Daubert Asks the Right Questions: Now Appellate Courts Should Help Find the Right Answers

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1 Daubert v. Merrell Dow Pharm., 509 U.S. 579, 588 (1993) (noting that FRE 702 superseded the Frye standard; under FRE 702, scientific evidence must be valid in the sense of being reliable and must “fit” the case and, even if the evidence does satisfy these requirements, it is subject to the possibility of exclusion under FRE 403) (citing Frye v. United States, 293 F. 1013, 1014 (D.C. Cir. 1923) (adopting what has come to be known as the “general acceptance” standard)).
practice," or "lacks sufficient basis in fact" or "lacks sufficient connection to the case at hand." A post-Daubert judge who admits such evidence is expected to say "the court finds that indeed this testimony is properly grounded in theory and practice, and adequately based on the facts and sufficiently related to the task at hand." To be sure, Daubert still leaves room to hide. Factors like "peer review" and "general acceptance" provide opportunities, as does the possibility of invoking FRE 403, and a judge can also distance himself by casting his decision in terms of "adequate assurances" or "inadequate assurances" of validity.3

The basic point, however, is that Daubert puts judges into the position of judging science. That makes Daubert revolutionary.4

Criticisms of Daubert abound, particularly in toxic tort cases. Perhaps such criticisms are inevitable when a single case so profoundly changes the legal landscape. The most serious criticisms are advanced from three perspectives: One is epistemological and structural (or "political" in the fine sense of the term). This criticism holds that judges are not much more able than juries to appraise proof offered as science, and that attempts to exercise the "gatekeeping" role infringe on the powers and responsibilities of juries to act as factfinders. Another criticism is pragmatic, substantive, and to some extent ideological. This criticism holds that judges applying Daubert are throwing out too much good evidence proffered by civil claimants in toxic tort cases. A third is philosophical. This criticism holds that Daubert misunderstands science.

What follows is a defense of Daubert against these criticisms, followed by a suggestion of my own, which is that the Daubert revolution would achieve more if appellate courts abandoned the

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2 For an argument that "peer review" is not what it is cracked up to be, see Joelle Anne Moreno, Eyes Wide Shut: Hidden Problems and Future Consequences of the Fact-Based Reliability Standard, 34 SETON HALL L. REV. ___ (upcoming in Fall 2003).


4 One crude measure of Daubert's impact can be seen just by glancing at citation history. In the 38 years between 1945 and the decision in Daubert in 1993, Frye was cited in approximately 260 reported federal cases and 800 reported state cases. As of this writing, Daubert is almost ten years old. As of January 25, 2003, Daubert has been cited in nearly 2,500 reported federal cases and 1,200 reported state cases, while Frye has been cited in approximately 270 reported federal cases and 1,000 reported state cases.
abuse-of-discretion standard in reviewing the rulings of trial judges in this area. To be fair, I should note that most critics have not called for the abandonment of Daubert and few would endorse a return to the Frye standard. Casting the criticisms in their best light, their aim is to improve Daubert, an undertaking that I gladly join. This essay addresses civil rather than criminal cases, and scientific evidence rather than "experiential expertise," even though this dichotomy is hard to draw and counts for less than it once did because the Daubert standard applies to all expertise.

1. JUDGES CAN DO BETTER THAN JURIES: DAUBERT GATEKEEPING MADE REAL

In a nutshell, Daubert is the right standard because it asks directly the question that Frye put only indirectly, and thus puts courts in a better position to arrive at satisfactory answers. The central issue is scientific "validity," and the criteria suggested by Daubert are useful in resolving that issue.

Here it is worth pausing to ask some pragmatic questions: Why have a validity standard to begin with? Why not simply approach science with the kind of openness suggested by FRE 702 on its face? In other words, why not simply admit scientific evidence if it seems relevant and helpful and the witness is qualified?

The answer given by the Court is more positivist than policy-based. FRE 702 requires science to satisfy a validity standard, so courts are bound to scrutinize such proof. That answer is unsatisfactory because it does not emerge from the "plain meaning" of FRE 702 or even a reasonable interpretation of the Rule’s language. The Court has acknowledged that the Rules did not displace all prior evidence.
doctrine, and this point is critical to another major holding. The conclusion in *Daubert* rests on the notion that the word "scientific" as used in FRE 702 is a rich or deep normative term that implies a standard of legitimacy. It is of course astonishing, if we suppose that this meaning really is to be found in FRE 702, that nothing in legislative background supports this reading (in fact the term seems merely descriptive). In truth, the Rules provide no compelling basis for discarding the old *Frye* standard. What we now call the *Daubert* standard is in reality judge-made law disguised as something else. That is not to say that I disapprove of the decision, for the opposite is true: I think *Daubert* represents an advance, that it is at least consistent with the elastic contours of FRE 702, and that it good lawmaking, even if disingenuous in its logic.

But there are questions that should be asked before reaching that conclusion: Should we have such a standard? Does it make the law better? Keeping the focus on civil cases, my answer is yes.

I think that three facts of modern life conspire to suggest that we need a validity standard: First, we ask courts to resolve difficult technical and scientific issues. Second, much scientific knowledge is fluid and contestable, inaccessible to laypeople, hard to understand, and qualified in ways that elude ordinary experience and intuitions. Third, our adversary system places primary responsibility for gathering and presenting evidence in the hands of the parties, and creates incentives that lead to risks.

What we see is something like this: Complex questions arise, which can be answered, if at all, only by calling on scientific expertise. A salient modern example is the question of causation in the toxic tort setting. Parties and courts look to science for the answer. Lawyers on each side find experts who agree to help for a price. The issue is joined, and we discover that there is no definitive answer and only partial information—studies and theories, usually involving

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7 See United States v. Abel, 469 U.S. 45, 50 (1984) (explaining that the drafters of the Rules could not have intended to "scuttle entirely" the practice of cross-examining to show bias, even though they offer no "express treatment" of the subject); see also Edward Cleary, *Preliminary Notes on Reading the Rules of Evidence, 57 Nev. L. Rev. 908, 915 (1978*) (stating that after adoption of the Rules, "no common law of evidence remains," at least "in principle," but "in reality" the situation is different, because "the body of common law knowledge continues to exist" in the "somewhat altered form of a source of guidance in the exercise of delegated powers") (internal citations omitted).

8 See generally RONALD DWORKIN, LAW'S EMPIRE 255 (1986) (contrasting, *inter alia*, law as pragmatism with law as integrity, meaning that judges seek to resolve hard cases by some "coherent set of principles" in order to make the "complex structure" of law and politics "the best these can be").
some combination of chemical structure analysis, animal tests, epidemiology, and/or “differential diagnosis.” None of the proof either does or can answer directly the question of individual causation (“specific” causation). Instead, such proof shows a possibility, and perhaps sometimes a probability, of causation in an individual case by suggesting that the substance in question can or does cause some ailments in some people (“general” causation) or by suggesting that no other explanation is likely (differential diagnosis testimony).

What can courts reasonably do? Here is one possibility: Courts can suppose that the data and conclusions presented by qualified experts reflect valid science, upon which our system can reasonably allow a jury to rely in rendering a verdict for or against recovery in some very substantial amount. That seems close to the view in jurisdictions that admit scientific evidence on the basis of a credentials test coupled with findings that the proof is relevant and helpful. But there is another possibility, which seems more realistic: We can make the judgment that not all evidence that is presented as science, even by qualified witnesses, is of such quality that it can be relied upon to make serious decisions of the sort required for civil judgments. We can believe that such evidence varies in quality, and that sometimes it is not reliable enough. We can suppose that gaps in scientific understanding create room for interpretive disagreement, and that financial incentives, whether arising from the involvement of scientists in commercial or other funded projects or from their involvement in litigation, can compromise expert testimony. We can believe that science, like law, leaves room for principled intellectual disagreement that reflects differences in technical understanding or personal philosophy. We can also suppose that these differences sometimes lead to errors or to conclusions that cannot be defended or would be condemned by most others of similar training. Obviously, Daubert reflects the latter view of science, and I think that is the more realistic view.

There is yet another question. Should we charge judges to be

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9 See, e.g., State v. Peters, 534 N.W.2d 867, 871-73 (Wis. Ct. App. 1995) (following neither Daubert nor Frye, in applying WRE 702, the trial judge is limited to considerations of relevancy, qualifications of the witness, whether the evidence is superfluous or will waste time or resources, whether probative value is outweighed by prejudice, whether jury can draw its own conclusions, whether the evidence is inherently improbable, and whether the area is suitable for expert opinion); Green v. Smith & Nephew AHP, Inc., 617 N.W.2d 881, 890 (Wis. Ct. App. 2000) (finding that unlike Daubert jurisdictions, where “the trial court has a significant ‘gatekeeper’ function in keeping from the jury expert testimony that is not reliable, the trial court’s gatekeeper role in [this state] is extremely limited”).
the gatekeepers, or should we fold the gatekeeping responsibility into
the factfinding responsibility, leaving the assessment of science to
juries? As others have suggested, the right approach is to ask this
question: Are judges more capable than juries of playing this role? In
his engaging contribution to this conversation, Professor Joseph
Sanders says yes. His conclusion rests on an examination of
empirical data (some published, some new and unpublished), and he
is cautious. Still, his conclusion is generally yes.

