CRS Report for Congress

FDA Regulation of Follow-On Biologics

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Summary

Legislation introduced in the 110th Congress (H.R. 1038/S. 623, H.R. 1956, H.R. 5629, S. 1505, and S. 1695) would expand regulatory activities of the Food and Drug Administration (FDA) by opening a pathway for the approval of follow-on biologics. A biologic is a preparation, such as a drug or a vaccine, that is made from living organisms. In contrast, most commonly used drugs are synthesized via a chemical process. A follow-on biologic is similar to the brand-name, or innovator, product made by the pharmaceutical or biotechnology industry.

The new regulatory pathway would be analogous to the FDA’s authority for approving generic chemical drugs under the Drug Price Competition and Patent Term Restoration Act of 1984 (P.L. 84-417), often referred to as the Hatch-Waxman Act. The generic drug industry achieves cost savings by avoiding the expense of clinical trials, as well as the initial drug research and development costs that were incurred by the brand-name manufacturer. The cost of specialty drug products, such as biologics, is often prohibitively high. For example, the rheumatoid arthritis and psoriasis treatment Embrel costs $16,000 per year. It is thought that a pathway enabling the FDA approval of follow-on biologics will allow for market competition and reduction in prices, though perhaps not to the same extent as occurred with generic chemical drugs under Hatch-Waxman.

In contrast to chemical drugs, which are relatively small molecules and for which the equivalence of chemical composition between the generic drug and innovator drug is relatively easy to determine, a biologic, such as a protein, is much larger in size and much more complex in structure. Therefore, comparing a follow-on protein with the brand-name product is more scientifically challenging than comparing chemical drugs. In many cases, current technology will not allow complete characterization of biological products. Additional clinical trials may be necessary before the FDA would approve a follow-on biologic.

This report provides a brief introduction to the relevant law, the regulatory framework at the FDA, the scientific challenges for the FDA in considering the approval of follow-on biologics, and a description of the proposed legislation. It will be updated as legislative events warrant.
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The 110th Congress is considering legislation (H.R. 1038/S. 623, H.R. 1956, H.R. 5629, S. 1505, and S. 1695) that would expand regulatory activities of the Food and Drug Administration (FDA) by opening a pathway for the approval of follow-on biologics. This pathway would be somewhat analogous to that which allowed for the approval of generic chemical drugs via passage of the Drug Price Competition and Patent Term Restoration Act of 1984 (P.L. 84-417), often referred to as the Hatch-Waxman Act. By offering an alternative to brand-name drug products, the Hatch-Waxman Act is credited with lowering the cost of drugs to consumers, as well as allowing for the expansion of the generic drug industry in the United States.

At the time that Hatch-Waxman was being debated by Congress and implemented by the FDA, the biotechnology industry was just beginning to develop its first human therapeutic agents. The first FDA approval of a biotechnology drug for human use, human insulin, occurred in 1982, followed by human growth hormone in 1985, alpha interferon in 1986, tissue plasminogen activator in 1987, and erythropoietin in 1989. Biotechnology products are expected to become a larger and larger share of the drugs sold by the pharmaceutical industry to U.S. consumers. However, with no equivalent to the generic alternatives to chemical drugs, the cost of therapeutic biologics is often prohibitively high for individual patients. For example, the rheumatoid arthritis and psoriasis treatment Embrel costs $16,000 per year, and biological drugs for multiple sclerosis range in price from $16,000 to $25,000 per year.3 Spending by Medicare in 2006 on just one such drug, Epogen,
was $2 billion, more than the entire FY2006 budget for FDA, which was $1.863 billion ($1.494 billion in direct appropriations and $369 million in user fees).  

In 2006, U.S. spending on such specialty drugs was $54 billion, or about 20% of total spending on pharmaceuticals. Speciality drugs are expected to comprise 26% of total pharmaceuticals purchased by 2010, almost doubling to $99 billion per year, a rate of increase that is second highest among all the components of health care spending, exceeded only by diagnostic imaging. From 2005 to 2006, the cost of non-speciality (i.e., chemical) drugs rose 6%, whereas speciality (mostly biologic) drugs rose 21%. Spending on all pharmaceuticals currently represents about 11% of health care spending in the United States.

In the case of chemical pharmaceuticals, before a generic drug can be marketed, the generic drug company must demonstrate to the FDA that the drug product is identical to the original product. For chemical drugs, some experts argue that “generic medications decrease prices 60% to 90% on branded oral-solid medications.” The Congressional Budget Office estimated the savings generated by generic drug use in 1994 was between $8 billion and $10 billion. The generic drug industry achieves these cost savings by avoiding the expense of clinical trials, as well as the initial drug research and development costs that were incurred by the brand-name manufacturer.

Even though patents for several specialty biotechnology drug products have expired, very few have had to face the same type of market competition that occurs with chemical drugs. In contrast to the relatively simple structure and manufacture of chemical drugs, follow-on biological products, with their more complex nature and method of manufacture, will not be identical to the brand-name product, but may instead be shown to be similar. The Generic Pharmaceutical Association (GPhA) has advocated that the FDA establish a regulatory system for the approval of follow-on

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3 (...continued)


8 Jonah Houts, testimony before the House Committee on Oversight and Government Reform, March 26, 2007.

biologics under its existing statutory authority.\footnote{Bill Nixon, President and CEO, Generic Pharmaceutical Association, letter to Daniel Troy, Chief Counsel, FDA, January 18, 2002, at [http://www.fda.gov/cder/ogd/GPHA_jan_21.htm].} However, the Biotechnology Industry Organization (BIO) has filed a citizen petition with the FDA requesting a number of actions that would inhibit the approval of follow-on biologics.\footnote{BIO Citizen Petition, Follow-on Therapeutic Proteins, April 23, 2003, at [http://www.fda.gov/OHRMS/DOCKETS/DOCKETSF/03p0176/03p-0176-cp00001-01-vol11.pdf].}

