May 1985

Failure to Identify the Defendant in Tort Law: Towards a Legislative Solution

Victor E. Schwartz
Liberty Mahshigian

Follow this and additional works at: https://scholarship.law.berkeley.edu/californialawreview

Recommended Citation

Link to publisher version (DOI)
https://doi.org/10.15779/Z383447

This Article is brought to you for free and open access by the California Law Review at Berkeley Law Scholarship Repository. It has been accepted for inclusion in California Law Review by an authorized administrator of Berkeley Law Scholarship Repository. For more information, please contact jcera@law.berkeley.edu.
Failure to Identify the Defendant in Tort Law: Towards a Legislative Solution

Victor E. Schwartz†

and

Liberty Mahshigian‡

Prior to the late 1970's, an essential element of any plaintiff's product liability case was identification of the named defendant as the manufacturer or supplier of the allegedly defective product.¹ But in the late 1970's, numerous claims began to arise which involved a "generic" product and in which the plaintiffs alleged a defect in design or failure to warn common to all such products, such as asbestos or diethylstilbestrol ("DES"). Even where these plaintiffs have been able to prove that a manufacturer breached a duty of care in manufacturing the generic product under either a negligence or strict liability theory, that the defect in the generic product was the cause-in-fact and proximate cause of the plaintiff's harm, and that the plaintiff has suffered some provable injuries, recovery has been uncertain. These cases are characterized by the existence of a generic product and a large number of potential defendants, and thus by an absence of evidence as to the identity of the manufacturer of the product that allegedly caused the injury. Moreover, in asbestos and DES cases, plaintiffs are exposed to products many years before the injuries become manifest, which increases the plaintiffs' burden of producing evidence identifying the manufacturer.

Courts have been wrestling with this problem of plaintiffs who are able to prove that their injuries were caused by some act that constituted

¹ See W. KEETON, D. DOBBS, R. KEETON & D. OWEN, PROSSER AND KEETON ON THE LAW OF TORTS § 103, at 713-14 (5th ed. 1984) [hereinafter cited as PROSSER & KEETON].
a breach of duty to them but are not able to identify the actual wrong-
doer. In a number of cases in which the plaintiff is able to prove all
elements of his or her cause of action except the identity of the wrong-
doer, courts have fashioned several theories to hurdle the plaintiff over
the barrier posed by the identification requirement. All of these theories
deviate from traditional and fundamental principles of tort law. Their
development is motivated by the courts' policy judgment that as between
an innocent plaintiff and defendants who are allegedly guilty of some
wrongful conduct, the plaintiff should prevail—even if the alleged (not
necessarily established) conduct in question did not cause the plaintiff's
injury.

Thus, in a number of recent cases plaintiffs have brought actions
against an entire group or a substantial number of manufacturers that
made such a generic product, seeking recovery against the joined manu-
facturers under several theories of liability. In a few cases, courts have
modified traditional requirements of alternative liability or concerted
action liability in order to afford recovery for plaintiffs, expanding these
theories beyond the point at which liability can be justified. In other
cases, courts have developed completely new theories, imposing liability
on many or all members of an industry while apportioning damages
according to any of a variety of factors. These latter theories are riddled
with many practical problems, often raising confusing or complicated
issues that render litigation slow, complex, and costly for all parties
involved. Furthermore, imposition of liability without regard to individ-
ualized fault is often unfair to certain defendants, and inconsistent with a
basic underlying notion of the tort system: that only one who is responsi-
ble for causing particular and identifiable harm should be held liable for
civil damages.²

The DES nonidentification cases present an unresolved dilemma.
On the one hand, courts can deny liability in all of these cases. Most trial
courts have done this and have thus remained in step with the existing
parameters of tort law. But denying all recovery to plaintiffs offends
some courts' sense of justice. On the other hand, courts can create a
novel judicial theory in order to impose liability. While this might satisfy
some courts' sense of justice, it leaves in its wake extraordinary legal
costs, delay, injustice, and the imposition of tort law liability on a party
who is, in fact, not responsible for plaintiff's harm.

This Article will begin by examining the various ways that courts
have struggled to deal with the identification problem in cases involving a
generic drug, DES. It will show that the tort litigation system is not able
to resolve the identification problem and thus that the problem can only

². J. FLEMING, THE LAW OF TORTS 179-81 (6th ed. 1983); PROSSER & KEETON, supra note
1, at 41, 263-68.
be resolved satisfactorily by a legislative solution. This Article will then provide, on the basis of the problems raised by these common law solutions, a framework for a legislative solution to the problem.

I

THE BACKGROUND: THE DEVELOPMENT AND MARKETING OF DES

In order to appreciate fully the identification problem as it applies to DES, it is essential to understand how the drug was developed and marketed. Let us begin with some basic scientific facts. Natural estrogens, the female sex hormones present in varying amounts in the body of every woman, were first isolated outside the human body in 1929. Physicians and scientists soon discovered that the administration of controlled doses of natural estrogens could alleviate medical problems associated with estrogen deficiencies. Natural estrogens, however, still very expensive to isolate, and were difficult and painful to administer by injection.

In 1937, two British scientists, Drs. E.C. Dodds and Leon Golburg, synthesized a compound that has most of the characteristics of estrogen. The synthetic estrogen, stilbestrol, also called diethylstilbestrol and more commonly referred to as DES, could be produced at a fraction of the cost of isolating natural estrogens and was effective when administered orally. Dr. Dodds did not apply for a patent, leaving the product available for marketing by all.3

Before DES could be marketed in the United States, approval of the Food and Drug Administration (“FDA”) was required.4 Federal law required that each company wishing to market DES submit a New Drug Application (“NDA”), and that this application include a description of the chemical composition of the drug, clinical data establishing its safety and efficacy, information regarding the methods for manufacturing, processing, and packing the drug, and the format of proposed labeling.5

By the end of 1940, ten pharmaceutical companies had filed NDA’s seeking approval to market DES in up to one milligram doses to treat various conditions, including menopausal symptoms, senile vaginitis, and gonorrheal vaginitis. None of these proposed uses involved medical conditions relating to pregnancy. A large amount of clinical data was independently collected and submitted to the FDA.6 On December 20, 1940, the FDA convened a meeting with the drug companies, at which it formally requested the companies to withdraw their NDA’s and submit

The FDA also requested the drug companies to use the same United States Pharmacopeia (U.S.P.) standard to ensure that the active ingredient in all DES products was the same, to develop uniform labeling regarding usage and dosage of the drug, and to include a "permission clause" in their NDA's authorizing the FDA to use the materials gathered by each firm in considering any other NDA's that might be filed. In response, representatives of firms that had filed NDA's formed a small committee to compile the studies and clinical data into the master file that was submitted to the FDA. The companies then filed individual NDA's and, in September of 1941, the FDA approved the marketing of DES for use in the treatment of menopausal symptoms, senile vaginitis, gonorrheal vaginitis, and suppression of lactation. The small committee disbanded in October 1941.

Throughout the late 1930's and the 1940's, independent physicians conducted research on other possible uses for DES, including the treatment of pregnancy problems. In 1944, the FDA approved the use of DES for treatment of prostate cancer in men. In 1947, following published reports of successful treatment, several companies filed new or supplemental NDA's for the use of DES as a miscarriage preventative. Although these NDA's did not refer to the master file of clinical data that had been submitted with the 1941 NDA's, it was the FDA's policy to consider all of the material that it had in support of the original NDA's. By 1952, the FDA had decided that DES was no longer a "new drug." This meant that companies wishing to market DES did not have to file NDA's.

In 1971, Dr. Arthur Herbst and two colleagues published a paper suggesting a statistical relationship between fetal exposure to DES and a specific form of cancer, called clear cell adenocarcinoma, in some of the daughters of women who had taken the drug during pregnancy. In the period of time between the FDA approval of DES and the publication of this article in 1971, almost 300 drug companies had marketed DES in

8. Id. at 374.
9. Id.
dosages suitable for use in the treatment of certain pregnancy problems.\textsuperscript{14} After much further study and discussion, the FDA required drug companies in their labeling to recommend against DES use in the prevention of miscarriages. DES was never banned by the FDA and is still used today for treatment of problems unrelated to pregnancy, such as cancer of the prostate, certain forms of breast cancer, and menopausal symptoms.\textsuperscript{15}

Women who were exposed to DES taken by their mothers during pregnancy have sought to recover damages from DES manufacturers. Some of these women have adenosis, which is a benign condition characterized by the presence of glandular tissue in the vagina where it does not usually appear. Adenosis has not been shown to be a precancerous condition and it seems to regress and disappear over time. It is a condition that also exists in women who were not exposed to DES.\textsuperscript{16} Because the condition is benign and usually disappears by the time the women reach their late twenties, no medical or surgical therapy is necessary and actual physical harm is minimal. For that reason, plaintiffs have seldom pursued adenosis damage claims in court.

