Patent Reform: Issues in the Biomedical and Software Industries

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Summary

Congress currently is considering reform of the existing patent system. This interest in patent policy reflects a recognition of the increasing importance of intellectual property to U.S. innovation. Patent ownership is perceived as an incentive to the technological advancement that leads to economic growth. As such, the number of patent applications and grants has grown significantly, as have the type and breadth of inventions that can be patented.

Along with the expansion in the number and range of patents, there are growing concerns over whether the current system is working efficiently and effectively. Several recent studies recommend patent reform. Other experts maintain that major alterations in existing law are unnecessary and that, while not perfect, the patent process can, and is, adapting to technological progress. Thus far in the 109th Congress, two bills, H.R. 2795 and H.R. 5096, have been introduced which, if enacted, would make significant alterations in current patent law.

At the present time, the patent laws provide a system under which all inventions are subject to the same requirements of patentability regardless of the technical field in which they arose. However, inventors and innovative companies in different industries tend not to hold identical views concerning the importance of patents, reflecting varying experiences with the patent system. Innovators in biomedical industries tend to see patent protection as critically important as a way to prohibit competitors from appropriating the results of a company’s research and development efforts. Typically only a few, often one or two, patents cover a particular drug. In contrast, the nature of software development is such that inventions often are cumulative and new products generally embody numerous patentable inventions. As a result, it may be expected that distinct industries might react differently to the various patent reform proposals currently under consideration by Congress.

This report will be updated if events warrant such action.
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Introduction

Congress currently is considering reform of the existing patent system. This interest in patent policy reflects a recognition of the increasing importance of intellectual property to U.S. innovation. Patent ownership is perceived as an incentive to the technological advancement that leads to economic growth. As such, the number of patent applications and grants have grown significantly as have the type and breadth of inventions that can be patented. In 1980, 104,329 utility patent applications were received at the U.S. Patent and Trademark Office (USPTO); by 2005, this number had more than tripled to 381,797 applications. During the same time period, the number of U.S. utility patents granted grew from 61,819 to 151,079.1

Along with the expansion in the number and range of patents, there are growing concerns over whether the current system is working efficiently and effectively. Several recent studies (including those by the National Academy of Sciences and the Federal Trade Commission2) recommend patent reform. Other experts maintain that major alterations in existing law are unnecessary and that, while not perfect, the patent process can, and is, adapting to technological progress. To date, two bills, H.R. 2795 and, more recently, H.R. 5096, have been introduced in the 109th Congress which, if enacted, would make significant alterations in current patent law. (The specific legislative changes contained in H.R. 2795 are discussed in CRS Report RL32996, Patent Reform: Innovation Issues, by Wendy H. Schacht and John R. Thomas.)

The issue of patent reform has led to the emergence of several, often opposing, points of view. While the patent laws provide a system under which all inventions are treated the same regardless of the technical field, the varying experiences of companies in different industries often give rise to differing views concerning the importance and role of patents. Innovators in biomedical industries tend to see patent protection as critically important as a way to prohibit competitors from appropriating the results of a company’s research and development efforts. Typically only a few, often one or two, patents cover a particular drug. In contrast, the nature of software


development is such that inventions often are cumulative and new products generally embody numerous patentable inventions. Acknowledging these differences, this report explores the relationships between patents and innovation and looks at the role of intellectual property in the biomedical and software industries, two sectors where U.S. investment in research and development (R&D) has led to market leadership, a strong export position, and contributed to the Nation’s economic growth.

**Patents and Innovation**

Patent law is based upon the Patent Act of 1952, codified in Title 35 of the United States Code. According to the statute, one who “invents or discovers any new and useful process, machine, manufacture, or any composition of matter, or any new and useful improvement thereof, may obtain a patent therefore, subject to the conditions and requirements of this title.”\(^3\) Patents are issued by the United States Patent and Trademark Office (USPTO), generally for a term of 20 years from the date of filing. The patent grants its owner the right to exclude others from making, using, selling, offering to sell, or importing into the United States the patented invention. To be afforded patent rights, an invention must be judged to consist of patentable subject matter, possess utility, and be novel and nonobvious. The application must fully disclose and distinctly claim the invention for which protection is sought.

