REVIEW ARTICLE

Demonstrating Value for Biosimilars: A Conceptual Framework

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BACKGROUND: The value proposition for biosimilars can be characterized as a concept that moves beyond the argument of cost reduction relative to the innovator biologic drug and into a framework that incorporates the diverse needs of key healthcare stakeholders during the transition from clinical development to commercialization in the marketplace.

OBJECTIVES: To identify factors that facilitate and inhibit the development, commercialization, and adoption of biosimilars, and to recommend modifications in program design that are likely to support the demonstration of the value of biosimilars for payers, providers, and patients.

METHODS: The primary data sources for this article include surveys conducted by Boston Healthcare Associates with payers and clinicians in the United States and the European Union 5 markets and blinded international protocol feasibility assessments completed by Worldwide Clinical Trials. Survey methodology used either convenience or purposeful sampling as appropriate, with participants extracted from diverse audiences, representative of those who generate or evaluate clinical data shaping the economic exchange and preferential status influencing physician adoption and patient access to biosimilars. Patient characteristics and psychosocial issues influencing patients’ perception of small-molecule generics were extracted from the available literature to inform exploratory hypotheses, given the relative absence of such information for biosimilars.

DISCUSSION: This article reviews the current evidence and summarizes results of surveys conducted with payers, providers, and drug investigation sites in the United States. Based on a review of published literature, as well as these survey results, conflicting and convergent demands exist for gathering data related to biosimilars. The motivations and data needs for these new agents are diverse, requiring adjudication of regulatory, economic, and clinical incentives beginning at program inception and extending through commercialization of the final biosimilar agent.

CONCLUSIONS: The development and commercialization of biosimilars represent an international activity that can encounter unanticipated challenges, as well as opportunities to achieve clinical and commercial success. Evolving regulatory guidance mapped in relation to payer, physician, and patient sentiments may inform the biosimilar development program designs, implementation, and positioning of the new drug.

KEY WORDS: biosimilars, biologics, reference drug, drug development, value proposition, generic drugs, payers, providers, patients, regulatory guidance, biosimilar adoption and commercialization

Although a range of regulatory definitions exist, a biosimilar drug generally is defined as a biological compound that is highly similar to the reference drug, with no clinically meaningful differences in safety, purity, and potency. In addition, biosimilars can be characterized by a value proposition centered on reducing healthcare costs while maintaining clinical efficacy and safety outcomes similar to the originator biologic. These objectives become particularly laudable for patient populations receiving biologic agents to treat chronic or life-threatening conditions.

In this article, the value proposition for biosimilars is characterized as one that moves beyond the cost reduction argument appropriately encountered for small-molecule generic drugs and into a framework that is more nuanced, incorporating the perspective of regulators, physicians, patients, and payers into an overall statement of value.

Using a nonprobability-based survey sampling from 17
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In March 2015, the FDA approved the first biosimilar in the United States.

As the US market for biosimilars transitions from clinical development to commercialization, multiple stakeholders who influence formulary placement, reimbursement, and the adoption of a biosimilar will shape the value proposition of biosimilars.

This article presents survey data from payers, providers, and international drug investigation sites showing that conflicting and convergent demands exist for biosimilars.

The authors build on and extend the current literature on biosimilars’ market entry, arguing that payers expect biosimilars to induce price competition leading to potential positive economic returns.

Addressing the barriers and challenges related to payers, providers, and patients during clinical trial development will help to ensure a successful adoption and commercialization of biosimilars into the US market.

Although the impetus for a biosimilar development originally might have been solely economic, the authors argue that manufacturers should devise a value proposition for biosimilar compounds that moves beyond price, demonstrating value to payers, physicians, and patients.

