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M. Stuart Madden*

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Abstract

The reach of Federal statutory preemption of inconsistent state law obligations has extended to numerous products liability subject matters, including most notably tobacco products, agricultural pesticides, medical devices and automobile air bags. The Supreme Court decision in Cipollone v. Liggett Group, Inc. countenanced a broad application of federal preemption when the subject statute contained an express preemption clause. Eight years later the Court appeared to back away from Cipollone, and held in Geier v. American Honda Motor Co. that even as to statutes with express preemption clauses, the simultaneous presence of a savings clause might trigger a narrow reading of the preemption provision. The potential effect of Geier upon lower court decisions in liability suits involving pesticides, medical devices, or even tobacco products, will necessarily be played out in litigation before lower courts for years to come.

I. INTRODUCTION

The significance of federal statutory preemption of inconsistent state tort law obligations was branded on the hindquarters of products liability law in the 1992 Supreme Court decision of Cipollone v. Liggett Group, Inc. Cipollone's emphasis on textual express preemption was followed by numerous lower court decisions holding that federal health and safety rules in product subject categories ranging from pesticides to

prostheses to propellers should be given sway over inconsistent state common law liability or statutory obligations.

The Cipollone-sparked romp over the historical federal hesitation to impose federal regulation in areas of health, safety and welfare traditionally ceded to the individual states was brought to a pause, if not a halt, by the Supreme Court’s 2000 decision in Geier v. American Honda Motor Co.2 In Geier, an air bag case, the Court gave new vitality to the application of ordinary preemption principles even in the context of a federal safety statute that contained an express preemption provision clause, should such statute also contain a “savings” clause provision essaying to preserve common law damage claims from any preemptive consequences. The implications of Geier and its explicit distancing from Cipollone will surely roil the waters of products liability litigation for years to come.

II. TYPES OF PREEMPTION

A. Express preemption

Perhaps the most important emerging issue in modern products liability law is the role of the federal statutory preemption of private civil suits brought under state statutory or common law. Characteristically, the question of the appropriateness of federal preemption is implicated when such state law claims for damages or equitable remedies trench upon the same safety objectives, which Congress has addressed directly, or indirectly through federal statute or administrative regulation. As discussed below, the Supreme Court’s 1992 decision in Cipollone v. Liggett Group, Inc.3 gave new definition to the constitutional doctrine of federal legislative preemption, and provided a basis for modern application of the tenets of that doctrine in a range of accident law and regulatory settings.

In general terms, federal safety-related statutes, or regulations pursuant to those statutes, that pertain to a particular field or subject matter may be deemed to preempt state regulation or common law that would impose design, performance, or informational requirements upon a seller that are inconsistent with the federal standard. Federal law may also preempt suits

brought under state common law or state codification of common law principles that create the risk that a seller who has satisfied federal safety-related requirements might nevertheless be found liable in money damages, or subject to equitable relief, in a products liability suit.\textsuperscript{4} The primacy of federal law in such subject areas as Congress may elect to regulate is grounded in the Supremacy Clause of Article VI of the Constitution, which provides that the laws of the United States “shall be the supreme Law of the Land; and . . . any Thing in the Constitution or Laws of any State to the Contrary notwithstanding.”\textsuperscript{5}

Preemption may be either express or implied, and “is compelled whether Congress’s command is explicitly stated in the statute’s language or implicitly contained in its structure and purpose.”\textsuperscript{6} Issues of express preemption are textual, while questions of implied preemption are contextual. By this it is meant that the question of whether federal law expressly preempts inconsistent or additional state common law, statutory or regulatory requirements may be deduced from the explicit language of a federal statute.\textsuperscript{7} Implied preemption, in contrast, must ordinarily be inferred from an evaluation of not only the language of the statute, which may or may not contain a preemption clause, but also from an assessment of the overall statutory objectives. As to the latter, a full understanding of statutory objectives will frequently be informed by pertinent


Also, the application of preemption principles to state statutory/regulatory and common law pursuits alike is explained by the Supreme Court’s observation in San Diego Building Trades Council v. Garmon, 359 U.S. 236, 247 (1959), that awards of money damages in suits for civil liability can act as a “potent method of governing conduct and controlling policy.”

\textsuperscript{5} U.S. Const. art. VI, § 2.

\textsuperscript{6} Jones v. Rath Packing Co., 430 U.S. 519, 525 (1977); See generally Oja v. Howmedica, Inc., 111 F.3d 782 (10th Cir. 1997) (held: negligent failure to warn claim in hip prosthesis products liability action is not preempted under 1976 Medical Device Act Amendments). Preemption issues associated with medical devices are discussed below in § II(H).

\textsuperscript{7} See generally Cipollone, 505 U.S. 504 (1992), discussed in detail at II(A) below.
legislative history and the interpretation of the statute given it by the regulatory agency charged with its effectuation.  

The approach taken by the Supreme Court in its interpretation of the Supremacy Clause as applicable to state law preemption "starts with the assumption that the historic police powers of the States [are] not to be superseded by . . . Federal Act unless that [is] the clear and manifest purpose of Congress." Express preemption is properly found "[w]hen Congress has considered the issue of preemption and has included in the enacted legislation a provision explicitly addressing that issue, and when that provision provides a 'reliable indicium of congressional intent with respect to state authority[.]'" In the words of one court concerning express preemption:

Federal preemption is a relatively simple concept, especially when Congress has explicitly provided the terms of preemption. It provides order. Instead of having 50 or more standards with respect to a given human pursuit, there is one. When a preemptive federal standard is applied evenhandedly, it further provides both the protection of the federal standard and some leeway to develop state standards where the federal standard does not apply.

A party advancing the defense of federal preemption must overcome an established presumption against federal preemption of state law. Thus any statutory provision forming the basis of a preemption defense will be narrowly construed, consistent with the tenet that state police powers, particularly state regulations relating to health and safety, should not be

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8. Implied preemption of state common law or statutory products liability litigation is discussed in § I(B) below.
9. Cipollone, 505 U.S. at 516 (citing Rice v. Santa Fe Elevator Corp., 331 U.S. 218, 230 (1947)).
10. Id. at 517.
superseded without a demonstration that such preemption is the "clear and unequivocal intent of Congress."\textsuperscript{13}

B. \textit{Implied preemption}

i. \textit{General implied preemption}

Where a federal safety-related statute does not contain language providing expressly for its preemptive effect, and, where pertinent, the preemptive effect of regulations or standards promulgated pursuant thereto, the products liability defendant will attempt to argue that the language of the statute, the context of the federal regulatory interest, and the wishes of the congressional authors, read together, require the conclusion that inconsistent or additional state law claims or requirements are impliedly preempted.\textsuperscript{14} This subsection and the two to follow describe briefly the three principal theories underlying the doctrine of implied preemption.\textsuperscript{15}

Defendant's defense of general implied preemption may succeed upon a showing that the scope of the safety-related statute, including a fact-specific appreciation of its comprehen-


As stated by the Supreme Court in \textit{Medtronic}, matters of the citizenry's health and safety "are primarily and historically matters of local concern[,]" and therefore "the States traditionally have a great latitude under their police powers to legislate as to the protection of the lives, limbs, comfort and quiet of all persons." \textit{Medtronic}, 518 U.S. at 475.


\textsuperscript{15} The three analytically distinct types of implied preemption that courts have recognized are (1) general implied preemption, (2) field preemption, and (3) conflict preemption. Ordered differently, the factor analysis employed by many courts in adopting these distinctions were summarized by the Supreme Court in this way: (1) "Congress' intent to supersede state law altogether may be found from a 'scheme of federal regulation so persuasive as to make reasonable the inference that Congress left no room to supplement it;'" (2) an act of Congress may touch a field in which the federal interest is "so dominant that the federal system will be assumed to preclude enforcement of state laws on the same subject;" or (3) because "the object sought to be obtained by the federal law and the character of obligations imposed by it may reveal the same purpose." \textit{Fidelity Fed. Sav. & Loan Assoc. v. de la Cuesta}, 458 U.S. 141, 153 (1982).
siveness and its particularity, taken together with the observations of its congressional authors and its regulatory legal experts, demonstrates that toleration of inconsistent or additional state common law or statutory requirements would frustrate the purpose of the federal act. As the Supreme Court has explained, in the absence of language revealing "an explicit congressional intent to preempt state law," courts examining the potential that federal law impliedly preempts state statutory or common law claims should consider "whether the federal statute's 'structure and purpose,' or nonspecific statutory language, reveal a clear, but implicit, pre-emptive intent."

ii. Field preemption

Even though concepts of federalism preserve a constitutional reluctance to permit federal law to supplant state regulatory prerogative, particularly in matters associated with the health, safety and welfare of citizens of that state, there are two additional theories by which a federal safety-related statute may be deemed to impliedly preempt an additional or inconsistent state claim or requirement. A second means of implied preemption is termed "field preemption." A conclusion that state statutory or common law remedies are preempted by a federal statutory initiative may be warranted when examination of the federal statute and allied regulations permit the conclusion that in passing the statute and creating the regulatory authority of the pertinent federal agency or agencies, Congress intended that federal law and regulation effectively and functionally occupy the safety field that the state law or regulation would purport to enter. Field preemption may be found when Congress or its administrative delegees have "so thoroughly occupied a legislative field 'as to make reasonable the inference that Congress left no room for the states to supplement it.'"

A finding of field preemption of state statutory enforcement or common law liability claims does not turn upon whether the federal agency has actually initiated regulation in the pertinent

subject matter, but rather upon whether it is empowered to do so. The Supreme Court so confirmed in Napier v. Atlantic Coast Line R. Co. in which, interpreting the Locomotive Boiler Inspection Act, it wrote: "The fact that the [Interstate Commerce] Commission has not seen fit to exercise its authority to the full extent conferred, has no bearing on the construction of the act delegating the power.

iii. Conflict preemption

Third, additional or inconsistent state safety common law or statutory obligations or requirements may be deemed to be impliedly preempted where the obligations or prohibitions imposed by state statute or regulation "actually conflict[ ] with federal law," which is to say, in instances in which "compliance with both state and federal law is a 'physical impossibility,' or where state law 'stands as an obstacle to the accomplishment and execution of the full purposes and objectives of Congress.'" This type of implied preemption is described as "conflict" preemption.

As is true in matters of potential express preemption, courts have adopted a rebuttable presumption against all types of implied preemption that would supplant state legislative authority, particularly where such preemption is urged in actions implicating accident law. In one court's words: "Where the state laws at issue affect health and safety issues, there is a presumption against implied preemption by congressional enactments."
III. PREEMPTION AND REGULATORY ISSUES

A. Tobacco Product Labeling

Preemption analysis as it relates to tobacco product matters derives almost exclusively from the Supreme Court decision in Cipollone v. Liggett Group, Inc.\textsuperscript{26} In Cipollone, the Court reiterated that "Congress' intent may be 'explicitly stated in the statute's language or implicitly contained in the structure and purpose.'"\textsuperscript{27} Consistent with the discussion of implied preemption, the Court confirmed that even absent express or explicit preemption, state law may be preempted where "that law actually conflicts with federal law, or if federal law so thoroughly occupies a legislative field 'as to make reasonable the inference that Congress left no room for the States to supplement it.'"\textsuperscript{28}

The suit in Cipollone was filed in the New Jersey federal trial court by Rose Cipollone, who would die of lung disease one year later, and her husband. The plaintiffs alleged that cigarette manufacturers: (1) sold a product that was defectively designed, in that there were safer alternative designs for cigarettes, and because the social utility of cigarettes was vastly outweighed by the dangers inhering in their use; (2) failed to provide adequate warnings; (3) "were negligent in the manner [that] they tested, researched, sold, promoted and advertised" cigarettes; (4) breached express warranties as to the lack of health risks from smoking; (5) fraudulently misrepresented health risks, thereby neutralizing federally mandated warnings; (6) "ignored and failed to act upon" scientific and medical data showing the health risks of cigarette smoking; and (7) by "conspiracy to defraud" combined with other manufacturers to deny the public of medical and scientific information.\textsuperscript{29}

Section 5 of the 1965 Cigarette Labeling and Advertising Act ("1965 Act"), captioned "Preemption," provided "(a) No statement relating to smoking and health, other than the [Section 4] statement . . . shall be required on any cigarette package," and "(b) no [such] statement . . . shall be required in the

\textsuperscript{26} 505 U.S. 504 (1992).
\textsuperscript{27} Id. at 516.
\textsuperscript{28} Id.
\textsuperscript{29} Id. at 509-10. The Cipollone decision itself did not address the issue of product defectiveness. See Richardson v. Phillip Morris, Inc., 950 F.Supp 700 (D. Md. 1997).

