PERSPECTIVE

Value-based evidence across health care sectors: a push for transparent real-world studies, data, and evidence dissemination

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Abstract
There is currently a heightened need for transparency in pharmaceutical sectors. The inclusion of real-world (RW) evidence, in addition to clinical trial evidence, in decision-making processes, was an important step forward toward a more inclusive established value proposition. This advance has introduced new transparency challenges. Increasing transparency is a critical step toward accelerating improvement in type, quality, and access to data, regardless of whether these originate from clinical trials or from RW studies. However, so far, advances in transparency have been relatively restricted to clinical trials, and there remains a lack of similar expectations or standards of transparency concerning the generation and reporting of RW data. This perspective paper aims to highlight the need for transparency concerning RW studies, data, and evidence across health care sectors, to identify areas for improvement, and provide concrete recommendations and practices for the future. Specific issues are discussed from different stakeholder perspectives, culminating in recommended actions, from individual stakeholder perspectives, for improved RW study, data, and evidence transparency. Furthermore, a list of potential guidelines for consideration by stakeholders is proposed. While recommendations from different stakeholder perspectives are made, true transparency in the processes involved in the generation, reporting, and use of RW evidence will require a concerted effort from all stakeholders across health care sectors.

Keywords: Health care system; peer review; real-world evidence; transparency; value assessment; value communication

1. Introduction
Across health care sectors, there is an accelerating recognition of the need for concrete action to secure a transparent flow of information on treatment effectiveness and value, given the specific and unique requirements for regulatory approval and, ultimately, access to new medications and health care technologies (Ross et al., 2012; Fraser et al., 2018). In parallel, the information available to inform health care decisions – especially decisions optimized for the individual – is expanding rapidly, and interest in the use of evidence beyond clinical trials has been growing for many years in step with technological and methodological advances. Besides patients and their caregivers, members of the general public are also entitled to transparency in health care decision-making. The inclusion of real-world (RW) evidence has been an important step forward in moving toward a more inclusive established value proposition (Dhruva et al., 2018). However, this advance has brought with it new transparency challenges.

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Society only truly benefits if the health care sector is universally transparent in inputs, processes, and outcomes, and integrates the participation and viewpoints of all key stakeholders (Henke et al., 2011). Transparency of methods, assumptions, and data is fundamental to improving the credibility and relevance of efforts to identify and pay for the highest-value health care. In turn, transparency increases the accountability of all stakeholders and constituencies regarding their actions, ensures visibility about the roles of stakeholders in decision-making processes, increases the awareness of patients and caregivers that the decisions being made reflect their preferences and priorities, and allows decision-makers to assess both the rigor and relevance of the evidence on which they base their choices in resource allocation (Henke et al., 2011; Leviton and Melichar, 2016). Transparency, in itself, does not equate to quality, but rather allows for the evaluation of quality. The achievement of true transparency in the processes involved in the generation, reporting, and use of RW evidence will require an intensification of the ongoing efforts from all stakeholders across health care sectors, taking concrete action to improve procedures and processes.

This policy perspective paper aims to highlight the need for wide-ranging transparency concerning RW studies, data, and evidence (see Figure 1) across the health care sector, and to identify areas for improvement and provide concrete recommendations and practices to help bring about substantive change for the future.

2. Methods
An initial virtual meeting was hosted to discuss aspects of transparency surrounding RW studies, data, and evidence dissemination. The participants consisted of a small but diverse panel of six experts, who could draw on specific expertise and experience, with backgrounds in relevant fields and professional organizations, including International Society for Medical Publication Professionals (ISMPP), global pharmaceutical companies, academia with health policy expertise, medical communications agencies, and a value framework agency. The participants agreed to participate in a roundtable discussion and act as authors of a subsequent manuscript. The roundtable consisted of two informal but structured virtual events during which participants discussed transparency issues from their individual perspectives and drawing on their professional experience. The discussions focused on identifying themes, deficiencies, and potential solutions, aiming to arrive at clear recommendations for improvements moving forward. Ideas generated during the first discussion were collected and presented to the participants prior to the second discussion, during which they were further refined. Further discussion and refinement of ideas occurred by email at various stages during the drafting of this manuscript.