KEY POINTS

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- There are significant international differences in the experience related to the development and clinical use of biosimilar drugs.
- In March 2015, the FDA approved the first biosimilar in the United States.
- As the US market for biosimilars transitions from clinical development to commercialization, multiple stakeholders who influence formulary placement, reimbursement, and the adoption of a biosimilar will shape the value proposition of biosimilars.
- This article presents survey data from payers, providers, and international drug investigation sites showing that conflicting and convergent demands exist for biosimilars.
- The authors build on and extend the current literature on biosimilars’ market entry, arguing that payers expect biosimilars to induce price competition leading to potential positive economic returns.
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payers (convenience sample in the United States [N = 7] and in the European Union [N = 10]), 50 practicing physicians (convenience sample within the United States [N = 15] and in the European Union [N = 35]), and 91 international investigative sites (purposive sample), as well as a review of the available published evidence, we developed a framework for assessing value for biosimilars.

These data provide milestones to guide critical development and commercialization decisions related to biosimilars.

An Evolving, Complex Environment

There are significant international differences between countries in the experience associated with the development and clinical use of biosimilars. This is particularly true when contrasting the European Union with the United States, where the regulatory climate is expected to change dramatically after the US Food and Drug Administration (FDA)’s approval of the first biosimilar, and the finalization of “interchangeability guidance.” These events have paved the way for the implementation of the Biologics Price Competition and Innovation Act of 2009.

To date, most of the experience with biosimilars is limited to 3 therapeutic classes—granulocyte colony-stimulating factors (G-CSFs), epoetins, and human growth hormones. Biosimilars in these therapeutic classes have been marketed in Europe since 2005 through the European Medicines Agency (EMA) regulatory framework (Figure 1).

In September 2014, the first biosimilar monoclonal antibody, infliximab (Inflectra), was approved in the European Union. On March 6, 2015, the FDA approved the first biosimilar in the United States—the G-CSF agent filgrastim-sndz (Zarxio).

Across the 3 major classes of biosimilars so far, historical data suggest that biosimilar penetration varies widely across biosimilar classes, with G-CSF biosimilars achieving an average market share of 42% in the European Union 5 (EU5) markets in 2011—that is twice the average market share of erythropoietin class and 4 times of the human growth hormone class. However, the differences in the adoption of biosimilars between countries are marked (Figure 2), suggesting a mosaic of different regional incentives, which are mirrored during the clinical development program leading to the regulatory approval of the drug.

As the US market for biosimilars transitions from clinical development to commercialization, multiple stakeholders who influence formulary placement, reimbursement, and, ultimately, the adoption of a biosimilar, will shape the value demonstration process, which must occur during clinical development, given the pharmaceutical industry incentives and bargaining power. Healthcare industry analysts forecast the US biosimilars market to grow up to $25 billion by 2020.

As in small-molecule generic markets, the growth of the biosimilars market is fueled by a series of patent expirations, such as in the case for blockbuster biologics in oncology, immunology, and inflammatory diseases, including rituximab (Rituxan), cetuximab (Erbitux), trastuzumab (Herceptin), and infliximab (Remicade), which will lose patent protection in the next 3 to 5 years. In 2015, branded biologics, specifically monoclonal antibodies, may generate $60 billion in revenue.

Because the adoption of any novel therapeutic agent reflects the interplay of the opinions of diverse stakeholders, the value proposition of a biosimilar—moving beyond cost reduction—should be payer-, physician-, and patient-centric. The key adoption factors, by stakeholder, are identified in Table 1 and are discussed below.
Value Framework for Biosimilars:
A Payer-Centric Perspective

In the US healthcare market, biosimilar drug manufacturers are likely to face significant challenges during the commercialization process. In contrast to small-molecule generic drugs, few drug manufacturers possess the complex research and development capabilities to advance a biosimilar to market; therefore, it is unlikely that the same competitive dynamics will exist as has been observed in the generic, small-molecule drug market. Considerable barriers, such as biologics' manufacturing capabilities (although drug developers may use contract manufacturing organizations to circumvent this problem), extend into the need for a more extensive, and, therefore, lengthier and more costly, clinical testing program, which effectively limits competition.

Nevertheless, experience to date with the commercialization efforts for biosimilars within the United States (payer and clinician survey conducted by Boston Healthcare Associates) suggests that drug manufacturers are challenged to devise strategies demonstrating the value of biosimilars moving beyond a narrow value proposition based on reducing direct healthcare costs through price competition.

