelines, which have not been updated since 1995.
  ○ ICH E3 guideline

- **Individual listings.** A colloquial term referring to a document or electronic dataset which contains data recorded at the level of the individual participant. In CSRs, individual listings are provided in section 16.2 (Patient Data Listings), 16.3 (Case Report Forms), and 16.4 (Individual Patient Data Listings). In written documents they generally come in the form of tabular data, but may also appear in other forms, for example as is the case of individual participant Serious Adverse Events narratives (ICH E3 guidelines section 12.3.2) and withdrawals. Some journal publications may include individual listings as supplementary online material. Contrast with Aggregate data.
  ○ Example: Paroxetine trial 329

- **Individual participant data (IPD).** Data for each participant in a trial. This contrasts with aggregate or summary data, which is produced by combining data from multiple participants. Individual participant data allows for the replication of all analyses in study reports and exploration of further analyses. IPD generally come in two forms: electronic datasets (that are therefore readily analyzable with software packages) and printed/paper listings (as in the type found in the sections of CSRs that contain individual listings).

- **Informed Consent Form (ICF).** An information sheet that is required by law to be provided to potential research participants to enable an informed decision regarding study participation. The information sheet is also accompanied by a form used to document study participants' understanding of the study and consent to participate. Major elements that information sheets should contain include a description of the study purpose, information on the study intervention(s), study procedures, potential side effects, risks and benefits, compensation, and participants' rights.

- **Integrated Summary of Effectiveness (ISE) and Integrated Summary of Safety (ISS).** Integrated summaries of effectiveness data and of safety (harms) data of more than one study, possibly including pooled/meta analyses, prepared for the FDA (required for New Drug Applications, and encouraged for Biologic License Applications). In the harmonized regulatory submission dossier, the Common Technical Document, the ISE and Integrated Summary of Safety (ISS) might be found in section “5.3.5.3 Reports of Analyses of Data from More than One Study (Including Any Formal Integrated Analyses, Meta-Analyses, and Bridging Analyses).” The Summary of Clinical Efficacy (section 2.7.3 of the CTD) and Summary of Clinical Safety (section 2.7.4 of the CTD) were meant to replace the ISE and ISS

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- **Integrated Summary of Safety (ISS).** Similar to the Integrated Summary of Effectiveness, but instead of clinical efficacy, the focus of an ISS is on safety (harms) of a product. See Integrated Summary of Effectiveness (ISE) and Integrated Summary of Safety (ISS).

- **Introduction, Methods, Results, and Discussion (IMRAD).** Acronym used to describe the typical structure of a scientific report, that begins with the Introduction section, followed by Methods, Results, and finally a Discussion section. Short reports, such as journal publications of clinical trials, and long reports, such as clinical study reports, are generally both structured similarly to IMRAD.

- **Investigator’s brochure.** A document written by a sponsor and intended for clinical investigators interested in becoming involved in a study. It summarises the current body of evidence about an intervention under investigation, typically based on preclinical and early phase human studies. The document is periodically updated in light of new information.
  - Example: [Rituximab Investigator’s Brochure](#)

- **Licence/License.** The formal permission a regulator grants a company to market a medical product in a given territory.

- **Marketing Authorization Application (MAA).** The complete dossier of information submitted to the European Medicines Agency when sponsors seek marketing authorization for a medicine throughout the European Union under the EMA’s centralised procedure.

- **Marketing Authorisation Holder (MAH).** (Also often referred to as a ‘sponsor’ or ‘manufacturer’.) The entity granted marketing rights for a given medicine in a given jurisdiction. The EMA uses the term MAH whereas the FDA uses the term “sponsor”.