Other women who were exposed to DES developed vaginal or cervical cancer and have brought suit. One important study has reported a statistical association between clear cell adenocarcinoma and DES.\textsuperscript{17} Many of these women, however, are unable to identify the manufacturer of the DES taken by their mother, in part because of the passage of time resulting in the absence of records, and in part because DES usually was prescribed by a generic and not a brand name. As a result of their inability to identify the manufacturers of the DES taken by their mothers, some women have brought suit against several DES manufacturers. Most courts, recognizing and following traditional principles of tort law, have denied recovery to plaintiffs unable to satisfy the identification requirement. A few courts have permitted these actions to proceed by applying various traditional theories of joint liability, but as discussed below, traditional tort theories of joint liability will not rationally extend to the circumstances of DES cases. Because of this, some courts have modified traditional joint liability theories and have developed new theories in order to provide causes of action for DES plaintiffs.

\textsuperscript{14} Martin v. Abbott Laboratories, 689 P.2d 368, 374 (Wash. 1984).

\textsuperscript{15} Id. at 373, 374; Ferrigno v. Eli Lilly and Co., 175 N.J. Super. 551, 565, 420 A.2d 1305, 1312 (1980).


\textsuperscript{17} Herbst, Ulfelder & Poskanzer, \textit{supra} note 13.
II

THE COMMON LAW THEORIES OF RECOVERY: INABILITY
OF THE TORT LITIGATION SYSTEM TO RESOLVE
THE DES IDENTIFICATION PROBLEM

A. Summers v. Tice Alternative Liability

The first modern theory of alternative liability was judicially created in *Summers v. Tice.*\(^{18}\) In that case, the California Supreme Court ruled that where two defendants had both acted negligently toward the plaintiff, each could be liable for the harm caused, although only one of the defendants could have been the cause-in-fact of the plaintiff's injury. The defendants in *Summers* negligently fired, at the same time, at a quail in the plaintiff's direction. The plaintiff was struck in the eye and face by a shot from one of the guns. Unable to prove which one of the two defendants had fired the injury-causing shot, the plaintiff filed suit against both.

The court recognized that while both defendants had acted negligently, only one of the two defendants had actually caused the plaintiff's harm. The court held that each defendant would be jointly and severally liable for the harm unless he could prove that he was not the person who fired the injury-causing shot. The court reasoned that it was preferable that the two wrongdoers, both of whom had acted negligently toward the plaintiff and had better access to the relevant evidence, should bear the burden of absolving themselves rather than leaving the innocent plaintiff remediless.\(^ {19}\)

The *Summers* alternative liability theory soon gained acceptance in a few courts and was adopted by the authors of the Second Restatement of Torts.\(^ {20}\) Today, the prevailing judicial view is that the *Summers* alternative liability theory should apply if (1) all the potential wrongdoers are joined as defendants, (2) all of the defendants acted tortiously, and (3) the defendants are in a better position to prove the identity of the actual wrongdoer.\(^ {21}\) The plaintiff is not required to show an express agreement or tacit understanding among the defendants; thus, unlike the concerted action theory discussed below, the *Summers* alternative liability theory can be used to impose liability on defendants who acted independently.

---
\(^{18}\) 33 Cal. 2d 80, 199 P.2d 1 (1948).
\(^{19}\) 33 Cal. 2d at 86, 199 P.2d at 4.
\(^{20}\) See, Restatement (Second) of Torts § 433B(3) (1964); see also F. Harper & F. James, The Law of Torts, § 20.2, at 1115-16 (1956). It should be noted that *Summers v. Tice* alternate liability differs from alternate liability in the strict sense in that the latter usually involves a situation where several parties were in a position to cause plaintiff's harm, but there is no proof that all defendants acted tortiously. See Prosser & Keeton, supra note 1, § 41, at 270-71; see also Restatement (Second) of Torts, § 433B(3) comment g (1964).
Courts have refused to apply the theory in DES cases, however, for several reasons. Most courts have refused to apply alternative liability because the plaintiffs as a practical matter are unable to bring all possible tortfeasors before the court. Over 300 drug companies marketed DES for use in connection with pregnancy problems. It would be very difficult for a plaintiff to determine which of the 300 companies may have marketed the DES taken by her mother, to locate these companies, to obtain jurisdiction over them, and to serve process on them.

Second, the alternative liability theory presupposes that all parties who could have been responsible for plaintiff's harm have been joined as defendants. Unless plaintiffs join every manufacturer of DES that was sold in the area during the time of exposure to the drug, the responsible manufacturer may not be one of the defendants. Where substantially less than all possible tortfeasors have been joined as defendants, it would be unfair to shift the burden of proving identification to the defendants, since it is probable that none of the defendants before the court the caused plaintiff's harm. Liability in such a case would shift the burden of proof to a group of defendants who may not have been the cause in fact of the injury.

Third, courts have pointed out that the Summers theory is inappropriate for DES cases because the DES manufacturers are not in a better position to offer evidence to establish who manufactured the drug ingested by plaintiff's mother. In Summers-like fact situations, the defendants are in a better position to prove which of them caused the injury, thus justifying a shift in the burden of proof. Summers and Tice were the best persons to testify as to the aim of their guns and the destination of their shots. In contrast, the DES manufacturers are not in control of all evidence relevant to the identification of the drug ingested by a plaintiff's mother—the pharmacy where it was purchased, the size and shape of the pills, the dates of purchase, and so on. Thus, the primary justification for shifting the burden of proving identification in Summers is not present in DES cases. To shift the burden of proving identification to DES manufacturers is not only unfair to the DES manufacturers but is also inefficient, because it shifts the burden of proof to a party not in control of the relevant evidence.

The Summers theory is inappropriate for application in DES cases for a fourth reason. A justification for imposing liability on both Sum-
mers and Tice, as well as for shifting the burden of proof, was that both defendants had acted negligently toward the plaintiff, although the negligence of only one of them was the cause-in-fact of the injury. Yet in almost every DES case decided to date, the plaintiff has failed to show that the manufacturers of DES acted negligently toward her, though they may have acted negligently toward DES users. Absent proof of any negligent conduct, the fact that a company produced a drug, but not the particular drug that allegedly caused the plaintiff's injury, by no means establishes that the company breached a duty of care toward the plaintiff.

Notwithstanding these difficulties, the Michigan Supreme Court has recently applied a variation of the *Summers* theory of alternative liability to a DES case. In *Abel v. Eli Lilly & Co.*, the Michigan Supreme Court ruled that the theories of alternative liability and concert of action may be applied in suits brought by DES daughters who cannot identify which specific company made the drugs that their mothers took when pregnant. The court modified the elements of the traditional theory of alternative liability, setting out three requirements that plaintiffs must meet in order to recover under this theory. To shift the burden of proof to defendants on the issue of causation in fact under the modified theory, a plaintiff must show that (1) all the defendants acted tortiously, (2) she has been harmed by the conduct of one of the defendants, and (3) she is unable, through no fault of her own, to identify which defendant caused her injury. As a prerequisite to recovery, the plaintiff is also required to show that she has made a genuine attempt to locate and identify the manufacturer responsible for her individual injury. The genuineness of this attempt will be determined by the standard of "due diligence".

In order to support the second requirement, the plaintiff must bring before the court all parties who promoted DES in Michigan at the time her mother took the drug. A plaintiff must prove that all the defendants manufactured one or more of the three chemically related drugs involved (DES, dienestrol, or diethylstilbestrol dipropionate), that her mother ingested one of these drugs, that the drug her mother ingested

---

25.  *Id.* at 331-32, 343 N.W.2d at 173.
26.  *Id.* Some of the plaintiffs in this case alleged that they were unable to identify the manufacturer of the product that harmed them because of the absence of pharmacy records and the defendants' use of a generic marketing scheme. Michigan law requires pharmacists to maintain prescription drug records for only five years. *Id.* at 321, 343 N.W.2d at 168. The court held that the pleadings were sufficient to withstand a motion to dismiss. *Id.* at 340-41, 343 N.W.2d at 176-77. Although the Michigan court called the motion a "motion for summary judgment," it was a motion based solely on the pleadings and not on the facts and accordingly, actually was a motion to dismiss. *Id.* Hereinafter, it will be referred to as a motion to dismiss. The remaining plaintiffs were permitted to offer proof of the identity of the manufacturers liable; if unable to identify the manufacturers, they could resort to the alternative liability theory.
27.  *Id.* at 331, 343 N.W.2d at 173.
was manufactured or distributed in Michigan, and that the three drugs are essentially identical in their allegedly injury-producing results. If the defendants are unable to exculpate themselves, they will be jointly and severally liable for plaintiff's damages.  

By requiring the plaintiff to bring forth all possible defendants and by requiring that all of the defendants to have breached a duty toward some potential plaintiff, the Michigan court solved some of the problems arising from the application of Summers to DES cases. Specifically, all defendants have allegedly breached a duty of care and are all present before the court, thus justifying to a limited extent a shifting of the burden of proof. But the Michigan court did not resolve the problem in administration caused by the requirement that the plaintiff bring all defendants before the court. Complex And the Michigan court failed to realize that since defendants are not in a better position than plaintiffs to come forward with identification information, the full policy justification of Summers is missing here, raising the spectre of an arguably unfair shift in the burden of proof that will in practice operate as an unfair finding of alternative liability. Finally, the adoption of the Summers doctrine in a context where no breach of a duty of care to the plaintiff by a defendant before the court has been established ignores, to a greater extent than in Summers, the traditional tort law requirement of cause-in-fact. In sum, the application of Summers to DES cases, even if qualified as in Abel, solves the DES problem by ignoring problems of proof for plaintiffs and problems of unfairness toward defendants.

