World Health Statistics 2020 has indicated that, although both life expectancy and healthy life expectancy have increased 8% between 2000 and 2016, non-communicable diseases, which account for 71% of all global deaths, remain a considerable health burden (1). As an example, breast cancer, the most common cause of cancer deaths among females in 2017 (2), was also responsible for an estimated 626,679 deaths globally in 2018 (3). Owing to contributions from researchers, health-care practitioners (HCPs), and stakeholders, significant progress has been achieved in breast cancer prevention and treatment. Critical to this success has been the use of randomized controlled trials (RCTs), which are considered to be at the pinnacle of the medical evidence pyramid. However, the uncertain external validity of RCTs and the extensive costs of conducting them have become a significant concern.

Due to these issues, real-world study (RWS), being more economically feasible and possessing greater external validity, has received increasing attention in the past decade. Still, many misunderstandings and uncertainties concerning RWS, including the opportunities it offers or its capacity to prove high-quality of evidence, remain unresolved. The current article aims to discuss and clarify definitions relevant to RWS, opportunities RWS brings, and ways to ensure high-quality evidence.

What is real-world study?

The “real world” concept is not a novel innovation, and its origins can be traced back to over 50 years ago. Many do not appreciate this relatively long history, as “real world” only began to receive more attention two decades ago, and its true value has only been appreciated in recent years. When using “(real world[Title]) OR (real-world[Title])” to search in the PubMed (search date: June 10th, 2020), we found 9,031 articles published from 1966 to 2020. The number of articles has increased year by year and surged to hundreds per year beginning in 2005 and thousands per year since 2017.

Speaking RWS, the two concepts of real-world data (RWD) and real-world evidence (RWE) should be clarified together. Table 1 summarizes the most authoritative definitions and classifications (4-7). Among them, we consider the definitions of the Food and Drug Administration (FDA) as appropriate and accurate. According to the FDA, RWS refers to a study design that includes, but is not limited to, randomized and non-randomized trials (such as pragmatic clinical trials and large simple trials) in addition to observational studies; RWD is “data relating to patient health status and the delivery of healthcare routinely collected from a variety of sources,
including registries, collections of EHRs (electronic health records), administrative and medical claims databases etc.”; RWE means “clinical evidence about the usage and potential benefits or risks of a medical product derived from analysis of RWD” (4). As described, the definitions of these three concepts refer to different contents: study design, data, and evidence. However, these concepts could be easily misunderstood or misinterpreted. In one study, the authors reviewed 53 documents and conducted 20 interviews with stakeholders. They found that 53% of definitions of RWD were classified as data collected in a non-RCT setting, 24% were data collected in a non-controlled or non-interventional setting, 13% were data collected in a non-experimental setting, and 11% were defined as something else (8).

It needs to be particularly emphasized and clarified that RWS is an umbrella term. RWS include studies with both randomized and non-randomized designs. In addition, the criteria for identifying RWS goes beyond its design or methodology, and also considers the use of RWD in a real-world implementation scenario (4-6). A common misunderstanding of RWS and RCTs is that RCTs do not reflect real-world settings and that all observational studies are real world (8). In fact, RCTs may include real-world settings, such as pragmatic trials, and observational studies, such as those with intensified follow-up, may not be situated in real-world care settings (7). Therefore, it is inappropriate to completely separate RWS from RCTs. Similar to conducting pragmatic trials, RWS could adapt tools and methods of traditional trials and apply these to real-world settings by selecting appropriate analysis methods and designs like prospective plans and randomization.

**What opportunities could real-world study bring?**

RWS not only offers benefits of potential economic feasibility and greater external validity but also provides precious opportunities for optimizing evidence generation and verification.

First, RWS can be used for post market surveillance to further confirm the efficacy and safety of the approved interventions based on pivotal RCTs (9). For instance, many of the recent cancer drugs investigated in pivotal RCTs have been approved more quickly than before, due to the Accelerated Approval or Designated Breakthrough programs and through the use of surrogate endpoints instead of the traditional gold standard overall survival (OS), or both (10-13). However, the association of treatment effects between surrogate endpoints and OS has always shown low or modest correlation (14), surrogate endpoints are still