The grant of a patent does not necessarily provide the owner with an affirmative right to market the patented invention. For example, pharmaceutical products are also subject to marketing approval by the Food and Drug Administration (FDA).\(^4\) Federal laws typically require that pharmaceutical manufacturers demonstrate that their products are safe and effective in order to bring these drugs to the marketplace. USPTO issuance of a patent and FDA marketing consent are distinct events that depend upon different criteria.\(^5\)

Patent ownership is perceived to be an incentive to innovation, the basis for the technological advancement that contributes to economic growth. Patent title provides the recipient with a limited-time monopoly over the use of his discovery in exchange for the public dissemination of information contained in the patent application. Award of a patent is intended to stimulate the investment necessary to develop an idea and bring it to the marketplace embodied in a product or process, although it does not guarantee that the patent will generate commercial benefits. The

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requirement for publication of the patent is expected to stimulate additional innovation and other creative means to meet similar and expanded demands in the marketplace.

Innovation produces new knowledge. However, innovation typically is costly and resource intensive. Studies demonstrate that the rate of return to society as a whole generated by investments in research and development (R&D) leading to innovation is significantly larger than the benefits that can be captured by the person or organization financing the work.\(^6\) Some estimate that the social rate of return on R&D spending is over twice that of the rate of return to the inventor. Ideas often are easily imitated as the knowledge associated with an innovation is dispersed and adapted to other products and processes that, in turn, stimulate growth in the economy. Patents permit novel concepts or discoveries to become “property” when reduced to practice and therefore allow for control over their use.

Issuance of a patent furnishes the inventor with a limited-time exclusive right, the benefits of which are mitigated by other factors, particularly the requirements for information disclosure, the length of the patent, and the scope of rights conferred. The process of obtaining a patent places the concept on which it is based in the public domain. In return for a monopoly right to the application of the knowledge generated, the inventor must publish the ideas covered in the patent. As a disclosure system, the patent can, and often does, stimulate other firms or individuals to invent “around” existing patents to provide for parallel technical developments or meet similar market needs.

Patents may also provide a more socially desirable outcome than its chief legal alternative, trade secret protection. Trade secrecy guards against the improper appropriation of valuable, commercially useful information that is the subject of reasonable measures to preserve its secrecy.\(^7\) Taking the steps necessary to maintain secrecy, such as implementing physical security and enforcement, imposes costs that may ultimately be unproductive for society.\(^8\) Also, while the patent law obliges inventors to disclose their inventions to the public,\(^9\) trade secret protection requires firms to conceal them. The disclosure obligations of the patent system may better serve the objective of encouraging the diffusion of advanced technological knowledge. Patents may also prevent unproductive expenditures of time and money associated with R&D that duplicates other work.

\(^6\) For a list of relevant research in this area see Council of Economic Advisors, *Supporting Research and Development to Promote Economic Growth: The Federal Government’s Role*, October 1995, pp. 6-7.

\(^7\) American Law Institute, Restatement of Unfair Competition Third §39, 1995.


The patent system thus has dual policy goals — providing incentives for inventors to invent and encouraging inventors to disclose technical information. Disclosure requirements are factors in achieving a balance between current and future innovation through the patent process, as are limitations on scope, novelty mandates, and nonobviousness considerations. Patents give rise to an environment of competitiveness with multiple sources of innovation, which is viewed by some experts as the basis for technological progress. This is important because, as Robert Merges (now at the University of California, Berkeley) and Richard Nelson (Columbia University) found in their studies, in a situation where only “... a few organizations controlled the development of a technology, technical advance appeared sluggish.”

Not everyone agrees that the patent system is a particularly effective means to stimulate innovation. Some observers believe that the patent system encourages industry concentration and presents a barrier to entry in some markets. They suggest that the patent system often converts pioneering inventors into technological suppressors, who use their patents to block subsequent improvements and thereby impede technological progress. Others believe that the patent system too frequently attracts speculators who prefer to acquire and enforce patents rather than engage in socially productive activity such as bringing new products and processes to the marketplace.