3. Results from the roundtable discussions
There are many concepts and definitions of what constitutes ‘RW studies’, ‘RW data’, and ‘RW evidence’. The authors feel that an initial step toward improving transparency would be a general acceptance of clearly defined terms. We suggest that a ‘RW study’ is an initiative to prospectively collect data on patients in a RW setting, or to retrospectively gather data from databases or health records that have been collected in a RW setting. ‘RW data’ are data originating from a RW study. ‘RW evidence’ is information from RW studies that has either been publicly communicated or assessed for relevance and quality (by peer review). These definitions are summarized in Table 1.

In the following sections, specific issues concerning RW study, data, and evidence transparency are discussed. Recommended actions for improved transparency are discussed, and those recommendations together with associated key stakeholders are presented in Table 2. As an aid to achieving enhanced RW data transparency among stakeholders, a number of guidelines and standards could be useful for consistency. Potential guidelines for consideration by governing societies (some already under consideration, and some new) are given in Table 3.
4. Health care in the real world and the need for transparency

Historically, health care decisions were primarily based on clinician expert opinion and were heavily reliant upon clinical trial data (Umscheid et al., 2011). Bias in the selective reporting of clinical trials, concealing relevant information from patients and health care providers, led to a call to action for clinical trial submission requirements. Publishers introduced ICMJE author-ship guidance (ICMJE 2018), the CONSORT (Schulz et al., 2010) and SPIRIT (Chan et al., 2013) guidelines, and other guidelines included in the EQUATOR network (2020). This resulted in obligatory registration of clinical trials and reporting of trial results (EU 536/2014, 2020; FDAAA 801 and the Final Rule, 2020), one of the first major advances aiming for greater transparency across health care sectors.

A driving force in the development of alternative research designs has been the need to generate evidence with greater external validity to complement the internal validity of RCT data.
Additionally, clinical trials are expensive and lengthy, tend to be relatively small and lacking in diversity, are difficult if not impossible to perform for ultra-rare disorders, and are sometimes precluded by ethical issues, such as the need for a cancer trial without a control arm or with a synthetic control arm. In response, RW studies were introduced, and they have been evolving in quality, availability, and size, as well as methods for data analysis (Berger et al., 2017; Khosla et al., 2018).

The combination of increasing health care costs, the drive toward patient centricity, and a growing interest in the effectiveness of treatments in RW settings have also contributed to the need for supplemental forms of data beyond clinical trials in relevant patient settings (Califf et al., 2016; Sherman et al., 2016; FDA, 2019). Decision-making should be based on a synthesis of data on RW effectiveness and clinical trial efficacy, combining external and internal

Table 1. Definitions of RW study, data, and evidence

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
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<tr>
<td>RW study</td>
<td>Designed to prospectively collect data on RW patients or retrospectively draw on existing RW databases and health records</td>
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<tr>
<td>RW data</td>
<td>Data originating from RW studies</td>
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<tr>
<td>RW evidence</td>
<td>RW studies that have either been publicly communicated or assessed for relevance and quality (peer review)</td>
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RW, real-world.

Table 2. Recommended actions for improved RW study, data, and evidence transparency and responsibilities of stakeholders

Pharmaceutical and biotechnology company responsibilities

- Transparency of engagement processes that include patient/caregiver experience and expertise in defining real-world outcomes of interest. Use of RW and appropriate/relevant PRO measures to allow assessment of evidence quality
- Transparent design and conduct of RW studies, through hosting of protocols and results on an online platform similar to requirements for clinical trials

Researchers, publishers, and professional society responsibilities

- Submission of transparent RW manuscripts by all researchers for peer review
- Investment by journals in understanding RW studies and endorsement of the peer review and publishing of such studies, and assurance by publishers that peer review includes reviewers with real-world study experience (at a scientific, clinical, and patient expertise level)
- Development and implementation of good RW communications practice by publishers and professional societies, through the development of specific RW-communication checklists (similar to the CONSORT checklist) including adherence to guidelines for conducting studies and reporting data, the use of study registration numbers, and access to study protocols
- Communication of evidence through peer-reviewed publications involving the cooperation of publishers, scientists, and societies
- Provision of no-cost, open-access, lay-audience summaries to enable patients, caregivers, and the general public to properly assess evidence