The reasons for this apparently counterintuitive position—in which less competition may warrant a more complicated demonstration of value—are manifold and include expectations regarding the level of discounting in comparison with the originator drug once the biosimilar is commercialized. For example, although US payers may recognize that research and development costs for biosimilars are multiples of the costs for small-molecule generics, and therefore should command higher acquisition costs, significant price discounts to the originator drugs may be anticipated as a spillover effect, based on the experience with generic, small-molecule compound drugs.

Commentary from a convenience sample of US commercial payers, obtained by informal interviews conducted in 2014 by Boston Healthcare Associates, suggests...
that a discount of 20% to 50% from the originator drug would be necessary to give the biosimilar preferential formulary placement status. Arguing further, payers in US and EU5 markets suggest that in the absence of a significant price discount, preference will be given to the reference biologic given existing contractual/pricing arrangements—demonstrating payers’ higher price sensitivity at biosimilar launch.

Biosimilar drug manufacturers may benefit from a clinically driven value proposition by demonstrating their commitment to improving patient outcomes and engaging with key opinion leaders to address current unmet needs. These data can be generated during the clinical development process.

Payers may nevertheless still negotiate on price, demanding a significant price discount over the branded biologics. However, developing data to support concepts that address current unmet needs are likely to allow favorable comparison of the biosimilar to the branded biologic agent. These data also may incentivize the adoption of a biosimilar among skeptical physicians who are concerned with immunogenicity and variability of efficacy in the absence of data.

In some markets, such as Italy, in which price is negotiated at the national level (ie, through the Agenzia Italiana del Farmaco), a differentiating concept based on clinically driven value proposition could support financial decision makers’ acceptability of the biosimilar drug in innovative pricing schemes, including payment for clinical outcomes. According to internal interviews (conducted by Boston Healthcare Associates) with members of the Agenzia Italiana del Farmaco, differentiation is particularly important, because in most markets, competition with biosimilars will result in a winner-takes-all market through tendering (ie, a competitive bidding process resulting in a sole-source contract at a contractually agreed price for a specified time frame with regional, local payers, such as sickness funds, or hospital administrators).

Table 2 provides a summary of all the elements that likely impact a manufacturer’s ability to realize economic returns for biosimilar development.

### Value Framework for Biosimilars: A Physician-Centric Perspective

Our experience in conducting clinical studies on biosimilar drugs, as well as feasibility assessments for the design of a biosimilar clinical development program, confirm that the interest of physicians in biosimilars is influenced by diverse factors, including the accessibility of the branded (reference) biologic agents; competing clinical trials for innovator drugs and other biosimilars; the level of scientific novelty engendered by the proposed clinical program; and the changing landscape of clinical care.

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**Figure 2**  
Biosimilar Adoption Differs Across European Union Markets

<table>
<thead>
<tr>
<th>Country</th>
<th>G-CSF</th>
<th>EPO</th>
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<tbody>
<tr>
<td>Austria</td>
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<tr>
<td>France</td>
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<tr>
<td>Germany</td>
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<td>80</td>
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<tr>
<td>Sweden</td>
<td>10</td>
<td>90</td>
</tr>
<tr>
<td>United Kingdom</td>
<td>50</td>
<td>50</td>
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</table>

EPO indicates epoetin; G-CSF, granulocyte colony-stimulating factor.

**Table 1**  
Key Biosimilars Adoption Factors, by Stakeholder

<table>
<thead>
<tr>
<th>Stakeholder</th>
<th>Key adoption factors</th>
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</thead>
</table>
| Payers           | • General perception on biosimilars—acceptability of clinical data package used for regulatory approval  
                  • Pricing power—ability to induce price competition, for example, through tendering processes and need for pricing negotiations at the national, regional, and local level  
                  • In the United States, regulation of therapeutic interchange and automatic substitution is controlled by state pharmacy boards and state laws, which may vary between states  
| Physicians       | • Noninferior versus equivalent better clinical outcomes to the originator drug; concerns highlighted include extrapolation of clinical data into other indications or to patients with different characteristics, variability of efficacy (batch-to-batch), immunogenicity, other safety concerns  
                  • Experience in switching patients from reference drug to biosimilars  
                  • Uncertainty in biosimilar performance resulting from shifts in standards of care that were used to evaluate the utility of the reference product  
                  • Absence of tools to assess clinical value in individual patients after biosimilar commercialization  
| Patients         | • Concern over manufacturers’ know-how and manufacturer capabilities, especially for brand-loyal patients  
                  • Uncertainty regarding longer-term efficacy and safety outcomes  
                  • Inability to distinguish and interpret evolving concepts, such as interchangeable biologic drugs  

