Patent-Eligible Subject Matter Reform: Background and Issues for Congress

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The statutory definition of patent-eligible subject matter under Section 101 of the Patent Act has remained essentially unchanged for more than two centuries. As a result, the scope of patentable subject matter—that is, the types of inventions that may be patented—has largely been left to the federal courts to develop through “common law”—like adjudication. In the 20th century, the U.S. Supreme Court established that three main types of discoveries are categorically patent-ineligible: laws of nature, natural phenomena, and abstract ideas.

A series of Supreme Court decisions in the 2010s broadened the scope of these three judicial exceptions to patent-eligible subject matter. Over a five-year period, the Supreme Court rejected, as ineligible, patents on a business method for hedging price-fluctuation risk; a method for calibrating the dosage of a particular drug; isolated human DNA segments; and a method of mitigating settlement risk in financial transactions using a computer. These cases established a new two-step test, known as the Alice/Mayo framework, for determining whether a patent claims ineligible subject matter.

The first step of the Alice/Mayo test addresses whether the patent claims are “directed to” a law of nature, natural phenomenon, or abstract idea. If not, the invention is patentable. If the claims are directed to one of the ineligible categories, then the second step of the analysis asks whether the patent claims have an “inventive concept.” To have an inventive concept, the patent claim must contain elements that transform the nature of the claim into a patent-eligible application of the ineligible concept, so that the claim amounts, in practice, to something “significantly more” than a patent on the ineligible concept itself. If the invention fails the second step of Alice/Mayo, then it is patent-ineligible.

The Supreme Court’s decisions have been widely recognized to effect a major change in the scope of patentable subject matter, restricting the sorts of inventions that are patentable in the United States. The Alice/Mayo test has been the subject of criticism, with some stakeholders arguing that the Alice/Mayo framework is vague and unpredictable, unduly restricts the scope of patentable subject matter, reduces incentives to invest and innovate, and harms American industry’s competitiveness. In particular, the Alice/Mayo test has created uncertainty in the computer technology and biotechnology industries as to whether innovations in medical diagnostics, personalized medicine, methods of treatment, computer software, and artificial intelligence are patent-eligible.

As a result, some patent law stakeholders—including academics, bar associations, industry representatives, judges, and former Patent and Trademark Office (PTO) officials—have called for the Supreme Court or Congress to act to change the law of patentable subject matter. Other stakeholders defend the legal status quo, arguing that the Alice/Mayo framework provides an important tool for combating unmeritorious patent litigation, or that the revitalized limits on patentable subject matter have important benefits for innovation.

Recently, there have been several substantial judicial, administrative, and legislative developments in patent-eligible subject matter law and potential reforms. On the judicial front, the Supreme Court has declined to hear further cases on this topic, despite calls by prominent stakeholders and judges on the U.S. Court of Appeals for the Federal Circuit. In 2019, the PTO issued and updated its guidance to clarify and improve predictability in how PTO patent examiners make Section 101 determinations, and in 2022 issued a new report on the topic. Following a series of hearings on the topic and draft legislative proposals in the 116th Congress, the 117th Congress saw several introduced bills seeking to reform the statutory standard for patentable subject matter.

Proposed changes to patent-eligible subject matter standards could have significant effects as to the types of technologies that are patentable. The availability of patent rights, in turn, affects incentives to invest and innovate in particular fields, as well as consumer costs and public access to technological innovation. Understanding the legal background and context of this complex issue may aid Congress as it debates the legal and practical effects that legislative Section 101 reforms would have if enacted.
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The statutory language governing patent-eligible subject matter—that is, the types of inventions that may be patented—has remained remarkably constant over the nearly 250-year history of U.S. patent law. Under the Patent Act of 1793, which Thomas Jefferson authored, “any new and useful art, machine, manufacture or composition of matter, or any new and useful improvement [of the same]” was patentable. Current law—Section 101 of the Patent Act of 1952—permits the patenting of “any new and useful process, machine, manufacture, or composition of matter, or any new and useful improvement thereof.” Through these four expansive statutory categories, Congress sought to ensure that nearly “anything under the sun made by man” is patentable if it meets all the requirements for patentability, such as novelty, enablement, and nonobviousness.

Consistent with its broad statutory language, Section 101 permits patenting in fields of applied technology such as pharmaceuticals, biotechnology, chemistry, computer hardware and software, electrical engineering, agriculture, mechanical engineering, and manufacturing processes. Even so, the Supreme Court has long read Section 101 as categorically prohibiting patents on three types of discoveries: “laws of nature, natural phenomena, and abstract ideas.” Even if “not required by the statutory text” of Section 101, the Court has held that these three judicial

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1 See generally Diamond v. Chakrabarty, 447 U.S. 303, 308–09 (1980) (tracing the history of statutory language on patentable subject matter). This observation—and this report more generally—is limited to traditional utility patents on useful inventions and discoveries. See 35 U.S.C. §§ 100–135. Congress did not provide patent protection for “original and ornamental designs for an article of manufacture” (design patents), id. §§ 171–173, and for “distinct and new variety[s] of plants” (plant patents), id. §§ 161–164, until 1842 and 1930, respectively. See An Act in addition to an act to promote the progress of the useful arts, and to repeal all acts and parts of acts heretofore made for that purpose, Pub. L. No. 27-263, 5 Stat. 543 (1842); An Act to provide for plant patents, Pub. L. No. 71-245, 46 Stat. 376 (1930).

2 See generally infra "Requirements for Patentability."


5 Chakrabarty, 447 U.S. at 308 (“In choosing such expansive terms as ‘manufacture’ and ‘composition of matter,’ modified by the comprehensive ‘any,’ Congress plainly contemplated that the patent laws would be given wide scope.”).


7 See 35 U.S.C. §§ 102–103, 112; see generally infra "Requirements for Patentability."


9 Diehr, 450 U.S. at 185.
exceptions “define[] the reach of the statute as a matter of statutory *stare decisis* going back 150 years.”

In a series of decisions in the 2010s, the Supreme Court relied on Section 101 to reject patent claims on

- a method for hedging price-fluctuation risks in commodity markets;
- a method for measuring metabolites in human blood to calibrate the dosage of particular drug;
- isolated human DNA segments; and
- a method of mitigating settlement risk in financial transactions using a computer.

These cases established a two-step test for patentable subject matter sometimes called the “*Alice/Mayo test*” or the “*Alice/Mayo framework*.” The Court’s decisions have been widely recognized to effect a major change in the scope of patentable subject matter, restricting the sorts of inventions that are patentable in the United States. The *Alice/Mayo* framework has thus shifted, for better or worse, the balance between encouraging innovation and the social costs of exclusive rights that is at the heart of patent law. The effects of this change have been particularly pronounced for computer technologies and biomedical technologies.

As a result, there is a significant and ongoing debate about the *Alice/Mayo framework*, with a number of patent law stakeholders questioning the Court’s patentable subject matter rulings. Critics argue that the *Alice/Mayo* framework is vague, unpredictable, and not administrable.

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11 *Id.* at 611–12.
17 See Bonito Boats, Inc. v. Thunder Craft Boats, Inc., 489 U.S. 141, 146 (1989) (“From their inception, the federal patent laws have embodied a careful balance between the need to promote innovation and the recognition that imitation and refinement through imitation are both necessary to invention itself and the very lifeblood of a competitive economy.”); Mark A. Lemley, Property, Intellectual Property, and Free Riding, 83 TEX. L. REV. 1031, 1031 (2005) (“[Traditionally,] the proper goal of intellectual property law is to give as little protection as possible consistent with encouraging innovation.”).
18 See PTO PSM REPORT, *supra* note 16, at 34–35 (finding “a general consensus that two industries have been most directly affected by the recent Supreme Court jurisprudence: life sciences and computer-related technologies”).
19 See generally *id.* at 27–34 (summarizing public comments that the *Alice/Mayo framework* is legally flawed, overly broad, unpredictable, and harmful to innovation).
20 *Id.* at 29–30 (describing public views that the Supreme Court “has failed to articulate objective, predictable criteria” for patentable subject matter). Hon. Paul R. Michel, The Supreme Court Saps Patent Certainty, 82 GEO. WASH. L. REV. 1751, 1758 (2014) (criticizing Court’s modern Section 101 jurisprudence as “subjective,” “indeterminate,” and “highly
muddies patent law by confusing patent eligibility with distinct patent law concerns, such as nonobviousness;\textsuperscript{21} reduces incentives to innovate and invest in particular industries, such as biotechnology;\textsuperscript{22} or puts U.S. industry at a disadvantage with international competitors.\textsuperscript{23} Other stakeholders defend the *Alice*/*Mayo* framework, arguing that the Court’s decisions are a part of the ordinary common law development of Section 101;\textsuperscript{24} an important tool for combating unmeritorious litigation\textsuperscript{25} or preventing overbroad or otherwise harmful patents;\textsuperscript{26} or beneficial to American consumers by lowering prices.\textsuperscript{27}

In response to stakeholder concerns, there have been several recent administrative and legislative developments that aim to clarify or reform the law of Section 101. In 2019, the Patent and Trademark Office (PTO) issued Revised Patent Subject Matter Eligibility Guidance designed to assist PTO patent examiners in determining patent eligibility with greater clarity and predictability.\textsuperscript{28} In the 116th Congress, Senators Thom Tillis and Chris Coons, along with Representatives Doug Collins, Hank Johnson, and Steve Stivers, released a “bipartisan, bicameral

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unpredictable”\textsuperscript{;} David O. Taylor, *Confusing Patent Eligibility*, 84 TENN. L. REV. 157, 158–60 (2016) (arguing that the Supreme Court’s Section 101 jurisprudence has created a “crisis of confusion” in patent law and that the doctrine “lacks administrability”).
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\textsuperscript{21} See PTO PSM REPORT, supra note 16, at 31–32; Michael Risch, *Everything Is Patentable*, 75 TENN. L. REV. 591, 598–606 (2008) (arguing that patentability criteria such as obviousness, novelty, utility, inventorship, written description, and enablement motivate the Supreme Court’s patentable subject matter decisions). But see Mark A. Lemley et al., *Life After Bilski*, 63 STAN. L. REV. 1315, 1319–32 (2011) (arguing that the preemption/overbreadth concerns driving Section 101 are not a part of the Supreme Court’s jurisprudence).

\textsuperscript{22} See, e.g., PTO PSM REPORT, supra note 16, at 32–33, 35–38; BCLT Report, supra note 16, at 582–84; Taylor, supra note 20, at 240 (“[The *Alice*/*Mayo* framework] substantially reduces incentives to invest in research and development, particularly in the biotechnology and software technology areas.”).

\textsuperscript{23} See PTO PSM REPORT, supra note 16, at 34; Ryan Davis, *Kappos Calls for Abolition of Section 101 of Patent Act*, LAW360 (Apr. 12, 2016), https://www.law360.com/articles/783604/kappos-calls-for-abolition-of-section-101-of-patent-act (quoting former PTO Director David Kappos as stating that international competitors “no longer have to steal U.S. technology in [biotechnology and software], since they can now take it for free”); Robert L. Stoll, *Courts Are Making Bad Patent Law*, THE HILL (July 16, 2015), https://thehill.com/blogs/pundits-blog/the-judiciary/248054-courts-are-making-bad-patent-law (“The courts’ focus on subject matter eligibility as a mechanism to deny patents for [inventions in diagnostics and personalized medicine] will drive investment into research in these technologies to other areas. We will lose our edge in the world . . . .”).

\textsuperscript{24} See PTO PSM REPORT, supra note 16, at 23–24.

\textsuperscript{25} See id. at 24; BCLT Report, supra note 16, at 555 (“Many technology companies that rely on software innovation . . . welcomed the tightening of patent eligibility standards on software claims and the opportunity to seek early dismissals of lawsuits.”); Paul R. Gugliuzza, *Quick Decisions in Patent Cases*, 106 GEO. L.J. 619, 652–53 (2018) (“The invigoration of the [patent] eligibility requirement can help courts resolve infringement disputes more quickly and efficiently by allowing validity to be resolved on the pleadings as a matter of law.”).

\textsuperscript{26} See *The State of Patent Eligibility in America: Part I: Hearing Before the S. Judiciary Comm., Subcomm. on Intellectual Property*, 116th Cong. (2019) (statement of Prof. Joshua D. Sarnoff, DePaul University College of Law), at 3–8, https://www.judiciary.senate.gov/download/sarnoff-testimony [hereinafter Sarnoff Testimony]; accord *Mayo Collaborative Servs. v. Prometheus Labs.*, Inc., 566 U.S. 66, 86 (2012) (“[E]ven though rewarding with patents those who discover new laws of nature and the like might well encourage their discovery, those laws and principles, considered generally, are the basic tools of scientific and technological work. And so there is a danger that the grant of patents that tie up their use will inhibit future innovation . . . .” (citations omitted)); Lemley et al., supra note 21, at 1329 (arguing that Section 101’s abstract ideas doctrine is “about encouraging cumulative innovation and furthering societal norms regarding access to knowledge”).

\textsuperscript{27} PTO PSM REPORT, supra note 16, at 27.

framework” for legislative Section 101 reform, and a draft bill to reform Section 101. After the release of the draft bill, the Senate Judiciary Committee’s Intellectual Property Subcommittee held three public hearings on Section 101 reform. These efforts did not result in formal legislation introduced by these Members during the 116th Congress. In the 117th Congress, Senator Tillis and Representative Thomas Massie have introduced bills on patent-eligible subject matter.

This report provides the necessary background and context to understand the legal and practical effects that these legislative reforms would have if enacted. First, the report reviews the basic legal principles of the U.S. patent system. Second, it examines the historical development and current state of patentable subject matter law. Third, it reviews several articulated rationales for Section 101 and potential options for Section 101 reform. Finally, it examines recent judicial, administrative, and legislative developments concerning patent-eligible subject matter, including the proposed legislative reforms to Section 101.

This report focuses on patent-eligible subject matter reform from a legal perspective. For an analysis of these issues as they relate to innovation policy, see CRS Report R47267, Patents and Innovation Policy, by Emily G. Blevins.

**Patent Law Background**

Congress’s authority to grant patents derives from the Intellectual Property (IP) Clause of the U.S. Constitution, which grants Congress the power “[t]o promote the Progress of Science and useful Arts, by securing for limited Times to . . . Inventors the exclusive Right to their . . . Discoveries.” Patents are generally available to any person who “invents or discovers any new

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34 U.S. Const. art. I, § 8, cl. 8.
and useful process, machine, manufacture, or composition of matter, or any new and useful improvement thereof.”

Patent rights do not arise automatically. Rather, to obtain patent protection under the Patent Act, an inventor must formally apply for a patent with the PTO, beginning a process called patent prosecution. During prosecution, a patent examiner at the PTO evaluates the patent application to ensure that it meets all the applicable legal requirements to merit the grant of a patent. To be patentable, an invention must be (1) directed at patent-eligible subject matter, (2) useful, (3) new, (4) nonobvious, and (5) adequately disclosed and claimed in the patent application. If the PTO finds these requirements met, it will issue (i.e., grant) the patent. Patents typically expire 20 years after the initial patent application.

