The role of biosimilars in value-based oncology care

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Abstract: Biopharmaceuticals (biologics) represent one of the fastest growing sectors of cancer treatment. They are recommended for treating underlying cancer and as supportive care for management of treatment side effects. Given the high costs of cancer care and the need to balance health care provision and associated budgets, patient access and value are the subject of discussion and debate in the USA and globally. As the costs of biologics are high, biosimilars offer the potential of greater choice and value, increased patient access to treatment, and the potential for improved outcomes. Value-based care aims to improve the quality of care, while containing costs. The Centers for Medicare & Medicaid Services (CMS) has developed value-based care programs as alternatives to fee-for-service reimbursement, including in oncology, that reward health care providers with incentive payments for improving the quality of care they provide. It is anticipated that CMS payments in oncology care will be increasingly tied to measured performance. This review provides an overview of value-based care models in oncology with a focus on CMS programs and discusses the contribution of biosimilars to CMS value-based care objectives. Biosimilars may provide an important tool for providers participating in value-based care initiatives, resulting in cost savings and efficiencies in the delivery of high-value care through expanded use of biologic treatment and supportive care agents during episodes of cancer care.

Keywords: biologics, biosimilars, oncology, patient access, value-based care, supportive care

Introduction

As of the early 1980s, biopharmaceuticals represent one of the fastest growing sectors of the drug industry worldwide¹ and are increasingly important in cancer care. Biologics (e.g., monoclonal antibodies [mAbs] and hematopoietic agents)² are recommended in oncology guidelines³ for treating underlying disease as well as for managing treatment side effects through supportive care agents such as granulocyte-colony stimulating factors (G-CSFs) and erythropoiesis-stimulating agents (ESAs).⁴

Biologics are produced from cells of living organisms and purified in complex, multi-step processes, including recombinant DNA technology, controlled gene expression, or antibody technologies.² Compared with chemically synthesized small molecule drugs, biologics have 100- to 1000-fold larger molecular weight and are relatively heterogeneous⁵; their physiochemical structure is complex and difficult to characterize. Furthermore, they are highly sensitive to changes in manufacturing conditions, and as a result, no two biological products can be identical,⁶ resulting in a complex production process. Biologic agents, including those used in cancer treatment and...
supportive care, have improved outcomes for patients, who often require ongoing treatment. As costs of biologics are high, long-term treatment of patients with biologics can be a chronic burden to health care systems.7

Biosimilars of reference biologic agents offer an alternative choice and value that has potential to open further patient access to treatment and associated outcomes. According to the United States Food and Drug Administration (US FDA) definition, a biosimilar is a biologic product that is highly similar to an already licensed reference biologic that has no clinically meaningful differences in terms of safety, purity, and potency.8,9 Biosimilars remain fairly new to the US market, particularly in the oncology space; however, this is anticipated to change rapidly with multiple biosimilar entrants expected in oncological treatment and supportive care in the upcoming years.10

Given the disproportionate burden of cancer in the elderly, understanding the intersection of the availability of biosimilars and the growing interest in value-based oncology care models, particularly within the Centers for Medicare & Medicaid Services (CMS), is of increasing importance in health care delivery.

The objective of this review is to provide an overview of value-based care models in oncology with a focus on the CMS programs and to discuss the potential contribution of biosimilars to CMS value-based care objectives. This review first describes the use of biologics in targeted and supportive oncology care, introduces biosimilars, and then examines the historical legacy and objectives of the CMS value programs with a focus on how biosimilars might support broader access to equitable, high-quality oncology care.

The high cost of cancer care
The increased prevalence of cancers, earlier treatment initiation, and improved patient outcomes all contribute to the growing use of oncology and supportive care biologic agents. These factors, coupled with the high costs of manufacturing biologics and macro- and micro-economic factors resulting in higher health care costs, have led to a rise in cancer care spending.7,11 In high-income countries, the costs of delivering cancer care are outstripping national budgets, and sustainability of health care financing remains a key public policy concern.12

Biologics in cancer treatment and supportive care
Biologics have been approved for use in primary cancer treatment and supportive care since 1989 (Figure 1). Primary treatment biologics include, but are not limited to, cetuximab,13 rituximab14 (chimeric mAbs targeting epidermal growth factor receptor and CD20, respectively),
trastuzumab, and bevacizumab (humanized mAbs that inhibit human epidermal growth factor receptor 2 and vascular endothelial growth factor A, respectively). These biologics have been shown to improve clinical, health-related quality of life (HRQoL) and hematological outcomes. Biologics are not exclusive to primary treatment, but they have been developed for supportive oncologic treatment as well. Supportive oncologic treatment addresses the adverse effects that are common with primary chemotherapy. Biologics in supportive oncology care include, but are not limited to, agents that help replenish hematologic components during and following chemotherapy. Epoetin alfa and darbepoetin are recombinant human erythropoietic proteins. Filgrastim and its analog, pegfilgrastim, are recombinant human G-CSF. The use of supportive care biologics with chemotherapy improves hematological response and has a positive effect on HRQoL.