Proposed legislation (H.R. 1038/S. 623, H.R. 1956, H.R. 5629, S. 1505, and S. 1695) would provide a mechanism for FDA approval of biological products that are similar to the brand-name product, thereby allowing for market competition and reduction in prices, though perhaps not to the same extent as with generic chemical drugs. An economic analysis released in February 2008 estimates savings in the United States of $67.2 billion to $107.7 billion over the 10 years from 2010 to 2019, and $235.7 billion to $377.7 billion over the 20 years from 2010 to 2029.\footnote{Robert J. Shapiro, Karan Singh, and Megha Mukim, “The Potential American Market for Generic Biological Treatments and the Associated Cost Savings,” February 2008, at [http://www.insmed.com/PDF/Biogeneric_Savings.pdf].} Previous economic studies on savings to the federal government over ten years due to the use of follow-on biologics ranged “between nothing and $14 billion.”\footnote{“CBO Weighs 2 Studies That Show Little Savings From Biogenerics,” Inside Health Policy, July 19, 2007.} A study by Avalere Health estimated “government savings at $3.6 billion in the first 10 years;” another study by Express Scripts estimated “10-year consumer savings at $71 billion and federal savings at $14 billion.”\footnote{Ibid.}

This report provides an overview of the FDA regulatory issues involved in the approval of follow-on biologics.\footnote{For patent issues, see CRS Report RL33901, *Follow-On Biologics: Intellectual Property and Innovation Issues*, by Wendy H. Schacht and John R. Thomas.}

### Relevant Laws

In general, biological products are regulated (licensed for marketing) under the Public Health Service Act — originally by the National Institutes of Health (NIH) and its precursors and later by the FDA — and chemical drugs are regulated (approved for marketing) under the Federal Food Drug and Cosmetic Act (by the FDA). This section provides a brief history of these two Acts and other relevant laws, as well as some of the important amendments that have occurred during the past 100 years.
The regulation of biologics by the federal government began with the Biologics Control Act of 1902, “the first enduring scheme of national regulation for any pharmaceutical product.” The act was groundbreaking, “the very first premarket approval statute in history.” It set new precedents, “shifting from retrospective post-market to prospective pre-market government review.” The Biologics Act was passed in response to deaths (many in children) from tetanus contamination of smallpox vaccine and diphtheria antitoxin. The act focused on the manufacturing process of such biologic products and required an inspection of the manufacturing facility before a federal license was issued to market the product.

The Biologics Act predates the regulation of drugs under the Pure Food and Drugs Act, which was enacted in 1906. The 1906 Act “did not include any form of premarket control over new drugs to ensure their safety ... [and] did not include any controls over manufacturing establishments, unlike the pre-existing Biologics Act and the later-enacted Federal Food Drug and Cosmetic Act (FDC Act).” The Pure Food and Drugs Act was replaced by the FDC Act in 1938. The FDC Act required that drug manufacturers submit a new drug application (NDA) prior to marketing that demonstrated, among other things, that the product was safe.

The Biologics Act was revised and re-codified (42 USC 262) when the Public Health Service Act (PHS Act) was passed in 1944. The 1944 Act specified that a biological product that has been licensed for marketing by the FDA under the PHS Act is also subject to regulation (though not approval) under the FDC Act. A biological product is defined under section 351(i) of the PHS Act, as

a virus, therapeutic serum, toxin, antitoxin, vaccine, blood, blood component or derivative, allergenic product, or analogous product ... applicable to the prevention, treatment or cure of a disease or condition of human beings.

Section 351(j) of the PHS Act states that “the FDC Act applies to a biological product subject to regulation under this section, except that a product for which a license has been approved under subsection (a) shall not be required to have an approved application under section 505 of such Act.” Most biological products regulated under the PHS Act also meet the definition of a drug under section 201(g) of the FDC Act:

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17 Ibid, p. 147.

18 Ibid.


20 For further information, see CRS Report RL32797, *Drug Safety and Effectiveness: Issues and Action Options After FDA Approval*, by Susan Thaul.
articles intended for use in the diagnosis, cure, mitigation, treatment, or prevention of disease in man or animals; and articles (other than food) intended to affect the structure or any function of the body of man or other animals.

The FDA Modernization Act of 1997 (FDAMA) amended the PHS Act to require a single biological license application (BLA) for a biological product, rather than the two licenses — Establishment License Application (ELA) and Product License Application (PLA) — that had been required between 1944 and 1997. The PHS Act provides authority to suspend a license immediately if there is a danger to public health.

As stated previously, biological products are, in general, regulated — licensed for marketing — under the PHS Act, and chemical drugs are regulated — approved for marketing — under the FDC Act. However, through a historical quirk, the FDA was given regulatory authority over certain natural source biological products; these products have been regulated as drugs under the FDC Act rather than as biologics under the PHS Act. Three years prior to the re-codification of the Biologics Act, Congress gave the FDA authority over the marketing of insulin.21 Insulin is a peptide hormone, a small protein that regulates carbohydrate metabolism.22 In the 1940s, insulin “was obtained in the same manner as many biologics, namely extraction from animals. Despite this similarity with biologics, insulin was regulated by FDA.”23 In addition to insulin, the distinction of a biological product regulated as a drug under the FDC Act rather than as a biologic under the PHS Act holds true for a small set of products that are mostly hormones: glucagon, human growth hormone, hormones to treat infertility, hormones used to manage menopause and osteoporosis, and certain medical enzymes (hyaluronidase and urokinase).24

This distinction is important because the Hatch-Waxman Act provides a mechanism for the approval of generic drugs under the FDC Act but not under the PHS Act. Specifically, Hatch-Waxman added two abbreviated pathways to the FDC Act for subsequent versions of already approved products: section 505(j) and section 505(b)(2).