The current law of patent-eligible subject matter will be discussed separately in detail below. The remainder of this section briefly reviews the other requirements for patentability, the scope and effect of patent claims, and the legal rights granted to the holder of a valid patent.

Requirements for Patentability

Section 101: Utility

Along with its subject matter requirements, Section 101 contains a requirement that a patented invention must be “useful.” In particular, courts have held that an invention must have both a specific and substantial utility to be patentable. The utility requirement derives from the Constitution’s command that patent laws exist to “promote the Progress of . . . useful Arts.” The constitutional purpose of patent law thus requires a “benefit derived by the public from an invention with substantial utility,” where the “specific benefit exists in currently available form.” This standard for utility is low, however, requiring only that the claimed invention have some “significant and presently available benefit to the public” that “is not so vague as to be meaningless.”

Section 102: Novelty

Perhaps the most fundamental requirement for patentability is that the claimed invention must be new. The PTO will not issue a patent if “the claimed invention was patented, described in a printed publication, or in public use, on sale, or otherwise available to the public before the

39 See id. §§ 101–103, 112.
40 Id. § 131.
41 Id. § 154(a)(2).
42 See infra “The Current Law of Section 101.”
46 Brenner, 383 U.S. at 534–35.
47 In re Fisher, 421 F.3d at 1371–72.
effective filing date of the claimed invention.” In other words, if every limitation of the claimed invention is already disclosed in the “prior art”—the information available to the public at the time of the patent application—then the alleged inventor “has added nothing to the total stock of knowledge,” and no valid patent may issue to her.

Section 103: Nonobviousness

Even if a claimed invention is novel in the narrow sense that it is not “identically disclosed” in a prior-art reference (such as an earlier patent or publication), the invention must further be nonobvious to be patentable. Specifically, an invention cannot be patented if “the differences between the claimed invention and the prior art are such that the claimed invention as a whole would have been obvious . . . to a person having ordinary skill” in the relevant technology. When determining obviousness, courts also evaluate secondary considerations (also known as “objective indicia”) of nonobviousness such as “commercial success, long felt but unsolved needs, [or] failure of others . . . to give light to the circumstances surrounding the origin of the subject matter sought to be patented.” By its nature, obviousness is an “expansive and flexible” inquiry that cannot be reduced to narrow, rigid tests. Nonetheless, if an invention merely combines “familiar elements according to known methods,” yielding only “predictable results,” it is likely to be obvious.

Section 112(a): Written Description, Enablement, Best Mode

Finally, the Patent Act imposes several requirements relating to the technical disclosures in the patent application. These provisions are intended to ensure that the patent adequately describes the invention such that the public can use the invention after the expiration of the patent term. Section 112(a) of the Patent Act requires that patents must contain a “specification” that includes

a written description of the invention, and of the manner and process of making and using it, in such full, clear, concise, and exact terms as to enable any person skilled in the art to . . . make and use the same, and shall set forth the best mode contemplated by the inventor or joint inventor of carrying out the invention.

This statutory language yields three basic disclosure requirements for patentability. First, to satisfy the written description requirement, the specification must “reasonably convey[] to those

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48 35 U.S.C. § 102(a)(1). There are certain exceptions to this requirement when, for example, the prior-art disclosure derives from the inventor and the patent application is made within one year of the disclosure. Id. § 102(b)(1).
49 Great Atl. & Pac. Tea Co. v. Supermarket Equip. Corp., 340 U.S. 147, 153 (1950); Graham v. John Deere Co. of Kan. City, 383 U.S. 1, 6 (1966) (“Congress may not authorize the issuance of patents whose effects are to remove existing knowledge from the public domain, or to restrict free access to materials already available.”).
51 Id. Patent law often relies on the concept of a “person having ordinary skill in the art,” a “hypothetical person” with a typical level of skill in the relevant technology who is “presumed to be aware of all the pertinent prior art” in the particular field. See Standard Oil Co. v. Am. Cyanamid Co., 774 F.2d 448, 454 (Fed. Cir. 1985).
52 Graham, 383 U.S. at 17–18; see also Apple Inc. v. Samsung Elecs. Co., 839 F.3d 1034, 1048 (Fed. Cir. 2016) (en banc) (“Objective indicia of nonobviousness must be considered in every case where present.”).
54 Id. at 416.
56 35 U.S.C. § 112(a) (emphases added).
skilled in the art that the inventor had possession of the claimed subject matter as of the filing date” of the patent application.58 Second, to satisfy the enablement requirement, the specification must contain enough information to teach a person skilled in the art how “to make and use the invention without undue experimentation.”59 Finally, to satisfy the best mode requirement, if the inventor knew of a preferred way of practicing her invention at the time of the patent application, the specification must disclose that “preferred embodiment[]” of the invention. 60

Patent Claims

Section 112(b): Definiteness

If granted, the legal scope of the patent is defined by the patent claims, a sequence of statements that formally set forth the patentee’s asserted rights. In essence, while the specification explains the invention in a technical sense, the claims set forth the legal effect of the patent.61 Much as a deed may describe the boundaries of a tract of land, the claims define the “metes and bounds” of the patent right.62 Patent claims must be sufficiently definite to be valid—that is, they must “particularly point[] out and distinctly claim[] the subject matter which the inventor . . . regards as the invention.”63 In other words, when the claims are read in context, they must “inform, with reasonable certainty, those skilled in the art about the scope of the invention.”64

Section 112(f): Functional Claiming

For the most part, the current Patent Act uses a system of peripheral claiming, in which the patent claims formally set out the outer boundaries of the patentee’s rights.65 However, the Patent Act

58 Ariad, 598 F.3d at 1351.
59 In re Wands, 858 F.2d 731, 735 (Fed. Cir. 1988).
60 Eli Lilly & Co. v. Barr Labs., Inc., 251 F.3d 955, 963 (Fed. Cir. 2001). Failure to disclose the best mode is not a basis on which a patent claim can be invalidated in subsequent patent infringement proceedings. 35 U.S.C. § 282(b)(3)(A).
61 See Ariad, 598 F.3d at 1347 (Fed. Cir. 2010); In re Vamco Mach. & Tool, Inc., 752 F.2d 1564, 1577 n.5 (Fed. Cir. 1985).
63 35 U.S.C. § 112(b); Laitram Corp. v. NEC Corp., 163 F.3d 1342, 1347 (Fed. Cir. 1998) (“[T]he claims, not the written description, which define the scope of the patent right.”).
65 See 35 U.S.C. § 112(b); Mark A. Lemley, Software Patents and the Return of Functional Claiming, 2013 Wis. L. Rev. 905, 911 (2013) (“Today, peripheral claiming is universal [in patent law]; patentees write claims in an effort to define the outer boundaries of their invention.”); Jeanne C. Fromer, Claiming Intellectual Property, 76 U. Chi. L. Rev. 719, 725–30 (2009) (explaining the distinction between peripheral and central claiming systems for intellectual property). Until the late 19th century, however, central claiming prevailed: the patentee had only to describe the core principle or an example of his invention, and courts would decide whether the accused infringer’s product or method was sufficiently similar to the patentee’s invention to infringe the patent. See Lemley, supra, at 910–11; Fromer, supra, at 731–33. Peripheral claiming began as a defensive strategy by patentees to describe their invention at a higher level of generality, and the gradual switch toward the modern patent claiming was eventually codified in the Patent Act in 1870. See An Act to revise, consolidate, and amend the Statutes relating to Patents and Copyrights, Pub. L. No. 41-230 § 26, 16 Stat. 198, 201 (1870) (requiring patent applicant to “particularly point out and distinctly claim the part, improvement, or combination which he claims as his invention or discovery”); see generally Fromer, supra, at 731–35 (reviewing American patent law’s historical shift from central to peripheral claiming); Dan L. Burk & Mark A. Lemley, Fence Posts or Sign Posts? Rethinking Patent Claim Construction, 157 U. Pa. L. Rev. 1743, 1766–71 (2009) (same). This account of patent-claiming history is somewhat simplified: notably, despite the 1870 statutory shift, the Patent Act retained (and retains) features of central claiming. See Burk & Lemley, supra, at 1771 (“[I]t may be fairer to say that during the twentieth century we had not a peripheral-claiming system, but a hybrid peripheral claiming system.”).
still retains elements of its former system of central claiming, in which the patentee would describe the core principles or examples of what he had invented, but need not formally delineate the outer boundaries of his rights.66 For example, under the doctrine of equivalents, an accused infringer may be found liable even if his product does not literally meet every element of the patent claims, if the differences between a claim element and its alleged equivalent in the accused product are “insubstantial.”67

A potential danger of a peripheral claiming system is that patentees may seek to claim more than they invented by couching the patent claims in broad, functional language—that is, by claiming a result or goal without limitation to any specific structure or device that accomplishes the result.68 In Halliburton Oil Well Cementing Co. v. Walker, the Supreme Court limited this practice, invalidating as indefinite a “functional” patent claim, in which the invention—an apparatus for determining the location of an obstruction in an oil well—was claimed not in terms of specific machinery, but instead as a “means for” performing various functions.69

Functional claims (also known as “means-plus-function” claims) such as those in Halliburton may be convenient for the patentee, who can express a claim element in terms of a general end, rather than an “exhaustive list” of every possible apparatus that could be used to perform that goal.70 On the other hand, as Halliburton recognized, functional claims may be overbroad and ambiguous, or permit the patentee to claim more than he actually invented.71 In the Patent Act of 1952, Congress enacted current Section 112(f) as a compromise for functional claims, overruling Halliburton72 but providing a standard to make functional claims more definite.73

Under Section 112(f), a patentee may opt to express a claim element as “a means or step for performing a specified function without the recital of structure, material, or acts in support thereof.”74 If the patentee chooses to claim functionally, however, the claim is construed not to cover all possible means of performing the function, but only “the corresponding structure, material, or acts described in the specification and equivalents thereof.”75 Courts have held that a

68 See Lemley, supra note 65, at 911–13. Such claiming should in theory be prohibited on novelty or enablement grounds, see 35 U.S.C. §§ 102, 112(a), but the problem persists, for example, in modern software patents. See Lemley, supra note 65, at 921–23 (citing examples).
69 See 329 U.S. 1, 8–9, 12–13 (1946).
70 Stephen Winslow, Means for Improving Modern Functional Patent Claiming, 98 Geo. L.J. 1891, 1892 (2010) (“A patent can be clearer, more concise, and more comprehensible when the patentee drafts her claims using language describing what a particular element does, rather than giving an exhaustive list of the various structures that could provide that function within her invention.”).
71 See Halliburton, 329 U.S. at 12.
72 See Williamson v. Citrix Online, LLC, 792 F.3d 1339, 1347 (Fed. Cir. 2015) (en banc) (“In enacting § 112(f), Congress struck a balance in allowing patentees to express a claim limitation by reciting a function to be performed rather than by reciting structure for performing that function, while placing specific constraints on how such a limitation is to be construed . . . .”); P.J. Federico, Commentary on the New Patent Act (West 1954), reprinted in 75 J. Pat. & Trademark Off. Soc’y 161, 186 (1993) (observing that “[t]he last paragraph of section 112” means that “decisions such as that in [Halliburton Oil] are modified or rendered obsolete . . . .”).
75 Id. (emphasis added).
The patentee is presumed to invoke Section 112(f) when the term “means” is used in the claims. Conversely, there is a presumption that the patentee does not invoke Section 112(f) if she does not use the term “means,” but that presumption may be overcome, such that Section 112(f) will apply to any claim that fails to recite a “sufficiently definite structure” for performing a function.

**Rights of Patent Holders**

With some exceptions, a patent is generally granted “for a term beginning on the date on which the patent issues and ending 20 years from the date on which the application for the patent was filed.” The Patent Act includes provisions that may modify the 20-year term, including to account for excessive delays in patent examination at the PTO, or delays associated with obtaining marketing approval from other federal agencies.

Once granted, a valid patent gives the patent holder the exclusive right to make, use, sell, or import the invention in the United States until the patent expires. Any other person who practices the invention (i.e., makes, uses, sells, offers to sell, or imports it) without permission from the patent holder infringes the patent and may be liable for monetary damages and injunctive relief if sued by the patentee. To obtain relief from infringement, the patentee must generally sue in court. Patent law is an area of exclusive federal jurisdiction, and the traditional forum for most patent disputes is federal district court. Although patent suits may be filed in any district court across the country with jurisdiction over the defendant and proper venue, a single specialized court, the U.S. Court of Appeals for the Federal Circuit (Federal Circuit), hears all appeals in patent cases.

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76 Williamson, 792 F.3d at 1348 (quoting Watts v. XL Sys., Inc., 232 F.3d 877, 880 (Fed. Cir. 2000)).
77 Id.
79 Id. § 154(b)(1).
80 Id. § 156. In the pharmaceutical context, patents claiming a drug product or medical device (or a method of using or manufacturing the same) may be extended for up to five years to account for delays in obtaining regulatory approval, if certain statutory conditions are met. See Eli Lilly & Co. v. Medtronic, Inc., 496 U.S. 661, 670–71 (1990); Merck & Co. v. Hi-Tech Pharmacal Co., 482 F.3d 1317, 1320–21 (Fed. Cir. 2007); Stephanie Plamondon Bair, Adjustments, Extensions, Disclaimers, and Continuations: When Do Patent Term Adjustments Make Sense?, 41 CAP. U. L. REV. 445, 460 (2013).
82 Id. §§ 271, 281, 283–85.
Defending Against Patent Suits

Parties accused of patent infringement may defend on several grounds. First, the accused infringer may claim an “absence of liability” because of noninfringement. In other words, even presuming the patent is valid, the patentee may fail to prove that the activities of the accused infringer fall within the scope of the patent claims—that is, the accused infringer is not making, using, selling, or importing the patented invention. Second, although patents benefit from a presumption of validity, the accused infringer may assert that the patent is invalid. To prove invalidity, the accused infringer must show, by clear and convincing evidence, that the PTO should not have granted the patent because it failed to meet the requirements for patentability.

Thus, for example, the accused infringer may argue that the invention lacks novelty, is obvious, or claims nonpatentable subject matter; that the patent fails to enable the invention; or that the patent claims are indefinite. Finally, the accused infringer may assert as a defense that the patent is unenforceable based on the inequitable or illegal activities of the patent holder, such as obtaining the patent through fraud on the PTO. While the patent holders bears the burden of proving infringement, the accused infringer bears the burden of proving invalidity or inequitable conduct.