Professor Sanders builds on what he calls a "counter-
revolutionary" Kansas decision and the work of Professor Alvin
Goldman. Goldman suggests that we should consider (a) the
characteristics of the audience (juries), (b) the characteristics of the
witnesses (experts, including scientists), (c) the criterion to be
applied (Daubert or some other standard), and (d) other alternatives.
As for juries, we have indications that they have trouble with complex
cases, and with scientific evidence, and we have reason to believe that
better-educated juries do better in these areas. We have indications
that juries approach expertise with skepticism. We have indications
that juries appraise expert testimony not by grappling with technical
issues, but by counting extraneous factors like qualifications, the
number of arguments (rather than quality), and personal
attractiveness. We understand that jurors give more credence to
messages framed in simple language, less to those framed in complex
language, and they pay close attention to demeanor. As for experts,
we have confirmation of what we have long suspected: They tailor
their testimony to please whoever pays them. They learn to
"perform" in court. As for judges, we have some mixed news: Data on
state judges suggest that many do not understand the "testability"
concept (can the evidence be "falsified")? or error rates, although

10 Joseph Sanders, The Paternalistic Justification for Restrictions on the Admissibility of
Expert Evidence, 33 Seton Hall L. Rev. 881, 937-38 (2003); see also Brian Leiter, The
Epistemology of Admissibility: Why Even Good Philosophy of Science Would not Make for Good
designed to substitute "the rulemaker's judgment about what is epistemically best for
agents for their own judgment" as "epistemic paternalism").

(refusing to apply the state's Frye standard to testimony by treating physician, based
on differential diagnosis, that Parlodel caused a new mother to suffer stroke)
(Parlodel is a drug taken by mothers, who prefer not to breast feed their babies, in
order to suppress lactation). The court's reasoning in Kuhn involves consideration
of factors similar to those advocated by Professor Goldman, except that the court
decided that those factors support the conclusion that juries can appropriately
evaluate such testimony, so the judge need not play screening role. The main work
by Professor Goldman on which Professor Sanders draws is Alvin Goldman, Epistemic
they do better with criteria of peer review and general acceptance. Surveys of federal opinions, however, suggest that judges are achieving a better understanding of science. As for alternatives, we have some indications that cross-examination does little to affect jury appraisals of expert testimony.

Looking at this data, Professor Sanders concludes that they provide "some support" for restricting admissibility by a standard applied by judges. He comments as well that his own reading of the cases indicates that courts are doing better than the survey of state judges suggests. I agree, and I too can report that reading many decisions leads me to believe that appellate judges are doing better in appraising scientific proof than they did in the days of Frye.

At the risk of being simplistic, I think four additional factors point toward the need for a validity standard in which trial judges screen out questionable science. First, on balance judges are better educated than juries and are selected with attention to merit and skill. Second, judges have experience with adversarial presentations and are likely to be better able to understand the substance of testimony and its relationship to the issues. Both of these points suggest that judges can do better than juries in separating what should count from what should not. Third, the complexity of scientific evidence suggests that the "relevancy" criterion that applies to other evidence is not adequate to deal with science. Although I cannot prove it, I suspect that the very fact that a court admits evidence that is daunting or complex conveys to jurors an unspoken message of invitation, suggesting that they can rely on it (even though they need not). Finally, in jury-tried civil cases, it seems wiser to have judges decide the validity point simply because it is better to separate the decision on this point from the decision on the merits. The point is not merely that "two heads are better than one" (counting one head for the judge and one for the jury), but that one can reasonably expect a better decision on validity by someone who is not also responsible to decide whether the plaintiff or the defendant has the stronger case.

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12 That jurors' expectations may affect the way they process the evidence they hear is recognized in a different context in **Old Chief v. United States**, 519 U.S. 172, 188-89 (1997) (stating that juror expectations may arise "from the experience of a trial itself," and that shifting from descriptions by witnesses "naturally" describing "a train of events" to a different kind of presentation may make jurors "wonder what they are being kept from knowing").

13 This point was raised during the Seton Hall Symposium, **Expert Admissibility: Keeping Gates, Goals and Promises**, in February 2003, but none of the various participants, whom I contacted, claims credit for it. I had not thought of it before.
Finally, I think that juries are even less likely than judges to conclude that credentialed witnesses with scientific expertise are mistaken. The eye-opening article in this Symposium by criminal defense lawyer James Shellow claims, on the basis of experience, that jurors cannot understand or follow cross-examination aimed at revealing "flaws in methodology," and that effective cross requires essentially peripheral tactics, such as attacks on character or a demonstration that the expert's opinion is contradicted by published texts. The burden of the examples cited by Mr. Shellow is that the cross-examiner should exploit any unwillingness of the witness to acknowledge the authority of texts by casting that very fact as a demonstration of mendacity. This practitioner's view supports empirical evidence described by Professor Sanders indicating that jurors do not effectively come to grips with scientific evidence.

II. *DAUBERT DOES NOT THROW OUT TOO MUCH EVIDENCE*

*Daubert* has been cast as one of the villains in toxic tort claims that fail, but I doubt that this claim is correct, and doubt even more that this claim shows that *Daubert* is in some serious sense misguided or mistaken. To start with, it was not clear on the day *Daubert* was decided whether the effect of the new doctrine was actually to raise or to lower the bar with respect to science (and now all expert testimony). Although a recent study concludes that *Daubert* subjects evidence proffered as science to increased scrutiny, *Daubert* itself threw out a ruling that excluded expert testimony, and some modern state decisions continue to declare that *Daubert* favors admissibility more than *Frye*.

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15 See generally Margaret A. Berger, *Upsetting the Balance Between Adverse Interests: The Impact of the Supreme Court's Trilogy on Expert Testimony in Toxic Tort Litigation*, 64 LAW & CONTEMP. PROBS. 289, 318 (2001) (pointing out that *Daubert* led to exclusion of epidemiological testimony, criticizing opinions requiring that such proof show a doubling of relative risk, and suggesting that federal courts should look to state requirements for proving causation); Lucinda Finley, *Guarding the Gate to the Courthouse: How Trial Judges are Using their Evidentiary Screening Role to Remake Tort Causation Rules*, 49 DEPAUL L. REV. 335, 375 (1999) (urging that courts should not place the burden of scientific uncertainty on plaintiffs).
16 See *Daubert v. Merrell Dow Pharm.*, 509 U.S. 579, 588 (1993) (contrasting the "liberal thrust" of the Rules and the "permissive backdrop" behind FRE 702 with the "austere standard" of *Frye*); *State v. Leep*, 569 S.E.2d 133, 143 (W. Va. 2002) (contrasting *Frye* with the "more liberal" *Daubert* standard and adopting the latter). But of course Judge Kozinski again excluded the evidence proffered in *Daubert*, this time applying the new standard. See *United States v. Daubert*, 43 F.3d 1311 (9th Cir. 1995); see also Lloyd Dixon & Brian Gill, *Changes In the Standards for Admitting Expert
A. Critical Look Approach

Properly understood, Daubert only asks courts to look critically at evidence proffered as science (or expertise more generally), and to determine whether it is valid and what it can prove. In modern decisions Daubert does not automatically block efforts to prove causation by expert testimony resting, for example, on such techniques as animal studies or differential diagnosis. The former brings questions of dosage or exposure and questions stemming from differences between humans and animals. The latter involves attempts by treating physicians to eliminate other causes until only one explanation is left. Proof of this sort can survive scrutiny under Daubert, although it may properly be excluded if it fails adequately to fit the case, its factual basis is inadequate, or the methods or laboratory protocols were not properly followed. The problems of rational inference raised by such evidence virtually invite attempts to prove cause by evidence that does not really do so, and one cannot seriously argue that all such proof is a reliable indicator of cause. There is room for difference of opinion in applying the Daubert standard, and no doubt room for mistakes. But in areas of such difficulty, why would anyone expect otherwise?

It is true that proof based on chemical structure analysis has had tough sledding, but skepticism is justified by the fact that the technique is not suited to this use, and serves better as a tool for mapping out future research. It is also true that cases applying
Daubert evince a preference for proving cause by means of epidemiological studies. The problem with too much enthusiasm for epidemiological evidence is that it is often unavailable: Claimants cannot come up with such proof because it is expensive and takes a long time to develop. Even if such evidence constitutes "the gold standard" in this setting, however, courts applying Daubert regularly allow causation to be proved in other ways.

B. Daubert as Source of Bad Rules

Some commentators suggest that Daubert is the cause of certain "rules" that block recovery in toxic tort cases. One is the supposed rule that epidemiological evidence is admissible only if it shows a doubling of incremental risk. The problem is as follows: Suppose a study of two groups of 500 people, one group exposed to agent X and one not exposed. In the exposed group, we find that thirty-six suffer ailment Y, but in the unexposed group only twenty people suffer ailment Y. The usual standard of statistical significance, in which \( p = 0.05 \), requires that we be able to say that pure chance would produce the observed result only one time in twenty (\( p = 0.05 \) refers to that low probability). Under this standard, our result is significant.

The calculations underlying the examples in this paragraph, and the text accompanying notes 68-71, infra, are the ones required in comparing the means of independent samples where standard deviation is not known (two groups of 500 people, one exposed to agent X, the other not exposed). The examples assume that prior research indicates a causal link between agent X and disease Y, which is important because the analysis must assume either that (1) it is unknown whether agent X might actually lessen the risk of disease Y or (2) it is known that agent X might increase the risk of disease Y and there is no reason to think it lessens that risk. In the latter situation, the analysis employs a one-tailed test, and more results survive scrutiny. See Russell T. Hurlburt, Comprehending Behavioral Statistics, 238-74
Scientists would take it seriously as an indication that agent X causes ailment Y.\textsuperscript{23} Actually, it is now recommended that researchers reporting statistically significant findings include the "confidence interval" with their report. The latter describes the range of outcomes that would be expected to occur by pure chance no more than five percent of the time. The narrower the interval, and the further up from critical value that the interval lies, the higher the quality of the reported result.\textsuperscript{24}

Still, if all we knew about the plaintiff was that he was exposed to agent X and suffers ailment Y, even our statistically significant result does not by itself indicate that agent X probably caused plaintiff’s ailment. We could say otherwise, however, if the result showed more than a doubling of incremental risk—let us say that 42 people in the exposed group suffer ailment Y, and only twenty people in the
unexposed group. In this case, combing the epidemiological study with our knowledge of the plaintiff would suggest that agent X probably did cause his ailment. The reason is that more than half the observed instances of ailment Y in the exposed population were caused by agent X, so any one person in the group is more likely than not to have become ill from exposure to agent X. The result of this test, by the way, would once again satisfy the conventional notion of statistical significance.