Accessibility of Branded Biologic Agents

Because branded biologic agents are generally associated with high cost, access and affordability vary greatly in different countries or regions of the world. Indeed, in prestudy assessments informing the operational footprint required for a proposed clinical development program, the level of interest expressed by physicians regarding participation in biosimilar trials is inversely associated with the affordability and availability of branded biologic agents. In addition, because of the perceived lack of benefit to patients in countries where branded biologic agents are available and accessible, the level of interest for physicians to participate in trials of biosimilars is usually much lower, as is routinely observed in the United States and in Western European countries.

In contrast, the interest of physicians in countries with limited access to expensive biologic agents can be apprecia-
Innovation
Physicians’ willingness to participate in clinical trials for biosimilars, and their success in recruiting patients, also can be driven by factors generally subsumed under the umbrella of professional satisfaction. In addition to providing patients with access to medications that are not usually available or affordable in their institution, and the financial incentives associated with study participation, other factors influencing decisions to participate in biosimilar trials include scientific interest; the possibility of defining improvements in other aspects of patient care; the need to be referenced in peer-reviewed quality publications for career advancement; and the prestige and publicity afforded for the individual or the institution as a result of participation in a biosimilar research program.

Subsequently, given the perception of a lack of innovation for biosimilar drugs, and the limited opportunities to publish on innovative research, many physicians often decline to participate in biosimilar studies, removing an invaluable center of influence for the transition from branded biologic to a biosimilar drug during the commercialization period. This is especially relevant within academic research centers in the United States and in Western Europe, where the need for professional and institutional recognition is marked.

For example, in the biosimilar feasibility assessment conducted by WCT, although academic and private practices (a total of 37 sites) were approached in the United States, no academic center and only 3 private practices responded favorably.

Changing Standards of Care
Based on the WCT survey experience that includes a purposeful sampling frame in which potential respondents are selected according to the diversity of location and practice demographics, a clinical study supporting commercialization regardless of the therapeutic class, is more acceptable to physicians if its design is closely aligned with the local standard of care. Although this concept is not unique to clinical investigations for biosimilars, it is accentuated by the lag time between the introduction of the innovator compound and the clinical development of a biosimilar, during which the standard of care may evolve.

This can be particularly notable in therapeutic areas with rapidly evolving standards of care, such as oncology and immunology and/or inflammatory disease. Consequently, physicians and other scientists may regard as unacceptable a study design mandating adherence to the original treatment paradigm used for regulatory approval for the branded drug.

As an example, significant regional differences in standard of care from site to site and from country to country have been noted in most feasibility assessments, particularly for indications within oncology for small molecules and biologics. The original regimen of docetaxel and doxorubicin used in the phase 3 registration studies for pegfilgrastim is no longer used as a neoadjuvant or adjuvant treatment for early-stage breast cancer. The National Comprehensive Cancer Network–preferred regimen, AC followed by paclitaxel, is the most widely used chemotherapy regimen.

Changing standards of clinical care also present a conundrum for the developers of biosimilars. If the original treatment paradigm is mandated within the study design for a new biosimilar, patient accrual rates for the proposed study may falter, because it is not aligned with the local practice, and the physicians may be reluctant to randomize otherwise acceptable patients to an investigational study or to subsequently transition from the reference drug to a biosimilar once it has been commercialized.

However, if the options within the treatment protocol acknowledge the evolving clinical care climate, there may be uncertainties regarding the anticipated effects for the branded drug, thus impacting the sample size required for the study. Finally, patients and ethics committees may question the justification of administering a treatment that may be either suboptimal or associated
with greater side effects solely on the prospect of a reduced cost of treatment for other patients who are not included within the current protocol, and possibly not within the country where the study has been conducted.