Following the passage of the 2011 Leahy-Smith America Invents Act (AIA), the Patent Trial and Appeal Board (PTAB) has become an increasingly important forum for patent disputes. The AIA created several new administrative procedures for challenging patent validity, including (1) post-grant review (PGR), which allows any person to challenge patent validity based on any of the requirements of patentability if the PGR petition is filed within nine months of the patent’s issuance; (2) inter partes review (IPR), which allows any person other than the patentee to challenge patent validity on limited grounds (novelty or obviousness based on prior patents or printed publications) at any time after nine months following the patent’s issuance; and (3) a transitional program for covered business method patents (CBM), a PGR-like process limited to

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89 35 U.S.C. § 282(a), (b)(2)–(3).
90 Id. § 282(b)(2)–(3); Microsoft Corp. v. i4i Ltd. P’ship, 564 U.S. 91, 95–96 (2011).
91 See supra “Requirements for Patentability.”
94 35 U.S.C. § 282(a); Therasense, 649 F.3d at 1291.
98 Id. §§ 311–319.
certain patents claiming “business methods” that was available only through September 2020.\footnote{100} Of these procedures, IPR is by far the most widely used.\footnote{101}

The Current Law of Section 101

At a general level, there are two basic requirements for an invention to claim patent-eligible subject matter. First, the invention must fit into one or more of the four statutory categories in Section 101—the claimed invention must be a (1) process, (2) machine, (3) manufacture, or (4) composition of matter.\footnote{102} Given the (intentionally) expansive nature of these terms, nearly all claimed inventions will satisfy this requirement.\footnote{103} Still, exceptions to this rule do exist. For example, in \textit{In re Nuijten}, the Federal Circuit held that a transitory electromagnetic signal was neither a process, machine, manufacture, or composition of matter, and was therefore not patent-eligible subject matter.\footnote{104}

Because most claimed inventions fit into one of the four statutory categories, the second requirement tends to be more practically important, and receives more attention.\footnote{105} The second patentable subject matter requirement is that the invention cannot claim one of the judicially created categories of ineligible subject matter. That is, the claimed invention must \textit{not} be a (1) law of nature, (2) natural phenomenon, or (3) abstract idea.\footnote{106} As explained below, the modern Supreme Court has articulated a two-step test for this second requirement, known as the \textit{Alice}/\textit{Mayo} framework.\footnote{107}

\begin{footnotesize}
\begin{itemize}
\item \footnote{100}{Pub. L. No. 112-29, § 18, 125 Stat 284, 329–30 (2011) (not codified in U.S.C.).}
\item \footnote{101}{\textit{See} 2018 Patent Dispute Year in Review, supra note 85 (finding that IPRs constituted 93.9\% of petitions submitted to the PTAB in 2018).}
\item \footnote{102}{35 U.S.C. § 101.}
\item \footnote{103}{\textit{See} Lemley et al., supra note 21, at 1328 (“[P]atent claims almost never fall outside of the four fundamental categories of § 101 . . . .”).}
\item \footnote{104}{500 F.3d 1346, 1354–57 (Fed. Cir. 2007).}
\item \footnote{105}{\textit{See} Kevin Emerson Collins, \textit{Patent-Ineligibility As Counteraction}, 94 WASH. U. L. REV. 955, 968 (2017) (“Contemporary debates over patent-ineligibility rarely parse the plain meanings of [the four statutory categories]. They focus instead on a set of judicial exclusions from patent-eligibility that are not expressly codified in the statute: laws of nature, products of nature, and abstract ideas . . . .”).}
\item \footnote{106}{Diamond v. Diehr, 450 U.S. 175, 185 (1981). \textit{Diehr}'s modern distillation of patentable subject matter doctrine to these three categories is a somewhat simplified version of the doctrine’s historical development, which often identified patent-ineligible categories in addition to these three. \textit{See}, e.g., Daniel J. Klein, \textit{The Integrity of Section 101: A 'New and Useful' Test for Patentable Subject Matter}, 93 J. PAT. & TRADEMARK OFF. SOC’Y 287, 288 (2011) (listing eight terms that the Court has used to denote patent-ineligible subject matter); Michel, \textit{supra} note 20, at 1757 (counting six categories of patent-ineligible subject matter); \textit{accord} Emily Michiko Morris, \textit{Intuitive Patenting}, 66 S.C.L. REV. 61, 66 n.31 (2014) (describing the Supreme Court’s patentable subject matter jurisprudence as “insolubly murky”).}
\end{itemize}
\end{footnotesize}
The Supreme Court has justified the three ineligible categories as necessary to prevent patent monopolies on the “‘basic tools of scientific and technological work,’” which “might tend to impede innovation more than it would tend to promote it.” Thus, the Court has explained that “a new mineral discovered in the earth or a new plant found in the wild is not patentable subject matter. Likewise, Einstein could not patent his celebrated law that E=mc²; nor could Newton have patented the law of gravity.” At the same time, the Court has said that even if a mathematical formula or law of nature is not patentable “in the abstract,” a practical application of such a principle or law “to a new and useful end” is patent-eligible.

Beyond such broad illustrations, it is not easy to define what an “abstract idea,” “law of nature,” or “natural phenomenon” is. Because these exceptions to patent-eligible subject matter are judicially created, they have no formal statutory definition; their meaning has instead been developed through two centuries of case-by-case “common law” adjudication in the federal courts. As a result, the scope of patentable subject matter has waxed and waned over time, depending on the trends in judicial decisions.

This section overviews the leading Supreme Court cases addressing patent-eligible subject matter, beginning with formative cases from the 19th century and culminating in the series of 2010s Supreme Court decisions that have led some to call for legislative reform of Section 101. Table 1 summarizes the facts and holdings of the major cases.

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111 See Morris, supra note 106, at 62 (describing the Supreme Court’s patentable subject matter jurisprudence as “insolubly murky”); Klein, supra note 106, at 289 (describing the three categories of nonpatentable subject matter as “metaphysically vague and extra-statutory”); Funk Bros., 333 U.S. at 134–35 (Frankfurter, J., concurring) (“It only confuses the issue, however, to introduce such terms as ‘the work of nature’ and the ‘laws of nature.’ For these are vague and malleable terms infected with too much ambiguity and equivocation. Everything that happens may be deemed ‘the work of nature,’ and any patentable composite exemplifies in its properties ‘the laws of nature.’”).
112 See, e.g., Peter S. Menell, Forty Years of Wondering in the Wilderness and No Closer to the Promised Land: Bilski’s Superficial Textualism and the Missed Opportunity to Return Patent Law to Its Technology Mooring, 63 STAN. L. REV. 1289, 1307 (2011) (“Since the founding of our nation, courts have evolved [patentable subject matter limits] within a hybrid constitutional/common law tradition.”); Lemley et al., supra note 21, at 1325 (describing the three judicially created ineligible categories as “common law exceptions” to patentable subject matter).
Historical Development of the Judicial Exceptions to Patent-Eligible Subject Matter

Nineteenth Century

The 1853 case of *Le Roy v. Tatham*, the “fountainhead” of American patentable subject matter jurisprudence,114 concerned a patent on machinery to manufacture metal pipes that exploited a newly developed property of lead.115 Although the Court ultimately did not decide the case on subject matter grounds,116 *Le Roy* relied on influential English patent cases117 to set forth a basic distinction between abstract “principles” and natural laws (which may not be patented) and practical applications of those principles (which may be patented).118 The Court stated that “[a] principle, in the abstract, is a fundamental truth; an original cause; a motive; these cannot be patented, as no one can claim in either of them an exclusive right.”119 On the other hand, a “new property discovered in matter, when practically applied, in the construction of a useful article of commerce or manufacture, is patentable,” for the “invention is not in discovering [the natural principles], but in applying them to useful objects.”120

In its next term, the Court applied this rule, in the famous case of *O’Reilly v. Morse*,121 to Samuel Morse’s patent on the telegraph. Although the Court found that Morse was the first inventor of the telegraph and sustained much of his patent,122 the Court rejected Morse’s eighth claim to any “use of the motive power of the electric or galvanic current . . . however developed for marking or printing intelligible characters, signs, or letters, at any distances, being a new application of that power of which I claim to be the first inventor or discoverer.”123 Observing that “the discovery of a principle in natural philosophy or physical science, is not patentable,”124 Chief Justice Roger Taney’s majority opinion held that Morse’s eighth claim was “too broad” because he had not discovered “that the electric or galvanic current will always print at a distance, no matter what may be the form of the machinery” used, but only that the specific machinery disclosed in the patent specification would do so.125

In the second half of the nineteenth century, the Court issued a series of important decisions on the patentability of processes. The result of these cases was a move away from an earlier rule that prohibited “pure” method patents as ineligible (i.e., a process claimed independently of the

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114 See, e.g., Lefstin, supra note 113, at 594 (describing *Le Roy* as “the fountainhead of subject-matter exclusion in American patent law”); Menell, supra note 112, at 1296 (describing *Le Roy* as “the foundation for much patentable subject matter jurisprudence”).


116 The dispositive issue in the case was the scope of the patent claims. See infra note 180; Lefstin, supra note 113, at 595 (“The outcome in *Le Roy* therefore turned entirely on the Court’s narrow construction of the claim.”).

117 For a full historical account of these English cases and how they shaped the Supreme Court’s jurisprudence, see Lefstin, supra note 113, at 577–644.


119 Id. at 175.

120 Id.

121 56 U.S. 62 (1853).

122 Id. at 111–12, 123–24.

123 Id. at 112–20.

124 Id. at 116.

125 Id. at 117, 119.
specific machinery used to accomplish the method) either by construing nominal process patents as claiming a machine or limiting the process patents to the machinery disclosed and its equivalents.\(^{126}\) In *Cochrane v. Deener*, which involved a patent on an improved manufacturing process for flour, the Court defined a patentable process as “a mode of treatment of certain materials to produce a given result. It is an act, or a series of acts, performed upon the subject-matter to be transformed and reduced to a different state or thing.”\(^{127}\) *Cochrane* held that such methods are patentable “irrespective of the particular form of the instrumentalities used.”\(^{128}\) Similarly, in *Tilghman v. Proctor*, the Court held that a method for separating fat into glycerin and fatty acids using water, pressure, and heat was patentable.\(^{129}\)

In *The Telephone Cases*, the Court distinguished *Morse* to allow Alexander Graham Bell’s patent claim on a “method of and apparatus for transmitting vocal or other sounds telegraphically, as herein described, by causing electrical undulations, similar in form to the vibrations of the air accompanying the said vocal or other sounds, substantially as set forth.”\(^{130}\) Chief Justice Edward Douglass White interpreted *Morse* as holding that “the use of magnetism as a motive power, without regard to the particular process with which it was connected in the patent, could not be claimed, but that its use in that connection could.”\(^{131}\) The Court found that Bell’s claim, unlike Morse’s, did not reach uses of electricity to transmit speech that are “distinct from the particular process with which it is connected in [Bell’s] patent,” and upheld the claim, so construed.\(^{132}\)

**Twentieth Century**

In the first half of the 20th century, the Court decided two major cases on the patentability of natural phenomena. In *American Fruit Growers v. Brogdex Co.*, the Court rejected patent claims on citrus fruit treated with a solution of borax to render it resistant to mold.\(^{133}\) The Court held that treated fruit was not a “manufacture” under Section 101, but a patent-ineligible “natural article”; treatment with borax did not effect a “change in the name, appearance, or general character of the fruit” or imbue it with a “new or distinctive form, quality, or property.”\(^{134}\) In *Funk Brothers Seed Co. v. Kalo Inoculant Co.*, the Court rejected patent claims on an inoculant for leguminous plants consisting of multiple species of bacteria, where the particular bacterial strains were selected to avoid inhibiting each other (as prior multispecies combinations had).\(^{135}\) Because the patentee’s combination “produces no new bacteria [and] no change in the six species of bacteria,” Justice

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\(^{126}\) See, e.g., *Corning v. Burden*, 56 U.S. (15 How.) 252, 268–70 (1853) (construing “equivocal” patent to claim a machine, and not a process, to save its validity because a “process” in the sense of “the function of a machine, or the effect produced by it” cannot be patented); see generally *Sarnoff*, supra note 113, at 67 (“[A]t the end of the eighteenth century, pure method patents—methods claiming all future applications and not merely those substantially similar to the disclosed implementing machinery and their equivalents—were ineligible for protection and remained so until the late nineteenth century.”) & id. n. 88 (collecting cases).

\(^{127}\) 94 U.S. 780, 788 (1876).

\(^{128}\) Id. at 787.

\(^{129}\) 102 U.S. 707, 728–30 (1880).

\(^{130}\) *Dolbear v. Am. Bell Tel. Co.* (The Telephone Cases), 126 U.S. 1, 531, 534–35 (1888).

\(^{131}\) Id. at 534.

\(^{132}\) Id. at 534–35.

\(^{133}\) 283 U.S. 1, 6, 11–12 (1931).

\(^{134}\) Id. at 11–12.

William Douglas’s majority opinion held that it was only “the discovery of some of the handiwork of nature and hence is not patentable.”

From 1972 to 1981, the Supreme Court decided four patentable subject matter cases. In *Gottschalk v. Benson*, the Court held that an algorithm for converting binary-coded decimal numerals into pure binary numerals (either by hand, or, more practically, on a computer) was patent-ineligible. Justice Douglas reasoned that “one may not patent an idea” and that upholding this patent would “wholly pre-empt the mathematical formula and in practical effect would be a patent on the algorithm itself.” Second, in *Parker v. Flook*, the Court rejected a patent on a method for updating alarm limits during catalytic conversion of hydrocarbons (such as petroleum), which relied in part on a mathematical formula, because the only novel feature of the method was the mathematical formula. Third, in *Diamond v. Chakrabarty*, the Court upheld a patent on a genetically engineered bacterium useful in breaking down oil (e.g., in cleaning up oil spills). Chief Justice Warren Burger distinguished *American Fruit Growers* and *Funk Brothers* because this bacterium, although a living organism, was human-made and possessed “markedly different characteristics from any [bacteria] found in nature.” Finally, in *Diamond v. Diehr*, the Court distinguished *Flook* to uphold a patent on a process for molding synthetic rubber that relied on a mathematical formula (the Arrhenius equation). Justice William Rehnquist’s majority opinion reached back to *Cochrane v. Deener*, holding that the process at issue was patentable because it transformed an article (uncured rubber) into a different state or thing. Even though the method used a mathematical formula, the patent in *Diehr* did not claim the formula itself and would not “pre-empt the use of that equation” in other fields.

After *Diehr*, the Court did not decide a major patentable subject matter case for nearly 30 years. Development of patent-eligible subject matter law was mainly left to the Federal Circuit,

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136 Id.
137 Three of these four (*Benson, Flook, and Diehr*), which concern the patentability of inventions relating to mathematical formulas and computers, are often referred to as a “trilogy.” See, e.g., Michel, supra note 20, at 1755; Menell, supra note 112, at 1290. This usage leaves out *Chakrabarty*, which was also decided in the same time frame, because that case concerned the exception for products of nature.
139 Id. at 71–72.
142 Id. at 310.
144 Id. at 184.
145 Id. at 187. In the view of many commentators, *Diehr* effectively overturned *Flook* (or at least some statements in *Flook*) without explicitly saying so. See, e.g., Michel, supra note 20, at 1756 (“*Diehr*, to my eye, overruled *Flook* five to four.”); Menell, supra note 112, at 1298 (“Justice Rehnquist [in *Diehr*] effectively overrode *Flook*’s statutory subject matter test.”); *BCLT Report*, supra note 16, at 554 (“*Flook* was effectively overruled three years later in *Diamond v. Diehr* . . . .”); *Athena Diagnostics, Inc. v. Mayo Collaborative Servs.*, 927 F.3d 1333, 1346 (Fed. Cir. 2019) (Chen, J., concurring in the denial of rehearing en banc) (“Given *Diehr*’s evident disagreement with *Flook*’s analysis, *Diehr*, as the latter opinion, was widely understood to be the guiding, settled precedent on § 101 for three decades.”); Dennis Crouch, *Revival of Parker v. Flook II*, PATENTLYO (Jan. 4, 2018), https://patentlyo.com/patent/2018/01/revival-parker-flook.html (presenting data showing that courts rarely cited *Flook* between 1982 and 2007).
146 See Lemley et al., supra note 21, at 1317; Menell, supra note 112, at 1298. There are two partial exceptions to this generalization. The first is *J.E.M. Ag Supply, Inc. v. Pioneer Hi-Bred Int’l, Inc.*, in which the Court held that human-made plant varieties were patentable under Section 101. 534 U.S. 124, 127 (2001). However, that case turned not on
whose decisions generally expanded patent-eligible subject matter, such that by the late 1990s Section 101 became perceived as “a dead letter.”