Medical evaluation and decision-maker responsibilities

- Implementation of solid methodology and good assessment practices to allow for proper evaluation of RW evidence
- Engagement with the patient community to inform the decision-making process

CONSORT, Consolidated Standards of Reporting Trials; PRO, patient-reported outcome; RW, real-world.
validity. Although different RW data sources have different strengths and limitations, the selection of appropriate RW data sources should be performed in a way that provides an appropriate level of validity, reliability, and transparency. So far, tangible improvements in transparency have been relatively restricted to clinical trials, and real progress in advancing standards of transparency concerning the generation and reporting of RW data has lagged behind (Loder et al., 2010; Szkultecka-Dębek and Drozd, 2015; International Society for Pharmacoeconomics and Outcomes Research, 2017; ISPE-ISPOR Special Task Force, 2020). Transparency surrounding RW study design and reporting of RW data is not regulated, and there remains a lack of clear guidance on best practices. Furthermore, there is not yet a consensus on appropriate uses of RW data, nor is there a system to evaluate the scientific rigor of RW evidence on a par with that of clinical evidence (e.g., GRADE) (Dhruva et al., 2018; Malone et al., 2018; Miksd and Abernethy, 2018).

Increasing transparency is a critical step toward accelerating improvement in type, quality, and access to data, regardless of whether those data originate from clinical trials or from RW studies (Henke et al., 2011). The imperative to improve transparency, if RW evidence is to be trusted and optimally utilized in decision-making processes, has been recognized by the ISPOR-led Real World Evidence Transparency Initiative (Orsini et al., 2020). Short-, mid-, and long-term recommendations have been made concerning RW secondary data studies, including the identification of the best site to register studies using secondary data, determination of a ‘good’ registration process, and provision of incentives for routine registration of studies (Orsini et al., 2020). For such calls for increased transparency concerning RW studies to result in real change, a concerted effort will be required from all associated stakeholders within the health care sector.

Fortunately, clearer guidance on conducting, reporting, and using RW data for regulatory purposes has been proposed by governmental bodies (Perfetto et al., 2015; US Government Information, 2016; Gabay, 2017; FDA, 2018), and professional societies, such as ISPOR (Berger et al., 2017; Orsini et al., 2020) and CONSORT (Husereau et al., 2013; Calvert et al., 2018).

Transparency in generating data and communicating robust evidence beyond clinical trials, across health care decision-making sectors, is an overall goal for the future. Throughout the health care sector, the drive toward this goal requires adaptation of current practices and careful considerations and collaborations with all stakeholders involved. Collectively, all health care sectors must work to overcome the challenges that prevent transparency (Figure 1).

Table 3. Summary of potential guidelines for consideration by stakeholders

<table>
<thead>
<tr>
<th>RW study registration and publication requirements (good real-world evidence practices)</th>
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<tr>
<td>• RW outcomes and patient registry protocol registration</td>
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<tr>
<td>• Identification of the most appropriate repository agency (or agencies) for registration</td>
</tr>
<tr>
<td>• Publicly available RW study outcomes and results disclosure</td>
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<tr>
<td>RW study reporting (good value-communication practices)</td>
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<tr>
<td>• Standardization of reporting RW studies, CONSORT-like adaptation</td>
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<tr>
<td>• ICMJE endorsement of RW study registration prior to publication</td>
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<tr>
<td>• Encouragement of open-access availability of publications</td>
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<tr>
<td>Proper assessment of RW studies (good value-assessment practices)</td>
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<tr>
<td>• Implementation of transparent assessment practices and methodology</td>
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<tr>
<td>• Open access to assessment methodology and results for appraisal, including list of identified studies, quality and bias assessment</td>
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5. Recommendations from the roundtable discussions