**Uncertain Clinical Utility After Approval of a Biosimilar**

After the approval of a biosimilar, physician uptake can be limited by factors that were embedded into the design of the study registration program. These factors are manifold and include perceived clinical differences across study designs using noninferiority, equivalency, or superiority hypotheses for comparisons between the biosimilar and the originator drug.

Concerns also may exist regarding the extrapolation of study data into clinical care because of variations from batch to batch in the biological properties of the drugs, or differences in patient characteristics or in standard of care from that permitted within studies evaluating the originator drug and the biosimilar. Unanticipated long-term safety concerns, such as immunogenicity, may be voiced regarding adverse events of clinical interest that could not be demonstrated in the trial’s duration that would otherwise be acceptable for regulatory approval.

Finally, the lack of real-world experience with switching strategies from innovator drugs to comparator (biosimilar) drugs introduces hesitancy into adoption of a biosimilar. For example, to gather real-world data on switching from an innovator drug to a comparator drug, Norway’s government is conducting the NOR-SWITCH Study to evaluate the safety and efficacy of switching from the innovator monoclonal antibody Remicade to its biosimilar Remsima in patients with rheumatoid arthritis, spondyloarthritis, psoriatic arthritis, ulcerative colitis, Crohn’s disease, and chronic plaque psoriasis.

The availability of a therapeutic monitoring tool that would enable a physician to determine clinical utility for an individual patient—rather than extracting guidance from group data obtained within a study—could obviate the need for this type of investigation.

**Value Framework for Biosimilars: A Patient-Centric Perspective**

Despite a wealth of clinical and scientific literature, regulatory documents, and expert opinion on the development of biosimilars, only recently have patient-related perspectives for this most important topic been addressed. Given the paucity of published literature in this area, factors dictating the perceived value of a biosimilar from a patient’s perspective are regarded as indeterminate; however, they may reflect the nature of clinical efficacy or safety measurements used during the development of a biosimilar, as well as difficulties in understanding the implications of a drug characterized as a biosimilar versus a drug characterized as a fully interchangeable biologic drug.

In contrast to biosimilar drugs, a wealth of published studies has described the variables that influence patient perspectives regarding the use of generic, small-molecule drugs. These data provide an informative framework for hypothesis generation for biosimilars in a postmarketing setting. For example, in national surveys in the United States, Japan, Australia, Portugal, and Malaysia, patients agree that generic drugs are less expensive and have a better value than brand-name drugs; however, the same patients are not eager to use generic drugs personally. The main factor associated with patients’ willingness to accept a generic drug substitution was identified as correct understanding of the characteristics of the generic drug relative to the brand drug after a detailed discussion of the drug’s attributes with the prescribing physician.

One of the few data points on this issue is provided through a recent survey of 3214 patients with type 1 or type 2 diabetes. The survey posed an open-ended follow-up question that addressed patients within the sample who said they would “definitely not use” biosimilars or were “unlikely” to use biosimilars (4% and 13%, respectively, of the sample) to provide a reason for their reluctance. The respondents mentioned the proved track record of brand-name insulin and the lack of such a record with biosimilars, their current personal satisfaction with a particular insulin, their past bad experiences with other types of generic medications, a lack of trust in generic medications in general and in biosimilars in particular, and allergic reactions to various forms of insulin. One respondent’s answer was particularly enlightening, stating, “It is not Humalog. I know how my body acts with Humalog. I do not trust things I do not know when it comes to my health.”

These concepts that are well-documented for small-molecule generic drugs prompt systematic inquiry for all biosimilars that are undergoing development. This sentiment suggests that because of side-effect concerns regarding biosimilars and the maintenance of adequate response, patient education will be crucial to secure a biosimilar acceptance developed in the context of its clinical trials or after its commercialization. Programs to ensure patient education on the use of biosimilars can serve as supportive activity for the clinical trial registration program of a drug.