The Modern Alice/Mayo Framework

In 2010, the Supreme Court reentered the field of patent-eligible subject matter, deciding four cases on the issue within five years. These cases established the two-step Alice/Mayo test for patentable subject matter.

The first step of the Alice/Mayo test addresses whether the patent claims are “directed to” an ineligible concept: a law of nature, a natural phenomenon, or an abstract idea. The inquiry at step one focuses on the “claim as whole.” To be “directed to” an eligible concept at step one of Alice/Mayo, the claims must not simply involve a patent-ineligible concept. Rather, the “focus on the claims” must be a patent-ineligible concept, and not the improvement of a technological process. If the patent claims are not directed to an ineligible concept, then the subject matter is patent-eligible.

If the claims are directed to an ineligible category, then the invention is not patentable unless the patent claims have an “inventive concept” under the second step of the Alice/Mayo test. Step two of Alice/Mayo considers the elements of each patent claim both individually and as an ordered combination in the search for an “inventive concept”—additional elements that “transform the nature of the claim” into a patent-eligible application of an ineligible concept. To have an “inventive concept,” the patent claims must contain elements “sufficient to ensure that

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148 Lemley et al., supra note 21, at 1318 (“[A]fter 1998, patentable subject matter was effectively a dead letter”).


150 Alice, 573 U.S. at 217.


153 Id.; see also Athena, 915 F.3d at 750 ("To determine whether a claim is directed to an ineligible concept, we have frequently considered whether the claimed advance improves upon a technological process or merely an ineligible concept, based on both the written description and the claim.") (citations omitted).

154 Alice, 573 U.S. at 217.

155 Id.

156 Alice, 573 U.S. at 217–28 (quotations omitted).
the patent in practice amounts to significantly more than a patent upon the [ineligible concept] itself.” 157 Claim limitations that are “conventional, routine and well understood,” such as generic computer implementation, cannot supply an inventive concept. 158

Bilski v. Kappos, the first in the series of Supreme Court cases that developed what became known as the Alice/Mayo framework, concerned a patent on a business method for hedging against price-fluctuation risks in energy and commodity markets. 159 The Federal Circuit had held that this method was not patentable as a “process” under Section 101 because it failed the “machine-or-transformation test”—that is, it was neither “tied to a particular machine or apparatus” nor “transform[ed] a particular article into a different state or thing.” 160 All nine members of the Supreme Court agreed with that result—that the business method at issue was not patent-eligible—but differed significantly as to their reasoning. Writing for five Justices, Justice Anthony Kennedy held that the machine-or-transformation test was not the “sole test” for determining whether a process is patent-eligible but still “a useful and important clue.” 161 While the majority rejected the “atextual” notion that business methods were categorically unpatentable under Section 101, 162 it relied on Benson and Flook to conclude that this particular patent attempted to claim an unpatentable abstract idea: the “concept of hedging risk.” 163 Concurring only in the judgment, Justice John Paul Stevens wrote for four Justices who would have held, based on the history of the Patent Act and its constitutional purpose, that business methods were always patent-ineligible. 164

In Mayo Collaborative Services v. Prometheus Laboratories, the Court addressed the scope of the “law of nature” exception. 165 The patent in Mayo claimed a method for measuring metabolites in human blood in order to calibrate the dosage of thiopurine drugs in the treatment of autoimmune disorders. 166 Writing for a unanimous Court, Justice Stephen Breyer’s opinion held that the patent claims were addressed to a law of nature: “namely, relationships between concentrations of certain metabolites in the blood and the likelihood that a dosage of a thiopurine drug will prove ineffective or cause harm.” 167 Because the claims were little “more than an instruction to doctors to apply the applicable laws when treating their patients,” the patent lacked any inventive concept and was held to be patent-ineligible. 168

The next case, Association for Molecular Pathology v. Myriad Genetics, Inc., concerned the applicability of the “natural phenomena” exception to the patentability of human DNA. 169 The

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157 Id. (quoting Mayo Collaborative Servs. v. Prometheus Labs., Inc., 566 U.S. 66, 73 (2012)).
158 Ariosa Diagnostics, Inc. v. Sequenom, Inc., 788 F.3d 1371, 1378 (Fed. Cir. 2015); accord Alice, 573 U.S. at 225; Mayo, 566 U.S. at 79 (“Purely ‘conventional or obvious’ ‘[p]re-[solution activity] is normally not sufficient to transform an unpatentable law of nature into a patent-eligible application of such a law.” (quoting Parker v. Flook, 437 U.S. 584, 590 (1978))).
159 Bilski, 561 U.S. at 598–99.
160 In re Bilski, 545 F.3d 943, 954 (Fed. Cir. 2008) (en banc) (Michel, C.J.).
161 Bilski, 561 U.S. at 604.
162 Id. at 609.
163 Id. at 609–12.
164 Id. at 626–57 (Stevens, J., concurring in the judgment).
166 Id. at 73–75.
167 Id. at 77.
168 Id. at 79.
inventor in *Myriad* had discovered the precise location and genetic sequence of two human genes associated with an increased risk of breast cancer. 170 Based on this discovery, the patentee claimed two molecules associated with the genes: (1) an isolated DNA segment and (2) a complementary DNA (cDNA) segment, in which the nucleotide sequences that do not code for amino acids were removed in the laboratory. 171 Justice Clarence Thomas’s unanimous opinion in *Myriad* held that isolated DNA segments were nonpatentable products of nature because the patent claimed naturally occurring genetic information. 172 The Court held, however, that cDNA, as a synthetic molecule distinct from naturally occurring DNA, was patentable even though the underlying nucleotide sequence was dictated by nature. 173

Most recently, *Alice Corp. v. CLS Bank International* examined the scope of the “abstract idea” category of nonpatentable subject matter. 174 *Alice* concerned a patent on a system for mitigating “settlement risk”—the risk that only one party to a financial transaction will pay what it owes—using a computer as an intermediary. 175 The Court first held, relying on *Bilski*, that the invention was directed at “the abstract idea of intermediated settlement.” 176 Although this idea was implemented on a computer (which is, of course, a physical machine), the patent lacked an inventive concept because the claims merely “implement[ed] the abstract idea of intermediated settlement on a generic computer.” 177

Table 1 summarizes the facts and holding of the Supreme Court’s major patentable subject matter cases, in reverse chronological order.

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<th>Case Citation</th>
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<td><em>Alice Corp. Pty. v. CLS Bank Int'l</em>, 573 U.S. 208 (2014)</td>
<td>Computer-implemented method and system for mitigating settlement risk in financial transactions using a third-party intermediary</td>
<td><strong>Ineligible</strong>: The claims are drawn to the abstract idea of intermediated settlement; implementation on a generic computer does not transform an ineligible abstract idea into a patent-eligible invention.</td>
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<tr>
<td>Ass'n for Molecular Pathology v. Myriad Genetics, Inc., 569 U.S. 576 (2013)</td>
<td>Isolated human DNA segments and exon-only complementary DNA (cDNA) segments corresponding to genes discovered to be linked to an increased risk of breast cancer</td>
<td><strong>Certain Claims Ineligible</strong>: Isolated human DNA segments are patent-ineligible because the nucleotide sequence is a product of nature and isolation from the rest of the genome is insufficient to render them patentable; however, cDNA is patentable because it is not naturally occurring.</td>
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170 *Id.* at 579.
171 *Id.* at 580–85.
172 *Id.* at 591–94. Justice Antonin Scalia joined the opinion save for the “fine details of molecular biology,” as he found himself “unable to affirm those details on my own knowledge or even my own belief.” *Id.* at 596 (Scalia, J., concurring in part and in the judgment).
173 *Id.* at 594–95.
175 *Id.* at 212.
176 *Id.* at 221.
177 *Id.* at 225.
### Case Citation | Claimed Inventions | Holding and Rationale
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Method for optimizing dosage of thiopurine drugs for treating autoimmune disease, by administering the drug, measuring a metabolite, and adjusting the dosage based on the measurement | Ineligible: The relationship between the concentration of particular metabolites in the blood and a drug’s effectiveness is directed to a law of nature, and the claims lack an inventive concept beyond conventional post-solution activity. |
Business method for hedging against price-fluctuation risks in energy and commodity markets | Ineligible: Although business methods are not categorically patent-ineligible, the process at issue was not patentable because it claimed the abstract idea of hedging risk. |
Human-developed inbred and hybrid corn plant varieties and seeds | Eligible: Newly developed plant varieties are human-made manufactures or compositions of matter, even though protection may also be available under the Plant Patent Act or the Plant Variety Protection Act. |
**Diamond v. Diehr, 450 U.S. 175 (1981)**  
Process for molding raw, uncured synthetic rubber into cured products, relying on the Arrhenius equation and a programmed computer to calculate the curing time | Eligible: The invention does not claim a mathematical formula or a law of nature as such, but applies a natural law to a particular industrial process that transforms an article into a different state or thing. |
Genetically engineered bacterium capable of breaking down components in crude oil | Eligible: The genetically engineered bacterium was not naturally occurring and possessed markedly different characteristics from any bacteria found in nature. |
**Parker v. Flook, 437 U.S. 584 (1978)**  
Method of updating alarm limits used in catalytic conversion of hydrocarbons (e.g., in oil refining) relying on a mathematical formula | Ineligible: The only novel feature of the invention was a mathematical formula, conventionally applied to a specific field. |
**Gottschalk v. Benson, 409 U.S. 63 (1972)**  
Method for converting binary-coded decimal numerals into pure binary numerals on digital computer | Ineligible: The patent claims cover all practical uses of a mathematical algorithm and would, in effect, amount to a patent on the algorithm itself. |
**Funk Bros. Seed Co. v. Kalo Inoculant Co., 333 U.S. 127 (1948)**  
Inoculant for leguminous plants comprising several strains of mutually noninhibitive species of bacteria to improve nitrogen fixation | Ineligible: Each bacterial strain is naturally occurring, and discovery of the noninhibitive qualities of certain strains was not invention but merely the discovery of a nonpatentable natural phenomenon. |
**Mackay Radio & Tel. Co. v. Radio Corp. of Am., 306 U.S. 86 (1939)**  
Radio antenna in which the angle of the wires and their length are determined by a mathematical formula | Assumed to be patentable: Although a mathematical expression of a scientific truth is not patentable, a novel and useful structure created with the aid of knowledge of scientific truth may be patentable. |
**Am. Fruit Growers v. Brogdex Co., 283 U.S. 1 (1931)**  
Citrus fruit treated with borax solution to render it resistant to mold | Ineligible: Treatment with borax did not transform the fruit (a product of nature) into a manufacture with a new or distinctive form, quality, or property. |

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178 Although *Mackay Radio* is widely quoted in subsequent jurisprudence for the proposition that useful applications of laws of nature are patentable, see, for example, *Mayo Collaborative Servs. v. Prometheus Labs., Inc., 566 U.S. 66, 71 (2012)*; *Diamond v. Diehr, 450 U.S. 175, 188 (1981)*, Justice Harlan Stone’s statement is dicta because the Court merely “assume[d], without deciding” that the invention was patentable, ruling instead on grounds of noninfringement, see *Mackay Radio*, 306 U.S. at 94, 101.
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<tbody>
<tr>
<td>The Telephone Cases,</td>
<td>Method and apparatus for transmitting method and apparatus for transmitting</td>
<td>Eligible: The patentee did not claim all uses sound telegraphically by causing of electricity to transmit speech at a distance, electrical undulations, similar to air but only the particular process and apparatus vibrations accompanying speech and disclosed in the patent. other sounds</td>
</tr>
<tr>
<td>126 U.S. 1 (1888)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tilghman v. Proctor,</td>
<td>Process for separating fat into glycerin and fatty acids using water, pressure, and heat</td>
<td>Eligible: New and useful manufacturing processes are “arts” that may be patented independently of the apparatus used.</td>
</tr>
<tr>
<td>102 U.S. 707 (1881)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cochrane v. Deener,</td>
<td>Improved industrial process for manufacturing flour</td>
<td>Eligible: A process (“a series of acts,</td>
</tr>
<tr>
<td>94 U.S. 780 (1877)</td>
<td></td>
<td>performed upon the subject-matter to be transformed and reduced to a different state or thing”) is patentable independent of the machinery used.</td>
</tr>
<tr>
<td>Rubber-Tip Pencil Co. v.</td>
<td>Rubber cap with cavity designed to be attached to lead pencils for convenient use as an eraser</td>
<td>Ineligible: An “idea of itself” (here, the idea</td>
</tr>
<tr>
<td>Howard, 87 U.S. (20 Wall.)</td>
<td></td>
<td>of attaching a piece of rubber to the end of a pencil for use as an eraser) is not patentable.</td>
</tr>
<tr>
<td>498 (1874)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Corning v. Burden, 56 U.S.</td>
<td>Machine for rolling puddle balls and other masses of iron used in the</td>
<td>Eligible: The patentee did not claim the function or abstract effect of a machine, but only the machine that produced the result.</td>
</tr>
<tr>
<td>(15 How.) 252 (1854)</td>
<td>manufacturing iron products</td>
<td></td>
</tr>
<tr>
<td>O'Reilly v. Morse, 56 U.S.</td>
<td>Any use of electro-magnetism for printing intelligible characters, signs, or letters, at a distance</td>
<td>Ineligible: The discovery of a scientific principle is not patentable, nor can a patentee claim a useful result in the abstract, apart from the particular process or machine by which the result is accomplished.</td>
</tr>
<tr>
<td>(15 How.) 62 (1854)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Le Ray v. Tatham, 55 U.S.</td>
<td>Machinery for manufacturing wrought metal pipes exploiting a newly discovered property of lead</td>
<td>Potentially patentable: Although a principle in the abstract is not patentable, a practical application of such a principle to a new and useful end is patentable.</td>
</tr>
<tr>
<td>(14 How.) 156 (1853)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Source: CRS.

179 The specific doctrinal basis of O'Reilly v. Morse is unclear, as the Court speaks in language that, when cast in modern terms, sounds at times like enablement and at times like patentable subject matter. Compare 56 U.S. at 113 (“The court is of opinion that the claim is too broad . . . .”) with id. at 116 (“[T]he discovery of a principle in natural philosophy or physical science, is not patentable.”). Many patent scholars regard Morse as a case not about Section 101 but about enablement under Section 112 of the modern Patent Act. See, e.g., Taylor, supra note 20, at 205 (“In modern terms, it is quite clear that the problem with Claim 8 in Morse’s patent was based on the enablement and written description requirements located in § 112 and not in § 101.”); Lefstin, supra note 113, at 597 (“Morse is about disclosure and scope, not patent-eligible subject matter.”). The Supreme Court, however, appears to regard Morse as primarily a subject matter decision. See, e.g., Mayo, 566 U.S. at 70, 73 (citing to Morse to support notion that “laws of nature” or claims that “preempt the use of a natural law” are “not patentable”).