5.1. Transparency of engagement processes that include patient/caregiver experience and expertise in defining RW outcomes of interest; use of RW and appropriate/relevant PRO measures to allow assessment of evidence quality

Generating data is a key step in gathering information on the clinical value of a treatment. Often pharmaceutical companies collaborate closely with clinical institutes or contract research organizations to develop clinical trials. While this process is transparent, it is mainly focused on meeting primary efficacy and safety objectives and endpoints. Regulatory bodies, such as the European Medicines Agency (EMA) and US Food and Drug Administration (FDA), have acknowledged that a deeper integration of the patient and caregiver experience is important and that integration of clinical outcome assessments [including patient-reported outcomes (PROs)] in clinical trial programs broadens the available evidence and can better describe the impact of interventions on the patient experience (FDA, 2018; European Medicine Agency, 2020). This guidance primarily focuses on the integration of PROs in a controlled clinical trial setting. However, health care trends, combined with increasing evidence linking the patient experience to clinical outcomes, are driving the need to integrate the patient/caregiver experience into drug development programs beyond the clinical trial setting. As the number of RW studies, including RW PRO studies, has been increasing over the last decade, the medical industry should actively engage with the move toward more transparent design and conduct of RW patient-orientated studies. Governmental and independent initiatives, such as the CAHPS surveys from AHRQ, and research supported by PCORI, do strengthen the validity of using RW patient-orientated outcomes (AHRQ CHAPS, 2020; Patient-Centered Outcomes Research Institute, 2020a). In addition, comparative-effectiveness research (CER) to assess relative effectiveness, using clinical endpoints that matter to the relevant (subset of) patients is becoming increasingly important to stakeholders such as health technology assessment (HTA) and value framework (VF) bodies and payers (Berger et al., 2017; Law et al., 2018), but is not yet a regulatory requirement of the FDA or the EMA (Patient-Centered Outcomes Research Institute, 2020b).

Additionally, although the patient perspective is becoming increasingly important, interpretation of results, and communication of findings, guidelines, and expectations for how the patient perspective should be incorporated into clinical research are currently lacking (Dhruva et al., 2018; Hoddinott et al., 2018; UK Standards for Public Involvement, 2020).

5.2. Transparent design and conduct of RW studies, through hosting of protocols and results on an online platform similar to requirements for clinical trials

The number of RW study registrations is increasing; however, many researchers currently do not provide RW study registration as there is no requirement for protocol registration. A key issue that drove the active implementation of clinical trial registration was the endorsement by the International Committee of Medical Journal Editors (ICMJE) of the CONSORT guidelines for reporting controlled trials as one of the requirements for publication of results in high-tier journals (DeAngelis et al., 2004). Subsequently, the FDA Amendments Act (FDAAA) of 2007, as well as agencies such as the EMA, required registration of summaries of trial protocols for ‘applicable clinical trials’ (FDA, 2007; EU 536/2014, 2020). Another important transparency measure was introduced through the requirement to report results within one year of completion of the clinical trial (with some provisions for delayed reporting) (Wood, 2009; Commission Guideline, 2012).

The lack of RW study protocol registration and reporting of results can potentially lead to significant bias in reporting positive/selective results, as studies that do not produce the expected data will probably not be completed or submitted for peer review. RW studies, regardless of the origin of RW data, need to be registered in a manner equivalent to that of clinical trials. It is, however, not always possible to fit RW study parameters, which include medical claims
data, EHR data, product and disease registries, patient-generated data (including from in-home-use settings), and data gathered from other sources such as mobile devices, into a CONSORT-style guideline. Consequently, as these CONSORT-style guidelines and officially recognized registration platforms with design features specific for RW studies to provide appropriate study registration information are not yet in place, a concerted effort will be needed with researchers registering RW studies, societies/publishers developing and endorsing RW publication guidelines and requirements, and governments signing measures into law. Additionally, a summary of the results should be made available, within a set time frame of 1 year (or 6 months for pediatric studies), as is now obligatory for clinical trials. Furthermore, there is often confusion concerning the definitions of ‘exploratory’ research vs confirmatory hypothesis testing, which can also contribute to a failure to report RW data, and can generate bias. With obligatory registration of RW study protocols and reporting of results, RW data would become more transparent and assessable (FDA, 2018; Katkade et al., 2018). The pharmaceutical industry, RW outcomes researchers, RW data database agencies, and patient advocacy groups, need to be prepared if study registration and reporting becomes mandatory in the future.