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advertising of any cigarettes the packages of which are labeled in conformity with” Section 4.  

Section 5 (b) was amended by the Public Health Cigarette Smoking Act of 1969 (“1969 Act”) to specify that: “No requirement or prohibition based on smoking and health shall be imposed under state law with respect to the advertising and promotion of any cigarette the packages of which are [lawfully] labeled.”

The Third Circuit Court of Appeals held that Congress “had impliedly pre-empted . . . claims challenging the adequacy of warnings on labels or in advertising or the propriety of [the manufacturers’] advertising and promotional activities.” The Supreme Court reversed in part and affirmed in part. The Court found that the preemptive scope of the 1965 Act and the 1969 Act was governed entirely by the express language of § 5 of each Act, and that, therefore, no justification existed for evaluating the possibility of implied preemption.

At the same time, the Cipollone Court determined that the appellate court erred in concluding that the preemptive language of the two respective Acts was operatively indistinguishable. In the Court’s view, § 5 of the 1965 Act “only preempted state and federal rule-making bodies from mandating particular cautionary statements and did not preempt state common law damages actions.” The Court reasoned that (1) the presumption against the preemption of state police power regulations required a narrow reading of the 1965 preemption provision; (2) label warning mandated by the 1965 Act’s § 4

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32. Cipollone, 505 U.S. at 517 (citing the Third Circuit Court of Appeals ruling in Cipollone v. Liggett Group Inc., 789 F.2d 181, 187 (3d Cir. 1986)). In so doing, the Court observed that the Third Circuit had placed unwarranted reliance upon the fact that the 1969 Act did not alter the statement of purpose from the 1965 Labeling Act. Id. at 517 n. 13.
34. See Cipollone, 505 U.S. at 517.
35. See id.
36. Id. at 519-20. Elsewhere in its opinion the court uses the terminology “positive enactment” synonymously with a “mandat[e] [as to] particular cautionary statements[,]” Id. at 520.
37. See id. at 518-19.
"does not by its own effect foreclose additional obligations imposed under state law,"38 because the mere fact "that Congress requires a particular warning does not pre-empt a regulatory field,"39 and (3) "there is no general, inherent conflict between federal preemption of state warning requirements and the continued viability of state common law damages actions."40 All of these considerations led the Court to conclude that § 5 of the 1965 Act "superseded only positive enactments by legislatures or administrative agencies that mandate particular warning labels."41

Disagreeing with the Third Circuit, the Supreme Court found that the preemptive language in the 1969 Act was materially different—and substantially broader in application—than the preemptive language in the 1965 Act. The Court explained:

First, the [1969] Act bars not simply 'statement[s]' but rather 'requirement[s] or prohibition[s] ... imposed under State law.' Second, the later Act reaches beyond statements 'in the advertising' to 'obligations with respect to the advertising or promotion' of cigarettes.42

The Supreme Court continued by noting that this broader reading was justified, in part, on the basis that the 1969 Act substantially altered the 1965 Act by "rewriting the label warning, banning broadcast advertising, and allowing the FTC to regulate print advertising."43

In addition, the Court found that the phrase "imposed under state law" is not limited to "positive enactments" obligating manufacturers to employ particular cautionary language or symbols.44 Focusing on the 1969 Act's broader language of "requirements or prohibitions,"45 the Supreme Court considered whether or not personal injury claims brought under state law might operate as a de facto imposition of a requirement or a prohibition. The question the Court posed could be put this way: "Does imposition of or vulnerability to a plaintiff's verdict

38. Id. at 518.
39. Cipollone, 505 U.S. at 518.
40. Id.
41. Id. at 518-19.
42. Id. at 520.
43. Id.
44. Cipollone, 505 U.S. at 522.
45. Id. at 521-24.
arising from a claim that the manufacturer should have provided warnings in addition to those required by the 1969 Act impose an obligation upon the manufacturer that can be considered a requirement or a prohibition within the meaning of the 1969 Act?" The Supreme Court answered its own question affirmatively.46

The Court stated that "the central inquiry" in the matter before it was straightforward: "We ask whether the legal duty that is the predicate of the common-law damages action constitutes a 'requirement or prohibition based on smoking and health . . . imposed under State law with respect to . . . advertising or promotion,' giving that clause a fair but narrow reading."47 Applying this "straightforward" analysis to the common law claims advanced by Mrs. Cipollone and her husband, the Supreme Court held that a products liability action alleging failure to provide adequate warnings was in essence a claim that the manufacturer's "post-1969 advertising or promotions should have included additional, or more clearly stated, warnings[.]"48 Such a failure to warn claim was, by the terms of the 1969 Act, preempted.49

A different result was appropriate, however, for such tort claims as were not associated with the advertising or promotion of cigarettes. Thus, the Supreme Court made clear that the 1969 Act did not operate to preempt claims in tort that might be based upon "the testing or research practices [of a defendant,] or other actions unrelated to advertising or promotion [of cigarettes.]"50 Further, the Supreme Court continued, the plaintiffs' claims for breach of an express warranty were not precluded insofar as they derived from the contractual nature of the relationship between the seller and buyer, rather than being imposed by state law.51 Thus, the Cipollone Court concluded "a common law remedy for a contractual commitment voluntarily

46. See id. at 524.
47. Id. at 523-24.
48. Id. at 524.
49. Cipollone, 505 U.S. at 524.
50. Id. at 524-25.
51. See id. at 525-26. The Court's evaluation regarding express warranties is of limited modern significance, as today, unlike the era in which Mrs. Cipollone began to smoke, no tobacco manufacturer is so incautious as to plump the health virtues of smoking.
undertaken should not be regarded as a 'requirement ... imposed under State law' within the meaning of § 5(b)."

The plaintiffs alleged that the manufacturer had engaged in two distinct types of misrepresentation. Taking each misrepresentation claim in turn, the Court found that a claim for fraudulent misrepresentation premised upon an allegation that the advertising practices of the cigarette manufacturers neutralized the effect of federally-mandated warning labels was preempted by the 1969 Act. Noting the interrelationship between regulatory prohibitions on advertising that downplay dangers of smoking and the associated statutory requirements for warnings, the Court found that such a theory of fraudulent misrepresentation was inextricably related to a theory of failure to warn and was similarly preempted.

The Supreme Court recognized, however, an important distinction between fraudulent misrepresentation claims associated with advertising and other forms or venues of misrepresentation. An example of the former might be a billboard with an image of young vibrant persons engaged in vigorous physical activity, serving to convey the message that cigarette smoking is compatible with cardiovascular fitness. Examples of the latter type of misrepresentation such as evidence that the cigarette manufacturers concealed material facts from administrative agencies, or that they included false statements of material fact, e.g., carbon monoxide or tar levels, in their advertising, would not be preempted by the 1969 Act. The Court reasoned that this second type of fraudulent misrepresentation claim was predicated on state law duties to disclose material facts, rather than on advertising and promotion. To further clarify the distinction, the Supreme Court stated that in the former type of alleged fraudulent misrepresentation, such as billboards, print media and the like, state laws obligating manufacturers to disclose safety or health risks to administrative agencies were "obligations with respect to advertising or promotion" and a state law claim premised upon the breach of

52. Id. at 526.
53. See id. at 527.
54. Cipollone, 505 U.S. at 528.
55. See id.
that duty would be preempted.\textsuperscript{56} Claims that an advertisement contains a false statement of a material fact, in contrast, are not "predicated upon a duty based on smoking and health, but rather on a more general obligation—the duty not to deceive."\textsuperscript{57} Regarding plaintiff's conspiracy claims, the Court found these, too, were not preempted, as the duty not to conspire to commit fraud is not a prohibition "based on smoking and health."\textsuperscript{58}

In summarizing its holding, the Court wrote:

The 1965 Act did not pre-empt state law damages actions; the 1969 Act pre-empts petitioner's claims based on a failure to warn and the neutralization of federally mandated warnings to the extent that those claims rely on omissions or inclusions in respondents' advertising or promotions; the 1969 Act does not pre-empt petitioner's claims based on express warranty, intentional fraud and misrepresentation, or conspiracy.\textsuperscript{59}

Mr. Justice Blackmun, joined by Justices Kennedy and Souter, concurred in part and dissented in part, arguing that the modified language of § 5(b) in the 1969 Act did not clearly exhibit the necessary congressional intent to preempt state common law damage actions.\textsuperscript{60} The three Justices concluded, therefore, that the plaintiffs' various failure to warn and fraudulent misrepresentation claims, as well as express warranty and conspiracy claims, should not be preempted by the 1969 Act.\textsuperscript{61} Conversely, Justices Scalia and Thomas argued that all of petitioner's common law claims were preempted by the 1969 Act.\textsuperscript{62}

i. \textit{Warnings or misrepresentation claims regarding second-hand smoke or nicotine addiction}

The legislative history is ambiguous as to whether Congress, in either the 1965 or the 1969 Acts, contemplated that the labeling provisions should cover risks that were then unknown, or as to which the scientific and medical data were in-

\begin{itemize}
\item \textsuperscript{56} \textit{Id.} at 528.
\item \textsuperscript{57} \textit{Id.} at 528-29.
\item \textsuperscript{58} \textit{See id.}
\item \textsuperscript{59} \textit{Cipollone}, 505 U.S. at 530-31.
\item \textsuperscript{60} \textit{See id.} at 531-44.
\item \textsuperscript{61} \textit{See id.}
\item \textsuperscript{62} \textit{See id.} at 544-56.
\end{itemize}
cipient, or as it is sometimes put, "immature." *Cipollone* did not address the issue of whether the 1969 Act should be interpreted to preempt warning or misrepresentation claims regarding risks allegedly associated with second-hand smoke or nicotine addiction. Read together, the statements of purpose and the committee reports underlying the 1965 and the 1969 Acts permit no confident conclusion. Evaluation of whether the 1969 Act ought properly to preempt claims that the manufacturers fraudulently concealed or purposefully misrepresented information regarding claimed risks of injurious exposure to second-hand smoke or nicotine addiction requires examination of the legislative tradeoffs, or bargained-for exchanges, if any, that were involved in the consideration of and drafting of the 1969 Act. In this connection, the legislative history of the 1965 Act also enjoys a role, as the metes and bounds of congressional consideration in both the 1965 and the 1969 Acts may, to a cautious extent, be considered collectively.

Section 2 of the 1965 Act declared that the statute's two purposes were "(1) adequately informing the public that cigarette smoking may be hazardous to health, and (2) protecting the national economy from the burden imposed by diverse, nonuniform, and confusing cigarette labeling and advertising regulations."63 Both the statement of the 1965 Act's purpose, and the language comprising its required labeling, make it clear that Congress intended the warnings to convey to the public only that cigarette "smoking" could be hazardous to health. Section 2 further states that the goal of the mandated warning language is that of "informing the public that cigarette smoking can be hazardous to health," while Section 4 of the 1965 Act required a specific label stating, among other things: "Caution: Cigarette Smoking May Be Hazardous To Your Health."64 Nowhere referenced in the statute's statement of purpose or its required labeling is the suggestion that a message was intended to be conveyed to nonsmokers, i.e., information suggesting that smoking by others could be hazardous to nonsmokers. It appears, therefore, that in the absence of any arguable revelation of congressional intent that this too was an objective of the 1965 or 1969 Acts, neither Act can logically be construed to apply to

63. *Id.* at 514.
64. *Cipollone*, 505 U.S at 514.
the health claims arising from the health risks of exposure to secondhand smoke.

The legislative history for the 1969 Act contains several references to the Congressional authors' awareness of (1) the risks of inhalation of environmental smoke and (2) the potential that with the passage of time, additional health risks of tobacco would be discovered and proved. Senate Report 91-566 and the accompanying Conference Report to Public Law 91-222, otherwise known as the Public Health Cigarette Smoking Act of 1969, contains several references to cigarette-related health risks. The fair inference of this is that, in passing the 1969 Act, (1) Congress was aware of health dangers other than the orthodox litany, i.e., respiratory disease, lung disease, pulmonary disease; and (2) Congress was further aware of health risks posed to individuals other than the smokers themselves.