Some experts argue that patents do not work as well in reality as in theory because they do not confer perfect appropriability. In other words, they allow the inventor to obtain a larger portion of the returns on his investment but do not permit him to capture all the benefits. Patents can be circumvented and infringement cannot always be proven. Thus, patents are not the only way, nor necessarily the most efficient means, for the inventor to protect the benefits generated by his efforts. A study by Yale University’s Richard Levin and his colleagues concluded that lead time, learning curve advantages (e.g. familiarity with the science and technology under consideration), and sales/service activities were typically more important in exploiting appropriability than were patents. That was true for both products and processes. However, patents were found to be better at protecting products than

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11 Kenneth W. Dam, “The Economic Underpinnings of Patent Law,” *Journal of Legal Studies*, January, 1994, pp. 266-267. Scope is determined by the number of claims made in a patent. Claims are the technical descriptions associated with the invention. In order for an idea to receive a patent, the law requires that it be “...new, useful [novel], and nonobvious to a person of ordinary skill in the art to which the invention pertains.”


processes. The novel ideas associated with a product often can be determined through reverse engineering — taking the item apart to assess how it was made. That information then could be used by competitors if not covered by a patent. Because it is more difficult to identify the procedures related to a process, other means of appropriation are seen as preferable to patents, with the attendant disclosure requirements.16

### Role of Patents in Biomedical R&D

The pharmaceutical industry perceives patents as critical to protecting innovation. Several studies over the years have demonstrated the important role patents play in the pharmaceutical sector. Of the 18 major manufacturing industries analyzed by Richard Levin and his colleagues, only drug companies rated product patents the most effective means of insuring that firms can capture the profits associated with their innovations.17 Later research by Professor Wesley Cohen (now at Duke University) and his colleagues demonstrated that patents were considered the most effective method to protect inventions in the drug industry, particularly when biotechnology is included.18 These studies reinforce earlier work by Edwin Mansfield that indicated 65% of pharmaceutical inventions would not have been brought to market without patent protection in contrast to the 8% of innovations made in other industries.19

Patents may be particularly important in the pharmaceutical sector because of the relative ease of replicating the finished product. Imitation costs vary among industries. For example, while it is expensive, complicated, and time consuming to duplicate an airplane, it is relatively simple to chemically analyze a pill and reproduce it.20 The degree to which industry perceives patents as effective has been characterized as “... positively correlated with the increase in duplication costs and time associated with patents.”21 In certain industries, patents significantly raise the costs incurred by nonpatent holders wishing to use the idea or invent around the patent — an estimated 40% in the pharmaceutical sector, 30% for major new

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17 Ibid., pp. 255 and 257.


21 Appropriating the Returns for Industrial Research and Development, p. 269.
chemical products, and 25% for typical chemical goods — and are thus viewed as significant. However, in other industries, patents have much smaller impact on the costs associated with imitation (e.g., in the 7%-15% range for electronics), and may be considered less successful in protecting resource investments.22

The costs associated with imitating pharmaceuticals “... are extremely low relative to the innovator’s costs for discovering and developing a new compound.”23 Studies by Dr. Joseph DiMasi of Tufts University and others indicate that the capitalized cost of bringing a new drug (defined as a “new molecular entity” rather than a new formulation of an existing pharmaceutical product) to the point of marketing approval is $802 million (2000 dollars).24 Additional research done by analysts at the Federal Trade Commission found the costs to be even higher; between $839 million and $868 million (2000 dollars).25 At the same time, the total capitalized costs appear to be growing at an annual rate of 7.4% above general price inflation.26

A large portion of new drug costs (in terms of money and time) are associated with the size and breath of clinical trials necessary to obtain FDA marketing approval. According to a study supported by the Federal Reserve Bank of Boston, only 10% of potential drug candidates reach the human trial phase and only a small portion of these actually reach the market.27 In research presented at a conference sponsored by the Federal Reserve Bank of Dallas, Duke University’s Henry Grabowski found that only 1% of drug compounds reach the human trial stage and 22% of those entering clinical trials receive FDA approval.28 Professor Iain Cockburn (Boston University) notes that “as drug discovery became more science-intensive, ... it became not just more expensive but also more difficult to manage.”29