180 Statements in Le Ray to the effect that a “principle, in the abstract” is not patentable, but a practical application of such a principle may be patentable, 55 U.S. at 174–75, are widely quoted and influential in subsequent American jurisprudence. See supra note 114. Nonetheless, because the result in Le Ray turned primarily on claim construction, see 55 U.S. at 176, these general statements were dicta and did not entail the holding of the case.
The Debate Over Alice/Mayo and Section 101 Reform

A substantial group of patent law stakeholders, including inventors, academics, industry representatives, patent attorneys, current and former Federal Circuit judges, and former PTO officials, has criticized the Alice/Mayo framework on various grounds. Other patent law stakeholders defend the Supreme Court’s Section 101 decisions.

Criticisms of the Alice/Mayo Framework

Generally, critics of the Court’s patentable subject matter jurisprudence raise four principal concerns. First, the Alice/Mayo framework is criticized as excessively vague, subjective, and unpredictable in application. For example, the Federal Circuit has stated that when determining whether a patent claim is “directed to” an ineligible concept at step one, courts must determine whether the “focus” of the claims is on that concept. At the same time, the Federal Circuit has cautioned that this “focus” must be articulated “with enough specificity to ensure the step one inquiry is meaningful.” The appropriate level of specificity can vary from patent to patent and from judge to judge.

Thus, in the view of many stakeholders, the Supreme Court’s patentable subject matter case law and the Federal Circuit’s implementation of the Alice/Mayo framework fail to articulate “objective, predictable criteria” for making patent-eligibility determinations. Key terms, such as what an “abstract idea” is, or precisely how claim elements can make an invention “significantly more” than an ineligible category (the “inventive concept”), are largely left undefined, making it difficult for patent applicants and litigators to know whether their patent claims will survive judicial scrutiny. Moreover, the Federal Circuit has explicitly recognized

181 See infra “Criticisms of the Alice/Mayo Framework.”
182 See infra “Defenses of the Alice/Mayo Framework.”
184 Thales Visionix Inc. v. United States, 850 F.3d 1343, 1347 (Fed. Cir. 2017).
185 See Visual Memory LLC v. NVIDIA Corp., 867 F.3d 1253, 1262 (Fed. Cir. 2017) (Hughes, J., dissenting) (disagreeing with the majority over whether characterizing the claims as directed to “categorical data storage” views the invention “at an unduly ‘high level of abstraction’”) (quoting Enfish, LLC v. Microsoft Corp., 822 F.3d 1327, 1337 (Fed. Cir. 2016)).
186 PTO PSM REPORT, supra note 16, at 29.
187 See id. at 30 (describing comments that the Alice/Mayo test “fails to define crucial terms, such as ‘abstract’ and ‘substantially more’”); Taylor, supra note 20, at 231 (“[N]o one really knows what an inventive concept is.”); Lemley et al., supra note 21, at 1316 (“[N]o one understands what makes an idea ‘abstract,’ and hence ineligible . . . .”); Morris, supra note 106, at 68 (arguing that the judicially created patentable subject matter decisions are “merely post hoc rationalizations”). Some Supreme Court Justices have echoed this criticism. See, e.g., Bilski v. Kappos, 561 U.S. 593, 621 (2010) (Stevens, J., concurring in the judgment) (“The Court . . . never provides a satisfying account of what constitutes an unpatentable abstract idea.”); Fred Funk Seed Bros. Co. v. Kalo Inoculant Co., 333 U.S. 127, 134–35 (1948) (Frankfurter, J., concurring) (“It only confuses the issue, however, to introduce such terms as ‘the work of nature’ and the ‘laws of nature.’ For these are vague and malleable terms infected with too much ambiguity and equivocation.”). To some extent, uncertainty in Section 101 is not a new phenomenon. See, e.g., Duffy, supra note 113, at 623–38 (reviewing history of failed patentable subject matter rules and observing that “instability in the law of patentable subject matter” is a recurring issue). However, at least in the decade before Mayo, uncertainty was less practically important for patentees because courts and the PTO only “rarely” rejected patents based on Section 101. See BCLT Report, supra note 16, at 575–76 (reviewing data showing a “dramatic” increase in the number of Section 101 district court decisions following Mayo, with a “10-fold” increase following Alice).
that the two steps of the analysis are not clearly defined and may overlap.\textsuperscript{188} As a result, many observers characterize the court’s Section 101 jurisprudence as a “highly subjective,” “I know it when I see it” approach.\textsuperscript{189} This subjectivity, in the view of critics, injects unpredictability and uncertainty into whether an invention is of a type that is patentable.\textsuperscript{190}

Second, the Alice/Mayo framework is criticized as legally flawed on various grounds. Some stakeholders argue that the Alice/Mayo framework misinterprets Section 101, imposing “extra-statutory” requirements for patent eligibility, contrary to congressional intent or the constitutional purpose of patent law.\textsuperscript{191} Others argue that Mayo’s requirement of an “inventive concept” rests on a historically inaccurate understanding of 19th century English patent law, first imported into American jurisprudence in cases such as Le Roy and Morse.\textsuperscript{192} Finally, many commentators and stakeholders argue that the Alice/Mayo framework confuses patent law by conflating eligibility under Section 101 with policy concerns—such as the obviousness of the invention and claim breadth—that are better addressed by other provisions in the Patent Act, such as Sections 102, 103, and 112.\textsuperscript{193} For example, patent claims have been found to lack an inventive concept at Alice/Mayo step two where they implement an abstract idea on conventional computer hardware.\textsuperscript{194} Issues about what was “conventional” or “well-understood” at the time of the invention, however, are questions usually reserved for novelty or nonobviousness analysis.\textsuperscript{195}

Third, the Alice/Mayo framework is alleged to have detrimental effects on incentives to innovate, especially in the biotechnology and computer software industries. Given the patent claims at issue in Alice (a computer-implemented business method), Myriad (an isolated human DNA segment), and Mayo (a drug dose optimization method), most observers agree that these two industries have

\textsuperscript{188} \textit{Elec. Power Grp.}, 830 F.3d at 1353 (“[T]he two stages are plainly related: not only do many of our opinions make clear that the two stages involve overlapping scrutiny of the content of the claims, but we have noted that there can be close questions about when the inquiry should proceed from the first stage to the second.”) (citations omitted).

\textsuperscript{189} See, e.g., PTO PSM REPORT, supra note 16, at 30 (quoting stakeholder view that Alice/Mayo is “hopelessly subjective”); Taylor, supra note 20, at 227–30 (arguing that Alice/Mayo framework has “no objective guidance” and “leaves the determination of eligibility to the unconstrained, subjective opinion of a patent examiner or judge”); Klein, supra note 106, at 288 (criticizing patentable subject matter case law as amounting to “an ‘I know it when I see it’ approach”).

\textsuperscript{190} See, e.g., BCLT Report, supra note 16, at 561 (describing “uncertainty and confusion resulting from the Court’s recent [patentable subject matter] jurisprudence”); accord PTO PSM REPORT, supra note 16, at 30–31 (describing views that the Alice/Mayo test yields “unpredictable” and “inconsistent” results).

\textsuperscript{191} See PTO PSM REPORT, supra note 16, at 28; Klein, supra note 106, at 289–91 (criticizing the three judicially created categorical exclusions as “extra-statutory” and proposing test that focuses on text of Section 101).

\textsuperscript{192} Lefstin, supra note 113, at 565 (arguing that Alice/Mayo test’s “inventive application” requirement rests on a “basic misapprehension” of the 19th century English case cited by the Supreme Court); PTO PSM REPORT, supra note 16, at 27–28 (same).

\textsuperscript{193} See PTO PSM REPORT, supra note 16, at 31–32; Taylor, supra note 20, at 157 (“[T]he current approach to determining patent eligibility confuses the relevant policy concerns underlying numerous discrete patent law doctrines.”); see also Risch, supra note 21, at 594 (arguing that the Court’s patentable subject matter doctrine would be more consistent and rigorous if replaced with a strict application of other patentability doctrines such as obviousness, novelty, utility, inventorship, written description, and enablement). This criticism has been echoed by Supreme Court Justices. See Parker v. Flook, 437 U.S. 584, 600 (1978) (Stewart, J., dissenting) (“[T]he majority] strikes what seems to me an equally damaging blow at basic principles of patent law by importing into its inquiry under 35 U.S.C. § 101 the criteria of novelty and inventiveness.”).

\textsuperscript{194} See, e.g., \textit{Elec. Power Grp.}, 830 F.3d at 1355.

\textsuperscript{195} See, e.g., Berkheimer v. HP Inc., 881 F.3d 1360, 1368–69 (Fed. Cir. 2018) (noting that Alice/Mayo step two determination of whether claims are “well-understood, routine and conventional” overlaps with Section 102 novelty inquiry).
been the most affected by the Supreme Court’s Section 101 rulings. In the biotechnology industry, stakeholders argue that the Alice/Mayo framework has limited their ability to obtain patents on diagnostic methods and kits, personalized medicine, and isolated natural substances. Views in the computer industry are “sharply divided,” but at least some stakeholders argue that Alice has devalued their patents and created uncertainty for their business. In both fields, some stakeholders argue that the law of Section 101 is reducing incentives to innovate in these areas and driving investment elsewhere.

Finally, the uncertainty and unpredictability caused by Alice/Mayo is alleged to put the United States at a disadvantage relative to international competitors. Some stakeholders argue that U.S. competitiveness may be harmed because a lack of patent availability will drive investment in certain industries to other countries where such inventions are more clearly patent-eligible. Others argue that one effect of Alice/Mayo is a loss of any patent protection for certain inventions, which will enable competitors to “free ride” off of American innovation.

**Defenses of the Alice/Mayo Framework**

Defenders of the current law of Section 101 respond that these criticisms of Alice/Mayo are overstated, or that the Supreme Court’s reinvigoration of Section 101 has important benefits for the patent system. As to the subjective or unpredictable nature of Section 101 doctrine, there is some indication that the Alice/Mayo framework is not quite as unpredictable as is sometimes claimed. Some commentators also observe that uncertainty in patentable subject matter law is hardly a new phenomenon, and may even be “inevitable.”

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196 PTO PSM REPORT, supra note 16, at 34–35 (“Among members of the public, there was a general consensus that two industries have been most directly affected [by the Alice/Mayo framework]: life sciences and computer-related technologies.”); see also BCLT Report, supra note 16, at 582–85 (examining the Alice/Mayo framework’s effects on diagnostics, personalized medicine, biosciences, software, and information technology).


200 See, e.g., Stoll, supra note 23 (“The courts’ focus on subject matter eligibility as a mechanism to deny patents for [inventions in diagnostics and personalized medicine] will drive investment into research in these technologies to other areas. We will lose our edge in the world . . . .”); accord PTO PSM REPORT, supra note 16, at 34; Kevin Madigan & Adam Mossoff, Turning Gold into Lead: How Patent Eligibility Doctrine Is Undermining U.S. Leadership in Innovation, 24 GEO. MASON L. REV. 939, 942–44 (2017) (expressing “concern about the U.S. conceding its gold standard patent system to China and Europe” because of the uncertainty of the Alice/Mayo framework).

201 See, e.g., Davis, supra note 23 (quoting former PTO Director David Kappos as stating that international competitors “no longer have to steal U.S. technology in [biotech and software], since they can now take it for free”).

202 See Jason D. Reinecke, Is the Supreme Court’s Patentable Subject Matter Test Overly Ambiguous? An Empirical Test, 2019 UTAH L. REV. 581, 583 (2019) (empirical study indicating that while “the [Alice/Mayo] test is likely not a beacon of absolute clarity, it is not completely amorphous,” as patent prosecutors correctly predicted judicial results 67.3% of the time based only on claim language).

203 See, e.g., Duffy, supra note 113, at 623–38 (reviewing 100-year history of failed rules and tests for patentable subject matter and observing that “instability in the law of patentable subject matter” is a recurring issue) & id. at 616 (citing 19th century treatise writers noting difficulty and complexity of the patentable subject matter); Risch, supra note 21, at 591 (criticizing, in 2008, the “currently confused and inconsistent jurisprudence of patentable subject matter”); Donald S. Chisum, The Patentability of Algorithms, 47 U. PITT. L. REV. 959, 992 (1986) (noting “confusion and arbitrary distinctions” in the law of the patentability of computer software resulting from the Benson decision).

204 Morris, supra note 106, at 107 (arguing that the Court’s “intuitive” approach to patentable subject matter determinations is “inevitable”).
approach to patentable subject matter, on this view, may have certain benefits, including flexibility and adaptability to new technologies.\textsuperscript{205} Moreover, even if one views the current state of the law as unacceptably vague, courts may eventually clarify or change Section 101 doctrine in line with the long history of common law development in this area.\textsuperscript{206}

As to the legal correctness of Alice/Mayo, defenders of the framework note that while the judicially created categories are not directly grounded in the text of Section 101, they have been treated as part of the law “as a matter of statutory stare decisis going back 150 years.”\textsuperscript{207} As to Mayo’s reliance on 19th century English patent law, some commentators defend the Supreme Court’s “inventive application” requirement as a faithful reading of this precedent.\textsuperscript{208} Finally, although the Alice/Mayo framework may overlap with other patent law doctrines, several commentators and judges of the Federal Circuit argue that Section 101 serves purposes distinct from Sections 102, 103, and 112.\textsuperscript{209} For example, even if the invention in Myriad—an isolated human DNA sequence discovered to be linked to increased breast cancer risk—was novel, nonobvious, and sufficiently disclosed, some commentators would still argue that the invention should not be patented based on harm to future innovation or moral concerns about patenting human DNA.\textsuperscript{210}

Regarding the alleged detrimental effects of the Court’s Section 101 decisions on innovation, some stakeholders point to countervailing benefits either generally or in certain industries. In particular, some stakeholders in industries (such as computer software) affected by litigation by patent assertion entities\textsuperscript{211} argue that Section 101 is a useful and important tool for weeding out overly broad or vague patents at the outset of litigation.\textsuperscript{212} Other commentators point to general utilitarian or moral benefits of robust exclusions for patents on basic discoveries in science and nature.\textsuperscript{213}

\textsuperscript{205} Id. at 107–09 (arguing that intuitive approach to Section 101 may be “desirable” because “there is simply no other more rigorous and yet durable way of identifying the proper boundaries for patentable subject matter” and “vagueness provides the flexibility necessary to adjust future technological developments”); Duffy, supra note 113, at 639 (“[T]he traditional doctrines of patentable subject matter—the prohibition against patenting abstract ideas, natural phenomena, and principles of nature—have survived because . . . they have been amorphous.”).

\textsuperscript{206} See PTO PSM REPORT, supra note 16, at 23–24 (expressing stakeholder views that the Court’s decisions are part of the normal common law development of Section 101, and that the Federal Circuit’s subsequent development of the law may be “headed in the right direction”).