As to the issue of nicotine dependence, the Conference Report to the 1969 Act cited conspicuously a 1967 FTC Report, filed pursuant to the 1965 Act § 5(d)(2), which stated that the very fact that "cigarette smoking is so strongly habit forming" was a basis for its statement that "it is unlikely that a mildly phrased cautionary statement will have any effect on confirmed cigarette smokers." The Senate Report later reiterated the FTC's observations that "another aspect of cigarette smoking that is... ignored, and has vital implications in terms of health hazard, is the fact that cigarette smoking is strongly habit forming." As a result of the reports, the FTC recommended the listing of nicotine content on cigarette packages.

Further reflective of Congress's awareness that not only tar levels, but also nicotine levels, contributed to the health risks, S. Rep. No. 91-566 referenced the Department of Health, Education and Welfare's (HEW) recommendation, filed pursuant to the 1965 Act, that "levels of 'tar' and nicotine in cigarette smoke should be published on cigarette packages[.]" That the 1969 Act contemplated the potential of new discernible health risks

of tobacco use was reflected in § 8 of the Act, “continu[ing] the requirement that the Secretary of [HEW] . . . transmit reports to Congress . . . providing current information on the health consequences of smoking[.]”71 In addition, the Senate Bill confirmed that the preemption entailed “is narrowly phrased to preempt only state action based on smoking and health. . . . It would in no way,” the Report points out, “affect the power of any state . . . [regarding] the prohibition of smoking in public buildings[.]”72

Commentators have urged that in the specific context of second-hand smoke claims, the labeling and advertising requirements do not extend to nonsmokers subjected to second-hand smoke; “reliance on the notice given to smokers via the labeling act[s] . . . cannot apply to nonsmokers, as they are not the ones who are warned through the contents of the cigarette label.”73 To further the argument that the warning provisions apply to smokers and not to nonsmokers is the fact that the language of the 1965 Act references the relationship between cigarette “smoking” and health, rather than a reference to the effects of “smoke” and health, a usage equally available to the legislative authors of the Act.

As to the question of whether at the very time of passing the 1969 Act Congress visualized that additional tobacco use health risks might be discovered, as previously noted, the pertinent Conference Report noted the inclusion of a new § 8 to the Act that directed HEW to report to Congress “current information on the health consequences of smoking . . .”74 One potential interpretation of this provision is that the congressional authors imagined that yet undiscovered health risks tied to tobacco use might be unearthed in the future and that Congress should not be considered to have preempted that which has not visualized. However, preemption advocates might respond that warnings or misrepresentation state law claims associated with such new risks would bear the unmistakable earmarks of “requirements or prohibitions relating to advertising or promotion

74. Id. at 33, 34.
of cigarettes" as those terms of art have been constructed by the Cipollone Court.\textsuperscript{75}

Various state disclosure laws have been held not to tram-mel preemption provisions under comparable tobacco-related acts.\textsuperscript{76} For example, the Court of Appeals for the Fifth Circuit affirmed the district court’s ruling that a state Disclosure Act was not preempted by either the 1969 Act or the Smokeless Tobacco Act.\textsuperscript{77} At issue was the Massachusetts Disclosure Act,\textsuperscript{78} that required “manufacturers of tobacco products to disclose the additives and nicotine yield ratings of their products to the state’s public health department.”\textsuperscript{79} The court in Philip Morris, Inc. v. Harshbarger\textsuperscript{80} held that the Disclosure Act did not ‘relate to’ advertising or promotion because it lack[ed] the requisite ‘reference to’ or ‘connection with’ the preempted realm.”\textsuperscript{81} The appellate court also declared that there existed no evidence to support defendant’s contention that Congress had intended to impose “national uniformity” with respect to disclosure or ingredient reporting regulations.\textsuperscript{82} To be compared is the decision of an Alabama federal trial court in Lacey v. Lorillard Tobacco Co. Inc.,\textsuperscript{83} in which the court held that common law claims regarding disclosure of cigarette ingredients were preempted by the 1969 Act.

B. Pesticide, Herbicide and Rodenticide Labeling

In the Federal Insecticide, Fungicide and Rodenticide Act (“FIFRA”), Congress established a comprehensive regulatory system for the registration and labeling of pesticides.\textsuperscript{84} Prior to being placed into the market, a manufacturer must register a

\begin{thebibliography}{9}
\bibitem{75} Cipollone, 505 U.S. at 506.
\bibitem{76} Philip Morris, Inc. v. Harshbarger, 122 F.3d 58 (1st Cir. 1997).
\bibitem{77} See id.
\bibitem{78} See \textit{Mass. Gen. Laws} ch. 94, § 307(B) (West 1996).
\bibitem{79} \textit{Harshbarger}, 122 F.3d at 58 (citing \textit{Mass. Gen. Laws} ch. 94, 307(B)).
\bibitem{80} \textit{Id.} at 74.
\bibitem{81} See \textit{id}.
\bibitem{82} \textit{Id.} at 85.
\bibitem{83} 956 F. Supp. 956 (N.D. Ala. 1997).
\end{thebibliography}
pesticide with the Environmental Protection Agency ("EPA"). As part of the registration process, the manufacturer must submit a proposed label to the EPA for approval. FIFRA requires that the label be "adequate to protect health and the environment" and be "likely to be read and understood." For purposes of this discussion, the preemption issues associated with FIFRA are limited to the ability of a state to permit common law damage actions based upon product labeling. The Supreme Court held in Wisconsin Public Intervenor v. Mortier (Ralph) that FIFRA does not preempt local governmental regulation of pesticide use.

FIFRA expressly prohibits states from imposing "any requirements for labeling or packaging in addition to or different from those required" under the Act. Although a state is permitted to regulate the sale or use of a registered pesticide to the extent that such regulation does not permit a sale or use prohibited by the Act, it is precluded from imposing labeling or packaging requirements which alter, in any way, those imposed by the EPA.

i. Decisional law prior to Cipollone v. Liggett Group

Prior to the Court's decision in Cipollone, the lower courts were divided with regard to the preemptive effect of FIFRA registration and labeling requirements upon suits brought under state common or statutory law for compensatory damages. In a pre-Cipollone decision finding no preemption, Ferebee v. Chevron Chemical Co., a trial verdict was rendered against Chevron in favor of an agricultural worker who died from pulmonary fibrosis allegedly contracted through long-term inhalation and dermal exposure to the herbicide Paraquat. Plaintiff claimed that the decedent's injuries were caused by Chevron's

85. See 7 U.S.C.A. § 136a(a) ("Except as provided in this Subchapter, no person in any state may distribute or sell to any person any pesticide that is not registered under this Subchapter").
90. 7 U.S.C.A. § 136v(b) (West 1999).
91. 7 U.S.C.A. §§ 136v(a) and 136v(b) (West 1999).
92. 736 F.2d 1529 (D.C. Cir. 1984)
failure to adequately label Paraquat to warn against the possibility that chronic inhalation and skin exposure could lead to lung disease and death. The Court of Appeals for the District of Columbia Circuit held that in FIFRA and its accompanying regulations (1) there was no express preemption of state common law actions; (2) there was no implied field preemption of such claims, as there was no evidence of Congressional intent to occupy the field; and (3) there was no conflict preemption, as compliance with both federal and state law was not mutually exclusive.

In holding that FIFRA § 136v(b) did not expressly preempt state tort recovery, the Court of Appeals stated:

While FIFRA does not allow states directly to impose additional labeling requirements, the Act clearly allows states to impose more stringent constraints on the use of EPA approved pesticides than those imposed by the EPA . . . . Given this provision, Maryland might well have the power to ban Paraquat entirely . . . . [I]f a state chooses to restrict pesticide use by requiring that the manufacturer compensate for all injuries or for some injuries resulting from the use of a pesticide, federal law stands as no barrier.93

In concluding that there was neither field nor conflict preemption, the court interpreted FIFRA as having primarily a regulatory aim—to ensure that, from a cost-benefit point of view, Paraquat, as labeled, does not produce "unreasonable adverse effects on the environment."94 The federal appeals court reasoned that state tort law had both a regulatory and a "broader, compensatory goal, [and that] conceivably, a label may be inadequate under state law if that label, while sufficient under a cost-benefit standard, nonetheless fails to warn against any significant risk."95 Although, the court continued, an award of damages for failure to warn imposed a dual obligation upon the manufacturer, Chevron could comply with both federal and state law by continuing to use the EPA approved label and by paying damages.96 The court also stated that tort recovery could promote legitimate regulatory aims by leading manufac-

93. 736 F.2d at 1541.
94. Id. at 1540; see 7 U.S.C.A. § 136(bb) (West 1999) (defining "unreasonable adverse effects").
95. 736 F.2d at 1540.
96. See id.
turers to petition EPA to allow more detailed labeling of their products, or by influencing the EPA to require revised labels in light of the new information brought to light through common law litigation.\footnote{See id. at 1541.} Several courts adopted the \textit{Ferebee} rationale and held that state common law remedies are not expressly preempted by FIFRA. For example, in \textit{Riden v. ICI Americas, Inc.},\footnote{763 F. Supp. 1500 (W.D.Mo. 1991).} the court held that FIFRA did not expressly or impliedly preempt state tort claims, that state common law remedies did not conflict with FIFRA’s purposes, and that FIFRA falls short of requiring uniform labeling.\footnote{See Roberts v. Dow Chem. Co., 702 F. Supp. 195, 199 (N.D.Ill. 1988) (adopting \textit{Ferebee} rationale and concluding that legislative history indicates FIFRA regulations not intended to be so comprehensive as to occupy field). See also Evenson v. Osmose Wood Preserving Inc., 760 F. Supp. 1345 (S.D.Ind. 1990); Stewart v. Ortho Consumer Prods., 1990 WL 36129 (E.D.La. 1990); Whitener v. Reilly Indus., Inc., No. 87-5224, slip op. at 4 (D.Ill. 1989); Cox v. Velsicol Chem. Corp., 704 F. Supp. 85, 87 (E.D.Pa. 1989).} In contrast, a Michigan federal trial court rejected the \textit{Ferebee} analysis in \textit{Fitzgerald v. Mallinckrodt, Inc.},\footnote{681 F. Supp. 404 (E.D.Mich. 1987).} and held that the common law claims for failure to warn were preempted. The \textit{Fitzgerald} court was not persuaded that a manufacturer had an authentic choice with respect to altering a pesticide’s label in response to an adverse jury award. The court stated that the \textit{Ferebee} “choice of reaction” analysis “seems akin to the free choice of coming up for air after being underwater. Once a jury has found a label inadequate under state law, and the manufacturer liable for damages for negligently employing it, it is unthinkable that any manufacturer would not immediately take steps to minimize its exposure to continued liability.”\footnote{Id. at 407 (quoting Palmer v. Liggett Group, Inc., 825 F.2d 620, 627-28 (1st Cir. 1987).}

In reaching its conclusion that FIFRA expressly preempted any state labeling or packaging requirements different from or additional to those mandated in FIFRA, the \textit{Fitzgerald} court relied heavily on \textit{Palmer v. Liggett}\footnote{825 F.2d 620 (1st Cir. 1987).} and the \textit{Palmer} court’s pur-
ported rejection of Ferebee. The Fitzgerald court held that whereas federal statute preempts any state regulation of labels, any recovery in tort is precluded. To hold otherwise "would have effectively authorized the state to do through the back door exactly what it cannot do through the front." Several courts criticized the Fitzgerald court's reliance on Palmer, but followed its conclusion and held that FIFRA preempts common law failure to warn claims.

103. Although the Palmer court noted its dissatisfaction with the Ferebee "choice of reaction" analysis as it applied to the 1969 Act, it did not question or criticize its application to FIFRA.


FIFRA, which applies to some 40,000 different herbicide and pesticide formulations, imposes upon an entirely different type of regulatory scheme from that established under the [1969] Act. Under FIFRA, each manufacturer drafts a warning label for each product for EPA approval. Thus, two manufacturers of the same regulated product may use different labels of their own choosing, provided only that they obtain prior EPA approval . . . . In contrast, the [1969] Act explicitly (i) applies to cigarettes only; (ii) mandates the precise language of the label; and (iii) prohibits any state from regulating any aspect of cigarette warnings.


See also Fisher v. Chevron Chem. Co., 716 F. Supp. 1283, 1284 (W.D.Mo. 1989), in which the federal trial court held that the claim that Paraquat was sold in an unreasonably dangerous, defective condition when put to its reasonably anticipated use was not preempted, in that there was no Congressional intent to "occupy the field" relating to pesticides and injuries arising from their use. Neither, the Fisher court held further, was plaintiff's claim that the aerial spraying of Paraquat was an inherently or abnormally dangerous activity.