Biologic therapies have improved treatment outcomes over previous standard-of-care chemotherapy, while biologic supportive care agents have been shown to be associated with reduced treatment side effects resulting in improved patient-reported HRQoL. However, patient access to biologics may be limited by availability, insurance coverage, and cost. As many available biologics reach the end of their patent protection periods (US patents for cetuximab expired in 2014, for rituximab in 2016, and for both trastuzumab and bevacizumab, they will reach the end of term in 2019), patient access has become an important consideration among the balance of high-quality care and costs. Within the context of this balance, biosimilars are being developed as alternative options with potentially lower costs and greater access. By 2020, a range of biosimilars of biologic agents used in oncology treatment are expected to receive US FDA approval and become available in the US market, providing increased treatment options and thus competition, with the potential for pricing reductions.

Costs of biologics cancer care in the USA

In the USA, total spending on cancer care has increased from $27 billion in 1990 to $124 billion in 2010, with spending projected to reach around $157 billion by 2020. Total costs of cancer care for the US population are predicted to increase across all phases of care (Figure 2). Cost drivers include technological innovation, rising costs of hospitalizations, and a population-level increasing susceptibility to malignancy due to an aging demographic and increasing life span. Global spending on oncology and supportive care drugs reached $100 billion in 2014, with targeted therapy expenditures accounting for almost 50% of this amount. In the USA, oncology drug expenditures, excluding supportive care agents, increased by 18.0% from 2014 to 2015. The

![Figure 2](https://www.dovepress.com/)

**Figure 2** Current and projected cost of cancer care in the USA by phase of care in 2010 and 2020, respectively, weighted to dollar values in 2010.

**Note:** Data from National Cancer Institute.27
fastest growing drug classes within oncology are mAbs and protein-kinase inhibitors, with mAbs accounting for 35% of US oncology spending. US sales figures in 2015 for three of the top 20 global products – bevacizumab, rituximab and trastuzumab – were $6.2 billion, $6.3 billion, and $5.6 billion, respectively. US patients are shouldering an increasing share of these rising costs as health plans restructure their benefit designs, including a transition to high-deductible health plans with higher patient out-of-pocket costs from traditional fixed copay plans. The financial consequences of cancer treatment on patients and their families can be substantial, which has been shown to be a substantial burden. Given the high costs of cancer care and the need to balance health care provision and associated budgets for the full range of conditions affecting population health, issues of patient access, value, and equity are the subject of global discussion and debate.

**Oncology biosimilars in Europe**

The European Medicines Agency (EMA) first introduced a regulatory framework for biosimilars in 2004, and by 2006, it had established a comprehensive set of guidelines for their approval. Since then, European countries have approved the highest number of biosimilars worldwide, having shown high similarity to their reference products via a series of studies (efficacy and safety), and on the basis of nonclinical and pharmaceutical quality data. For example, in the European Union, clinical guidelines were updated in 2009 to encourage the use of biosimilar filgrastim, which led to significantly increased consumption, enabling greater numbers of patients access to this treatment at earlier stages of the therapy cycle.

Approvals for supportive oncology care biosimilars in Europe have been ongoing. As of January 2018, nine filgrastim and five epoetin alfa/zeta biosimilar products were approved and have had tremendous impact on improving access to treatment. As of this publication, license applications for three pegfilgrastim biosimilars were under review by the EMA. For targeted therapies, five rituximab biosimilars, all licensed for the same or similar oncology indications, are authorized by the EMA, and five trastuzumab biosimilars are still under review.