- **Medical Officer Review.** Also known as a “Clinical review” or “Medical review”. As one part of the FDA’s process for evaluating marketing applications for new medicines (e.g. a New Drug Application or Biologics License Application), a medical officer, usually a physician, performs a review of and prepares a report regarding the clinical aspects of the application. These reports generally contain a listing of clinical studies included in the application and information about the design and results of those trials, including analyses conducted independently by the medical officer and additional commentary. Medical officer reviews are made public under FOIA and are posted to the FDA’s website (for drugs, under the Drugs@FDA database and for biologics, by searching the “Vaccines, Blood & Biologics” section of the FDA’s website).
  - Example: [Gardasil clinical review](#)
  - Example: [bevacizumab clinical review part 1](#)
  - Also see an example in the Screenshots Gallery

- **Medical Officer Report.** (Also referred to as a Medical Officer Review) See Medical Officer Review.

- **Medical Review.** See Medical Officer Review.

- **MedWatch.** The FDA safety information and adverse event reporting program. MedWatch encompasses both materials FDA makes available to the public and healthcare professionals as well as the three pharmacovigilance systems [FDA Adverse Event Reporting System (FAERS)], [Vaccine Adverse Event Reporting System (VAERS)], and [Manufacturer and User Facility Device Experience Database (MAUDE)].
  - Also see new web-based [FAERS Public Dashboard](#)

- **Module 2.5 (Clinical Overview) & Module 2.7 (Clinical Summary).** Common Technical Document module section 2.5 contains the Clinical Overview which is an accurate and exhaustive description of the evidence development plan. It contains the product development rationale, overview of biopharmaceutics, clinical pharmacology, efficacy, safety, benefit/risk conclusions, and literature references. The evidence development plan lists the completed, ongoing and planned studies by their study ID (which may or may not correspond to a register identifier). It is an invaluable overview and is relatively short (around 30 pages), and is complemented by Module 2.7, the Clinical Summary, which provides more detail of the same

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data (in around 50 to 400 pages), including the Summary of Clinical Efficacy (section 2.7.3) and Summary of Clinical Safety (section 2.7.4).

- **Module 5. Clinical Study Reports** and raw data (for regulators that require or request it) are included in Module 5 of the Common Technical Document. There is no space limitation for Module 5.

- **Narratives.** See Serious Adverse Event narratives.

- **New Drug Application (NDA).** According to the FDA, “The NDA application is the vehicle through which drug sponsors formally propose that the FDA approve a new pharmaceutical for sale and marketing in the U.S.” The application should contain sufficient information for the FDA to make a marketing decision. “The documentation required in an NDA is supposed to tell the drug’s whole story, including what happened during the clinical tests, what the ingredients of the drug are, the results of the animal studies, how the drug behaves in the body, and how it is manufactured, processed and packaged.” Compare with Marketing Authorisation Applications in the European Union. Also see Biologic License Application.

- **Patient Information Leaflet (PIL).** (Similar documents in the United States are called the ‘patient package insert’ or ‘medication guide’.) A document, typically a few pages in length, containing written medical information for patients that accompanies approved medicines. Information contained in the leaflet include active ingredient and indication, contraindications, warning and precautions, dosage and administration, possible side effects, storage of the medicine, marketing authorisation holder and manufacturer. Many forms of written medical information are reviewed and approved by regulators. However in the United States, patients may also receive a pharmacy leaflet when picking up a prescription medication. These documents are not regulator approved, but contain information that is similar in scope to official information and is written by third party vendors (not the manufacturer).
  - Example: Xydalba (Dalbavancin) PDF page 22
  - Also see an example in the Screenshots Gallery

- **Periodic Safety Update Report (PSUR).** EMA required pharmacovigilance document prepared by the marketing authorisation holder to provide an up to date evaluation of the benefit-risk-balance of a medicine. PSURs describe the worldwide safety experience with a medicine at a defined time after its authorization. Summarized data on the benefit-risk of a medicine and results of all studies of the medicine, authorised and unauthorised uses, are included.

- **Pharmacovigilance.** (Also often referred to as “drug safety”). The science and activities relating to the collection, detection, assessment, monitoring and prevention of adverse effects of medicines. Also see MedWatch.