In Villari v. Terminix Int'l, Inc., 692 F. Supp. 568 (E.D.Pa. 1988), the court held that FIFRA's prohibition of state labeling and packaging requirements did not preempt state law tort action against an exterminator and a manufacturer based on the exterminator's alleged failure to warn of dangers associated with termiticides used in the home. The plaintiff's injuries did not result from the de-
ii. Decisional law after Cipollone v. Liggett Group

The Court of Appeals for the Eleventh Circuit's decisions in *Papas I* and *Papas II* provide a temporal bridge between a pre-*Cipollone* FIFRA analysis and the post-*Cipollone* approach that seemingly governs today. In *Papas v. Upjohn Co. (Papas I)*, the court noted that the language in FIFRA, as well as its legislative history, could support a determination that state common law actions based upon inadequate labeling are expressly preempted, but declined to reach the issue. Instead, the court found that federal preemption of common law tort claims based upon labeling deficiencies could be implied from FIFRA and the labeling regulations promulgated pursuant thereto. The court reasoned (1) that FIFRA occupied the entire field of labeling regulation, leaving no room for the states to supplement federal law, even by means of state common law tort actions; (2) that "jury awards of damages in such tort actions would result in direct conflict with federal law"; and (3) that "allowing state common law tort suits for inadequate labeling would stand as an obstacle to the accomplishment and execution of the full objectives of Congress." The court stated further that permitting common law tort actions based on labeling claims "would permit state court juries to do what state legislatures and state administrative agencies were forbidden to do: impose requirements for labeling pesticides." In the subsequent proceeding before the Supreme Court, the Court vacated *Papas I*

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107. 926 F.2d 1019 (11th Cir. 1991), cert. granted and judgment vacated, 505 U.S. 1215 (1992), on remand, 985 F.2d 516 (11th Cir. 1993) (held: FIFRA expressly preempts state law claims based upon inadequate labeling or packaging). See also *Papas v. Upjohn Co.*, 985 F.2d 516 (11th Cir. 1993)(*Papas II*), cert. denied, *Papas v. Zoecon Corp.*, 505 U.S. 1215 (1993) (state law action for damages expressly preempted to extent dependent upon showing pesticide labeling or packaging failed to meet standards in addition to or different from those required by FIFRA).


109. *See id.* at 1024.

110. *See id.* at 1025.

111. *Id.*


113. *Id.* at 1026.
and remanded for reconsideration in light of its recent decision in Cipollone.\textsuperscript{114} With the Cipollone decision now before it, the Eleventh Circuit entered its ruling in Papas II, and held that FIFRA expressly preempted claims based upon inadequate labeling or packaging.\textsuperscript{115} A similar conclusion was reached in MacDonald v. Monsanto Co.,\textsuperscript{116} in which the Fifth Circuit Court of Appeals concluded that state common law judgments, including those predicated on a failure to warn theory, are “requirements” for the purposes of preemption.\textsuperscript{117}

In Shaw v. Dow Brands, Inc.,\textsuperscript{118} the Court of Appeals for the Seventh Circuit compared the preemption provisions in both the 1969 Act and FIFRA and found them sufficiently alike to compel the conclusion that FIFRA expressly preempted inconsistent state common law demands.\textsuperscript{119} To like effect, in King v. E.I. du Pont de Nemours & Co.,\textsuperscript{120} a Maine federal trial court considered whether FIFRA’s mandate barring states from imposing requirements on herbicide labels precludes state common law tort claims. After reviewing the legislative history of FIFRA, the regulations promulgated to govern the registration of pesticides, and guidance of Cipollone, the King court held:

\begin{quote}
[FIFRA § 136v(b)] says that states ‘shall not impose or continue in effect any requirements for labeling or packaging in addition to or different from those required under this subchapter . . . . Since the 1969 Cigarette Act, which prohibits states from imposing ‘requirements’ on cigarette advertising, was held to preempt com-
\end{quote}

\begin{footnotesize}
\begin{itemize}
\item \textsuperscript{114} 985 F.2d 516 (11th Cir. 1993).
\item \textsuperscript{115} Id. See also Clubine v. American Cyanamid Co., 534 N.W.2d 385 (Iowa 1995) (discussed and limited in Ackerman v. American Cyanamid Co., 586 N.W.2d 208 (Iowa 1998)) (held: claims associated with label-based warranties preempted by FIFRA as such claims constituted additional or different requirements than those imposed by FIFRA).
\item \textsuperscript{116} 27 F.3d 1021, 1025 (5th Cir. 1994).
\item \textsuperscript{117} Id. Accord Deshotel v. Rhone-Poulenc, Inc., 969 F. Supp. 397 (W.D. La. 1997) (held: label-based claims brought against pesticide manufacturer by farmer who experienced failure of sweet potato crop preempted). See also Ackles v. Luttrell, 561 N.W.2d 573 (Neb. 1997) (held: claims based upon either failure to warn or fraudulent labeling claims against pesticide manufacturer preempted by FIFRA).
\item \textsuperscript{118} 994 F.2d 364 (7th Cir.1993).
\item \textsuperscript{119} Id. (citing Cipollone v. Liggett Group, 505 U.S. 504 (1992)).
\item \textsuperscript{120} 806 F. Supp. 1030 (D.Me. 1992), aff’d, King v. E.I. DuPont de Nemours & Co., 996 F.2d 1346 (1st Cir. 1993), cert. dismissed, 510 U.S. 985 (1993) (FIFRA expressly preempts state tort law claims based upon alleged failure to provide adequate herbicide warning labels).
\end{itemize}
\end{footnotesize}
mon law damage actions with respect to inadequate warnings in *Cipollone*, we hold that the prohibition of 'requirements' under FIFRA preempts common law damage actions for failure to warn in the herbicide labeling context.  

Other courts having the opportunity to consider the preemptive effect of FIFRA in light of *Cipollone* have interpreted the Supreme Court’s decision more restrictively. For example, in *Burke v. Dow Chemical Co.*, the court undertook a detailed analysis of the statutory language of FIFRA to determine whether failure to warn claims would amount to a state imposed “requirement for labeling or packaging in addition to or different from those required [under FIFRA] and would therefore be expressly preempted.” The court found that the pre-emption provision in FIFRA lies “somewhere in between the [1965 Act] and [1969 Act] provisions.” The court stated that the prohibition upon the state imposing any “requirements” “for labeling or packaging” “different from” the EPA requirements, when viewed in conjunction with the general savings clause explicitly authorizing each state to regulate the sale or use of pesticides, “indicate[s] a congressional design to leave the states with expansive powers to ‘regulate’ pesticides.” Finding the doctrine of express preemption to be narrow in its scope and, moreover, that the failure to warn claims were not implicitly prohibited by FIFRA, the court wrote:

Applying the somewhat subtle distinctions of *Cipollone*, we hold that, if EPA-approved labels were in fact affixed to the relevant containers, plaintiffs may not claim that defendants’ products were mislabeled. If, however, warnings to the trade, warnings apart from labels or packaging, limitation on sales to professionals, or other protections falling generally within the ambit of warnings [which] should have been used when the content of the

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122. 505 U.S. at 504.


124. *Id.* at 1140.

125. *Id.* at 1140; *see* 7 U.S.C.A. §§ 136v(a)-(b) (West 1999).

126. 797 F. Supp. at 1140.

127. *See id.* at 1141.
label was fixed by EPA there remains a liability question for the trier of fact.\textsuperscript{128}

Similarly, in \textit{Couture v. Dow Chemical},\textsuperscript{129} the court held that "the narrow construction mandated by the preemption analysis utilized in \textit{Cipollone}\textsuperscript{130} supported an earlier interpretation of FIFRA in which the court had followed \textit{Ferebee}, and held that states are "free to regulate, through common law remedies, the use and sale of pesticides."\textsuperscript{131}

It is seen that most courts considering the matter have concluded that FIFRA expressly preempts state law claims that manufacturers should have used labels or warnings different from or in addition to those required by federal statute.\textsuperscript{132} Importantly, however, FIFRA's preemptive effects are limited to claims related to labeling or packaging. Other common law products liability or warranty claims may remain viable.\textsuperscript{133}

\textsuperscript{128} Id. at 1140.
\textsuperscript{130} Id. at 1302.
\textsuperscript{131} Id. at 1302. \textit{See also} Montana Pole & Treating Plant v. I.F. Laucks, 775 F. Supp. 1339 (D.Mont. 1991), aff'd, 993 F.2d 676 (9th Cir. 1993) (common law tort claims not preempted by FIFRA).


\textsuperscript{133} See \textit{DerGazarian}, 836 F. Supp. at 1429 (FIFRA did not preempt common law action premised upon insecticide manufacturer's alleged failure to use ordinary care in formulation, inspection and testing of insecticide); Levesque v. Miles, Inc., 816 F. Supp. 61 (D.N.H. 1993) (FIFRA did not preempt statutory cause of action for breach of warranty); Bingham v. Terminix Int'l Co., 850 F. Supp. 516 (S.D.Miss.1994) (preemptive scope of FIFRA extends only to claims relating to labeling and packaging; claims involving a failure to test or inspect product, non-labeling claims for breach of warranty, and claims for defective design or formulation not preempted).
C. Motor vehicle safety

In Geier v. American Honda Motor Company, the Supreme Court retreated from its Cipollone-grounded focus on express preemption, but in so doing the Court did not necessarily ease the complexity of federal preemption issues as they affect motor vehicle safety statutes and regulations. In Geier, the Court was asked to analyze the effect of the express preemption provision in the National Traffic and Motor Vehicle Safety Act ("NTMVSA" or "Safety Act") on a lawsuit alleging that a 1987 Honda was defective in design because it did not have a driver's side air bag. The NTMVSA, under which the Department of Transportation issued a federal motor vehicle safety standard ("FMVSS") which permitted automobile manufacturers in the late 1980s to choose among three options for passenger safety restraints, does contain a preemption provision, but also has a savings clause, described below.

Automobile design defect claims have been examined under the lens of federal law preemption for many years, and certain automobile design defect claims have been held preempted by the NTMVSA. The NTMVSA delegates to the Secretary of Transportation the authority to establish motor vehicle safety standards, and the Secretary of Transportation has in turn delegated the duty of standard promulgation to the National Highway Transportation Safety Administration ("NHTSA"). The NTMVSA contains a preemption clause that reads:

(b) Preemption.
(1) When a motor vehicle safety standard is in effect under this chapter, a State or a political subdivision of a State may prescribe or continue in effect a standard applicable to the same aspect of performance of a motor vehicle or motor vehicle equipment only if the standard is identical to the standard prescribed in this chapter.

While the Safety Act by its terms precludes state safety standards with respect to "any motor vehicle or item of motor vehi-

136. See 49 C.F.R. § 1.50 (current through September 29, 2000); See 49 C.F.R. § 501.2 (current through September 29, 2000).
cle equipment of any safety standard . . . which is not identical to the Federal standard,"\textsuperscript{138} it contains a savings clause that reads: "(e) Common law liability.—Compliance with a motor vehicle safety standard prescribed under this chapter does not exempt a person from liability at common law."\textsuperscript{139}

i. \textit{Decisional law preceding Geier v. American Honda Motor Co.}

Prior to the entry of the Supreme Court's decision in \textit{Geier}, courts in several jurisdictions had held that compliance with the criteria for one of the multiple passive restraint options set forth in the regulations impliedly preempted a tort claim targeting the manufacturer's election not to install airbags.\textsuperscript{140} For example, in \textit{Taylor v. General Motors, Corp.}\textsuperscript{141} the Court of Appeals for the Eleventh Circuit ruled that plaintiff's state law claims against the manufacturer of an automobile for failure to install an airbag was impliedly preempted by the NTMVSA insofar as the manufacturer had sold a product equipped with one of the other passive restraint options available to it. Because no Florida appellate court had been presented with such an issue, the role of the Eleventh Circuit was to predict the outcome should the matter be heard by the State's highest court.\textsuperscript{142} The federal appeals court focused on Supreme Court cases involving federal law preempting, by implication, state law claims, in which the Supreme Court had ruled that "under the principles of implied preemption, a state cannot impose common law damages on individuals for doing what a federal act or regulation 'authorized them to do.'"\textsuperscript{143} Insofar as personal injury claims brought under state law could, hypothetically but foreseeably, conflict with the passive restraint system approach permitted

\textsuperscript{140} 49 C.F.R. § 571.208 S5.1(a)-(b) (current through September 29, 2000).
\textsuperscript{141} 875 F.2d 816 (11th Cir. 1989).
\textsuperscript{142} \textit{See id.}
\textsuperscript{143} \textit{Id.} at 827 (citing Chicago & N.W. Transp. Co. v. Kalo Brick & Tile Co., 450 U.S. 311, 318 (1981)).
by the Safety Act, to permit such claims would "take away the flexibility provided by a federal regulation." 144

In Taylor, the Eleventh Circuit held that even absent express preemption, standards published by the NHSTA preempted, by implication, a common law claim for defect in design. The court held that the NHSTA as amended and regulations thereunder, authorizing the manufacturer a choice of three different methods for occupant crash protection, one of which being manual seat belts, impliedly preempted a common law claim of design defect for the manufacturer's conscious design choice not to equip an automobile with airbags. 145

A Wisconsin court of appeals also held that a state law claim for failure to install airbags in a 1980 Plymouth Horizon was preempted in circumstances of the defendant's compliance with the NTMVSA standard. 146 Observing that the phrase

144. Id. (citing Fidelity Federal Savings & Loan Ass'n v. de la Cuesta, 458 U.S. 141 (1982)). See also Wood v. General Motors Corp., 865 F.2d 395, 412-414 (1st Cir. 1988); Montag v. Honda Motor Co., 75 F.3d 1414, 1417 (10th Cir. 1996).