B. Concert of Action

Courts have also developed the concert of action theory to mitigate the identification problem. This theory imposes joint and several liability on all persons who acted in concert or pursuant to a common scheme or plan to commit a tortious act, even though fewer than all of those persons actually caused the harm.  

The paradigm concert of action case is a drag race in which all who participate in the race are deemed responsible should one of them cause harm to a bystander. The defendants who acted in concert were "joint tortfeasors" and "the act of one is the act of all." The rationale for imposing liability is that while the party on whom liability is imposed may not have actually caused the injury, his or her wrongful conduct actively encouraged the one who did.

28. Id. at 334, 343 N.W.2d at 174.
29. See PROSSER & KEETON, supra note 1, § 46, at 322-24.
30. See, e.g., Bierczynski v. Rogers, 239 A.2d 218 (Del. 1968); RESTATEMENT (SECOND) OF TORTS, § 876 (1977). Such "concerted" wrongdoers were considered "joint tortfeasors" even by the early common law. PROSSER & KEETON, supra note 1, § 52, at 346.
31. PROSSER & KEETON, supra note 1, § 52, at 346.
Unlike the theory of alternative liability, concert of action does not require the presence of the injury-inflicting party before the court.\textsuperscript{32} Thus, a plaintiff's inability to identify the defendant which caused his or her injury does not preclude recovery from all defendants. Similarly, a defendant joint tortfeasor who is able to prove that he or she did not cause the plaintiff's harm is not able to avoid liability. As applied to DES cases, and unlike the alternative liability theory, the concert of action theory does not permit a defendant to escape liability by showing that it did not manufacture the type of DES taken by the plaintiff's mother or that it did not market DES at the time or in the area in which the plaintiff's mother purchased the drug.\textsuperscript{33}

At least one court has imposed liability on DES manufacturers under a revised version of the concert of action theory. In \textit{Bichler v. Eli Lilly & Co.},\textsuperscript{34} the plaintiffs argued to a New York trial court that the "parallel and imitative" conduct of the DES manufacturers should be enough to show concerted action. The plaintiff made no showing that the defendant actively encouraged others into negligent conduct. The trial court instructed the jury that the "implied or tacit agreement or understanding" element of concerted action could be evidenced by the "conscious parallel" conduct of the drug companies.\textsuperscript{35} The court further stated that the concerted action could also be defined as "acting independently of each other in committing the same wrongful act, but although acting independently, their acts have the effect of substantially encouraging or assisting the wrongful conduct of the other, which, in this case, was the alleged failure to adequately test."\textsuperscript{36}

The trial court entered judgment in favor of the plaintiff, and the defendant appealed. The Appellate Division affirmed,\textsuperscript{37} and Eli Lilly & Company again appealed, seeking reversal on two grounds: the trial court's instructions on concerted action liability were erroneous, and the evidence before the jury was legally insufficient to support a verdict in the plaintiff's favor on the issue of concerted action. The New York Court of Appeals found that it could not review the instruction because the defendant had failed to preserve the challenge for review by appropriate objections.\textsuperscript{38} Thus, although it affirmed the Appellate Division, the Court of Appeals did not decide whether evidence of "conscious parallelism" among the drug manufacturers is sufficient to establish concert of

\textsuperscript{32} Id., § 47, at 327.
\textsuperscript{35} Bichler, 79 A.D.2d at 326, 436 N.Y.S. 2d at 631.
\textsuperscript{36} Id.
\textsuperscript{37} Id. at 336, 436 N.Y.S.2d at 636-37.
action. Therefore, it is not certain what type of conduct is necessary to constitute concert of action in New York.

Other courts have rejected the “conscious parallelism” approach to concert of action liability. The Wisconsin Supreme Court held that [the plaintiff’s allegations that the] “defendants failed to adequately test [DES] or to give sufficient warning of its dangers and that they relied upon the tests performed by one another and to the advantage of each others’ promotional and marketing techniques” . . . do not support the theory that the defendants tacitly agreed to produce and market DES without adequately testing the drug or warning of its potential dangers.\(^{39}\)

The court concluded that “[a]lthough there was a substantial amount of parallel action by the defendants in producing and marketing DES for use in pregnancy,” the extent of parallel action did not “rise to the level of ‘acting in concert.’”\(^{40}\) Similarly, the California Supreme Court determined that concerted action could not be established by evidence of “parallel or imitative conduct”\(^{41}\) and thus refused to apply concert of action liability to the drug companies’ conduct in “relying upon each others’ testing and promotion methods.” The court reasoned that such conduct was a “common practice in the industry”\(^{42}\) and that applying a concert of action theory to those circumstances “would expand the doctrine far beyond its intended scope.” “[V]irtually any manufacturer [could be held] liable for the defective products of an entire industry, even if it could be demonstrated that the product which caused the injury was not made by the defendant.”\(^{43}\) Courts in New Jersey, South Carolina, Florida, and Missouri have also rejected the New York Appellate Division’s “conscious parallelism” basis for concert of action liability.\(^{44}\)

In \textit{Abel v. Eli Lilly and Co.},\(^{45}\) the Michigan Supreme Court permitted the DES plaintiffs to proceed on a concert of action theory as well as an alternative liability theory. The court stated that the defendant manufacturers would be jointly and severally liable for the plaintiffs’ damages if the plaintiffs could establish that all defendants acted tortiously pursuant to a “common design.”\(^{46}\) In order to withstand a motion to dismiss,

---

40. Id.
42. Id.
43. Id.
46. Id. at 338, 343 N.W.2d at 176.
a plaintiff had only to allege that the defendants were “jointly engaged in tortious activity as a result of which the plaintiff was harmed.”47 The plaintiffs’ allegations that the defendants “acted together in negligently manufacturing and promoting drugs which were ineffective and dangerous, were inadequately tested, and were distributed without sufficient warnings,” thus could withstand a motion to dismiss.48 The court noted that the concert of action theory, unlike alternative liability, does not require that a plaintiff be unable to identify the specific defendant who caused his or her injury. The fact that some of the plaintiffs in this case had alleged that a specific DES manufacturer caused their injuries would not preclude them from recovering from all of the defendants under a concert of action theory.49

To summarize, all states except New York and Michigan have rejected the concert of action theory in DES cases. This may be due to the conceptual inapplicability of the theory in DES cases. In paradigm cases of acting in concert, such as the acting together to hold a drag race, the concerted activity was itself tortious. As pointed out by one district court, none of the drug companies’ concerted activities—the filing of wholly separate applications to the FDA for permission to market DES, the formation of a small committee, the resubmission of their applications to a master file—constitute tortious action.50 As pointed out by another court, the concerted action theory only applies to action such as express or tacit agreement, or a common plan to inadequately test DES or to fail to warn the public of the drug’s known dangers.51 Plaintiffs can rarely prove such conduct, and courts have been justifiably unwilling to assume that such conduct took place, and have generally looked elsewhere to solve the DES identification problem.

C. Enterprise Liability

The “enterprise liability” theory, first recognized by a federal district court in New York, presents a third possible basis for recovery in DES cases. In Hall v. E.I Du Pont De Nemours & Co.,52 thirteen children were injured in twelve separate blasting cap incidents over a period of four years. The plaintiffs were not able to identify the specific manu-

47. Id.; see also supra note 26.
48. Id.
49. Id.
50. Morton v. Abbott Laboratories, 538 F. Supp. 593, 596-97 (M.D. Fla. 1982); see also Lyons v. Premo Pharmaceutical Labs, 170 N.J. Super. 183, 406 A.2d 185 (App. Div. 1979) (no concert of action because drug companies’ act of seeking approval for DES was not tortious, nor shown to have encouraged a tortious act).
facturers of the caps that had injured them. They sought damages from six blasting cap manufacturers, that made up virtually the entire Ameri-
can industry, and from the manufacturers' trade association.

The court imposed liability on the entire industry, even though only one member of that industry had actually caused the harm, because all members jointly controlled the risk either by an express agreement or by parallel behavior in delegating safety functions and adhering to insuffi-
cient safety standards. Under this industry-wide liability theory, the existence of industry-wide standards or practices may support a finding of joint control of risk and shift the burden of proving identification to the defendants.