In fact, however, the situation is seldom so simple. The probative force of such proof turns on such things as levels of exposure (time and dose), adequate sampling techniques, specificity of symptoms measured, and controlling for extrinsic (or potentially confounding) variables. Moreover, taking seriously the conventions of statistical significance described above, even outcomes showing more than a doubling of incremental risk would not necessarily persuade scientists to draw any conclusions. Also, a relative risk in the neighborhood of two is not as high as it sounds (scientists often see far higher relative risks).

In this example (42 exposed people suffer ailment Y and 20 unexposed people), the result is statistically significant at $p = .05$. For this level of significance, the critical point value of $t$ is 1.65, and the observed point value of $t$, in the comparison of the two samples, is 2.98, which falls in the critical range. The confidence interval for the comparison is .019-.069. Again the null hypothesis is that exposure has no bearing on the number of ailing people. The null hypothesis assumes an observed value of $t$ below 1.65, and assumes that the confidence interval would span the number 0, or fall below it. Since the observed point value of $t$ exceeds the critical value of $t$ and since the confidence interval spans a range above 0, the result is statistically significant.

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26 See generally LEON GORDIS, EPIDEMIOLOGY 192-95, 204-17 (2d ed. 2000) (describing guidelines for studies of problems in causation, which require researchers to consider: (1) temporal relationship; (2) strength of association; (3) dose-response relationship; (4) replication of findings; (5) biologic plausibility; (6) consideration of alternative explanations; (7) cessation of exposure; (8) specificity of association; and (9) consistency with other knowledge). Other problems in epidemiological studies include: (a) selection bias; (b) information bias; (c) confounding factors; and (d) interaction. Id.; see also Barrow v. Bristol-Myers Squibb Co., No. 96-689-Civ-Orl-19B, 1998 WL 812318, at *23 (M.D. Fla. Oct. 29, 1998). This case describes the Bradford-Hill criteria for appraising epidemiological proof of causation, which include:

1. the strength of the association or how far above 1.0 is the relative risk; (2) the consistency of the association or its reproducibility; (3) the specificity of the signs and symptoms or whether they are unusual and distinctive; (4) the dose response; (5) the temporality; and (6) the biologic plausibility of the theory of causation.

27 See infra example 4, note 71 and accompanying text.

28 See, e.g., Hollander v. Sandoz Pharm. Corp., 289 F.3d 1193, 1212 (10th Cir. 2002) (noting study results that indicate that the relative risk of stroke in post-partum
Putting aside these problems, commentators are right that epidemiological studies should be admissible to show general causation even if relative risk is less than two, and a few decisions do miss this point. When the purpose is to prove specific cause, it makes sense to insist on relative risk exceeding two because such proof is mathematically sufficient to satisfy the preponderance standard. But even this restriction assumes that there is no other evidence of exposure: If there is other evidence, then even epidemiological proof that does not show a doubling of risk is still relevant as partial proof of specific cause. Many modern decisions (state and federal) approve epidemiological evidence showing a relative risk less than two, and many demonstrate good understanding of this idea. Decisions refusing to accept the proof when it shows a relative risk less than two often do so for other reasons, and not on the basis of simple misunderstanding.

women is 28.3, as compared with non-pregnant women); Falise v. Am. Tobacco Co., 94 F. Supp. 2d 316, 336 (E.D. N.Y. 2000) (noting in cigarette smoking case that risks of lung cancer are 5 times higher for asbestos workers than for other workers, 50 times higher if asbestos workers smoke cigarettes, and 87 times higher if asbestos workers smoke more than a pack a day); In re Joint Eastern and S. Dist. Asbestos Litig., 827 F. Supp. 1014, 1038 (S.D.N.Y. 1993) (reporting relative risk of lung cancer in cigarette smokers as compared to nonsmokers is “on the order of 10:1”).

Among federal cases, see In re Hanford Nuclear Reservation Litig., 292 F.3d 1124, 1133 (9th Cir. 2002), which held that the trial court erred in requiring that epidemiological evidence show a relative risk greater than 2.0 and further stated that, to show “generic causation,” plaintiffs only needed scientific evidence that radiation “was capable of causing” injuries such as those suffered by plaintiff. Among state cases, see McDaniel v. CSX Transportation, Inc., 955 S.W.2d 257, 265 (Tenn. 1997), which adopted the Daubert standard, and also rejected the defense claim that epidemiological evidence is admissible only if it shows relative risk exceeding 2.0.

32 See In re TMI Litig., 193 F.3d 613, 712 n.166 (3d Cir. 1999) (quoting passage from REFERENCE MANUAL); Allison v. McGhan Med. Corp., 184 F.3d 1300, 1315 n.16 (11th Cir. 1999) (finding a relative risk exceeding 2.0 permits “an inference that the plaintiff’s disease was more likely than not caused by the agent,” yet noting that no one doubts that smoking can cause heart disease even though relative risk in that setting is only 1.5); In re Joint E. and S. Dist. Asbestos Litig., 827 F. Supp. 1014, 1028 (S.D.N.Y. 1993) (holding that an epidemiologist might find cause where relative risk is less than 2.0, but a “more likely than not” test “is not satisfied by epidemiological evidence alone” unless relative risk exceeds 2.0).

33 See Allison, 184 F.3d at 1315 (approving exclusion of evidence showing relative risk of 1.24 because “it was so significantly close to 1.0 that the court thought the study was not worth serious consideration for proving causation” in a breast implant case).
Another supposed rule holds that evidence of differential diagnosis cannot be admitted to show that agent X caused plaintiff to suffer ailment Y unless there is additional proof of general cause—whether epidemiological or based on animal studies—that agent X does cause ailment Y in some people. Such proof is popular because it is less expensive, and it can be provided by treating physicians who are not toxicologists or epidemiologists. Essentially the physician testifies that she tried to account for the ailment in other ways, through testing or treatment regimens, and thus eliminated all other possible or likely causes except agent X. Here the cases conflict. Some hold that differential diagnosis can only eliminate other causes, and because the technique is necessarily uncertain (it is hard to know when one has eliminated all but one cause), it can only supplement affirmative proof that agent X sometimes causes such ailments. Other cases admit such proof without such preconditions, accepting it as relevant to show causation. There is no settled rule, and courts seem to be trying hard to distinguish between testimony that does eliminate other plausible risks, and testimony that does not. In other words, the decisions seem to draw sensible qualitative distinctions that connect with the task that Daubert asks courts to perform.

33 See e.g., Rider, 295 F.3d at 1194, 1199 (affirming summary judgment for defendant in a Parlodel suit after a Daubert hearing because the proffered evidence of causation was not sufficient to prove causation; also finding that case reports based on differential diagnosis could not by themselves prove causal link “because they report symptoms observed in a single patient in an uncontrolled context,” which can “rule out other potential causes” but cannot rule out the possibility that the observed effect “is simply idiosyncratic or the result of unknown confounding factors,” so such reports “may support other proof of causation,” but “ordinarily cannot prove causation” by themselves); Hollander, 289 F.3d at 1210-11; Glastetter v. Novartis Pharm. Corp., 252 F.3d 986, 988 (8th Cir. 2001) (containing an analysis similar to Rider).

34 Mattis v. Carlton Elec. Prods. 295 F.3d 856, 861 (8th Cir. 2002) (admitting differential diagnosis testimony in electrician’s suit against a maker of PVCs as sufficient to prove causation, where a physician “ruled out other possible causes,” including “smoking, asthma, or ammonia,” and concluded that plaintiff developed reactive airways syndrome “as a result of his exposure to Carlton cement fumes”).

35 See Joseph Sanders & Julie Machal-Fulks, The Admissibility of Differential Diagnosis Testimony to Prove Causation in Toxic Tort Cases: The Interplay of Adjective and Substantive Law, 64 LAW & CONTEMP. PROBS. 107, 137 (2001) (stating that Daubert has led to “greater skepticism” about differential diagnosis testimony, but that courts have reached a “fair degree of consensus” on questions such as whether the proponent must first offer “ruling-in” evidence before offering “ruling-out” testimony and on the sufficiency of proof resting largely on “temporal order”).

36 Cooper v. Smith & Nephew, Inc., 259 F.3d 194, 202 (4th Cir. 2001) (stating that differential diagnosis “normally should not be excluded because the expert has failed to rule out every possible alternative cause,” but that testimony may be
Of course both of these "rules" could be deployed unwisely to block just claims. It might even be true that a Daubert regime provides greater opportunity for courts to make such mistakes than Frye did. Certainly the most liberal "Rule 702" standard in current use, in which courts look at credentials and apply the "helpfulness" standard, would less often lead to exclusion of such evidence. But such proof should be excluded when it is thin, and looking directly at the science seems a good thing, not a bad thing.

These supposed "rules" also invite the criticism that courts are applying Daubert to measure the sufficiency of scientific evidence, rather than its relevancy. Some opinions appear to collapse notions of relevancy and sufficiency. That is not the fault of Daubert, however, and other modern decisions clearly understand the difference between relevance and sufficiency in this setting. In partial answer to this criticism, it is worth noting that a court asked to rule on an offer of proof sometimes should exclude the evidence because it is insufficient. In other contexts, it is perfectly conventional for courts to sustain objections on the ground that evidence does not suffice to prove the point for which it is offered. There is absolutely nothing wrong with doing so, at least in cases in which the proponent has no additional evidence on the point in question and has had an adequate opportunity to advise the court about the proof that he does have and to make the usual proffer.

excluded where it fails to consider other potential causes). The court in Cooper found that the record was "replete with evidence that smoking can cause non-unions to occur," and that plaintiff was a pack-a-day smoker for 25 years, a fact that expert "categorically dismissed."

37 See generally Edward J. Imwinkelried, Daubert Revisited: Disturbing Implications, 22 Champion 18 (May 1998) (criticizing federal courts that have invoked the "fit" or "relevancy" prong of Daubert to insist that epidemiological proof satisfy what amounts to a sufficiency standard).