145. 875 F.2d at 822 n.13 ("We conclude that a state common law rule that would, in effect, remove the element of choice authorized in Safety Standard 208 would frustrate the federal regulatory scheme."). See also Myrick v. Freuhauf Corp. (Myrick I), 795 F. Supp. 1139 (N.D. Ga. 1992) (held: state statutory or common law claims would impose stricter safety standards than established by federal regulations and were, therefore, preempted).

Accord Loulos v. Dick Smith Ford, Inc., 882 SW.2d 149 (Mo. Ct. App. 1994). In Loulos, a motorist's claim against Ford, among others, for failure to equip a 1979 vehicle with airbags, the court held that plaintiff's claim was preempted by the NTMVSA. The court noted specifically that regulations promulgated pursuant to the NTMVSA authorized manufacturers to adopt one of four permissible passive restraint system options, and that to permit a state tort law cause of action would, in effect, permit punishment of manufacturers for making a choice Congress specifically countenanced.

See also Gills v. Ford Motor Co., 829 F. Supp. 894 (W.D.Ky. 1993) (FMVSS 208, adopted pursuant to NTMVSA, preempted common law claim seeking damages for failure to install airbags); Pokorny v. Ford Motor Co., 902 F.2d 1116 (3d Cir. 1990). In Pokorny the Third Circuit held that state common law claims for failure to provide either air bags or automatic seatbelts were impliedly preempted by federal regulations authorizing the manufacturers their choice of safety mechanisms to be installed into the motor vehicle. However, the court held that common law claims for failure to install additional passive restraint systems not included among the options listed in Standard 208 were not preempted. See id. at 1126.

146. See Boyle v. Chrysler Corp., 501 N.W.2d 865 (Wis. 1993); see Panarites v. Williams, 629 N.Y.S.2d 359 (N.Y. App. Div. 1995) (state law claims arising from a manufacturer's election of a passive restraint system other than airbags both expressly and impliedly preempted insofar as judgment against manufacturer would impose safety requirements different from and in addition to those required by
"state law'... include[s] common law as well as statutes and regulations."

The state appellate court held that because the defendant complied with the federal act, state law claims, including common law claims, were preempted. To hold otherwise, the Wisconsin court noted, would have required judicial entertainment of state law claims that would have created a conflict with the federal standards. Lastly, the court applied a conservative interpretation to the Safety Act’s saving clause, and held that “[t]he savings clause does not operate to preserve common-law liability claims that conflict with the federal safety standards. Rather, the savings clause preserves only those common-law liability claims that do not conflict with the automobile safety equipment standards that Congress enacted.”

A harmonious conclusion was reached in the appeal of a suit bringing state law claims against a manufacturer for failure to install lap seat belts. A Minnesota state appeals court found the state law claims were impliedly preempted by the NTMVSA, reasoning that maintenance of potential state law liability, in effect, would have conflicted with the manufacturer’s prerogative under federal regulation to pick one from among a group of federally-approved passive restraint systems.

A limited preemption province was defined by the Seventh Circuit in Gracia v. Volvo Europa Truck, N.V., a claim against a truck manufacturer alleging that its windshield retention system was inadequate to prevent windshield ejection during collision. The federal appeals court stated the proposition broadly that the savings provision of the Safety Act “was not to preserve common law claims when they conflict with NHTSA standards, but to prevent a manufacturer from having

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federal government); Dykema v. Volkswagenwerk AG, 525 N.W.2d 754 (Wis. Ct. App. 1994) (state law claims preempted because they would have subjected manufacturers to different safety standards than established by the NTMVSA).

147. Id. (quoting Cipollone, 505 U.S. at 521).


150. See id.

151. 112 F.3d 291 (7th Cir. 1997).
a complete defense to a common law action not addressed by a
NHTSA standard by merely stating that it is in full compliance
with all federal safety standards."152 Observing that the Safety
Act's savings provision153 should not be interpreted as
"preserv[ing] conflicting or non-identical state common law ac-
tions from preemption,"154 the court held that common law
claims were not to be preempted by the NHTSA if, and only if,
they were identical to the federal standards.155

Regarding the windshield retardation standard specifically,
the court noted that in this instance, the agency had specifically
concluded that "given the design of forward control vehicles it
was both technically impracticable to design windshields which
would comply with the standards and impracticable to apply
the standard's barrier crash tests to these vehicles."156 Given
the NHTSA particularized conclusion that the standard would
not apply, the court held that "a state common law standard on
windshield retention would, accordingly, be a standard that is
not identical to the federal one[,]" and was thus preempted.157

Thus, even without backdrop of a specific standard, the
NHTSA dormant authority to create a safety standard may pre-
empt a personal injury claim brought under state law. Be the
question one of passive restraint systems or another vehicle
component within the NHTSA's NTMVSA purview, the
agency's decision to forego application of a standard to a partic-
ular type of product may still be given preemptive effect. For
example, regarding the safety standard for windshield retention
systems, and the exclusion from that standard of forward con-
trol vehicles with a gross vehicle weight exceeding 10,000
pounds, at issue in Garcia, the Ninth Circuit Court of Appeals
explained:

[H]ere there is a specific federal standard addressing windshield
retention for the truck at issue, in which the NHTSA determined
that this type of vehicle should be exempt from the affixing re-

152. Id. at 298.
154. 112 F.3d at 298.
155. See id.
156. Id. at 297.
157. Id. at 296.
to forego regulation in a given area may imply an authoritative federal determination that the area is best left unregulated, and in that event would have as much preemptive force as a decision to regulate."

In *Loulos v. Dick Smith Ford, Inc.*, a motorist's claim against Ford, among others, for failure to equip a 1979 vehicle with airbags, the Missouri appellate court held that plaintiff's claim was not preempted, and refused to consider an implied preemption theory, reasoning, according to *Cipollone*, an express preemption provision was "a reliable indicium of congressional intent," terminating implied preemption analysis.

Regarding Safety Act regulations for illumination, the Court of Appeals for the Third Circuit in *Buzzard v. RoadRunner Trucking, Inc.* held that the plaintiff's state common law tort claims of inadequate lighting were not preempted. The court, in its decision, noted the defendant's burden to overcome the generally applicable rebuttable presumption against a finding of Congressional intent to preempt state law or regulation. Emphasizing Congress’s stated goal of increasing transportation safety and the potential value of permitting states to enact more stringent regulations than are federally imposed to achieve that goal, the Third Circuit held that plaintiff's claims were not preempted, and wrote:

*Buzzard's action could encourage increased safety by enhancing motorists' ability to take in at a glance the size, location and movement of tractor-trailers encountered at night on the public highways. Encouraging manufacturers to act in a way that increases safety does not frustrate the primary purpose of the Safety Act. Nor does it make it impossible to comply with both federal and state law, as it does not suggest that illumination equipment mandated by state common law be used instead of that required by federal law, but only in addition to that specified in Standard 108.*

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158. *Id.* at 296 (emphasis added by appellate court).
160. *Id.* at 152.
162. *See id.* at 780.
163. *Id.* at 785.
A federal trial court in *Byrnes v. Honda Motor Co., Ltd.*,\(^{164}\) held that a products liability claim against the manufacturer of the 1990 Honda motorcycle's lighting system was not impliedly preempted by the NTMVSA, and explained that in the absence of any reference to the subject matter in the Safety Act, much less in its preemption provision, no further evaluation of implied preemption was necessary. Also finding no implied preemption is the decision of the Court of Appeals for the Eleventh Circuit in *Doyle v. Volkswagenwerk Aktiengeellschaft*,\(^{165}\) in which the plaintiffs challenged the restraint design of a Volkswagen Jetta. The model at issue had an automatic shoulder belt, but no lap belt for either the driver or front seat passenger. Rather, the design employed knee bolsters to restrain the driver or passenger from sliding beneath the belt upon collision. The Eleventh Circuit affirmed the trial court's determination that standards promulgated under the NTMVSA do not preempt common-law claims, citing the Supreme Court's decision in *Freightliner Corp. v. Myrick*,\(^{166}\) which noted specifically that there existed no federal safety standard creating a conflict of compliance.

In *Collazo-Santiago v. Toyota Motor Corp.*,\(^{167}\) a suit brought by a motorist claiming that deployment of her vehicle's airbag caused her facial injuries, a federal trial court found that the federal standards were not design standards at all, but rather constituted performance standards. As such, the court reasoned:

Manufacturers are apparently free to choose any airbag design, as long as the design meets the performance criteria[.] Thus, there is no conflict between the provisions of [federal law] and its implementing regulations, on the one hand, and tort liability for defective design, on the other, because such liability "[does] not remove or require any particular choice, or otherwise frustrate 'flexibility' that the federal scheme provides."\(^{168}\)

\(^{164}\) 845 F. Supp. 875 (S.D. Fla. 1994).

\(^{165}\) 114 F.3d 1134 (11th Cir. 1997).


\(^{167}\) 957 F. Supp. 349 (D.P.R. 1997).

\(^{168}\) Id. at 354 (quoting Perry v. Mercedes Benz, 957 F.2d 1257, 1264-65 (5th Cir. 1992); 49 U.S.C.A. § 30103(e) (West 1996)).
ii. Geier v. American Honda Motor Company

In the years following the Supreme Court's 1992 *Cipollone*\(^{169}\) decision, some courts and commentators have referred to the Court's preemption analysis as "schizophrenic"\(^{170}\) or "shaky."\(^{171}\) In *Geier v. American Honda Motor Company*,\(^{172}\) the Court seemed to retreat from its *Cipollone* focus on express preemption, but in so doing did not necessarily simplify matters for litigants and trial judges seeking the proper doctrinal analysis by which to answer preemption questions.

In *Geier*, the review of a lawsuit alleging that a 1987 Honda was defective in design because it did not have a driver's side air bag, the Court was asked to identify the specific species and phylum of the NTMVSA preemption provision. As noted earlier, the NTMVSA authorizes the NHTSA to promulgate and implement FMVS standards. The NHTSA did so, implementing a standard that permitted automobile manufacturers during the applicable time period to choose among three options for passenger safety restraints, while stating further that compliance with federal safety standards would not "exempt any person from any liability under common law."\(^{173}\) In *Grier*, the Court held that this statutory provision does not expressly pre-empt state common law damages actions, but, rather, that "ordinary preemption principles" do.\(^{174}\)

The express preemption clause of the NTMVSA provides that states may not maintain "motor vehicle safety standards" which conflict with federal performance standards on the same topic. In *Freightliner Corp. v Myrick*,\(^{175}\) which involved the effect of the absence, at that time, of a federal standard pertaining to anti-lock brakes, the Court, in a unanimous opinion written by Justice Thomas, concluded that since there was no federal standard in issue on the topic for eighteen-wheel trucks,

\(^{171}\) Wilson v. Bradlees of New England, Inc., 96 F.3d 552, 556 (1st Cir. 1996) (the Supreme Court's preemption analysis makes its application "shaky" in "a changing legal climate.").
\(^{172}\) ___ U.S. ___, 120 S. Ct. 1913 (2000).
\(^{173}\) *Id.* at 1915
\(^{174}\) *Id.* at 1913.
there was no express or implied preemption of state design defect claims based on the absence of such brakes. The Court did not reach the question of whether the Safety Act would preempt such claims if a federal standard did exist, but in the course of its opinion, the Court raised a question about the scope of Cipollone's express preemption analysis:

The fact that an express definition of the pre-emptive reach of a statute 'implies'—i.e., supports a reasonable inference—that Congress did not intend to pre-empt other matters does not mean that the express clause entirely forecloses any possibility of implied pre-emption. . . . At best, Cipollone supports an inference that an express preemption clause forecloses implied preemption; it does not establish a rule.176

The unanimous decision in Myrick, it is seen, foreshadowed a potential reinvigoration of the doctrine of implied preemption. Some commentators noted that the Court’s preemption analysis after Cipollone, Myrick, and its 1996 decision in Medtronic,177 was less and less a true express preemption analysis and more and more a veiled implied preemption analysis.178 Geier179 proved this observation to be true. Justice Breyer, writing for the majority in Geier, articulated a three-part preemption analysis: Does the express preemption provision explicitly preempt the lawsuit? If not, “do ordinary preemption principles nonetheless apply?”180 If so, does the lawsuit “actually conflict” with the federal statute? Of primary importance was how the Court answered the second question because if “ordinary,” also known as “implied,” preemption principles applied in the face of an express provision to the contrary, then the continued influence of Cipollone and Medtronic beyond their precise subject matter precincts—tobacco labeling and medical devices respectively—would arguably be ripe for recalibration.