Biosimilars provide an opportunity for cost savings. In Europe, individual member states are allowed to negotiate their own pricing on biosimilars. According to a 2016 IMS report, following the introduction of biosimilar competition varied substantially among countries: 25%–29% in Scandinavia; 39% in France; and 55% in Germany. Meanwhile, the observed price change in (2015) for biologic and biosimilar filgrastim, following the launch of the first approved filgrastim biosimilar, varied from 14% in France to 27% in Germany, based on gross ex-manufacturer price. A significant increase in consumption of therapeutic biologics and biosimilars has been shown in European countries upon entry of a biosimilar into the market and this increase is attributed to reduced costs.

It should be noted, however, that owing to substantial international differences in market forces, drug pricing, and health care policy (as well as in access and utilization), European pricing data are not applicable to the US market.

**Biosimilar use in other regions**

Biosimilars are used widely in the Middle East and in Asia. Reasons for use include lower price relative to reference biologics and bioequivalence of efficacy and safety. Regulatory approvals, where needed, are largely modeled after US FDA and EMA guidelines.

**Biosimilars in cancer treatment and supportive care in the USA**

The USA has a regulatory framework for biosimilars, which was enacted as the 2009 Biologics Price Competition and Innovation (BPCI) Act, part of the Patient Protection and Affordable Care Act. This created an abbreviated licensure pathway for biological products demonstrated to be biosimilar to, or interchangeable with, a US FDA-licensed biological product (or “reference product”), known as the 351(k) pathway. The Congressional Budget Office estimates that the sales-weighted market average discount on biosimilars would be 20%–25% relative to reference agents. The regulatory framework and evidence requirements of the US biosimilars program involve a stepwise approach that relies heavily on analytical methods to demonstrate, through the “totality of evidence” (i.e., all available analytical, nonclinical, and clinical data), that a proposed biosimilar functions the same way as its reference product.

Biosimilars have been available globally since 2008, with the first US biosimilar, the G-CSF filgrastim-sndz, available in the USA since September 2016. Wholesale acquisition costs (WACs) in the USA for 2017 show a 15% discount for biosimilar filgrastim-sndz over filgrastim; a recent cost-efficiency analysis determined that prophylaxis with filgrastim-sndz was associated with consistently significant cost savings over filgrastim and pegfilgrastim. The alternate filgrastim agent, tbo-filgrastim (which is not a biosimilar in the USA and is approved for only one filgrastim indication), is available at a 23% discount compared to the WAC for filgrastim.
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(Figure 3A).\textsuperscript{57} Based on CMS payment limits (i.e., average sales price plus 6\%) for fourth quarter of 2017, the payment limit for biosimilar filgrastim-sndz was 28\% lower than for filgrastim, while for alternate agent tbo-filgrastim, pricing was 36\% lower than for filgrastim (Figure 3B).\textsuperscript{58} Thus, the filgrastim biosimilar and alternate agent provide cost savings under the pricing available to commercial payers and CMS payment limits.

Multiple biosimilars are expected to obtain US FDA approval and enter the US market in the next 2–3 years as patent protection of reference supportive care drugs have come to the end of term recently or are expiring soon (Figure 1).\textsuperscript{59–61} As of this writing, filgrastim-sndz is the only approved biosimilar for supportive oncology care under the US FDA BPCI, but in May 2017, the US FDA Oncologic Drugs Advisory Committee recommended approval of an epoetin alfa biosimilar across all licensed indications of the reference product.\textsuperscript{62,63} Also regarding biosimilar supportive care, Biologics License applications for four proposed biosimilars of G-CSF pegfilgrastim have been under review by the US FDA as of June 2017\textsuperscript{64–66}; however, two of these applications were rejected and two others were still pending as of this writing. Biosimilars of targeted therapies bevacizumab (September 2017) and trastuzumab (December 2017) have recently been approved by the US FDA. Also in 2017, the US FDA has accepted a new Biologics License Application for a filgrastim biosimilar (September)\textsuperscript{67} as well as biosimilars of rituximab\textsuperscript{68} and trastuzumab.\textsuperscript{69–70} As more biosimilars become available after receiving regulatory approval, adoption in clinical practice is expected to increase.

**Figure 3** Commercial price comparison for short-acting G-CSFs (filgrastim-sndz, tbo-filgrastim, and filgrastim) based on the fourth quarter of 2017 (A) WAC/average wholesale price in the USA\textsuperscript{57} and (B) CMS payment limits (average sales price + 6\%).\textsuperscript{58}

**Abbreviations:** AWPC, average wholesale price; \( \mu g \), microgram; CMS, Centers for Medicare & Medicaid Services; G-CSFs, granulocyte-colony stimulating factors; WAC, wholesale acquisition cost.
therapies are expected to improve access and to reduce overall pharmaceutical expenditures.