\textsuperscript{209} See, e.g., Morris, supra note 106, at 113 (“To be sure, patentable subject matter overlaps with and serves some of the same purposes as the other patentability requirements . . . . But only patentable subject matter serves to distinguish patentable technology from unpatentable discoveries, information, and human thought and activity.”); Lemley et al., supra note 21, at 1330–32 (distinguishing Purpose of Section 101 from Section 112); accord Mayo, 566 U.S. at 90–91; Athena Diag., Inc. v. Mayo Collaborative Servs., 927 F.3d 1333, 1337–39 (Fed. Cir. 2019) (Dyk, J., concurring in the denial of rehearing en banc).

\textsuperscript{210} See generally infra “Potential Rationales for Section 101.”

\textsuperscript{211} A patent assertion entity, sometimes called a nonpracticing entity or (pejoratively) a “patent troll,” is a loose term for an individual or organization that seeks to license or litigate patents, but does not itself practice the patented invention. See Colleen V. Chien, From Arms Race to Marketplace: The Complex Patent Ecosystem and Its Implications for the Patent System, 62 HASTINGS L.J. 297, 326–27 (2010) (discussing distinction among various types of nonpracticing patent entities).

\textsuperscript{212} PTO PSM REPORT, supra note 16, at 24–26; BCLT Report, supra note 16, at 596; Gugliuzza, supra note 25, at 652–53.

\textsuperscript{213} Sarnoff, supra note 113, at 106–24 (reviewing asserted utilitarian and moral benefits of robust Section 101
Lastly, in response to concerns about the *Alice*/Mayo framework’s effect on international competitiveness, some commentators assert that these changes are good for the United States as a geopolitical matter. In particular, restricted patent-eligibility standards may benefit U.S. consumers if a lack of patent protection leads to increased competition and lower prices for certain products without harming innovation.

**Potential Rationales for Section 101**

More broadly, there is a long-running debate over the functions and purposes that Section 101 serves in the patent system. For its part, the modern Supreme Court has largely settled on the “preemption rationale” for the judicially created subject matter exclusions. These decisions assert that abstract ideas, laws of nature, and natural phenomena should not be patentable because permitting a monopoly on the “‘basic tools of scientific and technological work’ . . . might tend to impede innovation more than it would tend to promote it,” in that such patents would “significantly impede future innovation.” The gist of the preemption rationale is that Section 101 functions to prevent patents that reach so broadly that they “threaten downstream innovation” by preempting all uses of a natural law, abstract idea, or fundamental research tools.

The preemption rationale is not the only potential justification for Section 101, however. Although a complete survey of the various rationales proffered for Section 101 is beyond the scope of this report, at least four broad categories of rationales for Section 101 have been proposed.

First, some commentators argue that Section 101’s purpose is to identify certain patents or categories of patents that should not be granted because their economic harms exceed their benefits—that is, their net social costs are negative for innovation, or more generally, Preemption theory, which claims that certain overbroad patents should be denied patent protection under Section 101 because of their negative effects on downstream innovation, is an example from this group.

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214 PTO PSM REPORT, supra note 16, at 27.
215 Id.
217 Mayo, 566 U.S. at 91.
218 See, e.g., Lemley et al., supra note 21, at 1346–47; accord Benson, 409 U.S. at 72 (rejecting patent because it would “wholly pre-empt the mathematical formula and in practical effect would be a patent on the algorithm itself”). But see Katherine J. Strandburg, *Much Ado About Preemption*, 50 HOUS. L. REV. 563, 566 (2012) (critiquing preemption rationale’s “sole focus on broad downstream impact” as not providing a satisfactory explanation for the Supreme Court’s Section 101 case law).
220 See Anderson, supra note 219, at 284–85 (overviewing this group of theories); see, e.g., David S. Olson, *Taking the Utilitarian Basis for Patent Law Seriously: The Case for Restricting Patentable Subject Matter*, 82 TEMP. L. REV. 181, 184 (2009) (arguing that patentable subject matter doctrine should be driven by looking at when “granting a patent right for this type of innovation causes more loss to society than gain”).
221 See supra note 218 and accompanying text.
Second—in what is in some sense a special case of the first rationale—other commentators assert that Section 101’s purpose is to identify and deny patents to categories of inventions that would have been developed even without a patent incentive.222 For example, several commentators have argued the patents on business methods should be excluded under Section 101 either because they affirmatively harm innovation and the economy, or because they are simply unnecessary because sufficient incentives to create business methods would exist even if patents are unavailable.223

Third, some commentators assert that Section 101 (or elements of Section 101 doctrine) are based not on economic considerations but on moral or ethical concerns.224 For example, the judicial prohibition on patenting products of nature—such as human DNA sequences—may be motivated by noneconomic, deontological notions of human dignity, or the inviolability of natural creation.225

Finally, some commentators believe that Section 101 serves no independent purpose in patent law not already better served by other patentability requirements.226 On this view, Section 101’s judicially created exceptions to patentable subject matter should simply be eliminated as an independent requirement for patentability, in favor of a rigorous application of the other patentability requirements in Sections 102, 103, and 112 of the Patent Act.227

Potential Options for Section 101

Before examining the particular approaches used in PTO guidance and proposed legislative reforms, this section will review some of the general ways in which Section 101 may or may not be reformed. These different paths are introduced to contextualize the current Section 101 reform proposals within the universe of possible reforms. This list is not exhaustive, nor are each of these options necessarily mutually exclusive.

At a general level, most of the proposed paths forward for Section 101 fall into one of four categories.228 First, some oppose any legislative intervention, proposing instead to allow the

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222 See Anderson, supra note 219, at 285–86 (overviewing this group of theories); see, e.g., Pamela Samuelson, Benson Revisited: The Case Against Patent Protection for Algorithms and Other Computer Program-Related Inventions, 39 EMORY L.J. 1025, 1136 (1990) (arguing that software should not be patentable in part because “the fact that this growth [in the software industry] has occurred without the aid of patent protection is powerful evidence that patent protection is not necessary for the software industry to thrive”).

223 See, e.g., Rochelle Cooper Dreyfuss, Are Business Method Patents Bad for Business?, 16 SANTA CLARA COMPUTER & HIGH TECH. L.J. 263, 274 (2000) (arguing that business method patents are unwise because they “adversely affect innovation, and worse, the economy”); accord Bilski v. Kappos, 561 U.S. 593, 651 (2010) (Stevens, J., concurring in the judgment) (arguing that business methods should not be patentable because there are “ample incentives to develop business methods even without patent protection” (quoting Dan L. Burk & Mark A. Lemley, Policy Levers in Patent Law, 89 VA. L. REV. 1575, 1618 (2003))).

224 See Anderson, supra note 219, at 286 (overviewing this group of theories); see, e.g., Sarnoff, supra note 113, at 84–90 (surveying religious and deontological bases for prohibition on patenting science, nature, and ideas); Tun-Jen Chiang, Competing Visions of Patentable Subject Matter, 82 GEO. WASH. L. REV. 1858, 1860 (2014) (arguing that Section 101 determinations are “often about noneconomic moral values”).

225 Chiang, supra note 224, at 1873–81.

226 See Anderson, supra note 219, at 280 (overviewing this group of theories).

227 See, e.g., Risch, supra note 21, at 591–94 (articulating this view); Davis, supra note 23 (quoting former PTO Director David Kappos as calling for abolishing Section 101 and instead “faithfully applying other areas of patent law to ensure that patents are not obvious or anticipated or lacking in written description”).

228 See David O. Taylor, Amending Patent Eligibility, 50 U.C. DAVIS L. REV. 2149, 2189–2211 (2017) (listing proposed Section 101 reforms, including a European-style “laundry list” of exclusions, a new “workable eligibility standard,” or the elimination of the judicially created ineligible categories); PTO PSM REPORT, supra note 16, at 39–46 (reviewing
courts to continue to develop and refine the standards for patent eligibility.\textsuperscript{229} Second, some propose replacing the \textit{Alice}/\textit{Mayo} framework with an explicit list of subject matter that is patent-eligible or -ineligible, similar to the approach that is used for European patents.\textsuperscript{230} Third, some propose replacing the \textit{Alice}/\textit{Mayo} framework with a different, usually standard for patent eligibility, such as a requirement that the invention result from human effort, exist outside the human mind, or contribute to the technological arts.\textsuperscript{231} Fourth, some propose to do away with any limitations on patentable subject matter, beyond the four statutory categories and other existing statutory patentability requirements.\textsuperscript{232}

\textbf{Continued Common Law Judicial Development}

One option is for Congress to leave Section 101 as it is, and allow the courts and the PTO to continue developing the law of patent-eligible subject matter. Stakeholders and commentators may support this option for several different reasons. Some may disagree that the \textit{Alice}/\textit{Mayo} framework is as indeterminate or as harmful to innovation as the critics claim.\textsuperscript{233} Other commentators, even if they accept the criticisms directed at \textit{Alice}/\textit{Mayo}, believe that the courts will eventually refine, clarify, or otherwise improve the law of patentable subject matter given more time for judicial development.\textsuperscript{234} Still other commentators support the current law of Section 101 as affirmatively good for innovation and society because it precludes property rights in fundamental aspects of science, nature, and ideas,\textsuperscript{235} or serves as an important mechanism to weed out overly broad patents or obtain early dismissal of unmeritorious patent litigation.\textsuperscript{236}

Supporters of continued judicial development may point to the administrative guidance put forth by the PTO\textsuperscript{237} and significant Section 101 decisions of the Federal Circuit\textsuperscript{238} as promising steps in

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\textsuperscript{232} See, e.g., Risch, supra note 21, at 591–94; see generally “Requirements for Patentability” (reviewing requirements for patentability under Sections 102, 103, and 112 of the Patent Act).

\textsuperscript{233} See BCLT Report, supra note 16, at 566.

\textsuperscript{234} See PTO PSM Report, supra note 16, at 39.

\textsuperscript{235} Sarnoff Testimony, supra note 26, at 1.

\textsuperscript{236} See Patent Eligibility Hearings, supra note 31 (statement of Prof. Paul R. Gugliuzza, Boston University School of Law), at 1, https://www.judiciary.senate.gov/imo/media/doc/Gugliuzza%20Testimony.pdf [hereinafter Gugliuzza Testimony] (“[T]he eligibility requirement, though imperfect, plays a crucial role in reducing litigation costs by giving courts a mechanism to quickly dismiss infringement claims that plainly lack merit.”).

\textsuperscript{237} See infra “Administrative Developments: PTO Subject Matter Eligibility Guidance.”

the development of Section 101 after the *Alice*, *Mayo*, and *Myriad* decisions. Opponents of maintaining the legal status quo, for their part, observe that the Supreme Court has not shown much interest in revisiting its Section 101 jurisprudence despite many opportunities, and that the PTO and the Federal Circuit are bound by the Court’s decisions.

### Specific Statutory List of Included or Excluded Subject Matter Categories

Another potential route for reform would be to amend Section 101 to replace the *Alice*/*Mayo* framework with a more specific list of subject matter that is patent-eligible or ineligible. Currently, Section 101 contains a broad list of included subject matter categories (processes, machines, manufactures, and compositions of matter), and most of the doctrine focuses on the three judicially created ineligible categories: laws of nature, natural phenomena, and abstract ideas. The “laundry list” approach would seek to make Section 101 clearer and more predictable by more specifically defining categories of eligible or ineligible subject matter. Depending on how this sort of proposal is structured, it would retain the notion of ineligible classes of subject matter, but define such categories differently, more precisely, and perhaps more narrowly than the common law exceptions under the *Alice*/*Mayo* framework.

The European Patent Convention’s (EPC’s) approach to patent eligibility offers a potential model for this type of approach. Under EPC article 52(1), patent-eligible subject matter reaches “all fields of technology, provided that they are new, involve an inventive step and are susceptible of industrial application.” At the same time, EPC article 52(2) defines specific subject matter that is not patentable when claimed “as such”:

1. discoveries, scientific theories and mathematical methods;
2. aesthetic creations;
3. schemes, rules and methods for performing mental acts, playing games or doing business, and programs for computers;
4. presentations of information.

EPC article 53 further denies patents on inventions that are “contrary to [public order] or morality,” claim “plant and animal varieties,” or claim “methods for treatment of the human or animal body by surgery or therapy and diagnostic methods practised on the human or animal body.”

Assuming that the new statutory categories are more clearly defined than existing judicial categories, a potential virtue of the laundry-list approach is greater clarity and predictability in the sort of inventions that are patentable. This approach would also more firmly ground subject matter determinations in the statutory text. On the other hand, the list-of-specific-exclusions

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239 See infra “Judicial Developments.”
240 See supra “The Current Law of Section 101.”
241 See Taylor, supra note 228, at 2198, 2200 (coining this term).
242 BCLT Report, supra note 16, at 564.
244 Id. art. 52(2)–(3).
245 Id. art. 53.
246 See Taylor, supra note 228, at 2200.
approach might be less flexible and less able to adapt to unforeseen new technologies than other options. It might also, to some degree, replace case-by-case judicial judgments of eligibility with more categorical legislative ones, which may be a virtue or a vice depending on one’s perspective.

Replace Judicial Exceptions with a Different Standard

A third group of proposed Section 101 reforms seeks to replace the Alice/Mayo framework with a new statutory standard for assessing patent eligibility. Proposals in this category are fairly diverse, but common elements in proposed new standards would limit patent eligibility to inventions that

- result from human effort;
- contribute to the technological arts;
- have practical utility or application;
- cannot be solely performed in the human mind;
- do not preempt all practical uses of a law of nature, abstract idea, or natural phenomenon.

Usually, the proposed new patentability standard would supersede the three judicially created subject matter exclusions and the two-step Alice/Mayo test.

Several proposed new standards blend more than one of these elements. For example, the American Intellectual Property Law Association has submitted a Section 101 reform proposal that replaces the Alice/Mayo framework with a single exception to patent eligibility if an invention “exists in nature independently of and prior to any human activity” or “is performed solely in the human mind.” A 2017 proposal by the American Bar Association (ABA) would explicitly allow patenting “practical applications” of laws of nature, natural phenomena, and abstract ideas, so long as the patent claim does not “preempt the use by others of all practical applications of the law of nature, natural phenomenon, or abstract idea.”

It is difficult to generalize given the significant differences among the various proposals in this category, but stakeholders may wish to consider whether proposed new standards would provide

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247 See id. at 2201.

248 Compare id. at 2193–97 (arguing that judicial “policymaking” under Section 101 should be constrained), with Morris, supra note 106, at 107–17 (arguing that a subjective, intuitive, case-by-case, judgment-based approach to Section 101 is inevitable and “perhaps even desirable”).


250 See, e.g., Taylor, supra note 228, at 2202–05; BCLT Report, supra note 16, at 563.

251 See, e.g., PTO PSM REPORT, supra note 16, at 42, 64.

252 See, e.g., PTO PSM REPORT, supra note 16, at 43; BCLT Report, supra note 16, at 563–64; Taylor, supra note 228, at 2205–07.