**Perceived Asymmetries in Outcomes**

Patient-perceived differences in efficacy or safety may exist during the development of, or the commercialization process for, biosimilar drugs that are comparable with experiences with generic, small-molecule drugs. For
example, an investigational program for the biosimilar
filgrastim may be adequately characterized from a regula-
tory perspective based on a limited clinical program, in-
cluding pharmacokinetic and pharmacodynamic studies
in healthy volunteers, with one comparative study in-
volving patients with similar pharmacokinetic and phar-
macodynamic outcomes, followed by postmarketing sur-
veillance through the use of a patient registry.23 This is a
methodologically appropriate program. A patient’s deci-
sion to participate in the development of a biosimilar
would be framed in the context of short-term supportive
care, given the end point of neutropenia, and the easily
measurable cases of severe neutropenia that may occur
after a well-established chemotherapy regimen.23

By contrast, in studies of biosimilar monoclonal anti-
bodies in oncology, the use of a proxy for overall survival
(i.e., clinical end points) rather than overall survival itself
may be perceived as problematic by a patient, even if
fully acceptable from a regulatory perspective.24,25 These
end points speak to fundamental drug attributes that in-
fluence disease progression and morbidity, and can there-
fore weigh heavily on a patient’s decision to accept ex-
posure to a biosimilar rather than a branded biologic,
either as part of the development program or after the
drug’s commercialization.

In addition, the potential for long-term safety out-
comes that cannot be measured in short-term studies
become more clinically consequential and differentially
impact the informed consent process, either for trial
participation or for a switch in therapy. In conclusion,
although an acceptable risk for novel, interventional
therapy exists, the potential lack of clinical equivalency
between the reference biologic and the biosimilar jeop-
ardizes patient interest in a trial of an alternative drug if
a reference medication is commercially available and
accessible to that patient.

**Biosimilar, or Interchangeable Biologic Drug?**

Given the potential for differences in efficacy or safety,
characterization as either a “biosimilar” or an “inter-
changeable biologic drug” may obscure more than inform
the biosimilar adoption process. This is understandable,
given that even regulatory agencies use various terms to
define the characteristics of a biosimilar. For example,
under 351(k) of the Public Health Service Act, an “inter-
changeable” biologic drug has a more comprehensive
definition than a drug that has been shown to be biosim-
ilar to the reference drug: it can be expected to produce
the same clinical result as the reference drug across a
spectrum of various clinical applications.1

Because the difference between a biosimilar and an
interchangeable biologic drug may be difficult to appre-
ciate even for healthcare professionals, patients attempt-
ing to render an informed consent before randomization
in a clinical trial, or to engage in a new treatment option
suggested by a provider, are at a disadvantage.26

**A Unifying Concept**

Diverse stakeholders create a mosaic of conflicting
and compatible demands for clinical trial data to inform
the approval, commercialization, and adoption of bio-
similars. A fully integrated development program maxi-
mizing the value proposition of a biosimilar must ac-
knowledge all perspectives, and can be illustrated by
development of a biosimilar for an extensively used
monoclonal antibody, rituximab (Rituxan; MabThera).

Rituximab (a chimeric anti-CD20 monoclonal anti-
body) is indicated for several conditions, including
non-Hodgkin lymphoma, chronic lymphocytic leuke-
mia, rheumatoid arthritis, and severe granulomatosis
with polyangiitis. Rituximab’s largest revenue source is
attributed to non-Hodgkin lymphoma, yet biosimilar
comparability for approval purposes will pursue the most
efficient pathway to drug approval. As with small mole-
cules, this may be demonstrated in the most sensitive
and easily accessible patient population (such as in pa-
tients with rheumatoid arthritis) rather than in patients
representing all approved indications.

Characterized by the early engagement of key opin-
ion leaders and network organizations, patient recruit-
ment for the study would utilize emerging markets for
faster clinical trial completion resulting from differenc-
es in access to biologics, and include a planned post-
approval publication strategy for participating centers
highlighting the attributes of the new biosimilar during
clinical use, as well as a patient education program to
facilitate adoption.