**Value-based oncology frameworks**

With ongoing concerns about the escalating costs of cancer care, a number of US professional and private organizations have developed value assessment frameworks to define and measure the value of oncology drugs and other therapies.\(^{78,71-75}\) The overarching objectives of these frameworks differ, with some tailored to support physicians and patients in making informed, evidence-based treatment decisions, and others designed as tools to assist in coverage or reimbursement decisions. To date, however, these frameworks provide suggested guidance only; none has been implemented in US clinical practice or in a payer environment.

Outside of the USA, the UK’s National Institute for Health and Care Excellence (NICE) and decision-making bodies in and many other countries use a health technology assessment (HTA) approach, which includes cost-effectiveness models and incremental cost-effectiveness ratios, for health care reimbursement decision making. While it could be argued that countries using HTAs include some measure of value-based care, it should be noted that the National Health Service in England has studied incentive programs and value-based payments as a means to address inequalities in health care, but comprehensive programs have not been implemented as of this writing.\(^ {76}\)

**The CMS value-based care programs**

The CMS has developed value-based care programs that reward health care providers with incentive payments for improving the quality of care they provide to Medicare beneficiaries. In the future, it is anticipated that CMS payments will be increasingly tied to measured performance in oncology care.\(^ {77}\)

The CMS Quality Payment Program (QPP)

Fee-for-service (FFS) is a common US payment model in which medical services are not bundled, but paid for individually, thus incentivizing provision of high-quantity (but not necessarily high-quality) health care. An underlying tenant of value-based care is to move away from the FFS model and toward performance-based payments. In October 2016, the CMS finalized the Medicare Access and CHIP Reauthorization Act of 2015\(^ {78}\) that implemented the QPP (Figure 4). The QPP began in January 2017, with payment adjustments based on performance to be fully implemented by January 2019.\(^ {79}\) The QPP offers payment according to one of two tracks: 1) a Merit-based Incentive Payment System linked to performance including following defined, evidence-based clinical quality measures, and 2) Advanced Alternative Payment Models (APMs)\(^ {80}\) that give financial incentives to clinicians to provide high-quality and cost-efficient care (Figure 5).\(^ {78,81,82}\) One of the Advanced APMs is the Oncology Care Model (OCM).\(^ {81,83}\)

**The CMS OCM**

In response to rising cancer treatment costs, in June 2016, the CMS launched a new, voluntary OCM\(^ {84,85}\) as part of its broader initiative to improve the effectiveness and efficiency of specialty care; the program aims to provide higher quality, more coordinated oncology care at the same or lower cost to Medicare than traditional FFS payments (Figure 4A and B).\(^ {86}\) The OCM program ties payments to provider performance based on meeting specified quality metrics and practice reforms, with some practices already entering into payment arrangements that include financial and performance accountability for episodes of care involving chemotherapy administration to patients with cancer. As of this writing, the program is scheduled for 2017 through 2022. In July 2017, 192 practices and 14 commercial payers were participating in the OCM.\(^ {84}\)

The OCM incorporates a two-part payment system for physician practices: a per-beneficiary Monthly Enhanced Oncology Services (MEOS) payment and a performance-based incentive payment (Figure 5). The MEOS payment assists participating practices in effectively managing and coordinating episodes of care for oncology patients. The performance-based incentive payment is calculated retrospectively on a semi-annual basis, based on the practice’s achievements in quality measures and reductions in Medicare expenditures.