<table>
<thead>
<tr>
<th>Source</th>
<th>RWD</th>
<th>RWE</th>
<th>RWS</th>
</tr>
</thead>
<tbody>
<tr>
<td>FDA 2018 (4)</td>
<td>Data relating to patient health status and/or the delivery of health care routinely collected from a variety of sources</td>
<td>Clinical evidence about the usage and potential benefits or risks of a medical product derived from analysis of RWD</td>
<td>RDs, NRDs, OS</td>
</tr>
<tr>
<td>Sherman et al. 2017 (5)</td>
<td>Electronic health records, claims, billing data, product and disease registries, data gathered through personal devices and health applications</td>
<td>Information on health care that is derived from multiple sources outside typical clinical research settings</td>
<td>OS, IS</td>
</tr>
<tr>
<td>CTONG 2018 (6)</td>
<td>Data generated by routine healthcare</td>
<td>Evidence produced from the RWD after strict data collection, systematic processing, correct statistical analysis, and multi-dimensional interpretation of results</td>
<td>PCT, CR, CSS, NCCS, CS</td>
</tr>
<tr>
<td>CEMC 2018 (7)</td>
<td>Data derived from a wide range of sources, such as routine healthcare, traditional epidemiological studies, surveillance, administrative databases, or personal devices</td>
<td>A spectrum of studies that apply various epidemiological methods to data collected from real-world settings</td>
<td>PCT, OSR, OSP</td>
</tr>
</tbody>
</table>

RWD, real-world data; RWE, real-world evidence; RWS, real-world study; RDs, randomized designs using real-world data; NRDs, non-randomized, single-arm trials with external real-world data control; OS, observational studies; IS, interventional studies with or without randomization; PCT, pragmatic clinical trials; CR, case report; CSS, cross-sectional studies; NCCS, nested case-control studies; CS, cohort studies; OSR, observational studies using existing administrative data; OSP, observational studies involving prospective data collection; FDA, U.S. Food and Drug Administration; CTONG, Chinese Thoracic Oncology Group; CEMC, Chinese Evidence-based Medicine Center.
widely used in confirmatory trials (15), and the majority of approved oncology drugs eventually do not demonstrate desirable clinical benefit after several years of marketing authorization (16,17). The above situations make RWS become rational as confirmation study to further evaluate the efficacy and safety of the approved interventions based on pivotal RCTs. In addition, this use of RWS is further bolstered by its proven ability to replicate clinical trials (18). Moreover, with larger sample size RWS supplies sufficient statistical power to investigate the long-term impact of rare or suboptimal outcomes in pivotal RCTs, including adverse effects, patient-reported outcomes, and quality of life. According to these outcomes, RWS can simultaneously support post market digital pharmacovigilance, especially drug safety surveillance (19). Taken together, the above demonstrate the potential value of RWS in post market confirmation of approved interventions.

Conducting RWS also allows researchers, HCPs and stakeholders to investigate the effectiveness of interventions in the clinical practice population, including in excluded/under-represented subgroups or those with different molecular characteristics not investigated in previous RCTs (20,21). Similar research can be conducted in different geographic or economic contexts to verify the efficacy/effectiveness of the interventions among the specific population. Based on these efforts, cost-effectiveness studies tailored to the given geographic or economic contexts can be conducted, providing evidence and indications to optimize the allocation of resources in health-care services (21).

By using data linkage technology based on the datasets from health records (in primary and subsequent care), cancer registries, insurance claims, etc. (22,23), conducting longitudinal studies of patients’ cancer experiences becomes feasible. The investigations can examine any given point along the treatment pathway or any segment across the cancer continuum, including primary prevention (e.g., tobacco control and cessation, risk/preventive factors to the risk of cancer), secondary prevention and diagnosis (e.g., cancer screening, novel diagnostic techniques/strategies), treatments (e.g., advanced surgical strategies, radiotherapies, targeted therapies, immunotherapies), recovery or survivorship (e.g., prevention and early detection of recurrence, metastasis, or secondary cancer), and end-of-life care (e.g., palliative care). The vast quantity of these linked datasets can provide ample patient information to answer a diversity of research questions related to general practice, surgery, oncology, epidemiology, health services research, health policy, health economics, and even social science (6,9,19,20,22,23).

**How can high-quality evidence from RWS be ensured?**

Although RWS offers considerable research opportunities, whether it has played an essential role in decision-making is still controversial (24). The main reason for this doubt is that the nature of RWS precludes it from having the similarly rigorous study design and methodology as RCTs enjoy—rigor that can ensure the enhancement of internal validity. Accordingly, the strength of evidence from RWS has becomes a primary concern.