Unlike the concert of action theory, enterprise liability does not require the plaintiff to prove the existence of an explicit agreement to commit a tort. Furthermore, the plaintiff is not required to demonstrate that the industry is rigidly controlled through trade associations or that the purpose of this common control is some particularly reprehensible breach of duty. The plaintiff begins by showing that her injury was caused by a product made by one of the named defendants and that the particular manufacturer is unknown. Second, to shift the burden of proof to the defendants, the plaintiff must “demonstrate defendants' joint awareness of the risks at issue in [the] case and their joint capacity to reduce or affect those risks.” The court stated that the factors relevant to the existence of joint control of risk include

the size and composition of the trade association's membership, its announced and actual objectives in the field of safety, its internal proce-
dures of decision-making on this issue, the nature of its information-gath-
ering system with regard to accidents, the safety program and its implemen-
tation by the association and member manufacturers, and any other activities by the association and its members (such as legislative lobbying) with regard to safety during the time period in question.

The application of this theory is very limited. In fact, it has never been adopted by any other court and was not even applied by the trial courts in any other blasting cap case. The theory had been rejected by courts in California, South Carolina, and Missouri because of the requirement of a centralized industry. Similarly, in Collins v. Eli Lilly

---

53. Id. at 374.
54. Id.
55. Id.
56. Id. at 379.
57. Id. at 378.
58. Id. at 376.
& Co., the Wisconsin Supreme Court held that the enterprise liability theory is inappropriate in DES cases involving hundreds of potential defendant drug companies. The court pointed out that "the assumption that the defendants jointly controlled the risk of injury is necessarily weak given the fact that so many drug companies entered and left the DES marketplace from 1947 to 1971," and "[i]t is less clear in DES cases than in Hall that the defendant drug companies were each violating a standard of safety, whether federally imposed or industry imposed, in producing and marketing DES for use in pregnancy." The theory also has been expressly rejected in DES cases in Florida, Michigan, New Jersey, and Washington.

This widespread rejection of the enterprise liability theory is hardly surprising. As the Hall court explicitly recognized, it is "manifestly unreasonable" to apply the theory to a "decentralized industry composed of thousands of small producers." For as commentators have argued, the enterprise liability theory should not apply in cases involving a large number of defendants because as the number of manufacturers increases, joint control and awareness of the risks at issue, and joint capacity to reduce those risks, become more difficult to imply as a predicate to liability. Additionally, the drug industry is required to comply with the standards and conditions set by the FDA. Courts rejecting the enterprise theory have recognized that in industries where standards are mandated by the government, the manufacturer has no choice but to comply, and compliance thus should not be a basis for liability. To apply the enterprise liability theory absent a finding of industry-wide control of standards and conditions would, of course, provide compensation to DES plaintiffs, but would again raise serious problems of fairness to defendants.

D. Market-Share Theory of Liability

In Sindell v. Abbott Laboratories, the California Supreme Court

60. 116 Wis. 2d 166, 342 N.W.2d 37 (1984).
61. Id. at 186-87, 342 N.W.2d at 47 (footnote omitted).
65. Id.
adopted a new basis upon which defendants may be liable even though
the plaintiff cannot prove which defendant manufactured the injury-caus-
ing product. The court developed a theory of market-share liability first
proposed in a student law review note and based on the Summers doc-
trine, extended and modified in two key respects.

First, the defendants in Sindell had to shoulder the burden of proof
on causation although, in contrast to the circumstances of the Summers
case, the DES manufacturers were not shown to be in a “better position”
than the plaintiff to offer evidence to determine which of them caused the
injury. The court recognized that the absence of evidence as to identifi-
cation is due not to the fault of the defendants, but rather to the passage
of time. Nevertheless, the court determined that the fact that the
defendants did not have greater access to information to establish the
identity of the manufacturer of the DES that allegedly injured the plain-
tiff should not prevent application of the Summers rule on shifting the
burden of proof. Second, the Sindell court did not require joinder of all
possible defendants. In Sindell, only five out of the hundreds of DES
manufacturers were named as defendants. In contrast, in Summers all of
the parties who could have been responsible for the plaintiff's injury were
joined as defendants.

Under the Sindell approach, the plaintiff need only bring before the
court those manufacturers that produced a “substantial percentage” of
the drug. Because the possibility that any one of the five defendants
manufactured the DES ingested by the plaintiff's mother was so remote,
the court thought it unfairly severe to impose complete joint liability on
each defendant unable to exonerate itself. Rather, the court apportioned liability according to the “likelihood that any of the defendants
supplied the product which allegedly injured plaintiff.” This likelihood
was measured “by the percentage which the DES sold by each of them
for the purpose of preventing miscarriage bears to the entire production
of the drug sold by all for that purpose.” Damages thus were apportioned among the defendants based on their share of the market, with
exact damage amounts determined by whether the liability so apportioned is joint, or joint and several. Finally, the court held that each

69. Sindell, 26 Cal. 3d at 600, 607 P.2d at 929, 163 Cal. Rptr. at 137.
70. Id. at 601, 607 P.2d at 930, 163 Cal. Rptr. at 138.
71. Id. at 602-03, 607 P.2d at 930-31, 163 Cal. Rptr. at 138-39.
72. Id. at 611-13, 607 P.2d at 937-38, 163 Cal. Rptr. at 144-46.
73. Id. at 603, 607 P.2d at 931, 163 Cal. Rptr. at 139.
74. Id. at 611, 607 P.2d at 937, 163 Cal. Rptr. at 145.
75. Id. at 611-12, 607 P.2d at 937, 163 Cal. Rptr. at 145.
76. Id. at 612, 607 P.2d at 937, 163 Cal. Rptr. at 145.
77. See infra text accompanying notes 82-85.
defendant could escape liability only by proving that it could not have made the product which allegedly caused the plaintiff's injuries. In DES cases, this can be shown by evidence that the defendant did not market DES in dosages usable for treatment of problem pregnancies at the time or in the location in which the plaintiff's mother purchased the drug, or that the drug manufactured by the defendant was of a different size, shape or color than the drug used by the plaintiff's mother.

An Illinois trial court recently adopted the Sindell theory in a DES case. The court reiterated the substantial share justification by stating that the market-share theory encompasses the consideration of probabilities and, along with considerations of policy, justifies the shifting of the burden [of proof] from one who had no role whatsoever to play in the purchase and consumption of the drug to those who have placed a substantial volume of sales of these products into the stream of commerce.

The court noted that if a plaintiff pleads that the defendants' shares of the total market are substantial, there will be a substantial likelihood that the product actually consumed by the DES mother was made by one of the defendants. The court did not, however, clarify the many issues left unresolved in the Sindell opinion.

In sum, there are four requirements to a successful use of the theory of market-share liability in product liability cases:

1. injury or illness occasioned by a fungible product (identical-type product) made by all of the defendants joined in the law suit;
2. injury or illness due to a design hazard, with each defendant having sold the same type product in a manner that made it unreasonably dangerous;
3. inability to identify the specific manufacturer of the product or products that brought about the plaintiff's injury or illness; and
4. joinder of enough of the manufacturers of the fungible or identical product to represent a substantial share of the market.

If these four requirements are satisfied, each defendant will be liable for the plaintiff's damages in an amount determined by its share of the relevant market, unless a defendant can prove that it could not have manufactured the product that caused the plaintiff's harm.

The Sindell market-share theory is one court's attempt to find a solution to the DES problem. Its inadequacies, however, demonstrate that it is not substantially fairer than any of the common law theories already discussed. First, the Sindell theory presupposes that if there is a substantial share of the market represented in the case, then there is a substantial likelihood that the tortfeasor is actually before the court. But

79. Id., slip op. at 14-15.
80. PROSSER & KEETON, supra note 1, § 103, at 714.
the defendants who happen to be joined as the "substantial share" of the market may be no more responsible for the plaintiff's harm than manufacturers who were not joined as defendants.

The Sindell theory has two other significant inadequacies. First and foremost, not all of the possible defendants are before the court. Second, not all of the defendants have acted in a negligent manner toward this particular plaintiff. For both of these reasons, a Summers-like shift in burden of proof seems harsher here than it did in the paradigm Summers situation.

In addition, there are numerous unresolved administrative problems raised by the Sindell theory. First, it appears (but is not clear from the opinion) that the court intended that the defendants' liability be several, rather than joint and several. Second, the court did not specify how the market-share proof is to be offered and did not define "appropriate market" or "substantial share." Third, the court did not address the issue of the availability of punitive damages under the market-share theory.81

Under a theory of several (as opposed to joint and several) market-share liability, each defendant is liable only for the percentage of the plaintiff's damages that corresponds to the defendant's percentage of the relevant DES market. If the plaintiff does not join all manufacturers of DES, she will not recover 100% of her damages. Thus, the plaintiff has an incentive to join all defendants that could be responsible for her injuries. Such an approach would increase the likelihood that the manufacturer that made the DES taken by the plaintiff's mother eventually pays damages, and more fairly limits the liability of defendants to actual market share—at the expense, however, of limiting the plaintiff's recovery to less than 100% of damages suffered.

Under a theory of joint and several market-share liability, a plaintiff may recover 100% of her damages despite a significant discrepancy between the defendants' actual market-share percentages and the damage apportionment percentages. Since the plaintiff is required to join only those defendants that together represent a "substantial share" of the DES market, each defendant may be liable for a portion of the plaintiff's damages that substantially exceeds the defendant's market share.82

81. The California Court of Appeal recently held that no punitive damages may be awarded in a case based upon the market-share liability theory. Magallanes v. Superior Court, 213 Cal. Rptr. 547 (Cal. Ct. App. 1983).