5.3. Submission of transparent RW manuscripts by all researchers for peer review

Various methodology guidelines are available for submission of RW data for peer review, such as CHEERS for health economic evaluations, and STROBE for observational (RW) studies, but these guidelines are not suitable for every type of RW study (e.g., claims database studies, patient surveys, etc.), and they have a predominant focus on reporting of the study methodology alone. Regardless, we encourage submission of RW studies that adhere to the most relevant methodology guidelines, where available. Besides adhering to methodology guidelines, in the absence of official mandatory registration and reporting guidelines, it is recommended to register RW studies on existing platforms such as clinicaltrials.gov, to at least future-proof the publication for uptake in later value-based decisions.

Integrity, accountability, and responsibility for accurate, complete, and transparent reporting are key. The introduction of GPP and subsequent further development of GPP2, and GPP3 guidelines, as directed by the ISMPP, provided recommendations for individuals and organizations that contribute to the publication of research results sponsored or supported by the pharmaceutical industry, and has led to a more responsible and ethical manner of submitting trials and studies for peer review (Battisti et al., 2015). Nonetheless, there is still a lack of trust in the reporting of industry-sponsored studies (Fisher, 2008), which might only be expected to grow with sponsored RW studies. The challenge of achieving credibility for industry-sponsored research is heightened in the communication of RW evidence where there is often an inherent perception of bias and a corresponding lack of guidelines to effectively mitigate this. Thus, clear, transparent submission of data is warranted (Khosla et al., 2018).

5.4. Investment by journals in understanding RW studies and endorsement of the peer review and publishing of such studies, and assurance by publishers that peer review includes reviewers with RW study experience (at a scientific, clinical, and patient-expertise level)

Reporting through peer review is considered to be the gold standard to communicate outcomes and is regarded as a credible mechanism for assessing the quality and trustworthiness of research (Mayden, 2012). Most high-tier journals have robust peer-review processes in place with publication guidelines; these are, however, mostly focused on controlled clinical trials and lack clear guidance on RW study reporting. Moreover, existing RW reporting guidelines are not fully endorsed by most medical journals, thereby enabling the submission of less transparent RW studies for peer review. In some instances, even reputable peer-reviewed journals may struggle to find
appropriate peer reviewers, or may render a decision of non-acceptance due to evaluating RW studies through a clinical trial lens.

A report presenting the point of view of journal editors themselves identified some practical barriers, including the large volume of often confidential patient data and the lack of robust computational models with which to analyze it (Oehrlein et al., 2018). There may be restrictions in making the data and algorithms from RW publications publicly available. Additionally, identifying peer reviewers who have familiarity with RW data sources, as well as the specific analytic skills to evaluate the data, is a very real difficulty for journals. In an on-line article published by ESMO, Javier Carmona Sanz, Deputy Editor at Nature Medicine suggests that, “a set of ‘best practices’ for submitting RW [evidence], endorsed by both editors and researchers, to safeguard the confidentiality of patients’ data while also encouraging transparency about computational models, may facilitate the proliferation of these studies in the scientific literature” (ESMO, 2021).

With RW evidence becoming increasingly important throughout the health care sector, it is thus of great importance for journals to acknowledge the need to publish high-quality and transparent RW studies, but also to grow their expertise in allowing for appropriate peer review.

5.5. Provision of no-cost, open-access, lay-audience summaries to enable patients, caregivers, and the general public to properly assess evidence

Patient-centeredness and patient engagement with health care stakeholders has become an important part of health care decision-making. The paradigm has shifted from the more exclusive provision of information from a health care provider to a patient, toward a more collaborative relationship in which patients and their caregivers are involved and consulted with regards to health care decision-making processes. One way to increase the transparent transfer of information is for journals to intensify efforts to provide open access to publications together with the provision of freely accessible, researcher-provided, lay summaries of published research, especially with regards to RW studies, which may involve medications that are currently available to patients, rather than drugs being evaluated for clinical efficacy and safety.