In addition, to optimize the international regulatory
strategy during the development of a biosimilar, a step-
wise approach cited by predominant regulatory authori-
ties, such as the FDA and the EMA, would be used, in
which preclinical comparability was confirmed by stan-
dard parameters provided by regulatory guidance (e.g.,
state-of-the-art structural and analytic characterization,
functional characterization, pharmacology and toxicolo-
gy studies), followed by a clinical pharmacokinetic and
pharmacodynamic study in healthy volunteers (where
permitted) to demonstrate expected correlations. The
biosimilar development program would be concluded
through the incorporation of a clinical study with pa-
tients using either an equivalency or noninferiority hy-
pothesis (as appropriate) with a postapproval FDA Risk
Evaluation and Mitigation Strategies program and/or an
EMA’s Risk Management Plan strategy.

The efficiency of this stratagem would therefore be
dictated before the clinical program begins through a
regulatory quality comparability gap analysis. These analyses determine optimal countries and satisfaction of regulatory criteria, with staggered clinical trial initiation across countries to permit supplemental pharmacokinetic information on additional regulatory queries. Therefore, variables such as regulatory acceptance, a competitive environment, access to a relevant patient population, and operational knowledge of clinical centers would all be factored into consideration.

**Conclusion**

Although the impetus for a biosimilar development originally might have been economic, the value proposition for biosimilars can be enhanced by moving beyond cost reduction arguments that are often encountered for small-molecule generic drugs and into a framework that incorporates the regulatory, professional, and psychosocial concerns of diverse stakeholders. This process begins by acknowledging the marketing dynamics, evolving regulatory guidance, and the realities of the current environment for the clinical evaluation and commercialization of biosimilars in comparison with reference biologic agents.

Program strategies are diverse, including attempts to accommodate evolving standards of care into protocol design for drug registration trials; companion efforts addressing professional satisfaction during study participation; and the creation of abbreviated therapeutic monitoring strategies to facilitate the adoption of the biosimilar after its approval. For chronic illnesses characterized by a potential for significant morbidity as a consequence of the illness or as a reflection of treatment failure, the development of patient-specific outcome measures and a companion educational platform for the introduction of the drug are particularly important.

**Author Disclosure Statement**

The authors have no conflicts of interest to report.

**References**


**Stakeholder Perspective next page**
Challenges Surrounding the New Biosimilars Landscape

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The framework for understanding the evolution of a biosimilar marketplace outlined in the article by Dr Rompas and colleagues’ includes many of the most important biosimilar influencers but fails to reflect on the very competitive and economically constrained environment for biosimilar manufacturers. The key reasons for this are the lack of clarity and predictability regarding regulatory requirements, and uncertainty about the competitive biosimilar landscape. The authors highlight the long-term features of the stakeholder landscape that must be resolved to develop a robust biosimilar market, but fail to chart an evolutionary path to a new state, which is the fundamental challenge ahead.

The authors’ avoidance of a purely cost-centered value concept is important, because manufacturers cannot be expected to drive substantial cost-savings in the near-term. Ultimately, predictability around costs, risks, and regulatory requirements will encourage greater participation in biosimilar development and will support the increasing price competition. This, however, leaves out the continuing innovation by originator drug makers, which continues to address unmet clinical needs, and, as a consequence, supersedes earlier-generation biologics that are now, or will soon be open to biosimilar competition.

The broadly held belief that biosimilars will evolve to form a “generic biologic” market may be true in the long-term, but divergent stakeholder needs and perspectives indicate that generic biologic will not mean “vastly cheaper,” and will not entirely replace biologic drug originators within 6 to 8 weeks after patent expiration, as seen with traditional generics. Without recognition that the healthcare system will be paying high prices for biologics for the foreseeable future, why is this biosimilar concept the best approach? A greater focus on the personalization of care (ie, getting the right drug to the right patient at the right time) may address spending pressures more appropriately than a long-delayed effort to drive the adoption of biosimilars. It is possible that we should do both, but we should at least be clear about the savings that are possible.

**ORIGINATOR BIOLOGIC DRUG MAKERS:** The investment to develop an originator biologic drug is as substantial as in small-molecule innovations, and the long-term return on investment is dependent on the length of the patent protection. The end of a drug’s commercial life is not linked only to its patent life, because newer-generation medicines often supersede older drugs before patent expiration. All this has been part of the pattern of commercializing biologics for 20 years before the emergence of biosimilars.