Section 505(j) established an Abbreviated New Drug Application (ANDA) process for a generic drug that contains the same active ingredient as the brand-name innovator drug. In the ANDA, the generic company establishes that its drug product

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22 A protein is a large organic molecule composed of a long chain or chains of amino acids linked by chemical bonds. Insulin is a short chain of 51 amino acids. Examples of carbohydrates include sugars and starch.


is chemically the same as the already approved innovator drug, and thereby relies on the FDA’s previous finding of safety and effectiveness for the approved drug. The 505(j) pathway is used for the approval of most generic chemical drugs.

Under the second pathway, a drug that has a significant difference from an innovator drug, but is still sufficiently similar to that drug, may be the subject of a 505(b)(2) application. The company filing the application must submit additional non-clinical and clinical data to show that the proposed product is safe and effective. However, the application may rely on published literature or on the FDA’s finding of safety and effectiveness for the already approved product to support the approval of the proposed product. The 505(b)(2) pathway has been used to approve Omnitrope, a follow-on human growth hormone, and a few other follow-on protein products. All have been members of the small set of biologic products that were regulated as drugs.

### Regulatory Framework

Following enactment of the 1902 Biologics Act, regulatory responsibility for biologics was first delegated to the Hygienic Laboratory, a precursor of NIH. In 1972, regulatory authority for biologics was transferred from the NIH Division of Biological Standards to the FDA Bureau of Biologics, which eventually became the agency’s Center for Biologics Evaluation and Research (CBER).

Because biotechnology products frequently cross the conventional boundaries between biologics, drugs, and devices, determining the jurisdictional status of these new products has been difficult for both the FDA and industry. Some products have

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25 Janet Woodcock, testimony before the House Committee on Oversight and Government Reform, March 26, 2007.


27 Ibid., p. 148, and The NIH Almanac — Historical Data: Chronology of Events, at [http://www.nih.gov/about/almanac/historical/chronology_of_events.htm]. In 1937, the biologics control program was assigned to the newly established Division of Biologics Control. In 1955, the biologics control function was placed in the newly formed Division of Biologics Standards.

28 The NIH Almanac; Donna Hamilton, “A Brief History of the Center for Drug Evaluation and Research,” FDA History Office, November 1997, at [http://www.fda.gov/cder/about/history/Histext.htm]. During the early 1980s, the Bureau of Drugs and the Bureau of Biologics merged to form the National Center for Drugs and Biologics. In 1984, all of the National Centers within FDA were redesignated simply as Centers. In 1987, the Center for Drugs and Biologics was split into the Center for Drug Evaluation and Research (CDER) and the Center for Biologics Evaluation and Research (CBER). CBER continues to use NIH facilities and buildings until the expected move in 2012 to the new FDA headquarters in White Oak, Maryland (see [http://www.fda.gov/oc/whiteoak/projectschedule.html]).
had characteristics that met multiple statutory and scientific definitions. In 1991, the FDA published an Intercenter Agreement between CBER and the Center for Drug Evaluation and Research (CDER). In general, the agreement stated that traditional biologics (vaccines, blood, blood products, antitoxins, allergenic products), as well as most biotechnology products, would be regulated by CBER. The small set of biologics mentioned earlier that are regulated as drugs under the FDC Act would continue to be regulated by CDER, regardless of the method of manufacture.

In 2002, however, the FDA announced its intention to reorganize review responsibilities, consolidating review of new pharmaceutical products under CDER, thereby allowing CBER to concentrate on vaccines, blood safety, gene therapy, and tissue transplantation. On June 30, 2003, responsibility for most therapeutic biologics was transferred from CBER to CDER. Under the new structure, biological products transferred to CDER will continue to be regulated as licensed biologics under section 351 of the PHS Act. Examples of products transferred to CDER include monoclonal antibodies; proteins intended for therapeutic use (interferons, thrombolytic enzymes); immunomodulators (other than vaccines and allergenic products); and, growth factors, cytokines, and monoclonal antibodies intended to alter production of blood cells. Remaining at CBER are traditional biologics such as vaccines, allergenic products, antitoxins, antivenins, venoms, and blood and blood products, including recombinant versions of plasma derivatives (clotting factors produced via biotechnology).

As stated previously, the Hatch-Waxman Act added two abbreviated pathways under the FDC Act — 505(j) and 505(b)(2) — but not under the PHS Act, for the approval of additional products subsequent to the innovator product. Because of the complex nature of most biological products and their methods of manufacture, such products will not be identical to the brand-name product; therefore, the 505(j) pathway cannot be used for product approval. However, if a biological product is sufficiently similar to the innovator product, the 505(b)(2) pathway may be used by a company for the approval of its biologic. Following the enactment of Hatch-


30 The Intercenter Agreement is available at [http://www.fda.gov/oc/ombudsman/drug-bio.htm].


33 Transfer of Therapeutic Products to the Center for Drug Evaluation and Research, at [http://www.fda.gov/cber/transfer/transfer.htm]. Also of interest is Approved Products Transferring to CDER, at [http://www.fda.gov/cber/transfer/transfprods.htm], and Therapeutic Biological Products, at [http://www.fda.gov/cder/biologics/default.htm].
Waxman, the FDA published in 1999 a draft guidance on applications covered by section 505(b)(2); the guidance has never been finalized.34