5.6. Implementation of solid methodology and good assessment practices to allow for proper evaluation of RW evidence

Health care decision makers are involved at every level of the provision of care, whether it be governmentally/federally driven, state/regionally driven, or locally driven. The assessment of evidence and assessment of the value of a drug are key steps for health care organizations, whether at the approval level to assess the efficacy and safety of a drug, the appraisal of a drug with regard to comparative clinical effectiveness and cost-effectiveness (Figure 2), or by independent or commercial formulary review boards. Ultimately, these steps are required to allow access to a drug based on provided evidence (Rovira, 2008; Akehurst et al., 2017; EPHA, 2018; Paschke et al., 2018).

In parallel, the demand for transparent economic modeling is increasing (Eddy et al., 2012; Cohen et al., 2017; Cohen and Wong, 2017; Chapman and Kumar, 2019; Hay, 2019; Incerti et al., 2019; ICER, 2020). HTA and VF agencies require that models should be able to accommodate RW data in addition to randomized controlled trial data; however, established methods are not well suited to this. Current methods merely enable the narrative inclusion of qualitative aspects of the patient experience, but there is growing attention for the development of new methods to facilitate quantification (e.g., the potential of multi-criteria decision analysis) and incorporate alternative stakeholder perspectives (Leviton and Melichar, 2016; Hoddinott et al., 2018).

Sound clinical, access, and reimbursement decisions rely on proper appraisals, which include inputs from HTA and VF agencies. Many such agencies [e.g., National Institute for Health and Care Excellence (NICE), Haute Autorité de Santé (HAS), Canadian Agency for Drugs and
Technology in Health (CADTH), etc. are government-driven, as they are ultimately responsible for making decisions for an entire country. The US health care system is decentralized, and although assessment of value is becoming more commonplace, it is often performed by independent agencies (ICER, IVI, etc.) or societies [National Comprehensive Cancer Network (NCCN), American Heart Association (AHA), etc.] (Wilke et al., 2018) or through commercial health care plan review boards and local (hospital) formulary committees.

Approval and appraisal decisions are based on provided evidence, either through information provided by the drug manufacturer or through independent research assessment. HTA and VF assessment of a drug takes place at market approval, but ideally would take place later to allow for evaluation of RW evidence. Nonetheless, evidence synthesis methodology to allow for proper assessment of the evidence, whether from clinical trials or RW studies is widely variable among agencies (Desai et al., 2020), and reporting of methods is not uniform. Furthermore, although some agencies, such as ICER and NCCN, provide transparent assessment methodology, it is often unclear what the impact of RW evidence is on the overall assessment process. Clear guidance from those involved is warranted.

5.7. Engagement with the patient community to inform the decision-making process

The integration of RW and patient-centered outcomes is playing an increasingly important role in enabling better selection of treatments for those populations who benefit the most (and that show a favorable cost profile). This is reflected in the calls from leadership sectors, such as the FDA guidance on the use of RW evidence (FDA, 2019), and the white paper published by the Duke-Margolis Center for Health Policy exploring how and when studies producing RW evidence can inform FDA regulatory decisions concerning the effectiveness of a drug (Duke Margolis Center for Health Policy, 2020b).

With the integration of RW and patient-centered outcomes in the appraisal of drugs, questions arise regarding how these types of evidence weigh in the decision-making process. Stakeholders should shift their engagement from a more informing type of engagement toward a more collaborative involvement of the patient at every step of the process, and information provided should allow for patient evaluation of the evidence (EUPATI, 2016).
6. Conclusions

In conclusion, a truly transparent health care sector will only evolve if all stakeholders increase their efforts and bring about tangible changes resulting in greater transparency surrounding the generation and reporting of RW studies. Trust in the system can only be established if all evidence is generated meaningfully, reported uniformly, processed equally, and discussed openly. Ultimately, data and communication transparency will leverage the true value of a treatment or service for the benefit of the patient and society.

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