38 See Joint Dist. Asbestos Liab. Litig., 52 F.3d 1124, 1133, 1137 (2d Cir. 1995) (stating that "Daubert did not alter the traditional sufficiency standard" and reversing the trial court's judgment for defendant as matter of law because the trial court erred in ruling that plaintiff presented insufficient epidemiological and clinical evidence to prove causation).

39 See, e.g., Tennison v. Circus Circus Enters., Inc., 244 F.3d 684, 690 (9th Cir. 2001); Viking Theatre Corp. v. Paramount Film Distrib. Corp., 320 F.2d 285, 296 (3d Cir. 1963), aff'd, 378 U.S. 123 (both rejecting offers of proof because the evidence could not prove the point for which it was offered unless other evidence was offered, which offeror did not include or could not obtain).
C. Daubert and Erie

Professor Margaret Berger offers another criticism of what she takes to be the judge-made doubling rule. Imposing this rule in federal diversity suits, she argues, violates the Erie doctrine. Stated in its strongest terms, the argument is that this judge-made rule, created in a Daubert-inspired construction of FRE 702, is substantive and is intended to affect outcome in a particular class of cases. Drawing on a modern opinion by Judge Posner in the Healy case, Professor Berger suggests that federal courts must apply any state substantive rule that is “in actual conflict” with a Federal Rule, and any “state procedural rule” that applies to “a particular substantive area.”

Lest anyone think the decision in Hanna stands in the way because it puts the Federal Rules beyond Erie-based challenge, Professor Berger reminds us that a prominent modern decision requires federal courts to apply state substantive law even when a Federal Rule is in play. She concludes that when a state court interprets evidence law “so as to better a plaintiff’s odds of prevailing in toxic tort litigation,” the result is a state rule applying in a particular class of cases, so federal courts must observe it. Perhaps more importantly, the state rule is substantive because it creates an incentive for manufacturers “to take more care in testing their products.”

This is a brave and inventive argument. There is something to be said for the proposition that if the state and federal systems persistently produce different outcomes in similar cases, on account of what seems to be different standards of proof, while purporting to apply the same substantive principles, the result would be troublesome. I concur in Professor Berger’s argument that federal courts should consider state precedents on matters closely related to sufficiency standards, and on evidential conventions that seem closely

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40 See generally Berger, supra note 15.
42 The reference in this text is to Hanna v. Plumer, 380 U.S. 460 (1965). The modern decision is Gasperini v. Ctr. for Humanities, 518 U.S. 415, 419 (1996) (holding that, in diversity suits, federal courts must apply state statute controlling compensation awards for excessiveness or inadequacy; noting, however, that the statute directs appellate courts to exercise this power, while in the federal system, the trial judge must take this responsibility).
43 Berger, supra note 15, at 319.
related to substantive principles. Federal courts often do just that, but most of the time they do not view themselves as bound by state law, and it seems telling that Judge Posner himself goes to great lengths to avoid being bound by state rules relating to proof in tort cases.

In the end, the *Erie* argument seems misconceived. To begin with, federal decisions don’t impose a “doubling rule” for epidemiological evidence (instead they analyze and assess probative worth and sufficiency). More importantly, different results on this question seem epistemological, rather than policy-driven. What I mean by epistemological is that the differences look like variations in attempts by federal and state judges to implement the “sufficient evidence” standard by deciding “how much evidence is enough” to allow a reasonable juror to find that cause has been proved under the preponderance standard. If the results were policy-driven, one would expect to see opinions linking the discussion of “how much is enough” to particular substantive standards, or to the purposes of tort law as compensatory and loss-spreading or as shifting to manufacturers only actual costs of injury while keeping innovation alive and costs down. One would also expect policy-based decisions to generate bright-line rules, or even statutes governing recurrent situations.

Of course judges might mask their decisions, justifying them

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45 See Allison v. McGhan Med. Corp., 184 F.3d 1300, 1320 (11th Cir. 1999) (noting in a breast implant case that *Erie* requires application of state substantive standards and commenting, with reference to evidence of relative risk, that state law requires proof based on “reasonable medical probability,” interpreted to mean “the functional equivalent of preponderance of the evidence”); In re Simon II Litig., 211 F.R.D. 86, 157 (E.D. N.Y. 2002) (commenting that allowing “statistical proof” of causation in cigarette smoking litigation does not conflict with *Erie* because there is “no ruling New York case which holds that state substantive law will not permit the use of modern aggregation forensic tools to support a massive fraud action”).

46 See S.A. Healy Co., 60 F.3d at 309-10. Judge Posner, writing the opinion in *Healy*, cites two tort cases in which the *Erie* issue is “pretty easy” because the state rule is “limited to a particular substantive area.” Id. Judge Posner authored both prior opinions, and both times he avoided applying the state rule. See also Barron v. Ford Motor Co. of Canada, Ltd., 965 F.2d 195 (7th Cir. 1992) (holding that a state statute blocking proof that claimant was not wearing seatbelt was substantive, but that it did not apply where defendant claimed that seatbelts were a design element relevant to the question whether it was reasonable to make sunroof out of laminated glass); Flaminio v. Honda Motor Co., 733 F.2d 465, 471-72 (7th Cir. 1984) (refusing to apply a state rule letting plaintiff prove design change, because a federal rule blocks it; noting that the matter is both substantive and procedural, but that it would be “melodramatic” to label federal rule as substantive and require federal courts to apply state counterpart) (at the time, state and federal rule were textually identical, but state and federal courts diverged on question whether rule excludes such proof).
epistemologically rather than by reference to substantive policy. Indeed, humanly speaking it is hard to imagine deciding whether indirect proof of cause suffices unless one also thinks about substantive policies, and what is humanly at stake—a serious injury or ailment on one side, the future of a drug on the other. But if every judicial attempt to think epistemologically (hence procedurally) is viewed as a masked effort to implement policy choices, then the *Erie* doctrine is doomed. We would be forced to the conclusion that federal courts cannot at the same time operate as “an independent system for administering justice” as contemplated by Justice Brennan in *Byrd*\(^7\) while complying with *Erie*’s command to honor state substantive policy choices. Professor Berger does not make such an extravagant claim, but her contention that federal judges are sometimes implementing substantive policy choices comes close to that, since the opinions do not say that they are behaving in this way.\(^4^8\)

In any event, the federal decisions that have noticed this issue have mostly avoided concluding that *Erie* mandates following state practice on this point.\(^9\) On the root question whether a state or federal standard governs the sufficiency question in diversity litigation, there is stronger support for the proposition that federal law governs than there is for applying a state standard.\(^5^0\)


\(^{48}\) See Berger, *supra* note 15, at 301-06 (noting that in federal decisions “insisting on epidemiological proof” or those insisting that a plaintiff’s epidemiological evidence show a relative risk exceeding 2.0, judges “are not making value-free determinations that are the inevitable consequences of a system of rational proof”).

\(^{49}\) See *Bartley v. Euclid*, Inc., 158 F.3d 261, 272-73 & n9 (5th Cir. 1998) (finding that if state law requires the evidence to show more than doubled risk in the exposed population, the proof satisfied the standard; if this requirement defines burden of proof, it is arguably “procedural rather than substantive, and therefore controlled by federal rather than state law”) (dissent argues that state rule is substantive); Blue Cross & Blue Shield of N.J., Inc. v. Philip Morris, Inc., 178 F. Supp. 2d 198, 259 (rejecting defense claim, in smoking case, that *Erie* required federal courts to require “proof of individual injury,” and commenting that the question is “better posed as a question of legal sufficiency”; if mixture of statistical and individualized evidence can prove cause, then “no *Erie* question is presented by federal evidentiary procedures which allow for the use of aggregate proof”); Hall v. Baxter Healthcare Corp., 947 F. Supp. 1387, 1394-95 (D. Or. 1996) (rejecting a claim that *Erie* required the federal court to apply state rule relating to proof of causation in breast implant suit). *But see Nat’l Bank of Commerce v. Assoc. Milk Producers, Inc.,* 22 F. Supp. 3d 942, 948 n.4 (E.D. Ark. 1998) (stating that if the application of a “federal evidentiary rule” leads to dismissal, whereas the application of the state rule would not, “then, under *Erie*, the evidentiary ruling might be considered substantive rather than procedural”); Raynor v. Merrell Pharm., 104 F.3d 1371, 1376 (D.C. Cir. 1997) (suggesting that a “question of sufficiency would be a substantive rule under *Erie*”).

\(^{50}\) See generally 9A CHARLES ALAN WRIGHT & ARTHUR R. MILLER, FEDERAL PRACTICE AND PROCEDURE § 2525 (2d ed. 1995) (opining that many courts now agree that
The amendments to FRE 702 adopted in 2000 reinforce Daubert's message that courts are to take a close and independent look at evidence proffered as science, and increase the difficulties of arguing under Erie that rulings on sufficiency are substantive. In effect, these amendments say that everything that could affect validity and accuracy count. Not only should courts insure that scientific testimony "is the product of reliable principles and methods," which was the actual holding of Daubert, and not only should courts insure that scientific evidence rests on "sufficient facts or data," which was part of the sufficiency calculus that courts perform under FRCP 50 and sometimes in connection with rulings on offers of proof under FRE 103, but courts should also take steps to ensure that the witness "has applied the principles and methods reliably to the facts."

The latter provision resolves a conflict among the cases in favor of more judicial scrutiny. This language directs trial judges to consider issues of laboratory protocol in determining whether to admit or exclude expert testimony, meaning that these issues affect not merely weight, but admissibility. Some pre-amendment authority had pointed toward this conclusion, but other decisions pointed toward the opposite conclusion.