As things currently stand, and as discussed above, the 505(b)(2) pathway has been used only for those biologics that have been regulated as drugs under the FDC Act. However, the vast majority of biologics have been regulated under the PHS Act. The FDA’s position is that additional legislation is required to provide such a pathway under the PHS Act. For traditional biologics regulated under the PHS Act, the agency’s longstanding policy has been that a full BLA, including clinical testing, would be required for the licensing of each such product. In a 1974 Federal Register notice, the FDA stated that

 unlike the regulation of human and animal drugs, all biological products are required to undergo clinical testing in order to demonstrate safety, purity, potency and effectiveness prior to licensing, regardless whether other versions of the same product are already marketed or standards for the product have been adopted by rulemaking. Indeed, many of the existing standards require specific clinical testing before approval will be granted. This is required because all biological products are to some extent different and thus each must be separately proved safe, pure, potent, and effective.... There is no such thing as a “me-too” biologic.35

When publishing the final rule on the ANDA procedure that had been outlined in Hatch-Waxman, the FDA stated in 1992 that “these procedures are inapplicable to ... biological drug products licensed under 42 USC 262 (section 351 of the PHS Act).”36 Most recently, during hearing testimony on May 2, 2007, before the Subcommittee on Health of the House Energy and Commerce Committee, Janet Woodcock, Deputy Commissioner and Chief Medical Officer of the FDA, stated in response to questioning that there is no pathway under the PHS Act for the approval or licensing of follow-on biologics that is similar to the 505(b)(2) pathway under the FDC Act, and that the FDA would be willing to work with Congress in crafting a legislative approach to creating such a pathway.

Scientific Challenges

In prepared testimony, Dr. Woodcock outlined the scientific challenges involved in determining the safety and effectiveness of follow-on biologics. The FDA prefers to call these products follow-on protein products. In contrast to chemical drugs, which are relatively small molecules and for which the equivalence of chemical composition between the generic drug and innovator drug is relatively easy to determine, therapeutic proteins are much larger in size and much more complex in structure. A protein is a large organic molecule composed of a long chain of component parts, called amino acids, which are linked by chemical bonds. This

36 Federal Register, v. 57, no. 82, April 28, 1992, p. 17951.
amino acid chain folds into a complex three-dimensional structure. Slight changes in the chain or three-dimensional shape can influence the protein’s biological activity. Proteins can also be altered by the addition of other chemicals, such as sugar groups (glycosylation), at various points along the amino acid chain. Therefore, comparing a follow-on protein with the brand-name product is more scientifically challenging than comparing chemical drugs. In many cases, current technology will not allow complete characterization of biological products. Dr. Woodcock describes these technical problems in her prepared testimony:

Current technologies, such as peptide mapping, protein sequencing, and mass spectroscopy enable manufacturers to determine, with certainty, the amino acid sequence of a recombinant protein. However, the amino acid sequence is the most rudimentary characteristic of a protein. Conclusive analysis of other aspects of a protein’s structure requires much more sophisticated technologies and is fraught with uncertainties that are proportional to the size and complexity of the protein itself. Such complexities include folding of the protein’s amino acid chain into highly organized structures, post-translational modification of the protein with a broad range of biochemical additions (e.g., glycosylation, acetylation, phosphorylation, etc.), and association of multiple protein molecules into aggregates. It is the combination of the protein’s amino acid sequence and its structural modifications that give a protein its unique functional characteristics. Therefore, the ability to predict the clinical comparability of two products depends on our understanding of the relationship between the structural characteristics of the protein and its function, as well as on our ability to demonstrate structural similarity between the follow-on protein and the reference product. Although this currently may be possible for some relatively simple protein products, technology is not yet sufficiently advanced to allow this type of comparison for more complex protein products.

Several terms are important in the discussion of the follow-on proteins and their approval by the FDA. Products that are considered to be therapeutically equivalent “are approved drug products, usually made by different manufacturers, that are pharmaceutical equivalents and for which bioequivalence has been demonstrated. Therapeutic equivalents can be expected to have the same clinical effect and safety profile when administered to patients under the conditions specified in the labeling.”37 Pharmaceutical equivalents are products that contain the same active ingredient in the same strength, dosage form, and route of administration.38 Bioequivalence means that the products are absorbed into the body at a similar rate and extent.39 Interchangeability “is not defined by FDA and could have a number of different meanings. It could refer to products that are therapeutic equivalents, and thus could, in some circumstances, be substituted at the pharmacy level without a

39 Ibid.
physician’s intervention. Alternatively, the term could describe similar products that are not ‘substitutable’ but which, under a physician’s supervision, could be used to treat the same disease or condition in the same patient.”

Most drugs approved under section 505(j) are therapeutically equivalent to the already approved drug product. In her testimony, Dr. Woodcock explains the importance of a determination of therapeutic equivalence for a generic drug and the reasons why such a determination for a follow-on protein product may not be possible, at least at the present time:

In many jurisdictions, therapeutically equivalent drugs may be substituted at the pharmacy level, without a physician’s intervention.... Because of the variability and complexity of protein molecules, current limitations of analytical methods, and the difficulties in manufacturing a consistent product, it is unlikely that, for most proteins, a manufacturer of a follow-on protein product could demonstrate that its product is identical to an already approved product. Therefore, the section 505(j) generic drug approval pathway, which is predicated on a finding of the same active ingredient, will not ordinarily be available for protein products.