These issues related to study design and methodology are directly reflected in the current evidence-based system. Since the mid-19th century, evidence-based medicine has been the cornerstone in medical research (25), and, in the medical evidence pyramid, RWS (mainly observational studies) sit lower than RCTs. This judgement is reflected in the Grading of Recommendations Assessment, Development and Evaluation (GRADE) guidelines (26), a widely recognized approach to evidence grading. According to the guidelines, the quality of evidence is classified into four levels: high, moderate, low, and very low quality. Normally, the evidence from observational studies (like cohort studies and case-control studies, most common study design of RWS) are classified as low quality; however, the credibility of the evidence could be increased with the inclusions of a dose-response relation, plausible bias control, or other measures. On the contrary, the evidence from RCTs are generally classified as high quality, but its credibility can be reduced in cases of inconsistent results or reporting bias (26).

Apart from study design and methodology, the concerns in regard to the evidence from RWS also involve reporting. In academia, standards for creating transparent reports are a near-constant consideration. It is thus crucial to follow the principles established in the GRADE guidelines (26), the Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) statement (27), the extension of the CONSORT statement (28), and more relevant guidelines on the EQUATOR network (29). However, the quality of reporting in published studies is not always up to standard. For instance, studies which investigated the quality of reporting of observational studies after the introduction of the STROBE statement found that the
quality in these studies could still be improved (30,31).

Therefore, the guaranteeing a high quality of evidence from RWS is essential. In fact, quality control is critical in each stage of the research lifecycle. Specifically, it should be emphasized that conducting valid RWS requires data quality assurance, proper methodology design, and good reporting. Regarding data quality, prospective data collection of RWE should be foundational to preserving internal validity (19), especially for population-level studies. In addition, the items in datasets are subject to the design of the record systems, and the recorded contents are subject to the accuracy and completeness provided by clinical practitioners and patients. In terms of population-level research, the contents are also subject to the accuracy and completeness of data linkage and the feasibility of the linkage. Because of these limitations, vigilance, and administrative, financial, and technical support, are particularly appreciated and warranted to assure data quality. Also, proper study design and methodology are crucial.

RWS researchers encounter the risk of potential biases caused by confounders throughout the study and methodology design process (9). This risk can be mitigated through approaches logic-based [e.g., using pragmatic clinical trials (32), directed acyclic graphs (DAG) (33), stratification or matching (9)] to statistical (e.g., multivariable regression models (9), propensity-score matching (34)] (19), which can augment internal validity and external validity and better ensure generalizability.

As for reporting, it is strongly recommended that future RWS complies with the currently established principles from the guidelines/statements mentioned above. Adherence to these principles is required not only from article authors but from editors and reviewers as well. Indeed, more studies that investigate the quality of published RWS are expected (30,31,35), along with studies and explanations that educate article authors, editors, and reviewers on how to comply with quality of study design, methodology, and reporting guidelines/statements.

**Summary**

In summary, RWS is a type of study that is conducted through using RWD in a real-world implementation scenario. Its study design includes not only observational studies but also randomized and non-randomized clinical trials. RWS offers researchers and stakeholders unprecedented opportunities for optimizing clinical evidence generation and verification. To maximize the potential value of RWS, the quality of evidence is the key. Ensuring reliable data quality, proper research design and methodology, and standardized and transparent reporting all help increase the strength of evidence from RWS.

**Acknowledgments**

Funding: None.

**Footnote**

Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at http://dx.doi.org/10.21037/tbcr-20-30). KZ and DW declare that they are full-time employees of AME Publishing Company (publisher of *Translational Breast Cancer Research*). JZ is the section editor of *Annals of Translational Medicine*, *Annals of Cancer Epidemiology*, and *Journal of Hospital Management and Health Policy*, which are all managed by AME Publishing Company.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

Open Access Statement: This is an Open Access article distributed in accordance with the Creative Commons Attribution-NonCommercial-NoDerivs 4.0 International License (CC BY-NC-ND 4.0), which permits the non-commercial replication and distribution of the article with the strict proviso that no changes or edits are made and the original work is properly cited (including links to both the formal publication through the relevant DOI and the license). See: https://creativecommons.org/licenses/by-nc-nd/4.0/.

**References**

2. GBD Compare. Available online: https://vizhub.healthdata.org/gbd-compare/
3. GCO CANCER TODAY. Available online: https://gco.iarc.fr/today
4. FDA. Framework for FDA’s real-world evidence program.
Available online: https://www.fda.gov/media/120060/download
32. Groenwold RHH, Dekkers OM. Designing pragmatic
trials—what can we learn from lessons learned? J Clin Epidemiol 2017;90:3-5.
34. Reiffel JA. Propensity Score Matching: The 'Devil is in the Details' Where More May Be Hidden than You Know.

doi: 10.21037/tbcr-20-30