- **Protocol.** Generally refers to a Study Protocol document, but may also refer to a Study ID.

- **Regulatory document.** Colloquial term that generally refers to any document produced by, or held by, a regulatory agency. This may therefore include documents produced by regulators, such as an FDA Medical Officer Review, or a document submitted by a sponsor to a regulator, such as a Clinical Study Report.


- **Serious Adverse Event narratives.** Clinical Study Reports contain individual participant narratives of serious adverse events (ICH E3 section 12.3.2). They consist of unstructured free text and summarize information relevant to the serious adverse event. Each individual narrative is typically a paragraph to a page long.
  - Example: See PDF p.276 onwards of paroxetine study 329

- **Statistical Analysis Plan (SAP).** (Also known as a Reporting Analysis Plan.) Similar to a study protocol but focusing on the statistical methods and definitions to be used for data analysis. Like a study protocol any planned or actual changes from the original written SAP should be justified and documented with formal SAP amendments. The amendments should be dated.

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- **Statistical Officer Review.** Report similar to a Medical Officer Review, but written by an FDA statistician. The statistical review may include the statistician’s independent analyses using IPD submitted by the sponsor. Available as part of the Drug Approval Package on the Drugs@FDA website.
  - Also see an example in the Screenshots Gallery
- **Statistical Report.** See Statistical Officer Review.
- **Statistical Review.** See Statistical Officer Review.
- **Study ID.** (Sometimes referred to as Study Number. Pharmaceutical companies also often refer to a Study ID as a Protocol.) Identifier given to a single trial by its sponsor. Each trial may have multiple identifiers. For example: GlaxoSmithKline’s HPV vaccine study 580299/012 (GSK’s own ID) is also known as HPV-012 (part of the Cervarix programme) and NCT00169494 (registration ID), and may also be referred to by four known publications of the trial. The IDs may not be immediately reconcilable.
- **Study Protocol.** (Disambiguation: Protocol.) A document, written prospectively before recruiting participants into a study, which records the general rules and intended methods of conducting, analysing, and reporting the study. Detailed statistical methods are often recorded in a separate statistical analysis plan document, but the protocol should include the sample size calculation and an overview of the planned statistical analyses. Clinical trial protocols can be tens to hundreds of pages in length. A protocol may be required by the research ethics board, a data and safety monitoring board, or a funding body. Any planned or actual changes from the original written protocol in the conduct and/or analysis should be documented with formal protocol amendments.
  - Example: GSK Paroxetine Study 329 Protocol
  - Also see an example in the Screenshots Gallery
- **Summary Basis of Approval.** A document, according to the FDA, that contains “a summary of the safety and effectiveness data and information evaluated by FDA during the drug approval process.” (21CFR314.430) Sponsors may draft the Summary Basis of Approval; it may also be written by FDA scientists.
- **Summary of Product Characteristics (SmPC).** Term used in the European Union to refer to the drug label. The SmPC describes the properties and the officially approved conditions of use of a medicine, and is intended for use by healthcare professionals. The SmPC is part of the European Public Assessment Report (EPAR).
  - See an example in the Screenshots Gallery
- **Trial Master File (TMF).** A TMF is the collection of documentation that allows the conduct of the clinical trial, the integrity of the trial data and the compliance of the trial with Good Clinical Practice to be evaluated. It is also essential to allow the trial to be effectively managed by the sponsor as it allows the appropriate individuals access to the necessary trial documentation. The documentation contained within the TMF should be sufficient to adequately reconstruct the trial activities undertaken, along with key decisions made concerning the trial. Consideration should be given to the TMF being a stand-alone set of documentation that does not require additional explanation from the associated sponsor or site staff.
References


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32. Coyne DW. The health-related quality of life was not improved by targeting higher hemoglobin in the


42. All Trials Registered | All Results Reported [Internet]. AllTrials. 2017 [cited 2017 Oct 3]. Available from: http://www.alltrials.net/