Whether the 2000 amendments...
III. **DAUBERT PROPERLY CONCEIVES SCIENCE, AND TAKES THE RIGHT LEGAL STANCE TOWARD SCIENCE**

Critics complain that *Daubert* is incoherent, perhaps even internally conflicted, in two critical respects—in its view of science, and in its conception of the proper relationship between science and law. There is power in these observations, but I mean to say once again that *Daubert* is not at fault. Indeed, one of the strengths of the opinion is that its vision is broad enough to embrace internal tensions and difficulties in science, and in the relationship between law and science, that cannot be avoided. For us in the scholarly community, and for judges toiling in the vineyards, the “task at hand” is to make our way toward appropriate accommodations of

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[53] I am aware of one state that has apparently adopted the new language. See Miss. R. Evid. 702 (adopter the new federal language). In Colorado, the state Supreme Court rejected a recommendation by its Evidence Rules Advisory Committee to adopt the federal language, after I argued unsuccessfully in favor of its adoption. The expressed concern was that adopting language that seemed so closely related to *Daubert* would essentially adopt *Daubert* itself, a questionable position given that Colorado had just rejected the *Frye* standard, in order to adopt its own standard, which is similar to, but not identical with, *Daubert*. See Shreck, 22 P.3d at 73-79 (requiring trial courts to consider reliability of expert testimony, qualifications of witness, and usefulness of testimony, thus endorsing the *Daubert* factors without adopting *Daubert*).


[55] *Daubert*, 509 U.S. at 597 (characterizing the responsibility of trial judge to include ensuring that an expert’s testimony “rests on a reliable foundation and is
Let us consider first the charge that Daubert's view of science is incoherent. On the one hand, we can see in the core of the opinion (the part that adopts the validity standard and speaks of accuracy and propositions that can be tested or "falsified") an apparent belief that science is a static body of objective knowledge reflecting certainty. On the other hand, we also find in Daubert suggestions that (a) science is a process, hence anything but static; (b) scientific knowledge does not reflect certainty, but is uncertain and contingent; and (c) scientific expertise is affected by the forces that generate litigation, hence subjective in some respects, and socially constructed. This incoherent view, it is said, makes the task that Daubert sets for judges impossible to perform: In effect, Daubert charges them to apply static objective standards in appraising shifting subjective, contingent knowledge.

Let us consider the relationship between law and science, as Daubert envisions it. On the one hand, Daubert affirms that it is the job of courts to appraise science, and courts are not simply to defer to the scientific community on the question whether evidence presented as science is valid and reliable. This role for courts is what we mean by "gatekeeping." On the other hand, Daubert says courts are to judge science by the standards that scientists deploy in judging science. Kumho Tire adds an exclamation point in commenting that scientists are to bring to the courtroom "the same level of intellectual rigor that characterizes the practice of an expert in the relevant field." Again this incoherent view asks courts to do what they cannot do and fails to recognize that science and law have different agendas, goals and purposes, and operate under different constraints.

A. A Defective View of Science?

On the question whether Daubert has a defective view of science, I would begin by suggesting that the problem of objectivity has a familiar ring, perhaps because bridging the gap between human perceptions and the world has engaged philosophers for thousands of years, and the conversation is not over yet. How surprising is it to relevant to the task at hand"; see also Kumho Tire, 526 U.S. at 152 (explaining how judges apply Daubert factors "to the case at hand").

56 Daubert, 509 U.S. at 590, 596 (noting that "arguably, there are no certainties in science," and commenting that science is "a process for proposing and refining theoretical explanations about the world," making it qualitatively different from the law).

57 Kumho Tire, 526 U.S. at 152.
find that philosophers and historians of science report that science too is not the wholly objective edifice that we outsiders envision? Thomas Kuhn's salient work argues that the choice between what he called "competing paradigms" in science "cannot be determined merely by the evaluative procedures characteristic of normal science." The philosopher of science Karl Popper, of whom we have heard because Daubert draws on his work, takes a similar position. In defending the proposition that science is distinguished from other forms of knowledge by the fact that it can be "falsified," he argued that a proposition can be falsified only by a "basic statement," meaning a singular empirical statement that is accepted because it has been tested "inter-subjectively" rather than objectively. Yet views as skeptical as these cannot long survive unchallenged in a world that has seen such extraordinary accomplishments as laser surgery, the internet, space stations and jet airline travel. Obviously science has answers to critical questions, and Kuhn and Popper both recognized as much. What Kuhn called "normal science" proceeds, he wrote, out of random early "fact-gathering" toward something resembling "an accepted model or pattern" that he called a "paradigm," which succeeds because it is more successful than other paradigms "in solving a few problems that the group of practitioners has come to recognize as acute." "Mopping-up" operations that are "what engage most scientists throughout their careers," and these proceed after the adoption of a paradigm, and constitute "normal science." In the end, what counts most as a critical criterion of scientific paradigms is predictive accuracy. Popper was less direct in suggesting a positive account of scientific knowledge, but he did comment that scientists reach a kind of stopping point with "statements about whose acceptance or rejection the various investigators are likely to reach agreement," and he acknowledged that we must find such stopping points or end in a new "Babel of

59 KARL POPPER, THE LOGIC OF SCIENTIFIC DISCOVERY §§ 8, 22, 28, 29 (Routledge Classics 2002) (stating that a theory can be "falsified" only by means of "a reproducible effect which refutes the theory").
60 KUHN, supra note 58, at 15, 23, 25-27 (these "mopping-up" operations involve investigating those facts that "the paradigm has shown to be particularly revealing of the nature of things," as well as facts that lack "intrinsic interest" but "can be compared directly with predictions" from the paradigm).
61 THOMAS S. KUHN, Objectivity, Value Judgment, and Theory Choice, in INTRODUCTORY READINGS TO THE PHILOSOPHY OF SCIENCE 436 (Klemke et al. eds., 5d ed. 1998) (listing "as characteristics of a good scientific theory" the following factors: accuracy, consistency, breadth of scope, and fruitfulness in the sense of encouraging new phenomena or previously unnoticed relationships).
Tongues" in which scientific discovery "would be reduced to absurdity," and "the soaring edifice of science would soon lie in ruins."\(^{62}\)

Perhaps Daubert should be faulted for being too skeptical of science. As Professor Ron Allen argues, it seems odd to posit as a standard of scientific validity the question whether a proposition can be "falsified" as opposed to "verified" or "confirmed." Arguably such a standard is too demanding if we wind up accepting scientific knowledge only if it has been tested in every conceivable way, and Thomas Kuhn took issue with the very idea of "falsification" in an account that stresses more positive notions of verification.\(^{64}\) But I do not believe Daubert meant to erect a barrier as high as that. It may be comforting to consider that Popper, in explaining why he chose "falsifiability" as the central criterion of science, says his purpose was to distinguish science from myth and metaphysics. Specifically he had in mind the claims of Carl Marx, Sigmund Freud, and Alfred Adler, which he viewed as "pseudo-science" more closely resembling astrology than astronomy.\(^{65}\) Popper did not actually argue that nothing could be accepted until it was tested so exhaustively that nobody could doubt it. Rather, he argued that a scientific

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\(^{62}\) Popper, supra note 59, § 29 (characterizing this situation as "a failure of language as a means of universal communication").

\(^{63}\) See Ronald J. Allen, Expertise and the Daubert Decision, 84 J. CRIM. L. & CRIMINOLOGY 1157, 1169-71 (1994) (arguing that Daubert "adopted uncritically the view that Popperian falsifiability is at the heart of modern science," showing "no awareness" that that view is controversial and inadequate because it suggests that science produces knowledge only if it survives "all conceivable tests," but fails to account for the accomplishments of scientists, who "do not believe that all they know are negatives," and know "a lot of positive truths" too). For a reply to this criticism, see Sean O'Connor, The Supreme Court's Philosophy of Science: Will the Real Karl Popper Please Stand Up?, 35 JURIMETRICS J. 263 (1995).

\(^{64}\) Kuhn, supra note 58, at 145-47 (describing "probabilistic verification theories" that "compare the given scientific theory with all others that might be imagined to fit" the data, or construct by imagination "all the tests that the given scientific theory 'might conceivably be asked to pass,' and doubting that any test can falsify any theory because "no theory ever solves all the puzzles," and indeed "it is just the incompleteness and imperfection of the existing data-theory fit that, at any time, define many of the puzzles that characterize normal science").

\(^{65}\) Karl Popper, Science: Conjectures and Refutations, in INTRODUCTORY READINGS TO THE PHILOSOPHY OF SCIENCE, 38-40 (Klemke et al. eds.) (describing a conversation with Adler in which the author had mentioned a child, whom Adler had "no difficulty in analyzing in terms of his theory of inferiority feelings, although he had not even seen the child," because, as Adler said, he had "thousandfold experience," leading the author to conclude that that Adler's "previous observations may not have been much sounder than this new one," proving only that any case "could be interpreted in light of the theory").
proposition is one that can be tested. And of course the Popper account tracks a salient feature of the scientific method, which is to test a hypothesis to see whether experimental results refute it (testing the “null hypothesis” in common parlance).

In sum, what we might take as incoherence or internal conflict in Daubert’s view of science can also be understood more constructively as a kind of dualism that embodies a view of science similar to what we find in Kuhn and Popper. Rather than abandoning any search for a validity standard, this dualistic view should lead us to recognize, in words that Professor Nance might find congenial, that reliability is not an all-or-nothing concept, but a relative concept: Often it will be possible to insist on a kind of “certainty” of the sort that we have in mind when we speak of the tides or the hour of sunrise, but other times we can only expect the sort of “certainty” that we have when we say that asbestos causes some kinds of lung cancer.

B. Misconceived Relationship between Law and Science?

A serious criticism of Daubert is that courts are being led to demand a higher level of statistical significance than is appropriate. Epidemiological evidence might support the conclusion that exposure to agent X increases the risk of ailment Y, thus in turn supporting an inference of general causation (agent X causes some instances of ailment Y). Alternatively, it might support the conclusion that exposure more than doubles the risk, thus in turn supporting an inference of general causation and perhaps even specific causation (plaintiff was exposed and is ailing, and so agent X is the cause).