**Immunogenicity**, or the ability to elicit an immune response, is another important term in the discussion of follow-on proteins. An immune response to a therapeutic protein can range from detectable, but clinically insignificant, to one that can cause safety problems for the patient or limit the effectiveness of the product. For some biologics, such as vaccines, stimulating an immune response is the intended outcome. However, for other types of therapeutic products, an immune response can lower the clinical effect of a protein. Dr. Woodcock describes the implications at length in the prepared testimony:

Adverse safety events from an immune response could include hypersensitivity reactions such as anaphylaxis, rash, fever and kidney problems, to cross-reaction with an endogenous (naturally occurring in the body) protein (e.g., erythropoietin). Immunogenicity may be influenced by patient-related, disease-related, or product-related factors. Immune responses to administered protein products can be extremely serious or life-threatening; therefore, this issue requires significant attention. The ability to predict immunogenicity of a protein product, particularly the more complex proteins, is extremely limited. Therefore, some degree of clinical assessment of a new product’s immunogenic potential will ordinarily be needed. The extent of independent testing needed will again depend on a variety of scientific factors such as the indication, whether the product is to be administered chronically, the overall assessment of the product’s immunogenic potential, and whether there is the possibility of generating a cross-reaction with an important endogenous molecule.

Even if a follow-on protein product is found to be safe and effective by the FDA, this finding does not mean that the follow-on protein product would be interchangeable with, or substitutable for, the originally approved brand-name product. To establish that the follow-on protein product is substitutable for the

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41 Ibid.
brand-name product, the manufacturer of the follow-on product must demonstrate through additional clinical data that repeated switches from the follow-on product to the brand-name product (and vice versa) would have no negative effect on the safety and/or effectiveness of the products. In other words, there must be no problems with immunogenicity. “For many follow-on protein products, and, in particular, the more complex proteins, there is a significant potential for repeated switches between products to have a negative impact on the safety and/or effectiveness. Therefore, the ability to make determinations of substitutability for follow-on protein products may be limited.”

### Legislation

Initially, there were two competing legislative approaches for the approval by the FDA of follow-on biologics introduced in the 110th Congress: H.R. 1038/S. 623 vs. H.R. 1956 and S. 1505. The introduction of S. 1695 provided a third approach. H.R. 5629 provides an approach that is similar in some respects to S. 1695 with some important differences that are favored by companies that have developed the reference product, also referred to as the innovator or brand-name product.

In general, H.R. 1038/S. 623 is favored by the generic drug industry, whereas H.R. 1956 and S. 1505 are favored by the companies that have developed the reference products. H.R. 1038/S. 623 would allow the FDA to make a determination on interchangeability of a brand-name and follow-on biologic. H.R. 1956 and S. 1505 would not allow the FDA to designate a follow-on biologic as interchangeable with (or therapeutically equivalent to) the brand-name product. In addition, H.R. 1956 and S. 1505 would require the publication of a final product class-specific guidance document (H.R. 1956) or a final product class-specific rule (S. 1505) before an application for a follow-on biologic could be submitted to the FDA. H.R. 1038/S. 623 make no such requirement on the publication by FDA of guidance documents or final rules.

H.R. 1956 and S. 1505 would also set in place provisions governing the nonproprietary naming of biotechnology-derived biologics. The bills would amend the FDC Act to deem such a biologic to be misbranded if its labeling fails to meet these new requirements. H.R. 1038/S. 623 do not contain such product naming provisions. According to media reports, “the brand industry successfully pushed for different names [for brand-name and follow-on products] in Europe. The brand

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42 Ibid.

43 The nonproprietary name for a drug (also called a generic, common or established name) is provided in this country by the United States Adopted Names (USAN) Council. The USAN Council works with similar international groups to standardize drug nomenclature. While a drug’s chemical name can be lengthy and difficult to pronounce and the brand or trademark name is protected by intellectual property protections, the nonproprietary name is in the public domain and is used for all legal and regulatory matters and in all official correspondence. Dan Boring, “Names, Names, Names,” Modern Drug Discovery, v. 3, September 2000, p. 31-32; and, Dan Boring, “More Names,” Modern Drug Discovery, v. 3, October 2000, p. 35-36.
industry argues having different names helps pinpoint which drugs are hurting people, but the generic drug industry believes it is a ploy to thwart generic substitution.”

The brand name drug companies believe that “having different names would make it easier for FDA to tell when a brand or a biosimilar is the cause of side effects. However, FDA urged European regulators to not use this approach when they were debating it last winter. FDA says it already has ‘many alternative mechanisms’ to prevent inappropriate substitution.”

The bipartisan Senate authors of S. 1695 claim to have negotiated a compromise between the brand-name manufacturers and the generic drug industry. S. 1695 would allow FDA to make a determination on interchangeability. S. 1695 does not require the publication of guidance or rule prior to consideration of a follow-on biologic application and does not require a different nonproprietary name for the follow-on biologic than the brand-name drug. S. 1695 would provide, however, 12 years of exclusive marketing for the brand-name product prior to the approval of a follow-on biologic. H.R. 1956 and S. 1505 would provide at least 14 years of exclusive marketing for the brand-name product; H.R. 1038/S. 623 would not provide an exclusivity period.

Although the Bush Administration supports creating a pathway for the approval by FDA of follow-on biologics, it is opposed to several aspects of S. 1695, such as allowing FDA to make determinations on interchangeability, allowing the approval of follow-on biologics without prior issuance of guidance, the creation of the Biological Products Savings Fund, and the possible waiver by FDA of a requirement for clinical trials prior to the approval of a follow-on biologic. The Bush Administration also believes that a follow-on biologic should have a different nonproprietary name than the brand name product. A June 26, 2007, letter from HHS Secretary Michael Leavitt to the Chairman of the Senate HELP Committee outlines the Bush Administration’s position on S. 1695.