254 See, e.g., PTO PSM REPORT, supra note 16, at 60–61.


257 See PTO PSM REPORT, supra note 16, at 60.
greater clarity and predictability in patent-eligibility law, while still being flexible enough to adapt to new technologies.\textsuperscript{258}

**Eliminate Implied Patentable Subject Matter Limits**

A final option is to eliminate the *Alice*/*Mayo* framework and judicially created exceptions to patent eligibility altogether, without replacing them with a new standard.\textsuperscript{259} Several commentators have argued that patent-eligibility doctrine serves no purpose that is not already served by the existing statutory patentability requirements of utility, novelty, obviousness, written description, definiteness, and enablement.\textsuperscript{260} On this view, the appropriate course would be for Congress to simply eliminate the nonstatutory eligibility requirements (i.e., the judicial prohibitions on patenting laws of nature, natural phenomena, and abstract ideas) in favor of the application of the patentability requirements of Sections 102, 103, and 112 of the Patent Act.\textsuperscript{261}

Supporters of this approach argue that it advances the policy concerns motivating Section 101 law, but does so in a “more consistent and more rigorous” manner.\textsuperscript{262} Opponents argue that Section 101 serves important purposes that are distinct from the other patentability requirements, which would be lost if the judicial exceptions were eliminated.\textsuperscript{263}

**Recent Developments in Patent-Eligible Subject Matter Reform**

The Supreme Court’s modern patentable subject matter jurisprudence has led to responses from the courts, the PTO, and Congress. This section reviews recent judicial, administrative, and legislative developments on patent-eligible subject matter standards and proposed reform.

**Judicial Developments**

Since its 2014 decision in *Alice*, the Supreme Court has denied dozens of petitions for certiorari (i.e., requests that the Court hear an appeal) on Section 101 issues, despite calls from some patent law stakeholders asking the Court to revisit its patent-eligible subject matter jurisprudence.\textsuperscript{264} For example, in *Sequenom v. Ariosa Diagnostics, Inc.*\textsuperscript{265} the Supreme Court denied certiorari despite 22 amicus briefs supporting certiorari and calls from commentators, stakeholders, and Federal

\textsuperscript{258} See Taylor, supra note 228, at 2189–97 (articulating general principles for evaluating proposed Section 101 reforms).

\textsuperscript{259} See BCLT Report, supra note 16, at 565.

\textsuperscript{260} See Risch, supra note 21, at 594, 606–09; Taylor, supra note 228, at 2171–89.

\textsuperscript{261} Risch, supra note 21, at 606–09.

\textsuperscript{262} Id. at 594; accord Taylor, supra note 228, at 2211.

\textsuperscript{263} See, e.g., Mayo Collaborative Servs. v. Prometheus Labs., Inc., 566 U.S. 66, 91 (2012) (relying on concerns about preemption to “decline the Government’s invitation to substitute §§ 102, 103, and 112 inquiries for the better established inquiry under § 101”); see supra note 209 (citing academic sources); see generally “Potential Rationales for Section 101.”


\textsuperscript{265} See 788 F.3d 1371 (Fed. Cir. 2015), cert. denied, 579 U.S. 928 (2016).
Circuit judges urging the Court to take the case to clarify Section 101.266 Similarly, in opinions concerning rehearing en banc in *Athena Diagnostics, Inc. v. Mayo Collaborative Services*,267 all of the active judges on the Federal Circuit called upon the Supreme Court (or Congress) to change Section 101 law to clearly allow for the patenting of diagnostic methods.268 The Supreme Court nonetheless denied certiorari in *Athena* and again declined to revisit its Section 101 case law.269

The most prominent recent Section 101 case that the Court declined to hear was *American Axle & Manufacturing v. Neapco Holdings*.270 That case was thought by some observers to be an ideal vehicle for the Court because the patented technology—a method for manufacturing driveline shafts for automotive vehicles—was tangible and relatively straightforward, yet the lower courts held it ineligible as directed to a law of nature.271 As in *Athena*, the Federal Circuit was closely divided with respect to rehearing *American Axle* en banc, dividing 6-6, with 5 judges averring that “[Federal Circuit] rulings on patent eligibility have become so diverse and unpredictable as to have a serious effect on the innovation incentive in all fields of technology.”272 Many stakeholders again supported the petition for certiorari in *American Axle*, including a brief filed jointly by Senator Tillis, the Hon. Paul R. Michel (a former Chief Judge of the Federal Circuit), and David J. Kappos (a former PTO Director).273 The Supreme Court invited the views of the Solicitor General, who filed a brief supporting a partial grant of certiorari in *American Axle*.274 The Supreme Court declined to hear the case.275

In light of the Supreme Court’s apparent reluctance to revisit Section 101, some stakeholders have called for Congress to intervene on the issue.

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268 See Athena Diagnostics, Inc. v. Mayo Collaborative Servs., LLC, 927 F.3d 1333, 1335 (Fed. Cir. 2019) (opinions regarding the denial of rehearing en banc); CRS Legal Sidebar LSB10344, Judges Urge Congress to Revise What Can Be Patented, by Kevin T. Richards (reviewing the Federal Circuit’s opinions in *Athena Diagnostics*).

269 140 S. Ct. 855 (2020).

270 967 F.3d 1285 (Fed. Cir. 2020), cert. denied, 142 S. Ct. 2902 (2022).

271 Id. at 1292–99.


275 142 S. Ct. 2902 (2022).
Administrative Developments: PTO Subject Matter Eligibility Guidance

In 2019, the PTO issued Revised Patent Subject Matter Eligibility Guidance (the 2019 Guidance) to assist PTO patent examiners in determining subject matter eligibility for patent applications. The PTO noted that the “legal uncertainty” surrounding the Alice/Mayo framework “poses unique challenges” for the agency, which has thousands of patent examiners who must make patent-eligibility determinations on hundreds of thousands of applications each year. Accordingly, the PTO issued revised guidance to its patent examiners to provide “more clarity and predictability” in their Section 101 determinations.

The PTO subsequently incorporated the 2019 Guidance into the Manual of Patent Examining Procedure (MPEP), which guides PTO patent examiners in their review of patent applications. The 2019 Guidance guided at least two major changes to how patent examiners evaluate whether a patent application claims patent-ineligible subject matter. First, the Guidance seeks to provide a clearer definition of what constitutes an ineligible “abstract idea.” Previously, examiners would make that determination by comparing the patent claim at issue to those found to be ineligible “abstract ideas” in previous judicial cases. The PTO found that this approach had become “impractical” because of an expanding volume of sometimes contradictory Section 101 cases. The 2019 Guidance “distills” the case law into three categories that examiners will treat as “abstract ideas”:

1) Mathematical concepts—mathematical relationships, mathematical formulas or equations, mathematical calculations;

2) Certain methods of organizing human activity – fundamental economic principles or practices (including hedging, insurance, mitigating risk); commercial or legal interactions (including agreements in the form of contracts; legal obligations; advertising, marketing or sales activities or behaviors; business relations); managing personal behavior or relationships or interactions between people (including social activities, teaching, and following rules or instructions); and

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277 See 2019 Guidance, supra note 276, at 50 (“The legal uncertainty surrounding Section 101 poses unique challenges for the USPTO, which must ensure that its more than 8500 patent examiners and administrative patent judges apply the Alice/Mayo test in a manner that produces reasonably consistent and predictable results across applications, art units and technology fields.”); see also U.S. PAT. & TRADEMARK OFF., U.S. Patent Statistics Chart Calendar Years 1963–2015, https://www.uspto.gov/web/offices/ac/ido/oep/tal/us_stat.htm (last visited Nov. 21, 2022) (indicating that the PTO received 859,410 applications in 2015).

278 See 2019 Guidance, note 276, at 50.

279 See MPEP §§ 2103–2106.

280 Id. at § 2106.04(a).

281 2019 Guidance, note 276, at 51.

282 Id. at 52.
3) Mental processes – concepts performed in the human mind (including an observation, evaluation, judgment, opinion).\textsuperscript{283}

Under the Guidance, patent claims that do not recite matter that falls into one of these three groupings should not be treated as an “abstract idea” except in “rare circumstance[s].”\textsuperscript{284}

Second, the 2019 Guidance clarifies when examiners will treat a patent claim as “directed to” an ineligible category (abstract ideas, laws of nature, or natural phenomena) under step one of the Alice/Mayo test.\textsuperscript{285} In particular, the PTO will not treat a claim as “directed to” an ineligible concept if “the claim as a whole integrates the recited judicial exception into a practical application of the exception.”\textsuperscript{286} If the claim does integrate a practical application—such as improving the functioning of a computer, effecting a particular treatment for a disease, or implementing the exception into a particular machine or manufacture—then the PTO will treat the claim as patent-eligible, without having to examine the patent application for an “inventive concept” under step two of the Alice/Mayo framework.\textsuperscript{287}

The 2019 Guidance was generally perceived as lowering Section 101 barriers to patentability, especially for computer-related inventions.\textsuperscript{288} Some commentators praised the Guidance for providing greater clarity to patent examiners, while other stakeholders criticized the Guidance as inconsistent with the Supreme Court’s Section 101 decisions.\textsuperscript{289}

Although the PTO’s 2019 Guidance changes how PTO examiners review new patent applications, the Guidance is not binding on the courts when patents are challenged in litigation (unlike decisions of appellate courts or statutes). The PTO lacks general substantive rulemaking authority,\textsuperscript{290} and the Guidance itself states that it is only a “tool for internal [PTO] management” that lacks “the force and effect of law.”\textsuperscript{291} Although the Federal Circuit has issued somewhat

\textsuperscript{283} MPEP § 2106.04(a) (internal cross-references omitted).

\textsuperscript{284} Id.

\textsuperscript{285} Id. at § 2106.04. The PTO calls the Alice/Mayo test’s first step “Step 2A” of its Section 101 examination process. See id.

\textsuperscript{286} Id. at § 2106.04(d) (emphasis added).

\textsuperscript{287} Id. at §§ 2106, 2106.04(d).


\textsuperscript{289} See generally Stuart P. Meyer, No Shortage of Viewpoints on New USPTO Eligibility Guidelines, BILSKI BLOG (Mar. 26, 2019), https://www.fenwick.com/bilski-blog/no-shortage-of-viewpoints-on-new-uspto-patent-eligibility-guidelines (reviewing comments received by PTO on the 2019 Guidance and noting that “both the ‘new Guidance is great’ and the ‘new Guidance doesn’t follow Alice’ camps are very well represented”).

\textsuperscript{290} Merck & Co. v. Kessler, 80 F.3d 1543, 1549–50 (Fed. Cir. 1996) (holding that while the PTO may promulgate regulations directed to the conduct of its own proceedings, it lacks authority to “issue substantive rules” under the Patent Act); Ass’n for Molecular Pathology v. U.S. PTO, 689 F.3d 1303, 1357 (Fed. Cir. 2012) (Bryson, J., concurring in part and dissenting in part) (“As we have recognized, the PTO lacks substantive rulemaking authority as to issues such as patentability.”); see generally Melissa F. Wasserman, The Changing Guard of Patent Law: Chevron Deference for the PTO, 54 WM. & MARY L. REV. 1959, 1962 (2013) (“[The PTO] lacks robust substantive rule-making authority and receives no judicial deference for its legal interpretations of the Patent Act.”).

\textsuperscript{291} 2019 Guidance, supra note 276, at 51.
contradictory signals on this point, the Guidance would receive, at most, “some deference” if a court found its reasoning to be persuasive.

Following the 2019 Guidance, the PTO has continued efforts to increase clarity and consistency in its Section 101 determinations. In 2020, the PTO Office of the Chief Economist issued a report on patent examination outcomes following Alice. That study found that while Section 101 rejections in certain technological fields increased by 31% in the 18 months after Alice, the rejection rate decreased by 35% after issuance of the 2019 Guidance, with less variability in outcomes across examiners. In response to a 2021 letter from Senators Tillis and Cotton, the PTO launched the Deferred Subject Matter Eligibility Response Pilot Program, which invites selected patent applicants to defer consideration of subject-matter eligibility issues until other patentability issues (such as those under Sections 102, 103, and 112) are resolved.

In 2022, at the urging of a bipartisan group of Senators, the PTO solicited public comment and published a report for Congress summarizing stakeholder views on current patent-eligible subject matter law. While the report found a consensus that patent-eligibility law should be “clear, predictable, and consistently applied,” stakeholders differed on whether current Section 101 law achieved that ideal. Finding a “continuing divide” on the issue, the PTO report indicated that defenders of the Alice/ Mayo framework (primarily from the computer technology industry) found

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292 Compare Nat. Alternatives Int’l, Inc. v. Creative Compounds, LLC, 918 F.3d 1338, 1346 n.2 (Fed. Cir. 2019) (noting that “[t]he parties dispute the persuasiveness of this document and the weight we should afford it under [Skidmore],” but declining to decide whether the 2019 Guidance should receive any deference), with Cleveland Clinic Found. v. True Health Diagnostics LLC, 760 F. App’x 1013, 1020 (Fed. Cir. 2019) (“While we greatly respect the agency, and given the value of uniformity in its administrative and judicial understandings of what a national law requires.”) (citations omitted); Skidmore v. Swift & Co., 323 U.S. 134, 140 (1944) (“The weight of [an informal agency] judgment in a particular case will depend upon the thoroughness evident in its consideration, the validity of its reasoning, its consistency with earlier and later pronouncements, and all those factors which give it power to persuade, if lacking power to control.”).


296 Id. at 1.


301 Id. at ii, 41.
current law to be sufficiently clear and an important tool for addressing overbroad patents and abusive lawsuits. On the other side, critics of the Alice/Mayo framework (especially life-science industries) found the current law to be unpredictable and to have detrimental effects on innovation and investment in the development of new technologies.

### Legislative Developments in the 116th Congress

#### The First Tillis-Coons Proposal

In the 116th Congress, Senators Tillis and Coons, along with Representatives Collins, Johnson, and Stivers, released a “bipartisan, bicameral framework” for legislative Section 101 reform (the First Tillis-Coons Proposal). The framework’s release followed multiple roundtables with patent law stakeholders on Section 101 and the effect of the Alice/Mayo framework on, for example, innovation in artificial intelligence, medical diagnostics, and personalized medicine.

The First Tillis-Coons Proposal would have retained the four current statutory categories of patentable inventions, but removed the requirement that the invention or discovery be “new and useful” from Section 101. Patent eligibility would have instead been determined “by considering each and every element of the claim as a whole and without regard for considerations properly addressed by [Sections] 102, 103 and 112 [of the Patent Act].”

In place of the judicially created exceptions to patent eligibility, which the First Tillis-Coons Proposal would have abrogated by statute, the proposal listed five “exclusive” categories of patent-ineligible subject matter: (1) fundamental scientific principles; (2) products that exist solely and exclusively in nature; (3) pure mathematical formulas; (4) economic or commercial principles; and (5) mental activities. Effectively, this would have codified aspects of the judicial exceptions in a narrower form, with the first two ineligible categories roughly corresponding to the “law of nature” and “natural product” judicial exceptions, and the final three to the types of “abstract ideas” identified by the PTO in its 2019 Guidance. The Proposal would have narrowed the construction of these ineligible categories by creating a “practical application” test, akin to the ABA proposal, expressly permitting patenting of a practical application of ineligible subject matter. But “simply reciting generic technical language or generic functional language” would have been insufficient to “salvage an otherwise ineligible claim.”