To illustrate these points, consider some examples comparing 500-person samples (one group exposed to agent X, one not exposed). Example 1 (risk increase): We find that thirty-six exposed people have ailment Y, but only twenty unexposed people. These numbers suggest that ailment Y suffered by sixteen out of thirty-six

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66 LOGIC OF SCIENTIFIC DISCOVERY, supra note 59, § 22 (we must “clearly distinguish between falsifiability and falsification”); see also Conjectures and Refutations, supra note 65, at 43 (task is not to identify “meaningfulness or significance” or “truth” or “acceptability,” but rather to distinguish statements and systems belonging to “the empirical sciences” from all others, whether “psychoanalytic” or “myth” or something else; the latter are not “unimportant, or insignificant,” and “may contain important anticipations of scientific theories”; indeed, psychoanalytic theories “contain most interesting psychological suggestions, but not in a testable form”).

exposed people came from agent X, that relative risk is 1.8, and that there is a .444 probability that agent X caused any given case of ailment Y in the exposed population. Example 2 (risk increase): In the exposed group, twenty-nine people have ailment Y; again only twenty in the unexposed group. These numbers suggest that ailment Y suffered by nine out of twenty-nine exposed people came from agent X, that relative risk is 1.45, and that there is a .310 probability that agent X caused any given case of ailment Y in the exposed population. Example 3 (risk more than doubled): We find that forty-two exposed people have ailment Y, but only twenty unexposed people have it. These numbers suggest that the ailments suffered by twenty-two out of forty-two exposed people came from agent X, that relative risk is 2.1, and that there is a .542 probability that agent X caused any given case of ailment Y in the exposed population. Example 4 (risk more than doubled): We find that five exposed people have ailment Y, and only two unexposed people. These numbers suggest that the ailments suffered by five out of seven exposed people came from agent X, that relative risk is 3.5, and that there is a probability of .714 that agent X caused any given case of ailment Y in the exposed population.

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Footnotes:

69 In this example (36 exposed and 20 unexposed people suffer the ailment) recall from note 22 that the result is statistically significant at p = .05. The observed value of t exceeds the critical value of t, and the confidence interval for the comparison is .007 to .057, which lies above the value of 0 that the null hypothesis would assume.

69 In this example (29 exposed and 20 unexposed people suffer the ailment) the result is not statistically significant at p = .05. For this level of significance, the critical point value of t is 1.65, and the observed point value of t, in the comparison of the two samples is 1.28, which falls below the critical range. The confidence interval for the comparison of the two samples is -.005-.041. Again the null hypothesis is that exposure has no bearing on the number of ailing people. The null hypothesis assumes an observed value of t below 1.65, and that the confidence interval will span the number 0. Since the observed point value of t is less than 1.65, and since the confidence interval does span 0, the result is not statistically significant (we do not reject the null hypothesis).

70 In this example (42 exposed people suffer the ailment, and 20 unexposed people), recall from note 25 that the result is statistically significant at p = .05. The observed value of t exceeds the critical value of t, and the confidence interval for the comparison is .019-.069, which lies above the value of 0 that the null hypothesis would assume.

71 In this example (5 exposed people suffer the ailment, and 2 unexposed people), the result is not statistically significant at p = .05. For this level of significance, the critical point value of t is 1.65, and the observed point value of t, in the comparison of the two samples, is 1.2, which falls below the critical range. The confidence interval for the comparison of the two samples is -.002-.014. Again the null hypothesis is that exposure has no bearing on the number of ailing people. The null hypothesis assumes an observed value of t below 1.65, and assumes that the confidence interval will span the number 0. Since the observed point value of t is less
The usual approach to the question whether agent X causes ailment Y is to begin with the "null hypothesis" that there is no correlation. If that were so, then in the two samples (500 exposed people; 500 unexposed people) the number of observed instances of ailment Y would be the same. But the outcome of studies may suggest, as do Examples 1-4 above, that the null hypothesis should be rejected, and that agent X does indeed cause ailment Y. Let us begin by understanding the meaning of these outcomes. To start with, the suggested conclusions are all general and qualified, and each can be deployed to state a probability, but not a certainty. The conclusions are all general because they suggest that agent X is a causal factor in the mass of observed instances of ailment Y. The conclusions are qualified because they suggest that agent X causes certain percentages of observed instances of ailment Y in exposed populations. The conclusions can be deployed to state probabilities because they suggest that, among ailing and exposed people, there are certain probabilities, equal to the percentages suggested by the figures, that any given instance of ailment Y was caused by agent X.

In the description set forth above, Examples 1 and 2 indicate that exposure raises the risk but does not double it, thus supporting inferences of general cause. Examples 3 and 4 indicate that exposure more than doubles the risk, thus supporting inferences of both general and specific cause. Epidemiologists would likely accept the conclusions indicated by Examples 1 and 3, because they are statistically significant at the level of \( p = 0.05 \), meaning that there is but one chance in twenty that the numbers would appear by chance—by random and inevitable differences in 500-person samples taken from the general population. Epidemiologists would likely reject the conclusions indicated by Examples 2 and 4 because they are not statistically significant at this level. The effect of this convention is to discourage "false positives" in favor of "false negatives," or (as it is sometimes said) to discourage \( \alpha \)-errors by incurring more \( \beta \)-errors, or discourage Type I errors in favor of more Type II errors. In other words, the conventional standard is less tolerant of errors that would find a causal connection and more tolerant of errors that would fail to find one. Examples 2 and 4, which suggest respectively that twenty-nine exposed people have ailment Y (as opposed to twenty unexposed people), and that five exposed people have ailment Y (as opposed to two unexposed people), would not be accepted because the results do not satisfy the conventional standard of statistical

than 1.65, and since the confidence interval does span 0, the result is not statistically significant (we do not reject the null hypothesis).
significance.

It is important to note that the conventional standard does not mean that science usually accepts only results that are ninety-five percent certain. Statistical significance at the level of $p = .05$ means that there is but one chance in twenty that the observed results could happen by chance. Satisfying the standard means that there is one chance in twenty (or less) that mere accidental variation would produce such a result, not that we can be 95% certain that the outcome (twenty-two out of forty-two observed ailments were caused by exposure, or sixteen out of thirty-six, as Examples 1 and 3 indicate) is correct. We do not and cannot know that. All we know is that the observed outcome would rarely be produced by chance alone, which gives us some reason to believe that the indicated correlation is correct. Any suggestion, however, that the conventional standard produces results of which we are ninety-five percent certain is false.72

Hence, it still needs to be said, it would make no sense to suggest that the civil justice system should accept results that are statistically significant at, say, the level $p = .40$ since such results leave us sixty percent certain of the conclusion, thus easily satisfying the notion of a preponderance of the evidence. Suggestions of this sort are close akin to the “prosecutor’s fallacy,” to which courts and lawyers sometimes fall prey. That fallacy equates the inverse of a scarcity factor with the probability of guilt: “The evidence shows that only 1 in 1000 randomly-chosen people would have a DNA profile like the one found in the defendant’s blood and in the blood at the crime scene, so we must conclude that the probability is 99.9% that defendant is guilty.” It is no more the case that $p = .05$ means that we can be ninety-five percent sure that the indicated correlation exists than it is the case that a scarce sample common to the defendant and the crime scene makes for near-certainty that defendant is guilty. Of course it is possible to make a link between the one-in-twenty probability of reaching the observed outcome by chance and the probability of actual cause. Likewise, it is possible to link the “1 in 1000 randomly-chosen people” probability to the probability that defendant was at the crime scene. Doing so, however, involves use of Bayes’ Theorem, which brings new complications and raises the

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72 See David W. Barnes, Too Many Probabilities: Statistical Evidence of Tort Causation, 64 LAW & CONTEMP. PROBS. 191, 208-09 (2001) (there is “no convenient way to translate the .05 p-value into a ninety-five percent confidence that the fact probability is correct,” in part because that value “assumes that the hypothesis is true” and “does not measure whether it is true”) [hereinafter Barnes, Too Many Probabilities].
serious problem of assigning a prior probability to the point in issue.⁷³

Now the argument that Daubert is leading courts mistakenly to require of statistical outcomes the same level of significance that scientists normally require proceeds in this way: This strong scientific bias may be appropriate in the setting of science, but not in the setting of civil litigation. Science observes this strong bias because science can afford it. Science works incrementally and has "forever" to get it right. Here is the way that one modern text in statistics explains the strong scientific bias:

We want α to be small because Type I errors are expensive for the scientific (and the human) enterprise. Suppose, for example, we . . . report in a journal that doses of vitamin B₁₂ increase IQ. That will be an error that we made in good faith because we had no way of knowing that this particular result was a type I error . . . . As a result, our readers will alter their behavior, perhaps focusing on a B₁₂ diet while ignoring other avenues (such as reading enhancement programs) that might be effective in raising IQ. Sometime later, perhaps, someone will conduct many experiments and find that vitamin B₁₂ has no effect on IQ; that is, they will demonstrate that we had made a Type I error. Our readers who believed our earlier report were done a possibly uncorrectable disservice because they may have ignored other avenues. To prevent any further damage, we would want to find and contact the entire readership of our first report and inform them that our result was mistaken. That is clearly an expensive and difficult (if not impossible) thing to do.

In contrast, the same text continues, we do not need as much protection from Type II errors (β-errors):

Consider an investigator who conducts a single experiment to demonstrate that doses of vitamin B₁₂ increase IQ. After

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⁷³ Bayes’ Theorem describes the degree to which an item of evidence, when expressed as a datum of known frequency, affects one’s prior estimate of the issue on which the evidence bears. For accounts of Bayes’ Theorem, see David W. Barnes, Too Many Probabilities, at 208-09, and CHRISTOPHER B. MUELLER & Laird C. KIRKPATRICK, EVIDENCE § 7.18 (3d ed. 2003). One problem in utilizing Bayes’ Theorem in this setting (indeed any setting) is that it is necessary to quantify the prior estimate of the probability before using the theorem to find the new probability, after taking into account the datum of known frequency. If the prior estimate were .05 (very low probability of causation), then applying a statistical finding returned at the conventional level of statistical significance raises the odds to 1:1, meaning equilibrium, or a .5 probability that the degree of cause indicated by the finding (let us say 22 out of 42 instances) is true, or in other words even odds that 22 out of 42 instances of ailment Y in the exposed population exposed were caused by agent X.
the data are collected, she concludes that the null hypothesis should not be rejected; that is, she concludes that vitamin B₁₂ has no demonstrable effect on IQ. As a consequence, the investigator will not report findings in a journal. Instead, the investigation may be considered an exploration of a blind alley. She had thought vitamin B₁₂ was effective, but apparently it wasn’t.