176. Id. at 289.
179. ___ U.S. ___, 120 S. Ct. 1913 (2000). Geier was a five to four opinion. Justice Breyer, writing for the majority, was joined by Chief Justice Rehnquist, Justices O’Connor, Scalia and Kennedy. Justice Stevens, the author of both the Cipollone and Medtronic plurality opinions, dissented in an opinion in which Justices Souter, Thomas and Ginsburg joined.
180. Id. at 1918.
Neither in its analysis of the NTMVSA preemption provision nor the Safety Act’s preemption clause did the majority expend the time necessary to provide lower courts with such guidance as has been characteristic of the Court’s prior preemption decisions. The Grier Court declined to focus upon the meaning of the language of the NTMVSA preemption provision to determine its scope, including, without limitations, the question of what is meant by “standard,” as distinct from the term “requirement” emphasized in the Cipollone line of decisions. Instead, the Court concluded, with little fanfare, that the “savings clause” made that exercise unnecessary, and explained its reticence to interpret the term “standard” as based upon its conclusion that it should be read to include common law damages actions because the savings clause assumes “that there are some significant number of common-law liability cases to save.”

In essence, the Court found that the presence of the savings clause triggered a narrow reading of the express preemption provision, and that further, in order to give meaning to the savings clause, it operated to exclude common law damages actions from the preemption clause’s reach.

D. Foods, Drugs and Cosmetics

Manufacturers of pharmaceuticals are subject to the labeling and formulation standards articulated by the Food, Drug and Cosmetic Act (“FDCA”). Also, pharmaceutical sellers and manufacturers may be held strictly liable for injuries that result from their failure to warn of product dangers that are known or knowable within the scientific field.

In the realm of pharmaceuticals, the FDCA contains no language expressly preempting claims brought under state law. Moreover, courts considering the issue have held generally that with regard to health and safety issues, the statutory language

181. Id.
182. See id. stating, “We have found no convincing indication that Congress wanted to preempt not only state statutes and regulations, but also common-law tort actions.” Id. at 1918.
does not evince a congressional objective to so occupy the field of pharmaceutical regulation as to impliedly preempt state claims.186 Support for this conclusion can be found in MacDonald v. Ortho Pharmaceutical Corp.,187 in which the court rejected the manufacturer's preemption defense.188 The MacDonald court held that "[t]he regulatory history of the FDA requirements belies any objective to cloak them with preemptive effect."189

The Supreme Court has held that where Congress has not expressly preempted state tort law claims, there exists a strong presumption against implied preemption unless there are exceptional circumstances involved.190 The District Court in Mazur v. Merck & Co., Inc.191 inferred from the absence of express preemption that Congress, via the FDCA, had not intended to preempt state regulations imposed upon a vaccine’s manufacture, distribution and labeling.192

As stated by the District of Columbia Court of Appeals in Merrell Dow Pharmaceuticals v. Oxendine,193 a Bendectin suit, "FDA prescription drug regulations and safety determinations are intended to be minimum standards which 'do not conflict with state law which sets higher standards for due care and safety in the manufacture of drugs.'"194 One federal trial court described the underlying rationale for finding no necessary conflict between state tort remedies and FDA regulation in terms of the differing objectives of regulation and actions in tort:

[F]ederal regulation serves a very different purpose than state tort law. Essentially, federal regulation serves a deterrent pur-


188. See Ausness, supra note 186, at 218-25.

189. Id. at 219 (quoting MacDonald, 475 N.E.2d at 70).


192. See id. at 248.


194. Id. at 828 (citations omitted).
pose by limiting the manufacture of inherently dangerous products to those applicants who meet stringent safety standards, while state tort law serves the equally important purpose of compensating individuals injured by those very same products. Since compliance with FDA regulations will not insure that a manufacturer's products will not cause injury, compliance will not necessarily exempt a manufacturer from liability. When those products do cause injuries, the state tort system provides a means of compensation. State tort law is intended to supplement federal regulation by providing a vehicle for compensation of vaccine-related injuries.\textsuperscript{195}

Many individuals have brought actions against diphtheria, pertussis, and tetanus (DPT) vaccine manufacturers.\textsuperscript{196} The absence of an express congressional declaration as to whether federal regulation preempts state law concerning the labeling and design of the DPT vaccination has led to a split in the decisional law.\textsuperscript{197} Many courts have held that FDA approval of the DPT vaccine does not preempt more stringent state standards. These courts reason that the product's design may not necessarily be the safest, technologically achievable design, because the FDA is only able to approve those designs that are submitted by the manufacturers. Therefore, the reasoning continues, states are able to impose higher standards upon these manufacturers, in order to promote product and public safety.\textsuperscript{198}

While the FDCA contains no provision that would expressly preempt state food and drug requirements in general, there are several areas where the FDA has interposed product specific or

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\item \textsuperscript{196} See Ausness, supra note 186, at 219.

\item \textsuperscript{197} Compare Wack v. Lederle Labs., 666 F. Supp. 123 (N.D. Ohio 1987) (disallowing defendant's claim of implied "field" preemption based on FDA's extensive regulation of the subject matter); Patten v. Lederle Labs., 676 F. Supp. 233 (D. Utah 1987); MacGillivray v. Lederle Labs., 667 F. Supp. 743 (D.N.M. 1987) (state tort remedies allowed) with Hurley v. Lederle Laboratories (Hurley I), 651 F. Supp. 993 (E.D. Tex. 1986) (DPT design actions preempted by the comprehensive nature of FDCA's testing requirements); Morris v. Parke, Davis & Co., 667 F. Supp. 1332 (C.D. Cal. 1987). But see Hurley v. Lederle Labs. (Hurley II), 863 F.2d 1173 (5th Cir. 1988) (although plaintiff's failure to warn claims preempted by the FDCA, other state tort remedies permitted as Congressional intent to protect industry or product not sufficient to strip plaintiffs of claims against any manufacturer that has achieved regulatory compliance).

\item \textsuperscript{198} See id.

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subject matter specific rules which are not identical to state standards, or which would be vulnerable to a conflict with a later promulgated state standard. In some of these settings, the agency's regulatory initiatives have been held to impliedly pre-empt inconsistent state claims or regulations. The Supreme Court has explained that even conceding that most statutes and accompanying regulations administered by the FDA do not expressly preempt state statutory or decisional law, such FDA statutes may be found to impliedly preempt state law where application of state law would frustrate effectuation of "the full purpose and objective" of the FDA's broad ranging safety-related mandates. As is characteristic of agencies charged with implementing federal health and safety statutes, the FDA has interpreted both its governing statute and its own regulations as carrying a broad preemptive mantle. For example, in 1982, the FDA issued a Final Rule on Tamper-Resistant Packaging Requirements for Certain Over-the-Counter Human Drug and Cosmetic Products due to a tragic incidence of product tampering of over the counter (OTC) drugs. According to the final FDC rule, all OTC drug products that are subject to retail sale must be packaged in specific tamper-resistant packages (TRPs). The FDA stated that as the manufacturing and distribution of these products is national in scope, only national regulations are adequate to safeguard the interest of "the entire population." Its rule provided further that all local or state

199. See Geiger and Rosen, Rationalizing Product Liability for Prescription Drugs: Implied Preemption, Federal Common Law, and Other Paths to Uniform Pharmaceutical Safety Standards, 45 DePaul L. Rev. 395, 421 (1996) ("[S]tate tort law should be preempted because it may well discourage the development or marketing of beneficial drugs. The possibility of deterring development is contrary to the FDC Act's fundamental goal of making such drugs available.") (citations omitted).

200. Medtronic, 518 U.S. at 496.


203. See 21 C.F.R. § 211.132(a). There were certain exceptions to this rule, including dermatological, dentifrice, and insulin products that did not have to be packaged in compliance with the ruling. See 46 Food Drug Cosm. L.J. 629, 639.

204. 47 Fed.Reg. 50442-01, 50448 (1982), discussed in Mark B. Gelbert, State Statutes Affecting the Labeling of OTC Drugs: Constitutionality Based on Com-
packaging requirements that were not identical to the new federal regulation were specifically preempted by the new regulation.205

The FDA supplied three justifications for preemption in this area. First, if localities were permitted to have specific local requirements, the supply of specific OTC drugs might be diminished if those drugs did not comply with the local packaging requirements. Second, the federal requirements could be essentially negated if a large state could force a drug manufacturer to use that state's standards throughout the country. Third, the product cost for each drug would substantially increase if the drug manufacturers were forced to adopt the different packaging requirements for different states.206

The FDA maintains that its requirement of a pregnancy warning on a broad range of OTC drug products has been held to impliedly preempt all associated state regulations.207 In 1982, the FDA issued the rule that required “[a]ll over-the-counter drugs that are intended for systematic absorption, unless specifically exempted” to contain the warning: “As with any drug, if you are pregnant or nursing a baby, seek the advice of a health professional before using this product.”208 While some states, such as California, had adopted similar legislation, the FDA felt that “[p]roliferation of such state requirements may weaken FDA’s efforts to develop comprehensive national labeling and other requirements for OTC drugs . . . .”209 If each state had different warnings, the FDA suggested, such state by state warning requirements would “prevent the full purpose and objectives of the agency in issuing the regulation and that under the doctrine of implied preemption, these State requirements are preempted by the regulation as a matter of law.”210


206. See Gelbert, supra note 204, at 640.
207. 21 C.F.R. §§ 201.63(a), (330.2) (current through September 29th, 2000), cited in Gelbert, supra note 204, at 640.
In *Jones v. Rath Packing Co.*[^211] the Supreme Court held that inconsistent state food labeling regulations were impliedly preempted by the Fair Packaging and Labeling Act ("FPLA") and regulations thereunder. At issue in *Rath Packing* was a California statute that regulated the labeling of the net weight of food commodities.[^212] The record below was sufficient to show that the average net weight for a proportion of this seller's bacon and flour was less than the net weight stated on the packages, resulting in subsequent removal of those commodities from the shelves insofar as they were in violation of the California statute.[^213] The food packing company claimed that the state statute was preempted by federal laws which regulated labeling and the net weight requirements,[^214] drafted by Congress with the intent of informing consumers of accurate information about a package's quantity and contents.[^215] The Court found that the FDCA did not preempt the state requirements because the FDCA did not "contain a preemption clause with regard to its food misbranding requirements[.]"[^216] The Court explained that the issue of express preemption arises when the "state requirements are less stringent than or require information different from federal law,"[^217] and further found that the California law was not expressly preempted by any provision of the FPLA as the California law was more stringent than the federal requirements and did not require any information that was different from the federal law.[^218] Even absent express preemption, however, the Court concluded that the federal FPLA regimen impliedly preempted the state law inasmuch as effectuation of the state law would "prevent the accomplishment and execution of the full purposes and objective of Congress in passing the FPLA."[^219]

[^212]: See id. at 522.
[^213]: See id.
[^214]: See id. at 523-24.
[^216]: 430 U.S. at 538.
[^217]: Id.
[^218]: See id. at 540.
[^219]: Id. at 543.
E. Medical Devices

i. Medical Device Act Amendments of 1976 generally

The Food and Drug Administration is empowered by the 1976 Medical Device Amendments ("MDA") to classify and regulate medical devices. Medical devices have been divided into three categories: Class I devices are subject to "general controls" insofar as they represent only a low level of risk to public health and safety.