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The Advanced APM track of the QPP is designed to give Medicare providers greater flexibility in delivering value-based care tailored to the type of care they provide. As specific examples of Advanced APMs, the Medicare Shared Savings Program (MSSP) two-sided risk models and OCM were developed to deliver more effective and efficient specialty care through the provision of higher-quality, more coordinated care at the same or lower cost to Medicare as the traditional FFS model, utilizing participant-reported quality metric data to measure and reward high-value oncology care. Participants in the OCM and other similar incentives have an opportunity to play a key role in identifying clinical care practices to meet CMS program assessment goals including patient experience, reduced shared cost, and improved patient outcomes.
Biosimilars offer potential benefits under the OCM, including enhanced affordability and increased access to biologic treatments, along with the improved outcomes and HRQoL associated with biologics in both cancer treatment and supportive care. For example, the use of G-CSF as supportive care in 1,655 patients receiving standard-of-care chemotherapy for breast cancer reduced the incidence of neutropenia, which led to increased dose administration of the primary treatment and improved survival outcomes.87 Availability of biosimilars in the oncology setting in the European Union has expanded patient access to treatment that previously may have been unavailable due to cost.89 Economic modeling studies from Germany, France, Italy, Spain, and the United Kingdom have shown cost savings along with expanded access to supportive care treatments including biosimilar filgrastim and biosimilar epoetin alfa, compared with their respective reference biologics. This is illustrated by a budget impact analysis of real-world data for biosimilar rituximab (for rheumatology and cancer) in 28 European countries and by a recent Croatian study evaluating the budget impact of biosimilar trastuzumab for the treatment of breast cancer.92 These European examples provide insight into possible cost-savings scenarios with biosimilars in the USA; however, due to the unique nature
of the US health care market, it is unknown whether cost savings from biosimilars observed in Europe and elsewhere will manifest in the USA once a greater range of biosimilars used in cancer care are approved and available for use.

Biosimilars may offer a more affordable alternative to biologics as well as result in overall price decreases from market competition, which could result in substantial cost savings in the USA. These benefits have been demonstrated for biosimilar filgrastim, for which CMS (ASP + 6%) pricing shows a 28% discount over the reference biologic. The Rand Corporation reports that savings to the US health care system incurred from the use of biosimilars over biologics range from an estimated $13 billion to $66 billion over the 10-year period between 2014 and 2024. It is anticipated that the expanded treatment choices provided by biosimilars will open up new opportunities to improve value and care delivery. With biosimilar filgrastim available in the USA and biosimilar epoetin alfa expected to be available soon, the potential for clinicians to utilize these two supportive care biosimilars in oncology care, along with biosimilar targeted therapies, could be an important component in meeting MSSP and OCM objectives to improve the quality of care while reducing costs.

Realization of cost savings possible from biosimilars, however, will require that biosimilars are utilized. Results of a 2015–2016 survey led by the Biosimilars Forum showed that major knowledge gaps about biosimilars and their potential use in clinical practice still exist among US specialty physicians, including oncologists. Key gaps include defining biologics versus biosimilars in the context of biosimilarity, understanding the approval process and the use of the “total- ity of evidence” approach by the US FDA for biosimilar evaluation, understanding the evidence requirements for demonstration of safety and immunogenicity of a biosimilar versus its reference product, understanding the rationale for indication extrapolation, and defining interchangeability in the context of pharmacy-level substitution.

US physicians are now becoming more receptive to prescribing biosimilars, as a potentially important way to reduce drug costs and open access to effective therapies. A recent survey showed that efficacy (89%), safety (81%), and patient costs (71%) were the most important factors in determining whether US physicians would prescribe biosimilars overall, with discounts being a key influencer in anticipated prescribing patterns. As additional biosimilars are approved in the USA and awareness grows, it is anticipated that biosimilar uptake and utilization will increase. There is a need to educate US physicians about biosimilars and to raise awareness among US payers and patients as well as health care providers in order to increase utilization of these potentially cost-saving therapies.

**Conclusion**

The goal of value-based care in oncology is to improve the quality of care, while containing costs. Advanced APMs such as the MSSP two-sided risk models and OCM are examples of
a shift away from the traditional volume-based FFS model. For the OCM, this objective targets Medicare beneficiaries through an episode-based payment model that financially incentivizes high-quality, coordinated care. In moving toward a value-based specialized care system, payers recognize and reward providers who proactively seek to improve the patient experience and health outcomes. Biosimilars may provide an additional tool for providers participating in value-based care initiatives such as the MSSP and OCM, resulting in cost savings and efficiencies in the delivery of high-value care through expanded use of biologic treatment and supportive care agents during episodes of care. These savings may then be realized through the MSSP, OCM, or other incentive programs, with benefits passed on to health care providers, payers, and patients alike.

Acknowledgments
Medical writing support was provided by Robyn Fowler, PhD, Patricia McChesney, PhD, CMP, and Karen Smoyer, PhD, of Engage Scientific Solutions and funded by Pfizer Inc. Financial support for this review was sponsored by Pfizer Inc.

Disclosure
Dr Patel has been a consultant to Pfizer Inc at advisory boards. Dr Arantes Jr, Ms Tang, and Dr Fung are employees and stockholders of Pfizer Inc. The authors report no other conflicts of interest in this work.

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