If subsequent research indicates that the conclusion of the original investigation was a Type II error, what is the expense of that error and who bears it? One major expense is the time lost in the original investigation, but the bearer of that expense is the original investigator, not the scientific community at large. There is no necessity of informing the community of a previous mistake because there was no report of findings in the first place . . . .

Now, the argument continues, our civil justice system differs from science in its goals and social purposes. To start with, our civil justice system does not have unlimited time. We cannot and do not defer decision in the interest of becoming more certain, and correlative we hold that a lawsuit must reach a conclusion now. Moreover, we must leave the results reached in a lawsuit in place forever. Our legal system cannot be tentative about its conclusions. In a sense, our legal system views “getting it right” as less important than “getting it done.” Equally important, our civil justice system is premised on the principle that mistakes in denying relief are as harmful as mistakes in allowing recovery (plaintiff loses when the evidence is equally balanced only because we need a rule to resolve this case).

Thus our civil justice system is neutral as between “false positives” and “false negatives” (α-errors as opposed to β-errors, Type I as opposed to Type II errors). For our civil justice system, failing to

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74 HURLBURT, supra note 21, at 196 (acknowledging, however, that Type I and Type II errors can both be costly; citing the example of a Type I error in the form of a false report that a drug is effective against AIDS, the author recognizes that this error could have “cruel effects, such as raising false hopes or discontinuing the funding of some other research”; the author also acknowledges, however, that a Type II error in “failing to report a drug that is in fact effective” would “also have cruel results, depriving needy individuals of effective treatment,” concluding that there is “no statistical answer” to the question which kind of error is more costly; it is “a matter of complex human judgment”).

75 Obviously criminal cases are another matter. There our law strongly favors acquittals over convictions if the evidence is in close balance, which is somewhat akin to the idea of preferring false negatives (such as not finding cause when in fact there is cause) over false positives (like finding cause where none exists). Some would argue that the bias of science would be appropriate in discouraging the state’s use of thin scientific evidence and grotesquely inappropriate in discouraging the defense use of thin scientific evidence.
find a cause that exists is as bad as finding one that does not exist. Hence our civil justice system should not require the level of statistical significance required by science, with its heavy bias against false positives.

These differences between law and science do indeed suggest that we should consider carefully the possibility of accepting results in lawsuits that scientists are not yet prepared to accept. In this paper, I am not prepared to stake out a final position on this issue. Frankly, I'm not sure what the right answer is. However, I do want to address the question how this argument fits with Daubert, to talk briefly about who ought to resolve the argument, and to raise some cautionary points.

First, I think the Daubert framework can accommodate the view that courts ought to accept scientific evidence that does not satisfy the conventional standard that scientists require. Daubert recognizes that the enterprises of law and science differ, and adopts the view that judges must make their own decision, in context and with reference to the needs of the legal system, on admitting or excluding evidence proffered as science. It is true, as some have pointed out in criticizing Daubert, that there is language suggesting that courts should be more careful of science than is the professional community that produces science. But given the more basic premise of Daubert that the law must judge science for its own purposes, I don't think this language is a serious obstacle to arguments favoring the admissibility of careful studies showing, for instance, a causal connection between ailment Y and agent X even if the results do not satisfy the conventional significance standard.

Second, in this symposium Professor Cohen suggests that a scientist who has evidence indicating, for example, some marginal correlation between agent X and ailment Y, should be able to testify even if she says the results do not satisfy the conventional standard. In Professor Cohen's formulation, the scientist is seen explaining to the jury that science would not accept the indicated conclusion but that the scientist herself might do so for purposes of resolving a question that could not wait. This proposal merits consideration, but it also raises questions. To begin with, and I think Professor Cohen

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76 See Daubert, 509 U.S. at 596-97 (noting that scientific conclusions are “subject to perpetual revision,” and science advances through “broad and wide-ranging consideration of a multitude of hypotheses” that can “eventually” be thrown out if wrong, but “[c]onjectures that are probably wrong are of little use” to the law, that must reach a “quick, final, and binding” judgment”).

agrees here, *Daubert* requires judges to determine what level of certainty or confidence is high enough to merit consideration by a jury. It is not up to an expert or a jury to decide whether or not to accept evidence that does not satisfy the significance standard. More importantly, telling jurors they can base their verdict on evidence that science would not accept may not be the right thing to do, and it is certainly a strange message: In effect, it says that “you may conclude that defendant caused plaintiff’s cancer even when qualified experts think the case is unproved.” It is even questionable whether a scientist would feel comfortable (or able to comply with the oath required of witnesses) if she testified that while she does not professionally accept the indicated conclusion, she might personally do for purposes of resolving a lawsuit.

Third, how clear is it that differences between the agendas of science and law justify applying a much more lenient standard to science offered as proof in litigation? It is true that in any one lawsuit there is but one chance to get it right, but it is certainly not true that society as a whole, operating through the legal system, has but one chance. When apparent toxic agent X appears, the system provides many opportunities to resolve the question whether agent X causes ailment Y, and the common-law method of building step by step on experience actually bears some resemblance to the scientific method of moving incrementally and withholding judgment until proof comes that is persuasive. On the other side of the ledger, it is not true that the decision in one case affects only that case. Particularly with medicines and other substances believed to have toxic effect, every court judgment has ripple effects, encouraging or discouraging parallel suits and settlements, and sometimes having legal impacts on later judgments. Hence errors made in the judicial system, whether favoring claimants or defendants, can produce additional errors as lawyers, claimants, and defendants react to them.

Perhaps equally importantly, it is not really the case that our legal system is “neutral” with respect to errors. In single cases we may be “neutral” as between errors favoring claimants and errors favoring defendants, but we are not neutral in aggregate on such points. It is in part because courts recognize the perils of being wrong in huge cases that we have a series of decisions in the federal system that cut back

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78 For well-known reasons, new claimants are not collaterally estopped by judgments won by the same defendant against prior claimants, and usually new claimants cannot take advantage of collateral estoppel against a defendant stemming from defendant’s prior loss on similar claims. But decisions admitting or excluding scientific proof, or holding it sufficient or insufficient, are likely to have *stare decisis* effects in later suits.
on the use of class suits to resolve mass tort cases. Even on the supposition that toxic exposure cases are litigated individually, wrongly finding for the plaintiff would have considerable collateral effect, in likely reactions by the defendant and resultant changes in the availability of products. On the supposition that toxic exposure cases are litigated in aggregate fashion, these collateral effects are even clearer and more pronounced.

Finally, I want to suggest that the apparent caution of science may not be quite what it seems. The conventional standard, after all, is just numbers—just the product of analysis of the quantification of data. Behind the numbers are more and real uncertainties—the ones that go with designing tests, selecting cohorts, trying to eliminate differences apart from the factor in issue that might account for observed differences. Part of the reason science insists on impressive numbers may be the recognition that it is hard or impossible to eliminate confounding variables, and that even promising results might not be replicable. To the extent such apprehensions underlie the insistence on high numbers, the conventions of science are not conservative. And there is an additional factor to consider, which is that scientists (like the rest of us) want to be noticed, and have an incentive to maximize the importance of their findings, which suggests that the high conventional standard for statistical significance acts as a counterbalance against self-serving human motivations that are in play not only among lawyers and politicians, but among scientists too, and indeed the whole human species.

See, e.g., Ortiz v. Fibreboard Corp., 527 U.S. 815 (1999) (disapproving settlement of asbestos claims, largely on the basis of concern over adequate representation); Amchem Prods. v. Windsor, 521 U.S. 591 (1997) (refusing to allow certification of nationwide settlement class in asbestos case, largely on basis of concerns over adequate representation and because remedies under consideration cannot be created judicially and require legislative consideration); In re Rhone-Poulenc Rorer, Inc., 51 F.3d 1293 (7th Cir. 1995) (refusing to certify class in suit against maker of blood solids, largely because of reluctance to stake future of defendants on outcome of single trial).

See also Robert L. Park, The Seven Warning Signs of Bogus Science, 49 CHRON. OF HIGHER EDUC. 21, Jan. 31, 2003, at B20 (as “warning signs,” listing the fact that the scientist “pitches the claim directly to the media,” that she says “a powerful establishment is trying to suppress” her work, that the effect is “at the very limit of detection,” listing “anecdotal” evidence, stressing that the belief has “endured for centuries,” that the discoverer has “worked in isolation,” and that “new laws of nature” are required to understand the discovery).
IV. DAUBERT CAN BE IMPROVED: APPELLATE REVIEW MADE REAL

This paper defends the Daubert approach, but one troubling aspect of Daubert can and should be fixed. That is the misplaced emphasis on the discretion of trial judges, which appears particularly in the Court's two follow-up decisions in the Daubert line. Indeed, it is at least possible to speculate that the sequence in which Daubert, Joiner, and Kumho Tire were decided has much to do with the growth in the emphasis on discretion. The idea is hardly mentioned in Daubert itself, but it gained prominence in Joiner partly because the Ninth Circuit had adopted an implausible rule that trial judges have less discretion to exclude than to admit evidence proffered as science (the antidote was to hold that judges have discretion either way), and it made further gains in Kumho Tire because the project in that case entailed explaining how the standard for science could be applied usefully to experiential expertise (where a measure of discretion seems essential if the scheme is going to work at all). 81

I concur with commentators who say Daubert should be implemented by inviting appellate courts to take a close look at rulings by a trial court admitting or excluding evidence offered as science. 82 Having become accustomed to the refrain among federal appellate courts that they accord deference to trial court decisions applying Daubert, I was surprised to learn that nine states and the District of Columbia instruct appellate courts to review rulings admitting or excluding evidence presented as science by applying a de novo standard. 83 Some decisions apply this rigorous standard only to

81 In describing the gatekeeping function in Daubert, it is singular that the Court made no reference to discretion. By my count, the Court mentioned discretion ten times in describing this function in Joiner (making many additional passing references to the term), and six times in the same context in Kumho Tire. See Daubert, 509 U.S. at 594 (mentioning that its standards are “flexible”); General Elec. Co. v. Joiner, 522 U.S. 136, 141-47 (1997) (repeatedly stressing discretion); Kumho Tire, 526 U.S. at 151 (repeatedly stressing discretion, and pointedly saying that judge “must have the same kind of latitude in deciding how to test an expert’s reliability” that it has “when it decides whether or not that expert’s relevant testimony is reliable”) (emphasis in original).