H.R. 5629 would require the publication of product-class specific guidance prior to the approval of a biological product, as is the case with H.R. 1956 and S. 1505. H.R. 5629 would allow FDA to make a determination on interchangeability, like H.R. 1038/S. 623 and S. 1695. However, H.R. 5629 would require (1) the publication of final guidance on interchangeability prior to determinations on interchangeability, (2) the biological product must be biosimilar to the reference product and any licensed product that is interchangeable with the reference product, and (3) the biological product must produce the same clinical result for each condition of use on the reference product label. H.R. 5629 would provide at least 12 years and possibly up to 14½ years of exclusive marketing for the brand-name product. H.R. 5629 would require clinical studies of immunogenicity; these studies

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46 Ibid.

47 Letter found at [http://insidehealthpolicy.com/secure/data_extra/dir_07/he2007_2375.pdf].
may be waived only if final guidance on immunogenicity determinations has been published. In contrast, S. 1695 would require that clinical studies submitted in support of a biological product application must be designed to avoid needless duplication or unethical clinical testing; the requirement for clinical and other studies may be waived.

A summary of the key provisions in each of the bills is provided below.

**H.R. 1038 (Waxman), the Access to Life-Saving Medicine Act**, was introduced on February 14, 2007. H.R. 1038 was referred to the Committee on Energy and Commerce, and the Committee on the Judiciary. A companion bill, **S. 623 (Schumer)**, was introduced on February 15, 2007. S. 623 was referred to the Committee on Health, Education, Labor, and Pensions.

H.R. 1038 would amend section 351 of the PHS Act to establish a process for the approval of an abbreviated biological product application for products that contain the same or similar active ingredients as a previously licensed biological product (the reference product). The bill would allow a person to file an abbreviated biological product application with the FDA that includes (1) data demonstrating that the product is comparable to or interchangeable with the reference product; (2) information to show that the conditions or conditions of use prescribed, recommended, or suggested in the labeling proposed for the biological product have been previously approved for the reference product; and (3) information to show that the route of administration, the dosage form, and the strength of the biological product are the same as those of the reference product.

H.R. 1038 sets forth a number of conditions for approval of such an application by the FDA. The bill allows an applicant to request that the FDA make a determination as to the interchangeability of a comparable product and the reference product, based on whether a product can be expected to produce the same clinical result as the reference product in any given patient.

H.R. 1038 provides for a period of up to 36 months of market exclusivity for the first approved interchangeable product, during which time the agency is precluded from approving a second interchangeable product. H.R. 1038 requires the FDA to establish requirements for the efficient review, approval, suspension, and revocation of comparable biological product applications. The bill sets forth provisions governing patent infringement claims against an applicant or prospective applicant for a comparable biological product license.

**H.R. 1956 (Inslee), the Patient Protection and Innovative Biologic Medicines Act of 2007**, was introduced on April 19, 2007. H.R. 1956 was referred to the Committee on Energy and Commerce.

H.R. 1956 would amend section 351 of the PHS Act to provide for the approval of similar biological products. The bill would allow any person to submit an application to the FDA for approval of a biologics license for a biological product that is to be similar to an already approved biological product (the reference product). The application would be approved only if (1) the applicant demonstrates that the similar biological product conforms to the applicable final product-class specific
guidance and, on the basis of the data submitted in conformance with such guidance, the FDA concludes the product is safe, pure, and potent; (2) the facility in which the similar biological product is manufactured, processed, packed, or held meets standards designed to ensure that the biological product continues to be safe, pure, and potent; and (3) the applicant consents to the inspection of the manufacturing facility.

H.R. 1956 would allow FDA approval of an application submitted for a similar biological product (1) only for indications for which the reference product is approved and (2) only if, with respect to each such indication, the application conforms to the applicable final product-class specific guidance, and on the basis of non-clinical and clinical data submitted regarding such indication, the FDA concludes the product is safe, pure, and potent.

H.R. 1956 would not allow the FDA to designate a similar biological product as therapeutically equivalent to the reference product. Two years after enactment, and every two years thereafter, the bill would require that a report be submitted to Congress making recommendations on (1) whether it is feasible, in the current state of scientific and technical knowledge, to make therapeutic equivalence determinations for similar biological products, and (2) if so, the statutory criteria that should govern such determinations.

H.R. 1956 would not allow an application for a similar biological product to be submitted to the FDA unless (1) the FDA has published final product-class specific guidance applicable to the reference product and (2) not less than 12 years have elapsed from the date on which the reference product was approved or licensed. Under the bill, approval of an application would not be effective until at least 14 years after the date the reference product was approved or licensed. Approval would not be effective until 15 years after the reference product was approved or licensed if (1) during the 12-year period following the approval or licensing of the reference product, the FDA approves a supplement to the new drug or biologics license application for the reference product that seeks approval to market the reference product for a new indication and (2) the new indication provides a significant clinical benefit in comparison with existing therapies. The bill would allow any person to submit a request to the FDA for the issuance of product-class specific guidance, and the bill provides specific requirements on the issuance of such guidance documents.

H.R. 1956 includes provisions governing the naming of biotechnology-derived therapeutic protein products and other biological products. The bill would amend the FDC Act to deem a biotechnology-derived therapeutic protein to be misbranded if its labeling fails to meet these requirements.

S. 1505 (Gregg), the Affordable Biologics for Consumers Act of 2007, was introduced on May 24, 2007. S. 1505 was referred to the Senate Committee on Health, Education, Labor, and Pensions.