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302 Id. at 41.
303 Id.
305 Id.; see generally “The Debate Over Alice/Mayo and Section 101 Reform.”
306 First Tillis-Coons Proposal, supra note 29.
307 Id.
308 Id.
310 First Tillis-Coons Proposal, supra note 29.
311 See supra note 257 and accompanying text.
312 First Tillis-Coons Proposal, supra note 29.
The First Tillis-Coons Proposal thus blended elements of the PTO’s 2019 Guidance with a “laundry list” approach of specific ineligible categories, plus new statutory standards for how to apply the list of exceptions to patentable subject matter. The overall effect would be to lower Section 101 barriers to patentability, while still retaining more narrowly defined classes of ineligible subject matter.

Reactions to the First Tillis-Coons Proposal were mixed. Some commentators argued that the draft proposal was a promising start for much-needed congressional intervention. Indeed, some critics of the Alice/Mayo framework argued that the First Tillis-Coons Proposal did not go far enough, and urged elimination of any ineligible categories of patentable subject matter. On the pro-Alice side of the debate, the Electronic Frontier Foundation, for example, criticized the First Tillis-Coons Proposal as detrimental to innovation because it would eliminate a powerful tool to combat bad patents and patent troll litigation.

The Second Tillis-Coons Proposal

Following feedback on their first draft framework, the same group of Members released a “draft bill” to reform Section 101 (the Second Tillis-Coons Proposal). The Second Tillis-Coons Proposal was released before a series of three hearings held in the 116th Congress before the Senate Judiciary Committee’s Subcommittee on Intellectual Property, which solicited feedback on the draft legislative language. In these hearings, 45 witnesses testified over three days, with representatives from industry, academia, bar associations, and trade groups; former Federal Circuit Judges and PTO officers; and other patent law stakeholders expressing various views on Section 101 reform.

313 See supra “Specific Statutory List of Included or Excluded Subject Matter Categories”; “Administrative Developments: PTO Subject Matter Eligibility Guidance”; see also Nelson & Smith, supra note 309 (“[The First Tillis-Coons Proposal] includes some aspects of the proposals from several patent specialty associations, including those from the AIPLA/IPO, IPLAC, and the ABA-IPJL section.”).
314 See Daniel T. Taskalos, Returning to the Status Quo?—Proposed Outline for Section 101 Reform, NAT’L. L. REV. (Apr. 22, 2019), https://www.natlawreview.com/article/returning-to-status-quo-proposed-outline-section-101-reform (“In all, the proposed framework appears to focus on returning the 101 analysis to its previous status as more of a low hurdle to patentability, but a hurdle nonetheless.”).
320 Id.
321 See generally Coons & Tillis, supra note 31. For a succinct summary of the main views expressed at the hearings, see Bruce M. Wexler et al., Senate Hearing on “The State of Patent Eligibility in America”: Analysis of Viewpoints on Looming Section 101 Change, PAUL HASTINGS (June 25, 2019), https://www.paulhastings.com/publications-items/details/?id=c58c536d-2334-6428-811c-f00004cbded1. For a more detailed witness-by-witness breakdown, see...
As compared with the first proposal, the Second Tillis-Coons Proposal would have made more sweeping changes to Section 101 to expand patent eligibility. Like the First Tillis-Coons Proposal, the draft bill had several provisions that attempted to separate the Section 101 inquiry from other patentability requirements. Specifically, the draft bill would have struck the word “new” from Section 101 and established that patent subject matter eligibility must be determined “considering the claimed invention as a whole” and without regard to “considerations relating to section 102, 103, or 112 of [the Patent Act].”322 The Second Tillis-Coons Proposal provided that eligibility determinations would not depend on the “manner in which the claimed invention was made; whether individual limitations of a claim are well known, conventional or routine; [or] the state of the art at the time of the invention.”323 The draft bill also explicitly provided that Section 101 “shall be construed in favor of eligibility.”324

Rather than narrow the judicial exceptions to patentability, the Second Tillis-Coons Proposal would have eliminated those exceptions altogether. The draft bill provided that

No implicit or other judicially created exceptions to subject matter eligibility, including “abstract ideas,” “laws of nature,” or “natural phenomena,” shall be used to determine patent eligibility under section 101, and all cases establishing or interpreting those exceptions to eligibility are hereby abrogated.325

This language would have overturned by statute not only the Alice/ Mayo framework, but over two centuries of judicial decisions interpreting the “common law” exceptions to Section 101.326

The Second Tillis-Coons Proposal would have replaced the judicial exceptions with a new statutory definition of utility that incorporated elements of various prior proposals for a new Section 101 standard.327 To be patent-eligible subject matter under the Second Tillis-Coons Proposal, the invention would need to fit into one of the four statutory categories of eligible subject matter (which remain unchanged) and be “useful.”328 To be “useful,” an invention or discovery would need to provide “specific and practical utility in any field of technology through human intervention.”329

Finally, to combat overbroad patent claims, the Second Tillis-Coons Proposal would have altered the functional claiming rules under Section 112(f), which permits patentees to claim their invention in functional terms as opposed to reciting specific physical structures.330 In particular, the draft bill provided that if any patent claim element is “expressed as a specified function without the recital of structure, material, or acts in support thereof,” then that claim element will be limited to the “corresponding structure, material, or acts described in the specification” and


322 See Second Tillis-Coons Proposal, supra note 30 (proposed § 101(a)–(b) and “Additional Legislative Provisions”).

323 Id. (“Additional Legislative Provisions”).

324 Id.

325 Id.

326 See supra “Historical Development of the Judicial Exceptions to Patent-Eligible Subject Matter.”

327 See supra “Replace Judicial Exceptions with a Different Standard”; “Section 101: Utility.”

328 See Second Tillis-Coons Proposal, supra note 30 (proposed § 101(a)).

329 See id. (proposed § 100(k)). The draft bill did not further define “practical utility,” “field of technology,” or “human intervention.”

330 See Coons & Tillis, supra note 31 (indicating that the Section 112(f) amendments were intended “to guard against . . . overly broad, functional patent claims”); see generally “Section 112(f): Functional Claiming” (summarizing current law of functional claiming).
their equivalents.\textsuperscript{331} Consistent with decisions of the Federal Circuit,\textsuperscript{332} this language would have clarified that Section 112(f) applies to any claim element that fails to sufficiently recite a structure for performing a function.\textsuperscript{333} This change could have arguably made it tougher for a patentee to avoid the limiting effects of Section 112(f), even if the words “means for” are not used in the claim language.\textsuperscript{334}

As with the first proposal, reactions to the Second Tillis-Coons Proposal from patent law stakeholders were mixed.\textsuperscript{335} Critics of the Alice/Mayo framework generally applauded the draft bill as bringing much needed clarity and certainty to the law of patent eligibility,\textsuperscript{336} particularly for biotechnology innovation.\textsuperscript{337} Opponents of the draft bill expressed concern that changes to the Alice/Mayo framework would eliminate an important tool against unmeritorious patent litigation.\textsuperscript{338} Critics also questioned the necessity and advisability of such a sweeping change to Section 101 law.\textsuperscript{339} Both supporters and opponents raised concerns about potential ambiguities in the proposed definition of “useful,” particularly the terms “human intervention,” “practical utility,” and “field of technology.”\textsuperscript{340}

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\textsuperscript{331} Second Tillis-Coons Proposal, supra note 30 (proposed § 112(f)).

\textsuperscript{332} Williamson v. Citrix Online, LLC, 792 F.3d 1339 (Fed. Cir. 2015) (en banc).

\textsuperscript{333} Compare Second Tillis-Coons Proposal, supra note 30 (proposed § 112(f)), with 35 U.S.C. § 112(f). See also Patent Eligibility Hearings, supra note 31 (statement of Christopher A. Mohr, Vice President for Intellectual Property and General Counsel, Software and Information Industry Association), at 11, https://www.judiciary.senate.gov/download/mohr-testimony (“[The proposed § 112(f) language appears to do little more than cement the Federal Circuit’s Williamson v. Citrix decision . . . .”).

\textsuperscript{334} See Patent Eligibility Hearings, supra note 31 (statement of David W. Jones, Executive Director, High Tech Inventors Alliance), at 12, https://www.judiciary.senate.gov/download/06/05/2019/jones-testimony [hereinafter Jones Testimony] (“[The proposed Section 112(f)] amendment represents a modest improvement over the current language and will eliminate lingering arguments about the effect of inclusion or omission of the words ‘means for’ and whether particular terms should be interpreted as functional in the wake of [Williamson v. Citrix].”).

\textsuperscript{335} See generally Wexler et al., supra note 321 (summarizing arguments made by supporters and opponents of the Second Tillis-Coons Proposal).


\textsuperscript{337} See, e.g., Patent Eligibility Hearings, supra note 31 (statement of Laurie Hill, Vice President, Intellectual Property, Genentech, Inc.), at 8, 15–16, https://www.judiciary.senate.gov/download/hill-testimony (supporting the Second Tillis-Coons Proposal as “a strong step in the right direction” because of the “present uncertainty surrounding Section 101 [that] threatens to disrupt the development of a wide range of important medicines, diagnostics, treatments, and other innovations that benefit society”).

\textsuperscript{338} See, e.g., Gugliuzza Testimony, supra note 236, at 6–7 (arguing that “completely dismantling the eligibility requirement would take away a crucial tool courts can use to end, at relatively low cost, patent cases that plainly lack merit.”).

\textsuperscript{339} See, e.g., Jones Testimony, supra note 334, at 7 (“The evidence and arguments that have been advanced by proponents [of Section 101 reform] simply do not provide any reasonable justification for . . . . the complete abrogation of two centuries of eligibility case law.”).

\textsuperscript{340} See, e.g., Dickinson Testimony, supra note 336, at 33–34; Jones Testimony, supra note 334, at 10–11.
Stakeholders also debated the specific practical effects of the legislative changes at the hearings, such as the effect of elimination of the judicial exceptions on basic scientific research.\(^{341}\) One concern, raised by the American Civil Liberties Union in opposition to the draft bill, was that the Second Tillis-Coons Proposal, by abrogating the Myriad decision,\(^{342}\) would permit the patenting of human genes.\(^{343}\) Several witnesses denied that the draft bill would lead to that result because of the bill’s “human intervention” requirement or other patent law principles.\(^{344}\) For their part, Senators Tillis and Coons made clear that they had “no intention” of overruling the result in Myriad that no one may patent “genes as they exist in the human body.”\(^{345}\) Senators Tillis and Coons stated that the hearings in the 116th Congress reinforced their view that “patent eligibility is broken and desperately needs to be repaired,” and that there is a “necessity for Congress to intervene” to bring greater clarity to Section 101.\(^{346}\) Ultimately, the Members did not formally introduce a Section 101 reform bill during the 116th Congress.

### Legislative Developments in the 117th Congress

The 117th Congress to date has seen two introduced bills proposing reforms to Section 101, one in the Senate and one in the House.

### The Patent Eligibility Restoration Act of 2022

In the Senate, Senator Tillis introduced S. 4734, the Patent Eligibility Restoration Act of 2022 (PERA). PERA would retain the four statutory categories of eligible subject matter, but delete the word “new” in Section 101 and add a new definition of “useful.”\(^{347}\) PERA’s utility definition would require that “the invention or discovery has a specific and practical utility from the perspective of a person of ordinary skill in the art.”\(^{348}\) Moreover, PERA would change the definition of “process” to clarify that “a use, application, or method of manufacture of a known or naturally-occurring process” is patentable.\(^{349}\) PERA would also establish that patent eligibility determinations shall be made without regard to “any consideration in [35 U.S.C.] section 102, 103, or 112” including “whether a claim element is known, conventional, routine, or naturally occurring.”\(^{350}\)


\(^{342}\) See supra notes 169–173 and accompanying text (discussing the Supreme Court’s decision in Association for Molecular Pathology v. Myriad Genetics, Inc.).

\(^{343}\) See, e.g., Patent Eligibility Hearings, supra note 31 (statement of Kate Ruane, Senior Legislative Counsel, Washington Legislative Office, ACLU) at 3, https://www.judiciary.senate.gov/download/ruane-testimony (arguing that the Second Tillis-Coons Proposal “would clearly make human genes, isolated from the rest of the genome, patent-eligible again”).


\(^{346}\) Coons & Tillis, supra note 31.

\(^{347}\) S. 4734, 117th Cong. § 2.

\(^{348}\) Id. § 2(a)(1)(B).

\(^{349}\) Id. § 2(a)(1)(A).

\(^{350}\) Id. § 2(a)(2).
Like the First Tillis-Coons proposal in the 116th Congress, PERA contains a closed list of the types of inventions that are not patent-eligible when claimed “as such,” specifically:

(A) A mathematical formula, apart from a useful invention or discovery.

(B) A process that—
   (i) is a non-technological economic, financial, business, social, cultural, or artistic process;
   (ii) is a mental process performed solely in the human mind; or
   (iii) occurs in nature wholly independent of, and prior to, any human activity.

(C) An unmodified human gene, as that gene exists in the human body.

(D) An unmodified natural material, as that material exists in nature.

In effect, PERA would abrogate the Alice/Mayo framework, and replace the three judicially created ineligible categories with this closed statutory list of narrower ineligible categories.351

The Restoring America’s Leadership in Innovation Act of 2021

In the House, Representative Massie introduced H.R. 5874, the Restoring America’s Leadership in Innovation Act of 2021 (RALIA).352 Alongside provisions designed to reverse many of the changes in patent law enacted through the 2011 America Invents Act,353 Section 7 of RALIA responds to the Supreme Court’s Section 101 decisions. Expressing the view that the Court’s recent Section 101 jurisprudence “has harmed the progress of science and the useful arts,” the bill would “effectively abrogate[]” those decisions (specifically, Bilski, Mayo, Alice, Myriad, “[and] their predecessors”).354

To “ensure that life sciences discoveries, computer software, and similar inventions and discoveries are patentable,” RALIA would replace the three judicially created exceptions to patent-eligible subject matter with a single, relatively narrow statutory exception. Under RALIA, any new and useful process, machine, manufacture, or composition of matter is patent-eligible unless “the claimed invention as a whole, as understood by a person having ordinary skill in the art, exists in nature independently of and prior to any human activity, or exists solely in the human mind.”355 RALIA would therefore generally expand the types of inventions that are patentable even further than PERA would. Like PERA, RALIA abrogates the Alice/Mayo framework and provides that eligibility determinations under Section 101 shall be made “without regard as to the requirements or conditions of sections 102, 103, and 112 of this title, or the claimed invention’s inventive concept.”356

Conclusion

The Supreme Court’s 2010s decisions on patent-eligible subject matter have inspired a robust debate among patent law stakeholders as to whether the Court’s jurisprudence in this area advances or harms innovation. Recent actions by the courts, the PTO, and Congress have

351 See id. (providing that the four statutorily eligible categories would be “subject only to the [listed] exclusions”).
353 See, e.g., id. at §§ 4–5 (abolishing the PTAB and the IPR/PGR procedures).
354 Id. at § 7(b).
355 Id. at § 7(a).
356 Id.
responded to the Court’s decisions in various ways, including proposed statutory changes discussed in the 116th Congress and introduced in the 117th Congress.

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