82. Comment, supra note 64, at 631.
for example, the plaintiff joins two defendants, each with 35% of the market share, each defendant will be liable for 50% of the plaintiff's damages. This discrepancy decreases as the total market share of the defendants approaches 100%.83

In resolving this issue of interpreting Sindell, some commentators have suggested that "the court did not intend to provide for joint and several liability, but each defendant is liable for the proportion of plaintiff's damage equal to the defendant's proportionate market share."84 This might be inferred from the fact that the Sindell court rejected the concert of action and enterprise liability theories because the drug companies had not acted as joint tortfeasors.85 The first California trial court interpreting Sindell to address this issue also ruled, in a preliminary order determining issues, that liability should be several and not joint.86 Although this issue was raised again by defendant drug companies in a California Court of Appeal case,87 no California appellate court has yet addressed it.

The second issue unresolved in Sindell is interpreting the requirement that the plaintiff join drug companies representing a substantial share of the "appropriate market." The Sindell court defined that market in terms of the use of the product,88 but provided neither geographic nor temporal limits for the concept.89

Finally, the Sindell court did not indicate how market shares are to be determined and did not address the practical difficulties in proving

83. Id.
84. PROSSER & KEETON, supra note 1, § 103, at 714.
85. Sindell, 26 Cal. 3d at 606, 609, 607 P.2d at 933, 935, 163 Cal. Rptr. at 141, 143.
87. The defendants in Magallanes v. Superior Court, 213 Cal. Rptr. 547 (Cal. Ct. App. 1985), suggested that liability was several in their brief on plaintiff's petition for writ of mandate. See Whether Punitive Damages Can Be Awarded Under Sindell Is Issue in Dispute, DES LITIGATION REP. (Andrews Publications) 5137, 5139. Defendants argued that the language in the Sindell opinion clearly suggests several liability, and in addition note that Justice Stanley Mosk, author of the Sindell opinion, stated that "we felt that it would be only fair to limit liability by the percentage of the market by which any manufacturer was responsible," Trends in Product Liability Litigation, TRIAL, Nov. 1980, at 82, 84. DES LITIGATION REP. at 5139.
89. The court did explicitly indicate, however, that a defendant may exculpate itself by showing that it did not market DES until after the time in which plaintiff's mother purchased the drug. Id.

A California trial court interpreting Sindell has ruled that the "appropriate market" is the smallest market that fairly covers the plaintiff's involvement with the subject drug. Den Daas v. Boyle Drug Co., No. 73275 (Cal. Super. Ct., Ventura County, Oct. 1, 1984) (order determining issues). In this case, the court held that the appropriate market was the military market during the years 1955 through 1958.
market shares.\textsuperscript{90} The Sindell court did not even define the term "substantial share,"\textsuperscript{91} giving rise to a host of conflicting suggestions. It has been suggested that the substantial share requirement exists so that "more likely than not the culpable party is one of the defendants and will not escape all responsibility, and, therefore, substantial share ought to mean more than 50\% of the market."\textsuperscript{92} The commentator cited by the Sindell court suggested that 75 to 80\% of the market must be joined;\textsuperscript{93} a California trial court interpreting Sindell has held that a "substantial share" of the market is a minimum of 51\% of the relevant market, but that the percentage may be increased upon an appropriate showing in a particular case.\textsuperscript{94}

Perhaps because the Sindell theory has so many unresolved issues and remains only partially developed by the California courts, it has been widely rejected outside of California.\textsuperscript{95} Both courts and commentators have recognized that the market-share approach is not suitable for product liability cases where the products or the alleged defects are not identical.\textsuperscript{96}

The theory also has generated widespread criticism on related policy
grounds. It has been criticized as encouraging plaintiffs to perjure themselves by concealing evidence as to the identity of the manufacturers responsible for their injuries, where the companies have since gone out of business or are judgment-proof, in order to sue under the market-share theory.97 It also has been pointed out that imposing the market-share theory in a strict liability tort case "gives rise to a form of absolute liability by relieving the plaintiff of proving defendant's breach of duty and by guaranteeing plaintiff's proof of causation," which "forces an industry into the position of an insurer" of a product.98 The theory has also been criticized as not serving the goal of economic efficiency. By apportioning damages throughout an industry solely on the basis of market shares and irrespective of safety efforts, it enables unsafe manufacturers to spread the burden of their accident costs and thereby creates disincentives for safety. And as pointed out above, the theory raises formidable problems of both fairness to defendants and judicial administration by increasing the number of defendants in the courtroom, placing the burden of identification on the defendants without regard to the relative abilities of the parties to identify the actual responsible producer, and embroiling the court in the complexities of defining the relevant market.99

In short, the Sindell approach, while solving some problems, leaves a host of others unanswered. Not surprisingly, courts have further modified the Summers approach as glossed by Sindell—with improved, albeit still inadequate, results.

E. Variations on Sindell Market-Share Theory

1. Market-Share Alternative Liability

In the recent case of Martin v. Abbott Laboratories,100 the Washington Supreme Court adopted a modified version of the Sindell market-share approach, rejecting the Sindell theory because of "its inherent distortion of liability."101 This modified Sindell theory permits a plaintiff to bring suit against a single drug company or several drug companies by alleging the following elements: (1) plaintiff's mother took DES; (2) DES caused the plaintiff's subsequent injuries; (3) defendant produced or marketed the type of DES taken by the plaintiff's mother," ("e.g., dosage, color, shape, markings, size, or other identifiable characteristics"), and

97. Note, supra note 96, at 676.
100. 689 P.2d 368 (Wash. 1984).
101. Id. at 380.
(4) defendant's conduct in producing or marketing DES constituted a breach of a legally recognized duty to the plaintiff. Moreover, "the plaintiff need not allege or prove any facts related to the time or geographic distribution of the subject DES."

Under this theory, defendants may escape liability by showing that they did not manufacture the type of DES taken by the plaintiff's mother or did not market DES at the relevant time or in the relevant geographic area. Those defendants who cannot exculpate themselves will be presumed, initially, to have equal market shares, and will be liable in equal shares for the plaintiff's total damages. Each defendant may reduce its liability by proving that its actual market share is less than its presumed share. The defendants who are not able to produce evidence of their actual market shares will be liable for the remainder of the plaintiff's damages in excess of the other defendants' actual market shares.

The court illustrated how this rule of apportionment would work in a hypothetical case where the plaintiff's damages are $100,000. If the defendants, X and Y, are unable to establish their actual market shares, each will be liable for 50% of the judgment. If X, on the other hand, establishes that its actual share of the relevant market is 20%, X is liable for 20% of the plaintiff's damages and Y is liable for the remaining 80%. If, however, X establishes that its actual market share is 20% and Y establishes that its actual market share is 60%, then X is liable for 20% of the damages and Y is liable for 60%. In this situation, the plaintiff will not recover 100% of her damages because the remaining 20% of the market consists of manufacturers who were not joined in the suit. In contrast to Sindell, under which it is unclear whether the plaintiff recovers 100% of her damages, this market-share alternative liability theory awards a plaintiff 100% of her damages unless all the named defendants are able to prove their actual market shares. Further, "defendants may implead third party defendants in order to reduce their presumptive share of the market or in order to establish an actual reduced market share."

The Martin court specifically rejected the Sindell requirement that the plaintiff join defendants representing a "substantial" share of the market. The court stated that "[n]ot only does the Sindell court fail to define 'substantial' share of the relevant market, the theory distorts market liability by providing that the 'substantial' market share bears joint

---

102. Id. at 382.
103. Id.
104. Id. at 382-83.
105. Id. at 383.
106. Id.
107. Id. at 382.
responsibility for 100 percent of plaintiff's injuries."\textsuperscript{108}

The court justified its theory in part on the basis that all the defendants contributed to the risk of injury, even though they might not have contributed to the actual injury of a given plaintiff.\textsuperscript{109} Furthermore, the court assumed that a "drug company is in a better position to absorb the cost of the injury" because it "can either insure itself against liability, absorb the damage award, or pass the cost along to the consuming public as a cost of doing business."\textsuperscript{110} And the court pointed out that while "elimination of individual causal responsibility as an element of plaintiff's case is liability enhancing," the apportionment of damages according to respective market shares and the possibility that the plaintiff might not recover her entire damages if all defendants can prove their actual market shares is "liability limiting."\textsuperscript{111}

The Martin approach refines the Sindell market-share theory and provides a fairer result. It is based on the presumption that DES manufacturers should pay damages because they contributed to the risk of injury to the public and not because they are actually responsible for a particular plaintiff's injuries. Aside from its inconsistency with traditional principles of tort law, the Martin theory leads to a fairer result than the other theories discussed above. It clearly establishes that the defendants will be severally liable, which ensures that the damages they pay will be limited to the risks they actually caused. It permits the defendants to exonerate themselves, which ensures that the defendants who clearly are not responsible for a plaintiff's harm will not pay damages.