For the vast majority of biologics, a biosimilar pathway offers a unique commercial threat. The likelihood of investing in a biosimilar for a 3rd to 7th biologic in a class is extremely low; therefore, the challenge for biologic originators can be split into those that will be reference brands for a biosimilar and the ones that will be part of a next tier of therapeutic options that may not be directly affected by biosimilars. If costs are not substantially lower for biosimilars, the economic factors that drive therapeutic choices may not materialize. Thus, many originator biologic makers may be most concerned with traditional market issues. The key will be the type of reimbursement offered for new biosimilars, and whether there are changes to reimbursement for originator drugs (whether they are reference drugs or not).

**BIOSIMILAR MANUFACTURERS:** The uncertainties about market share capture for a biosimilar manufacturer are a key driver of early trends in marketing investments and price setting.

In Europe, biosimilars have most often captured volumes of the molecule equivalent to small-molecule specialty injectable drugs in the same markets. That achievement is mostly because of the reimbursement and tendering processes that are in place for hospital-administered drugs, which include most of the biosimilars to date. As biosimilars begin to be marketed in the United States, it is important to note that although filgrastim biosimilars achieved a 60% to 70% volume share in Germany and in some other European markets, the US share for the biologic tbo-filgrastim (Granix) has achieved a 10% share after 1 year. This disconnect raises a note of caution and will likely result in a strategic choice to either escalate the commercial efforts to drive greater biosimilar uptake or to a more cautious level of restraint in the choice of molecules and commercial targets. Already fewer drug manufacturers are targeting biosimilars than when the US Biologics Price Competition and Innovation...
tion Act of 2009 was passed, and ultimately price competition can only be effective with more players.

**REGULATORS:** The goals of safe, effective, and sufficiently numerous biosimilars to drive economic savings in the healthcare system as a whole is at least partly at the root of the US biosimilars pathway legislation. That it took 5 years for the first biosimilar approval to make its way through the pathway is a serious issue, and the reasons why a company may choose to file an application under a 505(b)(2) or a biologic pathway seriously hamper the rapid evolution of the biosimilar pathway. The next catalyst event will likely be if the biosimilar version of filgrastim, which was approved in 2015, does vastly better in terms of volume uptake than the biologic license application–approved version of the originator drug launched in 2013. Whether greater incentives are needed to use one pathway or another remains to be seen.

**PATIENTS:** Depending on the therapy, patients’ perspectives on biosimilars may differ vastly. With some therapies, such as insulin, patients’ share of the cost, and their perception of the drug’s similarity, may make the choice a “no brainer.” In other cases, where the physician or the health system sets the protocols, the patient may not have a specific choice, and in still others, the patient may be reluctant to use a biosimilar.

Reluctance may come from concerns about similarity, or the price of the biosimilar may not be substantially lower. As with many cost issues in the US healthcare system, the patient’s perspective can be seen as an afterthought.

**PAYERS:** Although the US healthcare system retains medical and pharmacy benefit silos, payers’ ability to influence biosimilar utilization will remain bifurcated. The evolving incentive structures regarding accountable care organizations and bundled payments may encourage greater biosimilar use in institutional- or provider-administered treatments, but these structures are still only applicable to 33% of the US population. The earliest opportunities to influence controls over biologics have been related to administered treatments where payers have historically had the least direct influence. As their influence grows in these areas, and as new biosimilars begin to target self-administered treatments, the influence of payers on the adoption and pricing of biosimilars will most likely grow.

**PROVIDERS:** The earliest biosimilars, as seen in Europe, have demonstrated a decade of cost-savings, as well as remarkable safety. Many discussions about biosimilars have focused, at least in part, on similarity, which is, at minimum, an oblique reference to the show-stopping issue that many people fear. Providers have a strong interest in whether their patients can afford the treatments they prescribe. So, in some sense, they will look to a potential cost-savings with hope. As noted earlier, the lack of substantial cost-savings, the