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29 Iain Cockburn, “The Changing Structure of the Pharmaceutical Industry,” Health Affairs, (continued...)
Furthermore, returns to new drug introductions vary widely and the median new drug does not bring in sufficient profits to cover the costs of bringing the product to the marketplace.\(^{30}\) According to research by Professors Grabowski, John Vernon (Duke University), and DiMasi, only 34% of new drugs (new chemical entities) introduced generated profits that equaled the industry average R&D cost.\(^{31}\)

The significant costs of pharmaceutical R&D, coupled with the uncertainty of the clinical trial process, lend consequence to patents in this area because “… the disparity between the investments of innovators and those of imitators is particularly large in pharmaceuticals — almost as large as when software pirates simply copy the diskettes of an innovator.”\(^{32}\) While the capitalized cost of developing a new drug to the point of market approval is over $800 million, it takes only between $1 million and $2 million to obtain approval for a generic version of the pharmaceutical.\(^{33}\) This difference is a result of the costs associated with clinical trials needed to demonstrate the safety and efficacy of a new drug, data that could be utilized by generic companies if not protected by a patent.\(^{34}\) A generic company does not have to fund these studies to get FDA marketing approval; under the provisions of the Hatch-Waxman Act generic firms only have to prove that their product is “bioequivalent” to the innovator drug.\(^{35}\)

While patents are designed to spur innovation, some experts maintain that certain patents, particularly those on research tools\(^{36}\) in biotechnology, hinder the innovation process. Professors Rebecca Eisenberg (University of Michigan) and Richard Nelson (Columbia University) argue that ownership of research tools may “… impose significant transaction costs” that result in delayed innovation and possible future litigation.\(^{37}\) It also can stand in the way of research by others:

\(^{29}\) (...continued)

January/February 2004, p. 15.


\(^{31}\) *Returns on Research and Development for 1990s New Drug Introductions*, p. 23.

\(^{32}\) *The Economics of Human Gene Patents*, p. 1352.

\(^{33}\) *Patents, Innovation, and Access to New Pharmaceuticals*, p. 852.

\(^{34}\) *The Economics of Human Gene Patents*, p. 1352.


\(^{36}\) A biotechnology research tool is a cell line, reagent, or antibody used in research.

Broad claims on early discoveries that are fundamental to emerging fields of knowledge are particularly worrisome in light of the great value, demonstrated time and again in history of science and technology, of having many independent minds at work trying to advance a field. Public science has flourished by permitting scientists to challenge and build upon the work of rivals.38

Eisenberg and her colleague at the University of Michigan Law School, Michael Heller, contend that in the future scientists might need to obtain numerous patent licenses in order to undertake basic research.39 Similar concerns were expressed by Harold Varmus, President of Memorial Sloan-Kettering and formerly the Director of the National Institutes of Health. In July 2000 prepared testimony, he spoke to being “...troubled by widespread tendencies to seek protection of intellectual property increasingly early in the process that ultimately leads to products of obvious commercial value, because such practices can have detrimental effects on science and its delivery of health benefits.”40

However, other experts dispute this assertion. A study by Professors John Walsh (University of Illinois, Chicago), Ashish Arora (Carnegie Mellon University), and Wesley Cohen found that although there are now more patents associated with biomedical research, and on more fundamental work, there is little evidence that work has been curtailed due to intellectual property issues associated with research tools.41 Scientists are able to continue their research by “... licensing, inventing around patents, going offshore, the development and use of public databases and research tools, court challenges, and simply using the technology without a license (i.e., infringement).” According to the authors of the report, private sector owners of patents permitted such infringement in academia (with the exception of those associated with diagnostic tests in clinical trials) “... partly because it can increase the value of the patented technology.”