82 See David L. Faigman, Appellate Review of Scientific Evidence Under Daubert and Joiner, 48 HASTINGS L.J. 969 (1997) (for ordinary decisions to admit evidence, where preliminary facts “depend on the testimony of witnesses,” appellate deference is warranted, but scientific evidence is “quite different,” and the trial judge is not in a “preferred position” in evaluating it). See also Richard D. Friedman, Squeezing Daubert Out of the Picture, 38 SETON HALL L. REV. 1047, 1065 (2003) (appellate courts should play more of a role in reviewing Daubert issues than Joiner suggests).

83 I looked at modern decisions from all fifty states, most of which endorse an abuse-of-discretion standard. But a de novo standard has been adopted in Arizona, Florida, Kansas, Maryland, Minnesota, New Jersey, Oregon, Oklahoma, and Washington, as well as the District of Columbia. See generally Goeb v. Tharaldson, 615
the basic question whether the theory and method are valid—or in *Frye* terms, to the proxy question whether they are generally accepted—and leave related questions like "helpfulness" under FRE 702 for the trial judge to decide under an abuse-of-discretion standard.

There are four reasons for preferring a more exacting standard in review.

First, issues relating to the validity of theories and techniques transcend the facts of individual cases. This observation applies, for example, to the question whether DNA profiling can reliably identify a blood or fluid sample as having very likely come from one person or another (the validity of the theory), to the question whether particular methods of analysis (such as RFLP, PCR, and STR) accurately measure the attributes of blood or fluid, and whether a particular laboratory protocol adequately guards against missteps and laboratory error. It applies to the question whether proffered

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N.W.2d 800, 814-15 (Minn. 2000) (whether proffered expertise satisfies state's *Frye-Mack* "general acceptance" standard "is a question of law that we review de novo," but questions of "foundational reliability" are reviewed under "abuse of discretion" standard, as are matters of witness qualification); Kuhn v. Sandos Pharm. Corp., 14 P.3d 1170, 1179 (Kan. 2000) (adopting de novo standard of review for proof of medical causation); Jennings v. Baxter Healthcare Corp., 14 P.3d 596 (Or. 2000) (rejecting argument that appellate court should accord deference to trial court's decision on scientific validity, and concluding that the issue is reviewed as for "errors of law"); Hadden v. State, 690 S.2d 573, 578 (Fla. 1997) (review of *Frye* issues is de novo); State v. Harvey, 699 A.2d 596, 619 (N.J. 1995) (in applying *Frye* standard, question whether scientific community generally accepts a method or test "can transcend a particular dispute," and to the extent that *Frye* focuses on "issues other than a witness's credibility or qualifications, deference to the trial court is less appropriate"); Taylor v. State, 889 P.2d 319, 331 (Okla. 1995) (decision by trial court to admit novel scientific evidence should be subject to "an independent, thorough review," and appellate court should "not simply ask whether an abuse of discretion was committed"); State v. Tankersley, 956 P.2d 486, 464 (Az. 1994) (rejecting *Daubert* and staying with *Frye*, and announcing that *Frye* issues are subject to *de novo* review); Schultz v. States, 664 A.2d 60, 64 (Md. App. 1994) (question of reliability of scientific technique "does not vary according to the circumstances of each case," so it is inappropriate to apply abuse of discretion standard on review); State v. Cauthron, 846 P.2d 502, 505 (Wash. 1993) (stating that court would "review the trial court's decision to admit or exclude novel scientific evidence *de novo*"); United States v. Porter, 618 A.3d 629, 634 (D.C. 1992) (questions of general acceptance of new scientific techniques invite court "to establish the law of the jurisdiction for future cases," so court would "engage in a broad review").

84 The initials cited above refer to three of the more common methods for conducting DNA profiling. RFLP refers to "restriction fragment length polymorphism," and it is the first broadly useful approach that made its way into courtroom use. PCR refers to "polymerase chain reaction," a later development that allowed small samples to be "extended" so that the inevitable consumption of such materials in laboratory testing did not destroy the whole sample. The drawback of PCR was that it could only extend some of the attributes of the original sample, so
statistical proof should satisfy the standard that scientists would
require, to the question whether differential diagnosis can or cannot,
standing alone, prove specific cause, to the question whether animal
studies of any particular drug or chemical can prove causation in the
human population, and to the question whether similarities between
the chemical structures of a particular drug or other substance, on
the one hand, and some other agent known to cause certain
consequences, such as disease, on the other hand, can prove
causation. Questions of this magnitude need steadier guidance than
the abuse-of-discretion standard provides, and the answers that courts
reach should be applied in similar cases, rather than left to vary with
the differing views of trial judges exercising discretion.

Second, appellate courts are better situated than trial courts to
resolve such questions. To start with, three or more minds are likely
to do better than one in appraising such technical issues. And
appellate review goes forward in a setting less subject to severe
schedule pressures. Furthermore, Daubert issues are likely to benefit
from thorough appellate briefings. And appellate review can involve
consultation with technical materials and expert advice by means of
amicus briefs or affidavits, or even live testimony. Reviewing courts
can even take judicial notice of technical books, articles, and other
materials.\textsuperscript{86} Some sense of proportion is clearly warranted: It is one
thing to supplement the arguments and briefs of counsel with
references to additional material, and another thing to decide the
case on grounds never considered by the lawyers who briefed the case
without giving them any opportunity for input.

Third, the Daubert standard needs elaboration in the variety of
settings in which it is to apply, and trial judges need help and
guidance beyond that provided by the standards themselves. The
problem of causation in toxic tort cases is a prime area in which
appellate courts could play useful roles, and in which trial courts
clearly want and need guidance.

There is of course one countervailing concern, and that is that

\textsuperscript{86} FRE 201 governs only judicial notice of "adjudicative" facts, and most technical
material that might be noticed in this setting involves "evaluative" facts utilized by
courts in their attempts to formulate wise rules of law. The fact that FRE 201 does
not cover evaluative facts does not mean they cannot be noticed. Instead, the
omission from coverage simply means that judicial notice of evaluative facts is not
regulated by any formal rule. See State v. Jones, 922 P.2d 806, 809 (Wash. 1996) (\textit{de
novo} review includes "sources outside the record such as scientific literature, law
articles, and the decisions of other jurisdictions") (applying state's \textit{Frye} standard).
the *Daubert* standard is both wide-ranging and case-specific. It is wide-ranging now, if it was not when *Daubert* was decided, because amended FRE 702 indicates that judges are to consider “principles and methods” and the sufficiency of underlying “facts or data,” and *also* the question whether the expert “has applied the principles and methods reliably to the facts,” and *Kumho Tire* makes it clear that the focus is the “task at hand,” as Professors Denbeaux and Risinger remind us. To the extent that the admissibility decision actually focuses, for example, on the question whether a particular laboratory protocol was or was not followed in the case at hand, or on the question whether a particular lapse or discrepancy in the data materially affected the outcome, some degree of deference to the decision of the trial judge is in order. It is with larger questions, including those of theory and technique, and the appropriateness of the technique to the issue being decided, that closer scrutiny is warranted.

**CONCLUSION**

The Court’s decision in *Daubert* changed the relationship between law and science. Critics have argued that judges cannot act constructively in the way that *Daubert* envisions, but there are good reasons to think that indeed judges can rise to the task. Critics have argued as well that *Daubert* has led to the exclusion of too much scientific evidence, particularly in toxic tort cases, and have adopted restrictive rules that are out of place and, in the federal system, infringe on the *Erie* doctrine. But *Daubert* does not require adoption of such rules, and the cases show that courts are in fact working hard in very challenging areas to achieve appropriate outcomes in appraising science. The *Erie* doctrine is not offended by federal efforts to implement a sufficiency standard.

Critics have also argued that *Daubert* misconceives science, and the relationship between law and science. But the dualism visible in *Daubert’s* account of science is also visible in the accounts of philosophers and historians of science, and the task is to reconcile notions of objectivity and subjectivity in scientific undertakings. The truly difficult question whether scientific evidence proffered in civil

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86 See amended FRE 702 (described in the text accompanying note 51, supra); see also *Kumho Tire*, 526 U.S. at 154 (stressing that the question for the court is not “the reasonableness in general” of a particular technique, but “the reasonableness of using such an approach . . . to draw a conclusion regarding the particular matter to which the expert testimony was directly relevant”) (emphasis in original); Mark P. Denbeaux & Michael Risinger, *Kumho Tire and Expert Reliability: How the Question You Ask Gives the Answer You Get*, 34 SETON HALL L. REV ___ (upcoming in Fall 2003).
cases should achieve a level of certainty that scientists themselves would require has not yet been resolved, but *Daubert* leaves room either to require that level or to admit scientific evidence on a lesser showing of significance. Which choice should be made here remains open to debate.

In the federal system, reviewing courts speak highly of the discretion that trial judges have in applying the *Daubert* standards, but a handful of states follow a different approach in allowing reviewing courts to appraise claims of error in applying *Daubert* on a *de novo* basis. These courts are doing the right thing, as trial judges need more extensive appellate guidance in handling science in civil cases under the *Daubert* standard.