On the other hand, the lack of a requirement that the plaintiff join the defendants representing a substantial share of the market creates the possibility that the joined defendants who are unable to prove their actual market shares will be forced to pay damages greatly disproportionate to their contribution to the risk of injury to the public. In addition, problems in judicial administration remain. The court did not define the relevant market for the purpose of apportioning market shares, either in terms of time or geography. Moreover, while a defendant is permitted to produce evidence to reduce its presumptive market share, the court did not indicate whether a plaintiff may produce evidence showing that a defendant's actual market share is greater than its presumptive market share. Furthermore, the theory raises a possibility that a plaintiff will recover less than 100% of her damages.

Most significantly, the presumption that manufacturers should pay

\textsuperscript{108} Id. at 381.
\textsuperscript{109} Id. at 382.
\textsuperscript{110} Id.
\textsuperscript{111} Id. at 383.
damages for contributing to the risk of injury to the public overrides the basic principle of tort law that one who is responsible for causing a harm should pay for the resulting injury.112 Creating a risk that does not result in specific damage is simply not a tort. This type of conduct, when wrongful, is more in the bailiwick of criminal law, as with a drunk driver whose recklessness does not result in an injury.

2. Risk Contribution Theory

In Collins v. Eli Lilly Co.,113 the Wisconsin Supreme Court rejected the plaintiff's proposed theories of alternative liability, concerted action, enterprise liability, and market-share liability, and framed a new method of recovery for DES plaintiffs in Wisconsin, purportedly based on the state's existing comparative negligence laws. The court noted that this method of recovery could apply to other product liability actions in situations which are factually similar to DES cases.114 The plaintiff must allege the following elements: (1) plaintiff's mother took DES; (2) DES caused the plaintiff's subsequent injuries; (3) defendant produced or marketed the type of DES taken by the plaintiff's mother (e.g., color, shape, markings, size, or other identifiable characteristics); and (4) defendant's conduct in producing or marketing the DES constituted a breach of a legally recognized duty to the plaintiff.115 With respect to the third element, the plaintiff may satisfy the requirement even if she cannot prove what type of DES her mother took; she need only "prove that the defendant . . . produced or marketed the drug DES for use in preventing miscarriages during pregnancy."116

If the plaintiff is able to prove these four elements, she may recover her entire damages from a single defendant. In this key respect, the Collins rule is tougher for defendants than the Sindell rule.117 If, however, the plaintiff joins more than one drug company, the damages will be apportioned among the defendants by the jury. In determining the percentage of liability to assign to each defendant, the jury may consider a

112. See authorities cited supra note 2.
113. 116 Wis. 2d 166, 342 N.W.2d 37 (1984).
114. Id. at 181, 342 N.W.2d at 49.
115. Id. at 193, 342 N.W.2d at 50. A plaintiff may proceed under theories of negligence and strict liability. Under a negligence theory, the plaintiff will have to prove that a drug company breached a duty of care when it made or sold DES. Under a strict liability theory, the plaintiff must show that the DES was defective, that it was unreasonably dangerous, that the defect was the cause of plaintiff's injuries, that the company engaged in making or selling the drug, and that the company expected the DES to reach the consumer without substantial change. Id. at 195-96, 342 N.W.2d at 51.
116. Id. at 193-94, 342 N.W.2d at 50.
117. Id. at 194, 342 N.W.2d at 50-51. "[A]ny defendant may . . . implead as third-party defendants other drug companies which it can allege produced or marketed the type of DES taken by the plaintiff's mother." Id. at 195, 342 N.W.2d at 51.
number of factors, one of which is whether the defendant had a small or large share of the relevant market. The other factors include, but are not limited to, the following:

- whether the drug company conducted tests on DES for safety and efficacy in a use for pregnancies; to what degree the company took a role in gaining FDA approval of DES for use in pregnancies; . . . whether the company took the lead or merely followed the lead of others in producing or marketing DES; whether the company issued warnings about the dangers of DES; whether the company produced or marketed DES after it knew or should have known of the possible hazards DES presented to the public; and whether the company took any affirmative steps to reduce the risk of injury to the public.\(^{118}\)

The court noted that a "DES plaintiff, who was a fetus at the time the DES was taken, could [not] be attributed with . . . negligence," and "that in all DES cases [presumably] 100 percent of the liability will be apportioned only among the defendants."\(^{119}\)

The Collins theory is even less fair to defendants than the other theories discussed above. A manufacturer who is the sole defendant in an action will pay damages far in excess of its share of the market and far in excess of the risk of injury it created. Moreover, it will incur all the litigation costs in defending the case. Under this theory, a manufacturer who did not manufacture the actual injury-inflicting drug could be forced to pay 100\% of the plaintiff's damages as well as its own transaction costs for the action. Unless the manufacturer impleads other drug companies that it can allege produced the type of DES ingested by the plaintiff's mother, it will be liable for the plaintiff's entire damages. The procedure seems unfair particularly because the plaintiff, who is in a better position to prove the type of DES ingested by her mother, need only prove that the defendants she joins produced or marketed DES for use in pregnancy problems. Thus, the plaintiff easily could join a number of defendants. Once again, the court's attempt to provide compensation to an innocent plaintiff demonstrates that a judicial scheme for recovery in DES cases may be developed only at the cost of unfairness to defendants.

III

A Framework for a Legislative Solution to the Identification Problem

The above analysis of the various theories courts have developed to provide compensation for persons suffering injuries allegedly caused by

\(^{118}\) *Id.* at 200, 342 N.W.2d at 53.

\(^{119}\) *Id.* at 200 n.14, 342 N.W.2d at 53 n.14. The court stated that punitive damages may not be awarded to a plaintiff who recovers under this theory, however, because the concept of punitive damages is based on the notion "that the wrongdoer being punished because of his conduct actually caused the plaintiff's injuries." *Id.* at 202, 342 N.W.2d at 54.
DES shows that the judicial system is unable adequately to resolve the DES identification problem. In many cases, particularly those involving clear-cell adenocarcinoma, the plaintiffs can allege all elements of a tort cause of action except the identity of the actual wrongdoer. Some courts appear to be motivated by a desire to "compensate" injured persons but are restrained by the traditional rules that impose liability only on those who are the cause-in-fact of a harm. To the extent that courts have overridden the traditional rules, problems of fairness to defendants have arisen.

The majority of courts have followed the traditional approach, requiring a DES plaintiff to produce evidence of the identity of the manufacturer of the particular pills that allegedly caused her injury. For instance, courts in Missouri, South Carolina, Texas, New Jersey, Florida, Massachusetts, Connecticut, and the District of Columbia have denied recovery to plaintiffs who could not identify the manufacturer of the DES that allegedly caused their injuries. The courts that have permitted recovery for the plaintiffs who are unable to identify the manufacturer of the drug have reached outside the traditional tort system to do so. They have modified traditional joint liability theories and have developed new theories, all of which conflict with the basic tort notion that one who is responsible for causing a harm should pay. On their faces, these theories are unfair to defendants. In addition, the theories raise substantial problems of judicial administration. Theories such as the Sindell market-share theory and the Martin market-share alternative liability theory were not fully developed or explained by the courts, creating many problems for other courts wishing to follow these approaches. Furthermore, application of these theories involves practical problems in apportioning damages according to the defendants' market shares. The process of determining the relevant market and the defendants' market shares in individual cases will involve lengthy litigation and huge transaction costs for all parties.

A number of commentators have recognized the inability of the tort litigation system to resolve the identification problem and have advocated various alternative systems of recovery. The alternatives pro-

---


121. See, e.g., Biebel, DES Litigation and the Problem of Causation, 51 INS. COUNS. J. 223 (1984); Delgado, Beyond Sindell: Relaxation of Cause-In-Fact Rules for Indeterminate Plaintiffs, 70 CALIF. L. REV. 881 (1982); Downey & Gulley, Theories of Recovery for DES Damage: Is Tort Liability the Answer?, 4 J. LEGAL MED. 167 (1983); Fischer, Products Liability—An Analysis of
posed include legislative and administrative schemes, such as a limited no-fault product liability fund for the plaintiffs unable to identify the manufacturer of a generic product that produced a latent injury, suits against the federal agency responsible for regulating the particular industry using the Federal Tort Claims Act and the Administrative Procedure Act; ad hoc congressional responses to mass injuries caused by products of unidentifiable manufacturers; legislation designed to hold certain industries liable through trade associations for all injuries caused by those industries' products whenever the manufacturer of an injury-causing product is not identifiable; a no-fault compensation system for persons injured by DES which would be funded by a tax imposed upon all manufacturers who produced DES for use as a miscarriage preventative; and a toxic tort compensation system not limited to a single industry or a single type of product-injury but designed to deal with the toxic tort problem as a whole.

Ideally, the identification problem in DES cases should be solved by legislation. The legislation's goal should be to provide a speedy and efficient process for dealing with claims where the injured person cannot identify the manufacturer of the allegedly injury-causing product. It should solve the problems created by the court-made solutions to the nonidentification cases, while providing fair and adequate compensation to injured persons. Before examining the broad framework of a legislative solution to accomplish these two goals, however, it is necessary to define precisely the limited scope of a solution to the DES problem.