Role of Patents in the Software Industry

Over the past 25 years, there has been a demonstrable and sustained increase in the number of software patents granted in the United States. Research by James Bessen and Robert Hunt for the Federal Reserve Bank of Philadelphia noted that the 1,000 software patents issued annually in the early 1980s42 had increased to an annual total of 5,000 by 1990. Today over 20,000 software patents are granted each year.

38 Ibid.
42 There is no official USPTO category for “software” patents; Bessen and Hunt use their own definition.
While software patents comprised approximately 2% of all patents awarded in the early 1980s, they now account for approximately 15% of the total number of U.S. patent issued each year.43

Experts differ as to their assessment of the role of patents in promoting innovation in the computer software sector. This discussion centers around the issue of whether the increase in the number of patents is a result of inventive behavior generated by intellectual property protection or a result of changes in law during the 1980s and 1990s that made patents on software easier to obtain. Some experts argue that patent protection is not a significant factor in the development of computer software programs. Other analysts maintain that they play an important role in generating new technologies, particularly for small firms in the marketplace.

The nature of software development is such that inventions often are cumulative and new products generally embody numerous patentable inventions. This has led to what has been described by some observers as a

... poor match between patents and products in the [software] industry: it is difficult to patent an entire product in the software industry because any particular product is likely to include dozens if not hundreds of separate technological ideas.44

This situation may be augmented by the multiplicity of patents often associated with a finished computer product that utilizes the software. It is not uncommon for thousands of different patents (relating to hardware and software) to be embodied in one single computer. In addition, ownership of these patents may well be fractured among hundreds or thousands of different individuals and firms.

Studies by Bessen and Hunt explored the characteristics of software patents and determined that most are not owned by software companies but by large manufacturing companies. They found that

Firms in just three manufacturing industries (machinery, electronics, and instruments) alone accounted for 66 percent of software patents [yet]...Firms outside the manufacturing sector employed 90 percent of computer programmers, but together they accounted for only 25 percent of software patents.45

This data leads the authors to the conclusion that patents may not be closely tied to the development of new software technologies. Ownership of such patents is concentrated in sectors that have large patent portfolios and use them for strategic


purposes. Instead, they believe that companies are utilizing patents as a means to protect or leverage their investments rather than to generate more innovation through R&D spending.

In industries where innovation is sequential and complementary, as with software and computers, some experts argue that strong patents interfere with the innovation process. Inventions in these sectors typically are built upon earlier technologies and are integrated into existing systems. Commentators pose that patents inhibit or prevent enhancements to existing products because the patent owner may not have the interest or capability necessary to generate improvements at the same time that other firms cannot advance the technology without infringing on the original patent.

Not everyone agrees with this assessment. Professor Robert Merges maintains that patents have not hindered innovation in the software industry and that the significant ownership of title to inventions by large companies in this sector has not resulted in the demise of small firms developing new technologies. Analysis of software companies by Professor Ronald Mann (University of Texas, Austin) indicates the importance of software patents to small companies, particularly later-stage start-ups firms. He notes that the software industry is comprised primarily of small businesses and “the data suggests a different picture, one in which software R&D is impressively robust.” Mann’s research indicates that small firms spend proportionally more on software R&D than large companies. Research and development spending by software firms “...tends to be relatively stable over time as a percentage of sales. Indeed, company size seems to be more important in explaining variations in R&D spending within the industry.”

Mann’s research also indicates that the importance of software patents is dependent on where the firm is in its development process. Patents play a more significant role in later-stage start-up companies when firms can generate revenues through licensing. At that point, “... patents are useful as “barter” in cross-licensing agreements that the firm enters if it reaches a sufficiently mature stage to be a

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46 An Empirical Look at Software Patents, p. 4.
50 Do Patents Facilitate Financing in the Software Industry?, p. 1002.
51 Ibid., p. 1003.
52 Ibid., p. 985.
significant player in the industry. Patents may allow a firm to differentiate its areas of expertise and innovative activity.