Class II devices are governed by an order of federal regulation known as "special controls," a higher level of superintendence because "general controls by themselves are insufficient to provide reasonable assurances of the safety and effectiveness of [such] devices." Lastly, Class III devices are subject to the most rigorous MDA controls as they represent "a potential unreasonable risk of illness or injury."

As to devices denominated as Class III, the MDA provides: "Before a new Class III device may be introduced to the market, the manufacturer must provide the MDA with 'reasonable assurance' that the device is both safe and effective." The process of providing the FDA with "reasonable assurance" is known as "premarket approval" or "PMA." The process by which a new class III device may gain premarket approval was described by the Supreme Court in Medtronic, Inc. v. Lohr as involving submission of detailed safety and efficacy information about the device followed by equally meticulous FDA review. Upon receipt of PMA, marketing of the device may begin. Sub-

221. See 21 U.S.C. § 360k(a) (West 1999).
225. Id.
226. 21 U.S.C. § 360e(d)(2)(West 1999). See Steele v. Collagen Corp., 63 Cal. Rptr. 2d 879 (Cal. Ct. App. 1997), stating, "With respect to medical devices that represent the highest risk to human life, the federal government imposes standards specific to each of those devices, and Congress has declared that the federal standard is preemptive." Id. at 880.
229. See id. at 477.
sequent changes in the product trigger a requirement that the manufacturer submit a PMA supplemental application.\footnote{230} Moreover, the regulations require annual post-approval reports detailing changes in the device, clinical investigational results, or pertinent scientific literature.\footnote{231}

Two exceptions exist as to the general requirement that Class III medical devices obtain premarket approval. First, the MDA includes a “grandfathering” provision that permits devices that were on the market prior to the 1976 passage of the MDA to remain on the market until the FDA undertakes and completes the required PMA.\footnote{232} Second, as the Supreme Court would later summarize, and “to prevent manufacturers of grandfathered devices from monopolizing the market while new devices clear the MDA hurdle, and to insure that improvements to existing devices can be rapidly introduced into the market,” the MDA “also permits devices that are ‘substantially equivalent’ to preexisting devices to avoid the PMA process.”\footnote{233}

ii. \textit{Medical Device Act Amendments of 1976 preemption provision}

The MDA’s preemption provision states:

\textbf{[N]o State or political subdivision of a State may establish or continue in effect with respect to a device intended for human use any requirement (1) which is different from or in addition to any requirement applicable under this chapter to the device, and (2) which relates to the safety or effectiveness of the device or to any other matter included in a requirement applicable to the device ...} \footnote{234}

The FDA has interpreted this preemption provision in these terms:

State or local requirements are preempted only when the [FDA] has established specific counterpart regulations or there are other specific requirements applicable to a particular device under the Act, thereby making any existing divergent state or local require-

\begin{thebibliography}{9}
\footnote{230. \textit{See} 21 C.F.R. § 814.39 (current through September 29th, 2000).}
\footnote{231. \textit{See} 21 C.F.R. § 814.84 (current through September 29th, 2000).}
\footnote{234. 21 U.S.C. § 360k (West 1999).}
\end{thebibliography}
ments applicable to the device different from or in addition to, the specific [FDA] requirements.235

iii. *Medtronic, Inc. v. Lohr and the decisional law thereafter*

*Medtronic, Inc. v. Lohr*236 was a products liability claim against the manufacturer of a failed pacemaker. On the issue of potential preemption, the manufacturer argued that plaintiff's claims were preempted by the FDA's "general 'good manufacturing practices' regulations, which establish general requirements for most steps in every device's manufacture, and by the FDA labeling regulations, which require devices to bear various warnings."237 The Court was unanimous in finding that these general FDA strictures did not constitute specific FDA requirements applicable to a particular device.238 Rather, the court concluded that as to MDA § 510(k) devices, state products liability claims would not be preempted, as the § 501(k) process "does not constitute FDA approval of the safety or effectiveness of the device, but was merely the preservation of the pre-1976 status quo, which included potential liability under state law."239

The *Medtronic* Court adopted a two-pronged inquiry to determine if a state regulation was preempted by regulations or policies issued by the FDA. First, the federal requirement had to be "applicable to the device" in question,240 i.e., the federal requirement would have preemptive effect only if it was "specific" to a "particular device."241 Second, the state requirement had to be "different from, or in addition to" the federal requirement,242 and thus "[s]tate regulations of 'general applicability'
are not preempted except where they have ‘the effect of establishing a substantive requirement of a specific device.’”

Speaking finally of the relationship between the PMA process and the limited MDA 510(k) procedure, the Supreme Court stated:

Thus, even though the FDA may well examine § 510(k) applications for Class III devices (as it examines the entire medical device industry) with a concern for the safety and effectiveness of the device, . . . it did not ‘require’ Medtronic’s pacemaker to take any particular form for any particular reason; the agency simply allowed the pacemaker, as a device substantially equivalent to one that existed before 1976, to be marketed without running the gauntlet of the PMA process. . . . There is no suggestion in either the statutory scheme or the legislative history that the § 510(k) exemption process was intended to do anything other than maintain the status quo with respect to the marketing of existing medical devices and their substantial equivalents. That status quo included the possibility that the manufacturer of the device would have to defend itself against state-law claims of negligent design.

In Medtronic, the Court appeared to build on its analysis in Cipollone, focusing on express and not implied preemption analysis. It concluded that the FDA regulations did not expressly preempt damages actions based on the design of the pacemaker in question because the language of the express preemption provision, which preempted state “require[s] . . . different from or in addition to” any federal requirement related to safety or effectiveness, was not intended to include common law damages actions based on design defects in instances where there was no device-specific federal requirement with which such a claim actually conflicted.

Of the various Medtronic opinions, that of Justice Stevens garnered the plurality. Redolent of the express preemption analysis articulated in Cipollone, the plurality wrote:

243. Id. (quoting 21 C.F.R. § 808.1(d)(1) (current through September 29th, 2000)). See also discussion in Oja v. Howmedica, Inc., 111 F.3d 782, 787-788 (10th Cir. 1997).

244. Id. at 493. See also Reeves v. Acromed Corp., 103 F.3d 442 (5th Cir. 1997). See generally Robert J. Katerberg, Patching the "Crazy Quilt" of Cipollone: A Divided Court Rethinks Federal Preemption of Products Liability in Medtronic, Inc. v. Lohr, 75 N.C. L. Rev. 1440 (1997).
We have long presumed that Congress does not cavalierly preempt state law causes of action. . . . We "start with the assumption that the historic police powers of the States were not to be superseded by the Federal Act unless that was the clear and manifest purpose of Congress." . . . We used a "presumption against the preemption of state police power regulations" to support a narrow interpretation of such an express command in Cipollone. That approach is consistent with both federalism concerns and the historic primacy of state regulation of matters of health and safety.245

A majority of the justices would have included common law damages actions within the scope of the preemption provision but, again, differed on whether that particular preemption provision was to be interpreted narrowly or broadly. The plurality found no preemption by interpreting the scope of the statute and regulations narrowly, using the legislative history and the FDA's own interpretation as support. None of the justices applied an implied preemption analysis.246 The concurring opinion of Justice Breyer confirmed the importance of Congressional intent in determining the statute's preemptive scope and complained of the "highly ambiguous" nature of the preemption provision in issue, requiring that courts look elsewhere for help as to "just which federal requirements preempt just which state requirements, as well as just how they might do so."247 Justice Breyer's frustration over Congress's inability to clearly identify and plainly state the purpose and the scope of this and other preemption provisions, together with his explicit dissatisfaction with the task of interpreting ambiguous language, foreshadowed the Court's return, in Grier v. American Honda Motor Company, to a focus on the implied preemption doctrine.248


246. Id. But see Justice Stevens's plurality opinion, which ended with the following cryptic notation: "Until such a case [announcing a device specific requirement which might require preemption] arises, we see no need to determine whether the statute explicitly preempts such a claim. Even then, the issue may not need to be resolved if the claim would also be preempted under conflict preemption analysis, see Freightliner Corp. v. Myrick, 514 U.S. 280 (1995)." Id. at 503 (italics omitted).

247. Id. at 505.

In *Martin v. American Medical Systems, Inc.* the Court of Appeals for the Fourth Circuit held that the MDA did not preempt common law tort and implied warranty claims brought, under Virginia law, by a penile implant recipient alleging that he suffered an injurious infection due to a lack of the device's sterility. Explaining its decision, the federal appeals court stated: "Because the 1976 amendments so abruptly changed the status quo, Congress was compelled to take the existing market into account. Any device on the market at the time was permitted to stay on the market until and unless the FDA, after conducting a review like that for new devices, ordered otherwise." Simple identification or classification regulations have been held not to preempt state regulation. As such, federal regulations do not "relate to the safety or effectiveness of the device."

Another departure from the PMA process is for investigational devices. The MDA exempts investigational devices from the PMA process "to encourage, to the extent consistent with the protection of public health and safety and with ethical stan-

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249. 116 F.3d 102 (4th Cir. 1997).
250. Id. at 105.
251. 116 F.3d at 103 (citing 21 U.S.C. § 360e (b) (1) (A) (West 1999)); see also Sylvester v. Mentor Corp., 663 So.2d 176 (3d Cir. 1995), in which the state appellate court followed *Medtronic* and held that the MDA § 360k did not apply to those medical devices which had been allowed to enter the stream of commerce subjected to only the premarket notification process by being "substantially equivalent" to a product already on the market. Id. at 178. See also Dutton v. AcroMed Corp., 691 N.E.2d 738 (Ohio Ct. App. 1997), which held that devices allowed to enter the market via the "substantial equivalence" standard were not subject to the blanket preemption under the MDA. Id. at 740. The appellate court also held that Plaintiff's claim of fraudulent misrepresentation was not preempted because the MDA preempts claims raised as to the device's safety and effectiveness, not fraudulent misrepresentation. Id. at 742. But see English v. Mentor Corp., 67 F.3d 477 (3d Cir. 1995), holding that the fact that the inflatable penile implants had been introduced to the market via the "substantial equivalence" test preempted state law claims against the manufacturers. Id. at 482. The court reasoned that the MDA preempted the state law claims because the "substantial equivalence" test or standard was related to the product's safety and effectiveness, barring the implementation of state standards of the same. The Supreme Court vacated the judgment and remanded for reconsideration in light of *Medtronic*. Id. at 482-83.
252. Kealoha v. E.I. Du Pont De Nemours & Co., 82 F.3d 894, 898-99 (9th Cir. 1996) (action brought by recipient of temporomandibular jaw (TMJ) implants against supplier of raw polytetrafluoroethylene (PTFE) used in the manufacture of the implants). *Accord* Anguliano v. DuPont, 44 F.3d 806, 809-10 (9th Cir. 1995); LaMontagne v. DuPont, 41 F.3d 846, 853 (2d Cir. 1994).
dards, the discovery and development of useful devices intended for human use.” Unlike the “substantial equivalent” § 510(k) process, however, which can be completed in an average of only 20 hours, investigational device status only is granted after a comprehensive and particularized procedure in which the applicant must set forth a report of all prior investigations, as well as a “description of the methods, facilities, and controls used for the manufacture, processing, packing storage, and, where appropriate, installation of the device, in sufficient detail so that a person generally familiar with good manufacturing practices can make a knowledgeable judgment about the quality control used in the manufacture of the device.” It is in the context of these exacting standards and others that FDA approval of investigational devices has been held to preempt state law products liability claims. Thus, in Martin v. Telectronics Pacing Systems, Inc., a case involving an implantable cardioverter-defibrillator-demand pacemaker, one of only 50 such devices, the Sixth Circuit Court of Appeals held that “where the FDA has specifically approved the design of the device for investigational purposes,” to permit a state law design defect claim “would thwart [the federal] goals of safety and innovation[,]” and, accordingly, investigational device approval would be held to preempt the common law claim.