S. 1505 would amend section 351 of the PHS Act to provide for the approval of biosimilars. The bill would allow anyone to submit an application to the FDA for approval of a biologics license for a biosimilar that is to be similar to an already approved biotechnology-derived therapeutic biological product (the reference
product). The application would be approved only if (1) the applicant demonstrates that the biosimilar conforms to the applicable final product class-specific rule and, on the basis of the data submitted in conformance with such rule, the FDA concludes the product is safe, pure, and potent; (2) the applicant demonstrates that the biosimilar is as similar to the reference product as may be achieved given the state of scientific knowledge and technology capabilities at the time of submission of the application; (3) the applicant demonstrates that the biosimilar has the same route of administration, dosage form, mechanism of action, and strength as the reference product; (4) the facility in which the biosimilar is manufactured, processed, packed, or held meets standards designed to ensure that the biological product continues to be safe, pure, and potent; and (5) the applicant consents to the inspection of the manufacturing facility.

S. 1505 would allow FDA approval of an application submitted for a biosimilar (1) only for indications for which the reference product is approved; (2) only if, with respect to each such indication, the application conforms to the applicable final product class-specific rule, and on the basis of non-clinical and clinical data submitted regarding such indication, the FDA concludes the product is safe, pure, and potent; and (3) only if the applicant agrees to provide to the FDA, on an ongoing basis, all written documents it prepares for any purpose (including any patent litigation) that characterizes the difference between the biosimilar and the reference product.

S. 1505 would not allow the FDA to designate a biosimilar as interchangeable with (or therapeutically equivalent to) the applicable reference product. Two years after enactment, and every two years thereafter, the bill would require an assessment of the state of scientific and technical knowledge regarding the ability of the FDA to make a determination that a biosimilar is interchangeable with (or therapeutically equivalent to) a reference product on a product class basis. If the assessment finds that the state of scientific and technical knowledge enables the FDA to make a determination of interchangeability (or therapeutic equivalence) with respect to one or more product classes, a report would be submitted to Congress that describes such findings and recommendations for statutory criteria that should govern such a determination.

S. 1505 would not allow an application for a biosimilar to be submitted to the FDA unless (1) the FDA has published a final product class-specific rule applicable to the reference product and (2) not less than 12 years have elapsed from the date on which the reference product was approved or licensed. Approval would not be effective until at least 14 years after the date on which the reference product was approved or licensed. Approval would not be made effective until at least 16 years after the reference product was approved or licensed if (1) during the 12-year period following the approval or licensing of the reference product, the FDA approves a supplement to the new drug or biologics license application for the reference product that seeks approval to market the reference product for a new indication and (2) the new indication provides a significant clinical benefit. The bill would allow any person to submit a request to the FDA for the issuance of a product class-specific rule, and the bill provides specific requirements on the issuance of such a rule.
S. 1505 includes provisions governing the naming of biotechnology-derived therapeutic protein products and other biological products. The bill would amend the FDC Act to deem a biotechnology-derived therapeutic protein to be misbranded if its labeling fails to meet the requirements.

S. 1695 (Kennedy), the Biologics Price Competition and Innovation Act of 2007, was introduced on June 26, 2007. S. 1695 was referred to the Senate Committee on Health, Education, Labor, and Pensions. On June 27, 2007, the bill was ordered to be reported with an amendment in the nature of a substitute.

S. 1695 would amend section 351 of the Public Health Service Act to establish a pathway for the licensure of biosimilar biological products. Under the bill, such an application would be required to include, unless the Secretary determines that it is unnecessary, information demonstrating that the biological product is similar to the reference (brand-name) product based on data derived from (1) analytical studies that show the two products are highly similar, notwithstanding minor differences in clinically inactive components; (2) animal studies, including a toxicity assessment; and (3) a clinical study or studies, including an assessment of immunogenicity and other factors. S. 1695 would require that the application include information demonstrating that (1) the biological product and reference product have the same mechanism of action; (2) the condition of use in the proposed labeling for the biological product has been previously approved for the reference product; (3) the route of administration, the dosage form, and the strength of the biological product are the same as those of the reference product; and (4) any facility in which the biological product is manufactured or held meets standards that ensure the product is safe, pure, and potent. The application may also include information demonstrating interchangeability of the biological product with the reference product.

S. 1695 would allow the Secretary to license the biological product if the information submitted in the application is sufficient to show that the biological product is either (1) biosimilar to the reference product or (2) interchangeable with the reference product. The biological product would be interchangeable with the reference product if (1) it can be expected to produce the same clinical result as the reference product in any given patient and (2) the risk, in terms of safety or diminished efficacy, of alternating or switching between use of the biological product and the reference product is not greater than the risk of using the reference product without such alternation or switch.

S. 1695 would allow for a one-year period of exclusive marketing for the first interchangeable biosimilar biological product to be approved as interchangeable for a particular reference product. The bill would not allow for the approval of any biosimilar application until 12 years after the reference product was first licensed.

S. 1695 would allow the Secretary, after the opportunity for public comment, to issue general or specific guidance on the application process for licensure of a biosimilar biological product. The issuance (or non-issuance) of guidance would not preclude the review of, or action on, a submitted application.

S. 1695 would establish a new process for identifying patents that might be disputed between the brand-name company and the company submitting a biosimilar
application. The bill would provide a multistep, but according to the bill’s sponsors, expedited, patent resolution process.

S. 1695 would require all applications for the approval of a biological product to be submitted under section 351 of the PHS Act rather than section 505 of the FDC Act. The bill would provide an exception for the class of biological products that have traditionally been approved under section 505 of the FDC Act; the exception would terminate 10 years after enactment of S. 1695. All approved applications under section 505 of the FDC Act would then be deemed to be a license for the biological product under section 351 of the PHS Act.

S. 1695 would make preliminary provisions for the collection of user fees for the review of biosimilar biological products.

S. 1695 would direct the Secretary of the Treasury to determine the amount of savings to the federal government as a result of enactment of S. 1695 and would transfer the amount to a special reserve fund, the Biological Product Savings Fund, to be expended by the Secretary of HHS on activities authorized under the Public Health Service Act.