A. Limiting the Scope of a Legislative Solution to the Identification Problem

A legislative system should serve only as an alternative method of providing compensation to injured persons who are unable to prove the identity of the responsible manufacturer. It is important that the legislative solution not preempt tort suits where identification is possible. It would be unfair to require nonresponsible manufacturers to bear the burden of compensating injured persons, through the legislative system, in cases where the responsible manufacturer can be identified. Unfairness also would result if the legislative solution were to permit manufacturers responsible for causing harms to escape tort liability. Furthermore, it would be administratively inefficient to overload the legislative compen-
sation system with cases that currently are handled appropriately by the judicial tort system.

To accomplish this important goal, legislation could encourage those persons who truly are able to identify the manufacturer to do so by requiring identification of the manufacturer for recovery under traditional tort principles. Thus, persons who can prove the identity of the responsible manufacturer could sue that manufacturer for tort damages, including pain and suffering. Where persons are unable to prove the identity of the responsible manufacturer (i.e., “nonidentification claims”), they could be compensated under the legislative system for out-of-pocket losses. In nonidentification cases, the claimant should be required to show fault, causation, damages, and a good-faith, genuine attempt to identify the manufacturer. Proof of these elements could trigger compensation through the legislative mechanism. Moreover, the legislation should impose penalties on claimants and unscrupulous counsel who falsely identify a defendant. This may occur because a greater recovery would be available to a claimant who can make a proper identification. If a court determines that a false identification has been filed, the defendants’ legal costs should be borne by the plaintiff and, in appropriate cases, a civil fine should be imposed on the plaintiff’s counsel.

A separate but related problem exists in cases where the claimant can identify a defendant but is unable to recover in the tort system because the defendant is insolvent or judgment-proof. These claimants may be denied any recovery, while claimants who cannot identify a defendant may recover under the legislative solution. The legislative solution should address this issue to avoid this unfair result and, again, to forestall false claims. One solution may be to permit these claimants to recover under the legislative system for out-of-pocket losses.

Moreover, legislative energy should be focused on addressing the identification problem. The identification requirement presents a unique problem in cases involving latent injuries linked to a generic product such as DES. The elements of fault and causation are not peculiar to these cases. Thus, a basic premise of the legislative solution must be that an injured person is required to prove that the product actually caused his or her injury.

In addition, and also in connection with the scope of the legislative solution, consideration should be given to the amount of payment to be received by claimant. The system should provide a smooth and efficient means whereby persons who (in most jurisdictions) are excluded from the possibility of payment would receive an award. Even in jurisdictions that currently permit such persons to secure an award under the tort system, delays and other deficiencies would be mitigated by a legislative solution. The system is “standby equipment” for the tort system, a vehi-
cle to provide relief where the tort litigation system fails. The damage award should reflect that fact and be limited to the claimant's true excess economic losses. This would include amounts lost and not otherwise recovered through collateral sources, but would not include damages for pain and suffering.

Finally, careful consideration should be given to how claimants enter the system. It is contemplated that the system would serve only those who have a problem identifying the defendant who made the product; individuals with problems relating to basic causation (i.e., whether the product caused the injury) or nonliability (i.e., whether the defendant, if he can be identified, owed no duty to the plaintiff) would be excluded. The simplest approach is to let the current court system handle these matters, but they are part of an area that must be addressed.

B. The Legislative Framework

Given the narrow focus of an appropriate legislative solution, it is important to sketch a framework for construction of the solution. First, let us quickly recall the problems such a solution can face, as revealed by the courts' difficulties in fashioning a common law solution to the DES problem. The difficulties presented by the judicial theories create three undesirable results—unfairness to defendants, incomplete compensation for the plaintiffs, and inefficiencies in judicial administration. The following list summarizes the most significant problems of the judicial theories and their adverse results:

1. Alternative liability theory, Sindell theory, Martin theory, Collins theory: Not all parties who may have been responsible for the plaintiff's harm are required to be joined as defendants. Thus, the responsible party may escape liability and the joined defendants may be forced to pay damages even though none of them caused the plaintiff's harm. Result: Unfairness to some joined defendants and imperfect deterrence of tortious conduct.
2. Alternative liability theory: The burden of proving identification is shifted to the defendants. In DES cases, the defendants are not in a better position to provide all relevant evidence. As to some of this information, it would be fairer and more efficient to place the burden of proving identification on the plaintiff, who has better access to the type of drug taken by her mother, her mother's geographic location at the time of ingestion, the location, etc. Result: Unfairness to the defendants, inefficiency in judicial administration, and impairment of the deterrence function.
3. Alternative liability theory: The theory presupposes that all defendants have acted negligently toward the plaintiff and thus
shifts the burden of proof unfairly. Manufacturers of drugs not taken by the plaintiff's mother have not acted negligently toward the plaintiff, so no shift in the burden of proof should follow. Result: Unfairness to defendants and imperfect deterrence of tortious behavior.

4. Concert of action theory: The theory applies only where there has been a tacit agreement to act wrongfully and, thus, the parties to the agreement are joint tortfeasors and all are responsible for the acts of the others. There was no tacit agreement among the manufacturers of DES. Accordingly, one manufacturer is not responsible for the acts of the others. Result: Unfairness to defendants.

5. Enterprise liability theory: There are no centralized industry standards and, thus, no basis for enterprise liability on a theory of joint control through uniform industry standards. Result: Unfairness to defendants.

6. Sindell theory: Appointment of damages throughout an industry solely on the basis of market shares, irrespective of safety efforts, is economically inefficient and puts those manufacturers who have made little or no effort to market a safe product in the same position as those who did make safety efforts. Result: Unfairness to some defendants and impairment of the deterrence function.

7. Sindell theory: The requirement that the plaintiff join a substantial share of the market increases the number of defendants in the courtroom in each case. This increases administrative costs. Result: Inefficient judicial administration.

8. Sindell theory, Martin theory, Collins theory: The definitions of substantial share and the relevant market are not clear. Thus, the court is forced to become embroiled in the complexities of defining these terms in each case. Result: Inefficient judicial administration.

How might a legislative solution resolve these difficulties? Problems 4 and 5 are insoluble, because they indicate that the concert of action and enterprise liability theories do not justify imposing liability. Thus, the only justifiable forms of recovery are those within the confines of Summers, and the modifications thereof.

Problem 1 could be solved if all defendants could be identified in advance of litigation. A legislative solution could achieve this result through a one-time centralized set of findings of fact.

Problems 2 and 3 could be solved if the burden of producing evidence were placed on defendants only when information was in their control, and only if and to the extent that the defendants had breached at
the very least some duty of care to someone. This could be accomplished legislatively by predetermining which issues will be resolved on a centralized basis among defendants, and by apportioning costs of administration among defendants in proportion to their tortious behavior and market shares. Problems 6, 7, and 8 would also be solved by such a centralized set of findings of fact and a liability and market-share apportionment of administrative costs. Thus, if a legislative solution adheres to the basic guidelines relating to centralized fact-finding and apportionment of costs, it could achieve what each of these theories of recovery attempts to achieve, but in fact achieves only imperfectly.

With the lessons of the common law solutions clearly in mind, let us proceed to delineate a set of guidelines for a legislative solution to the DES problem.

First and foremost, careful consideration should be given to whether, how, and to what extent legislation will require manufacturers to contribute to compensation paid to plaintiffs. To require all manufacturers of DES (or the product in question) to contribute to the compensation paid to plaintiffs in nonidentification cases would be fairer than the random targeting of defendants that occurs under current judicial theories. A major criticism of courts’ approaches is that they impose liability on manufacturers that are not responsible for causing a plaintiff’s harm. There are over 300 DES manufacturers who may have been responsible, yet the judicial theories impose liability only on the handful of manufacturers that have been singled out as defendants. The defendants may be no more responsible than other DES manufacturers who were not joined in a lawsuit, yet the defendants are forced to pay damages. The unfairness results because some nonresponsible manufacturers pay damages and legal costs while other responsible manufacturers do not. Requiring joinder of all DES manufacturers in every court case would often be impossible, would always be inefficient, would impose great burdens on plaintiffs, and would create problems in judicial administration. A legislative solution, on the other hand, could avoid both an unfair result and problems in inefficiency of administration because it could require payment of nonidentification claims by all DES manufacturers.

Thus, consideration should be given to whether contribution to compensation for plaintiffs should be made solely by industry or also by government. Some commentators have called for development of compensatory funds to be financed by special taxes on an industry or by the federal government, or have called for trade associations to establish their own compensation funds through a self-imposed excise tax. The legislative approach recommended in this Article, however, focuses on

---

122. See, e.g., Note, Legislative Alternatives, supra note 80, at 1037-42.
payment of compensation to claimants, either directly or indirectly, by
the manufacturers themselves, thus furthering the cause of effective
deterrence of (and where appropriate, penalty for) tortious behavior. In
sum, the source of payment is a key issue that legislation drafters must
resolve. If there is active participation in the development of a drug by
the government, serious consideration should also be given to utilizing
the tax base as a part of the compensation fund. In such a situation,
responsibility is, in part, public.