Patents enable a company to transform ideas into a tangible form of property that can provide value. This can be useful in negotiations for the acquisition of the firm. While intellectual property is important to some investors but not to others, it is considered a significant factor when a company is involved in acquisition negotiations or in an IPO. It can prevent large companies from appropriating a small firm’s technology. Bradford Smith and Susan Mann, writing in the University of Chicago Law Review, concur with the argument that patents are beneficial for small, software firms. They maintain that patents prevent larger companies from utilizing the technologies developed by small businesses while allowing these companies to attract venture capital.

The multiplicity of patents involved in computer-related products has resulted in the extensive use of cross licensing in these industries such that one commentator argues: “licensing of software patents has become an industry unto itself.” Instead of promoting innovation, some experts maintain that the ownership of intellectual property has become an obstacle to the development and application of new ideas. The expansion in the number of patents associated with software is a consequence of the changes in patent law that make these patents easier to obtain, rather than an indication of increased innovative activity. There are indications, according to Bessen and Hunt, that patents are being substituted for increases in R&D. The substitution occurs in industries that patent strategically but not in other sectors. The propensity to patent software appears to be related to the utilization of the software by companies rather than to the R&D resources expended in developing the product. This is of interest because a rationale behind the patent system is that it provides incentives for the additional investments necessary to bring a product to the marketplace.

Concerns have been expressed in the academic community that the propensity to patent and the extensive use of cross licensing has resulted in a “patent thicket” where ownership of patent title is used to block others from innovating. According to Bessen and Hunt, “This may have increased the attractiveness of a strategy that

53 Ibid., p. 990.
54 Ibid., p. 985.
55 Ibid., p. 978.
58 The Software Patent Experiment, pp. 28-29.
59 An Empirical Look at Software Patents, p. 34.
60 The Software Patent Experiment, p. 27.
emphasizes patent rights over a strategy based on R&D.”61 However, other experts maintain that this might not be a true assessment of the situation. In an article for the Virginia Journal of Law and Technology, David Evans and Anne Layne-Farrar argue it is not clear that a patent thicket exists. “Other industries with longstanding histories of patenting could be categorized as having cumulative and sequential R&D, yet they do not display signs of innovation gridlock.”62 There are additional ways to prevent the use of patents to block innovation including the use of pro-competitive patent pools and antitrust enforcement.

Others agree that innovation in the software industry is not hindered by a patent thicket. In one study where actual software companies and investors were surveyed, the analyst found new companies were not concerned with existing patent portfolios as a barrier to their work as “none of the startup firms [interviewed] suggested a practice of doing prior art searches before beginning development of their products.”63 Because the software industry is so diverse, it is “... difficult for any single patent or group of patents to control a major part of the whole industry.”64

Concluding Observations

Innovators in the biomedical and software industries tend to view patents differently and thus may exhibit divergent positions on the issues surrounding patent reform. Patent protection is critically important to the pharmaceutical and biotechnology sectors as a way to prohibit competitors from appropriating the results of a company’s research and development efforts. However,

...patents are not among the key means used to protect innovations in either the computer or semiconductor industries. In those two industries, firms rely more heavily on secrecy, lead time and complementary capabilities to protect their inventions.65

A difference between the role of patents in the biomedical community and their role in the computer software sector lie with the dissimilar composition of the respective products. Typically only a few, often one or two, patents cover a particular drug. In contrast, the nature of software development is such that inventions often are cumulative and new products generally embody numerous patentable inventions. While few companies other than those that manufacture drugs need to deal with the relevant pharmaceutical patents,

61 Ibid., p. 30.
64 Ibid., p. 1007.
...computers are ubiquitous — and as a result, so is software authorship...Thus, a patent on a drug creates potential liability for those companies in the pharmaceutical business, while a software patent creates potential liability for any company with its own website or software customizations, regardless of its business.66

At the present time, the patent laws provide a system under which all inventions are subject to the same requirements of patentability regardless of the technical field in which they arose. However, as discussed in this paper, inventors and innovative companies in different industries may not hold identical views concerning the role of patents and may have had varying experiences with the patent system. As a result, it may be expected that distinct industries might react differently to the various patent reform proposals presently under consideration by Congress.