S. 1695 would require the Government Accountability Office to conduct a study and report to Congress, not later than three years after enactment, on the extent to which pediatric studies of biological products are being required under the FDC Act and any pediatric needs not being met under existing authority.

Lastly, S. 1695 would specify that biosimilars to orphan drug products (i.e., reference products that have been designated under section 526 of the FDC Act for a rare disease) would be licensed only after the expiration for such reference product, the later of either (1) the seven-year period specified in section 527(a) of the FDC Act or (2) the 12-year period described in S. 1695.

H.R. 5629 (Eshoo), the Pathway for Biosimilars Act, was introduced on March 13, 2008. H.R. 5629 was referred to the Committee on Energy and Commerce, and the Committee on the Judiciary.

H.R. 5629 would require the biological product license application to include information demonstrating that the biological product is biosimilar to the reference product based on data from (1) analytical studies, (2) animal studies, and (3) a clinical study or studies (of immunogenicity) for each condition of use for which the reference product is approved. The requirement for analytical studies and animal studies may be waived. Immunogenicity assessment may be waived only if draft and final guidance has been published. The bill would not allow such an application to be submitted until the later of (1) the start of a proceeding for issuing applicable product class guidance for that product or (2) four years after the reference product was licensed. The bill would not allow the Secretary to accept an application until the Secretary has initiated a proceeding for issuance of applicable product class guidance for that biological product.

H.R. 5629 would require that a license application for such biological products would be reviewed by the division that reviewed and approved the reference product.
Risk evaluation and mitigation strategies under the FFDCA would apply. The bill would allow the approval of an application only if the biological product is biosimilar to the reference product with respect to each condition of use for which the reference product is approved, and the applicant consents to the inspection of the manufacturing facility. If the biological product is, bears, or contains a select agent or toxin, the application would not be approved. An application may not be approved until the Secretary has completed the proceeding for issuance of guidance with respect to the product class within which the biological product falls.

H.R. 5629 would not allow the approval of an application for a follow-on product to become effective until at least 12 years after licensing of the reference product. For pediatric applications, application approval would not become effective until 12 years and 6 months after approval of the reference product. Pediatric exclusivity must be determined by the FDA no later than nine months prior to the expiration of the marketing exclusivity period or no additional pediatric exclusivity will be awarded.

H.R. 5629 would not allow the approval of an application for a follow-on product to become effective until 14 years after the reference product was first licensed if, during the 8-year period following licensure of the reference product, approval of a new indication for the reference product would provide a significant improvement (compared to current marketed products) in the treatment, diagnosis, or prevention of disease. For pediatric applications in this situation, approval of a follow-on product application would not become effective until 14 years and 6 months after approval of the reference product. Pediatric exclusivity must be determined by the FDA no later than nine months prior to the expiration of the marketing exclusivity period or no additional pediatric exclusivity will be awarded.

H.R. 5629 would require a determination on interchangeability if the application shows that the biological product (1) is biosimilar to the reference product and any licensed biological product that has been determined to be interchangeable with the reference product, (2) can be expected to produce the same clinical result in any given patient for each condition of use on the reference product label, and (3) can be alternated or switched between use of the reference product without risk to the patient in terms of safety or diminished efficacy compared with use of the reference product alone. The bill would require that determinations on interchangeability would not be made prior to publication of draft and final guidance advising that it is feasible to make interchangeability determinations on products in that product class, and explaining the data that will be required to support such a determination.

H.R. 5629 would require the Secretary to publish proposed guidance for public comment prior to publication of final guidance with respect to licensure of a biological product or product class. The Secretary must establish a process to allow public input regarding priorities for issuing guidance. For a reference product that was licensed more than seven years prior to enactment, a person may petition the Secretary to commence the process for issuing final guidance for the reference product’s product class. The petition must include a description of the scientific feasibility and rationale for the request. The Secretary must issue final product class guidance within two years of such petition. A guidance may state that the Secretary will not license a product or product class (not including any recombinant protein)
because the science and experience, as of the date of the guidance, does not allow licensure. The bill would require the product class-specific guidance to include a description of (1) the criteria that will be used to determine whether a biological product is biosimilar to a reference product in such product class; (2) the criteria, if available, that will be used to determine whether a biological product meets the standards for interchangeability; and (3) the criteria, if available, that will be used to assess immunogenicity. The bill would allow the Secretary to issue subsequent guidance to modify or reverse previous guidance.

H.R. 5629 would require the Secretary to ensure that the labeling and packaging of each biological product bears a unique name that distinguishes it from the reference product and any other biological products that are evaluated against such reference product.

H.R. 5629 would allow a period of market exclusivity for the applicant that is the first to establish that its product is interchangeable with the reference product for one or more conditions of use. This period of market exclusivity would be 24 months after the later of either the date of the first commercial marketing of the product that is interchangeable with the reference product, or if marketed before interchangeability is determined, the date the product is determined to be interchangeable.

H.R. 5629 would establish a process for identifying patents that might be disputed between the brand-name company and the company submitting a biosimilar application. The bill would provide a multistep patent resolution process.

H.R. 5629 would require all applications for the approval of a biological product to be submitted under section 351 of the PHS Act rather than section 505 of the FDC Act. The bill would provide an exception for the class of biological products that have traditionally been approved under section 505 of the FDC Act; the exception would terminate 10 years after enactment of H.R. 5629. All approved applications under section 505 of the FDC Act would then be deemed to be a license for the biological product under section 351 of the PHS Act.

H.R. 5629 would allow for the collection of user fees for the approval of a biological product licensed under this newly created section of the PHS Act, section 351(k).