Assuming responsibility for causing a claimant's harm would not be
the sole basis for requiring payment, the legislation could rely on some
other fair basis for requiring all DES manufacturers to pay compensation
in nonidentification cases. One possible solution is to require all manu-
facturers of DES to make contributions in each nonidentification case.
Alternatively, the legislation could require periodic payments by all DES
manufacturers to a central account or fund that would distribute com-
ensation to successful claimants in nonidentification cases. Assessments
against all DES manufacturers in nonidentification cases could be based
on some combination of the market-share theories developed by the
courts. For example, legislation could apportion compensation claims
among all manufacturers of DES based on a number of factors, including
the following: (1) the manufacturer's share of the market for the prod-
uct, (2) the degree of the manufacturer's culpability, (3) the manufac-
turer's good faith or bad faith in remedying the product and in
responding to claims.123

Second, the above discussion of the market-share theories applied by
courts concluded that consideration of market share and other factors in
individual cases is inefficient and creates problems in judicial administra-
tion.124 Furthermore, the judicial market-share theories require proof of
the relevant market and joinder of a substantial share of the relevant
market. Legislation requiring apportionment of claims in nonidentifica-
tion cases according to market shares and other criteria, however, can
and should strive to reduce these problems. Specific guidelines in the
legislation itself for defining the relevant market or markets would
resolve that issue; the question of what constitutes a "substantial share"

123. One commentator has suggested a "latent technological injury compensation" system for
plaintiffs who suffer from injuries not by their nature discoverable within the running of the statute
of limitations. These plaintiffs would apply for relief to an administrative tribunal and be compen-
sated by funds from a tax paid by all (presumably by all DES) manufacturers. Note, Industry-Wide
Liability, supra note 121, at 1015-22. This solution suffers from a defect of the common law solu-
tions: it imposes a tax on all DES manufacturers regardless of market share, fault, or responsiveness
to claims. For further criticisms of this scheme, see Note, Legislative Alternatives, supra note 81, at
1023-24. (The note proposes a legislative solution requiring a trade association-funded compensation
pool, id. at 1037-42, which suffers from the same defect).
124. See supra notes 67-119 and accompanying text.
of the market would be immaterial because the legislation would apportion compensation claims among all manufacturers in the relevant market or markets.

A further problem inherent in the judicial market-share theories is the litigation of market shares in every case. Under a legislative solution, however, the apportionments for the relevant market or markets need not be made repeatedly in each nonidentification case but could be made only once or on a defined schedule on the basis of administrative or arbitration procedures. Any acceptable legislation must develop and set forth carefully drafted provisions embodying such procedures. The legislation should require participation in the hearings by all manufacturers in the relevant market or markets. Consideration should be given to whether a method could be developed for nonadversarial hearings; the hearings’ sole purpose could be to receive evidence on the factors that will determine the apportionments. In situations where the manufacturers are in a better position to prove the factors relevant to determining apportionment, it would be appropriate for the manufacturers to do so. Provisions could be included in the legislation to permit modification of the apportionments if new evidence emerges at a later time. Such administrative or arbitration procedures could of course be displaced by a suitably structured, suitably publicized civil quasi-class action—another possibility for a legislature to consider.

The legislation could also establish a mechanism to make assessments on manufacturers based on the apportionments developed in such hearings that would provide a method for transferring payment to claimants in nonidentification cases. This could be accomplished by making assessments, in accordance with the apportionments developed in the hearings, for each nonidentification case. On the other hand, a careful legislative examination of the facts may show that it would be best to have the assessments made periodically and transferred to an account or fund for distribution to individual claimants.

A fourth, related guideline for a legislative solution is that the legislation should fairly apportion transaction costs among all manufacturers of a product. The legislation should address the issue of when, how, and by whom counsel participating in the system should be paid. This aspect of the legislation focuses primarily on the role of counsel representing defendants. The legislation should provide a method for spreading the costs of nonidentification cases among all manufacturers of the product. It would be inefficient to require joinder of all manufacturers of a product in every lawsuit where the plaintiff is unable to identify the manufacturer. Thus, the legislation should include a set of provisions that address the issue of who should defend the action (for the purpose of litigating causation, fault, and damages) when a plaintiff is unable to
prove that one of the named defendants manufactured the product alleged to have caused her injury. Defense counsel that litigate these issues should be paid through the legislative system.

Assessments for litigation costs could be made on all manufacturers of the product in some fair way, perhaps based upon the apportionments made in the centralized findings of fact. Counsel's rates should be computed in proportion to the rates in the market in which they operate. Counsel should be required to keep detailed hourly records subject to review at the discretion of judges. A staff of defense counsel representing the legislative compensation system might be set up that would, in effect, represent all manufacturers of the product. Another possibility is the development of a method for coordination of counsel who represent the named manufacturers in nonidentification cases.

C. Other Legislative Considerations

A few other problems and considerations, of a lesser degree of importance than these first four guidelines, should be mentioned here. First, any proposed legislation must address the problem of how current insurance coverage for manufacturers under the present tort litigation system can be transferred and (in effect) captured by the new legislative system. Although this is an extremely complex problem with little precedent, the goal is clear: the legislation should contain a provision that all insurance coverage applicable in litigation and settlement of current DES cases should apply to manufacturers' payments of assessments or compensation claim apportionments under the legislative system. Similar provisions have been included in other types of legislation, such as Senate Bill 2708 in the 98th Congress, which proposed an asbestos compensation fund.125 The legislative solution would be fair to manufacturers only if it ensured that manufacturers retain the insurance coverage that they have paid for and are entitled to. Of course, the legislation should not retroactively create broader coverages for carriers beyond the contractual agreement between insurer and policyholder. In sum, it is essential to develop the application of present insurance coverage in legislation that facilitates the payment of claims in nonidentification cases.

Second, consideration must be given to the role and method of payment to claimant's counsel. The normal contingent fee approach may not work well in this context. To assure fair payment, but also to minimize transaction costs, claimant's counsel should be required to keep records of their time and services, which should be reviewed before payment is made.

Third, careful consideration should be given to whether legislation

should be state or federal. A persuasive argument can be developed that the identification cases are still in the experimental stage. The well-worn argument of Justice Brandeis that the states should be given an opportunity to act as "little laboratories" to address new problems is particularly persuasive here. Unlike the situation with respect to other areas of general product liability—such as standards for design and warning, or punitive damages—the identification problem is still new and solutions are in the experimental stages. Also, unlike federal product liability legislation, the legislation in this area involves both the creation of new mechanisms for apportionment of responsibility and, perhaps, facts and issues that are not common to product liability cases in general. The most appropriate resolution to these problems may be best left to the states. On the other hand, the products involved have been distributed in interstate commerce. It may be best to have one uniform method of apportioning responsibility. Finally, it would not be desirable to encourage forum shopping by plaintiffs. In sum, two things are clear: this is a matter that deserves much more study, and it is a problem distinct from the current needs for, and development of, uniform product liability law in more traditional areas of litigation.126

Finally, there must be a determination as to whether the legislation is to be product-specific or should address the more general problem of nonidentification. As this Article has demonstrated, DES litigation has a unique history and background. In structuring a legislative system addressing the identification problem, this history and background must be considered. On the other hand, a wholly separate approach for nonidentification problems that have some similarities (i.e., asbestos) may be inefficient and wasteful. This is a problem that can be most efficiently explored by legislative study and examination.

There are many other issues regarding the specific provisions of a legislative solution for the identification problem: the forum for filing and hearing nonidentification claims, the mechanism for transferring payment from manufacturers to nonidentification claimants, hearings for determining apportionment of compensation in nonidentification claims, and so on. This Article has suggested the basic role and structure of a legislative solution. Development of the specific provisions for implementation of a legislative solution requires further careful study and consideration.

CONCLUSION

The case in which a person injured by a product can prove that his

or her injuries result from culpable conduct of the manufacturer but is unable to identify the manufacturer is a peculiar problem requiring a unique legislative solution. Courts have attempted to develop solutions in order to provide recovery to plaintiffs in litigation, but in overriding and modifying the tort system, they have created needless and extensive litigation costs. Furthermore, these judicially imposed approaches do not facilitate prompt and appropriate recoveries for injured persons and are unfair to defendants.

The general problem can be solved by legislation that would promptly provide out-of-pocket losses to injured persons. Those persons should nevertheless be required to prove all other elements of a product liability action. Such a legislative system should balance the interests of injured persons and manufacturers. The legislature, through the hearing process, is well equipped to do this. In this Article, we have attempted to present some of the major considerations and alternatives a legislature should address if it attempts to venture down this path. While these problems are not easy to resolve, their very nature demonstrates the appropriate forum in which they ought be addressed. The legislature, if it follows these guidelines and parameters, can resolve a problem which the judicial system is not equipped or designed to handle.