In other settings, such as claims relating to super absorbant tampons, there have been numerous cases concerning tampons and toxic shock syndrome which have held that general FDA requirements of warning statements do, in actuality, constitute particularized FDA regulations that are specific to a particular device. As tampons have been classified as Class III medical devices, the FDA requires that certain information and warnings be affixed to the product and that the average

254. Medtronic, 518 U.S. at 479.
255. 21 C.F.R. § 812.20(b)(3) (current through September 29th, 2000).
256. See 21 C.F.R. § 812.25 (current through September 29th, 2000) (descriptions of methodology, protocols, controls, written procedures, etc.).
257. 105 F.3d 1090 (6th Cir. 1997).
258. Id. at 1099.
individual be able to understand that information. For example, the district court in *Krause v. Kimberly-Clark Corp.* held that the state law claims as to the sufficiency of warnings or labels were preempted, but that claims of negligence or claims of breach of an implied warranty were not. In *Papike v. Tambrands, Inc.* the Ninth Circuit Court of Appeals reached the same conclusion, holding that "[t]he tampon labeling regulation is device—and disease—specific and preemption is warranted in this case." In making its decision, the court noted the highly individualized tampon labeling requirements set forth in FDA regulations.

Upholding the principle that FDA regulations preempt state regulations concerning inadequate warning claims with regard to tampons, there is authority to the effect that "FDA labeling requirements, which establish a uniform standard, were intended to strike a balance between product safety and protecting interstate commerce from the undue burdens imposed by non-uniform standards." Because of the detailed warning requirements concerning toxic shock syndrome, many plaintiffs fight an uphill battle when litigating a negligence or breach of implied warranty claim. The court in *Krause* noted that the "plaintiff may have a difficult time establishing and proving her negligence and breach of warranty claims." The plaintiff in *Haddix v. Playtex Family Products, Corp.* was unable to maintain her burden of proving that the tampon was unreasonably dangerous because of the warnings concerning toxic shock syndrome that were affixed to the product's package, as mandated by federal law.

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260. *See Ausness, supra* note 186, at 227-28 (citing 21 CFR § 884.5460(b)) (current through September 29th, 2000)).
262. *See id.* at 169.
263. 107 F.3d 737 (9th Cir. 1997).
264. *Id.* at 742.
265. *Id.* at 739-40.
266. *Ausness, supra* note 186, at 228 (citing Lindquist v. Tambrands, Inc., 721 F. Supp. 1058, 1063 (D. Minn. 1989)).
268. 138 F.3d 681 (7th Cir. 1998).
269. *Id.* at 686.
F. Miscellaneous Product and Subject Classifications

Congress has included preemption-like language in numerous statutes in addition to those highlighted in the previous sections. Some of these legislative initiatives are fairly old, while others are quite modern. As has been seen, the law of federal preemption “to a large extent defies useful generalization[.]”270 Rather, “[t]he cases are very specific to the regulated subject [matter].”271

i. Federal Boat Safety Act

The Federal Boat Safety Act (“FBSA”)272 is intended to promote boating safety through various means that include, inter alia, requiring manufacturers of certain boating equipment to comply with safety standards promulgated by the Secretary of Transportation.273 The Coast Guard is the federal agency to which this regulatory function is delegated.274 In 1988 the Coast Guard “directed” the National Boating Safety Advisory Council to evaluate whether or not the Coast Guard should promulgate a regulatory standard requiring propeller guards. The Council recommended that no such regulatory action be taken.275 The Coast Guard adopted the Council’s recommenda-

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271. id.

The regulatory process is very structured and stringent regarding justification. Available propeller guard accident data do not support imposition of a regulation requiring propeller guards on motorboats. Regulatory action is also limited by the many questions about whether a universally acceptable propeller guard is available or technically feasible in all modes of boat operation. Additionally, the question of retrofitting millions of boats would certainly be a major economic consideration. The Coast Guard will continue to collect and analyze data for changes and trends. . . . The Coast Guard will also review and retain any information made available regarding development and testing of new propeller guard devices or other information on the state of the art.
tion, and in official correspondence recognized and detailed its ongoing supervision of this subject matter.276

The issue of whether the Coast Guard's decision not to implement a standard requiring propeller guards on recreational boats preempted state law personal injury claims arising from a claimed causal connection between the absence of such guards and injuries suffered was litigated in Ryan v. Brunswick Corp.277 The suit was brought by the wife of a swimmer who was killed following an accident in which he came into contact with a revolving propeller on a recreational watercraft.278 The FBSA preemption provision reads:

[A] state . . . may not establish, continue in effect, or enforce a law or regulation establishing a recreational vessel or associated equipment performance standard or imposing a requirement for associated equipment . . . that is not identical to a regulation prescribed under [this title].279

The Michigan Supreme Court, evaluating the FBSA language "law or regulation" with the preemptive language "requirement or prohibition" in the statute interpreted in Cipollone v. Liggett Group,280 found the difference between the two phrases to be "insignificant."281 Finding that the federal regulatory decision that propeller guards should not be required preempted state tort law claims based upon the absence of such devices, the Ryan court noted that it was "join[ing] numerous other courts that have held that 'common law causes of action may constitute state regulation and [impermissibly] impose a requirement on manufacturers to have propeller guards through an award of damages'"282

277. 557 N.W.2d 541 (Mich. 1997).
278. See id. at 543-44.
281. Ryan, 557 N.W.2d at 547-48.
Reaching the opposite conclusion, the Missouri Court of Appeals in *Ard v. Jensen* was persuaded that the FBSA savings clause was properly considered *in pari materia* with legislative history indicating a legislative purpose that state statutes and common law continue to provide a floor for appropriate safety measures, or, in the court's words, "the common law [should] be the minimum standard to be built upon by the Secretary's regulations." In finding no preemption of an injured skier's suit, the Missouri appellate court found particularly telling the language of the accompanying Senate Report that "[t]he purpose of the [savings clause] is to assure that in a product liability suit mere compliance by a manufacturer with the minimum standards promulgated under the Act will not be a complete defense to liability."

ii. **Consumer Product Safety Act**

In *Moe v. MTD Products, Inc.* the Court of Appeals for the Eighth Circuit ruled that the Consumer Product Safety Act (CPSA) preempted a state common law damages claim. The court examined the preemptive effect of the CPSA's lawn mower safety standard, which requires (1) that each power lawn mower have a blade control system that stops the blade rotation within three seconds after the operator's hands leave the handle; and (2) that each mower contain a label warning "DANGER, KEEP HANDS and FEET AWAY" and showing a blade cutting into the forefinger of a hand. Plaintiff injured his hand when reaching into his mower's side chute to remove some grass, and his hand came into contact with the blade, which had unexpectedly begun to turn.

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283. 996 S.W.2d 594 (Mo. Ct. App. 1999) (A suit was brought against the boat operator and its manufacturer when a waterskier suffered injuries when the boat backed over him as he prepared to ski.).

284. *Id.* at 599.

285. *Id.* at 594 (citing S. REP. No. 92-248 (1971), reprinted in 1971 U.S.C.C.A.N. 1333, 1352, stating "This section is a Committee Amendment and is intended to clarify that compliance with the Act or standards, regulations, or orders promulgated thereunder does not relieve any person from liability at common law or under State law." *Id.*)

286. 73 F.3d 179 (8th Cir. 1995).


288. *See id.* at 181. *See Moe, 73 F.3d* at 181.

289. *See* 16 CFR § 1205.5(a) (current through September 29th, 2000).
The CPSA preemption clause provides:

Whenever a consumer product safety standard under this chapter is in effect and applies to a risk of injury associated with a consumer product, no State or political subdivision of a State shall have any authority either to establish or to continue in effect any provision of a safety standard or regulation which prescribes any requirements as to the performance, composition, design, finish, construction, packaging, or labeling of such product which are designed to deal with the same risk of injury associated with such consumer product, unless such requirements are identical with the requirements of the Federal standard.290

The Act's "savings clause" reads: "Compliance with consumer product safety rules or other rules or orders under this Act shall not relieve any person from liability at common law or under state statutory law to any other person."291 The Court of Appeals for the Eighth Circuit held that plaintiff's warning claim was preempted, but that the design claim was not.292

iii. Federal Hazardous Substances Act

The majority of courts considering the matter have concluded that compliance with Federal Hazardous Substances Act ("FHSA")293 labeling regulations preempt state common law claims premised upon a party's argument that the manufacturer should have provided better hazard warnings.294 The legislative history of the FHSA announced that its purpose was to "provide nationally uniform requirements for adequate cautionary labeling on packages of hazardous substances which are

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292. See Moe, 73 F.3d at 183.
sold in interstate commerce and are intended or suitable for household use."295 The Act's preemption clause reads:

[If a hazardous substance or its packaging is subject to a cautionary labeling requirement . . . [under this title] designed to protect against a risk of illness or injury associated with the subject, no State or political subdivision of a State may establish or continue in effect a cautionary labeling requirement applicable to such substance or packaging and designed to protect against the same risk of illness or injury unless such cautionary labeling requirement is identical to the labeling requirement [imposed pursuant to this title].296

_Gurrieri v. William Zinsser & Co._297 was a personal injury suit arising from the injurious inhalation of and contact exposure to a stain remover.298 The plaintiff claimed that the label on the defendant's product, which contained about 2% methyl alcohol, should have had more vigorous language or hazard signage, such as a skull and crossbones, that would be mandated had the product contained more than 4% methyl alcohol.299 The court found that the product's label complied with the warning label standard set pursuant to the FHSA,300 and, affirming judgment for the defendant, wrote:

[T]o the extent that the plaintiff proposes additional, different, or more clearly-stated warnings, [these claims are preempted by the FHSA].301 . . . In the present case, both [the statute and the regulations] address the labeling of hazardous or dangerous substances. . . . A finding that specific local warnings pursuant to state law must apply to products containing less than 4% of methyl alcohol would create system of possibly fifty or more different labeling requirements throughout the country, contrary to Congress' obvious intent in passing the FHSA to 'provide nation-

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298. See _id_. at 833.
299. See _id_. at 841.
300. See 15 U.S.C. § 1261(p) (West 1997); _but see_, 16 C.F.R. § 1500.14(b)(4) (current through September 29th, 2000) (imposing different labeling requirement for household products containing 4% or more ethyl alcohol).
301. _Gurrieri_, 728 A.2d at 841 (citing _Salazar v. Whink Products Co._, 881 P.2d 431 (Colo. Ct. App. 1994)).
ally uniform requirements for adequate cautionary labeling.\textsuperscript{302}

We reject this result.\textsuperscript{303}

In \textit{State ex rel. Jones Chemicals, Inc. v. Seier},\textsuperscript{304} a Missouri appeals court found similarly that the FHSA statutory language pertaining to product labeling preempted a plaintiff's products liability claim under state law and that the labeling of a container of muriatic (hydrochloric) acid contained inadequate notice of the product's risks.\textsuperscript{305} In reaching this conclusion, the Missouri court found that the FHSA reflected congressional intent to establish uniform nationwide standards for labeling of hazardous substances, and to avoid the "impracticality of having states produce potentially fifty different labels for a particular hazardous substance."\textsuperscript{306}

\section*{IV. CONCLUSION}

\textit{Cipollone v. Liggett Group, Inc.}\textsuperscript{307} ended the long reign of judicial deference to state common law liability and regulatory obligations, and intimated that the existence of an express preemption provision might, standing alone, suffice to permit a finding of federal preemption. In so doing, the Court stifled the effect of equivalently specific savings clauses that purported to preserve state products liability or regulatory actions from federal suffocation.

In \textit{Geier v. American Honda Motor Co.},\textsuperscript{308} the Court refined its \textit{Cipollone} analysis, and in so doing gave breathing room to extant and future savings clauses in federal safety-related statutes. The \textit{Geier} court expanded little in suggesting a means for lower courts to proceed with confidence in reconciling express preemption provisions with similarly explicit savings clauses. The result of the Court's recalcitrance will surely be played out in the decisions of lower courts which are now left to decide if savings and preemption clauses may be interpreted in a comple-

\begin{footnotesize}
\textsuperscript{302} Id. at 841 (quoting H.R.Rep. No. 86-1861 at 1 (1960), reprinted in 1960 U.S.C.C.A.N. 2833.)

\textsuperscript{303} Id. at 841.

\textsuperscript{304} 871 S.W.2d 611 (Mo.App. 1994).

\textsuperscript{305} See id. at 614.

\textsuperscript{306} Id. at 612-13, quoting Moss v. Parks Corp., 985 F.2d 736 (4th Cir. 1993).

\textsuperscript{307} 505 U.S. 504 (1992).

\textsuperscript{308} ___ U.S. ___, 120 S. Ct. 1913 (2000).
\end{footnotesize}
mentary way. The inevitable consequence of the Supreme Court's failure to harmonize *Cipollone* and *Geier* will be a bumper crop of conflicting decisions brought about by the inability of courts to determine in a consistent way whether the polar magnetic field of express preemption clauses, or that of savings clauses, is the stronger.