ABSTRACT The Institute of Medicine report, Unequal Treatment: Confronting Racial and Ethnic Disparities in Health Care, claims that medical studies document a systematic causal relationship between race and disparities in health inputs and outcomes among individuals of different races. This article argues that the majority of studies are not powerful enough to establish a causal link, since they do not sufficiently control for differences among patients that happen to correlate with race, and it outlines a powerful audit study that could isolate any effect of race on health care decisions. Even if there are race-based disparities in health inputs, evaluations of welfare and policy prescriptions should be based on health outcomes, since the relationship between care and health is, at least in some cases, weak.

WHY IS THERE UNEQUAL TREATMENT between groups? The Institute of Medicine (IOM) report, Unequal Treatment: Confronting Racial and Ethnic Disparities in Health Care (Smedley, Stith, and Nelson 2003), makes the claim that a negative attitude—namely, “bias,” “prejudice,” and “discrimination”—on the part of nonminority physicians is one significant reason. This conclusion, in our view, is premature. In various sections of the IOM report, it appears to be regarded as an established fact, though it should remain a hypothetical factor.
This is not to say that clinical uncertainty and miscommunication within the doctor-patient relationship springing from racial differences might not exist; the possibility is quite plausible. The differences, theoretically, might even be based on invidious discrimination, though we doubt that. But their relative importance in determining health disparities remains unclear, especially when compared to factors such as access to care, quality of care, and health literacy—which have undisputed and sizeable influence (Aaron and Clancy 2003; Mechanic 2002). In military health settings, where access to care is excellent and quality and patient characteristics are relatively homogeneous, there are negligible disparities (Dominitz et al. 1998; Swift 2002; Taylor et al. 1997).

Words such as *prejudice*, *bias*, and *discrimination* represent charged and divisive language that is needlessly provocative and potentially counterproductive at this stage of our knowledge. This paper will discuss the kinds of evidence that would bestow a fuller picture of the dynamics involved in treatment disparities, allowing us to be more confident in our policy recommendations.

First, consider the data presented in the IOM report. The committee looked mainly at peer-reviewed retrospective studies. These studies often, but by no means always, revealed that minority patients were less likely to undergo complex procedures such as cardiac catheterization. The most rigorous studies tried to control for the many clinical or economic variables that might explain the differential (e.g., co-morbid illness, supplemental insurance, patient interest in procedure). But because the majority of the studies were retrospective and relied upon chart review or large Medicare databases, there were many variables that they could not capture, such as details of clinical presentation, EKG subtleties, patient preferences, and so on. Nor could doctors or patients be queried about their decision-making process. It is possible, too, that certain procedures and medications are used selectively because they are observed to yield differential outcomes by race. For example, black patients with chronic heart failure have been reported to derive less benefit from angiotensin-converting enzyme inhibitors, on average, than do whites (Exner et al. 2001; Wood 2001). A sizeable literature on pharmacogenomics suggests differential response to various medications and susceptibility to adverse effects by racial group (Holden 2003; Merikangas and Risch 2003; Wood 2001).

As the IOM report acknowledges, the more confounding variables that are identified, the smaller the differential between whites and minorities becomes: “almost all of the studies reviewed here find that as more potentially confounding variables are controlled, the magnitude of racial and ethnic disparities in care decreases” (Smedley, Stith, and Nelson 2003, 50).

In many cases, whatever residual remains between care of whites and minorities may be so small as to not be worth worrying about, but in other situations what is left may be important, and it represents fertile ground for serious investigation. For example, new research published since the release of the IOM report suggests that the profit status of a patient’s health plan can significantly
affect access to certain procedures. Schneider, Zaslavsky, and Epstein (2004) find that Medicare beneficiaries enrolled in for-profit health plans are more likely to receive some high-cost operative procedures than similar patients enrolled in not-for-profit health plans. (Once case-mix and demographic variables were controlled for, however, although for-profit enrollees had higher rates for all procedures, only partial colectomy and closed cholecystectomy were significantly higher.)

If it were in fact the case that enrollment in for-profit versus not-for-profit health plans is systematically related to race, then this research suggests that the IOM study suffers from an omitted-variables bias. That is, the observed racial differential might be an artifact of health plan type. Even if there were no discrimination at the physician decision level, a treatment differential would exist if minority patients were overrepresented in not-for-profit health plans. On the other hand, if minority patients are underrepresented in not-for-profit plans, the IOM differentials are biased downward.

Given that we are in the relatively early stages of research on race differences in medical treatments, there are likely to be scores of other factors that the IOM neglected to take into account (Zuvekas and Taliaferro 2003). Many of those factors could significantly alter the eventual policy prescriptions needed to alleviate any problem. Proceeding before we have more information about the sources of any race-based differential almost guarantees that we will not fix a problem if it does exist, and leaves the very real possibility that we could worsen health outcomes or increase health costs for everyone in the process.

**What Kinds of Investigations Could Illuminate the Recesses of the Treatment Gap?**

**Prospective Studies**

We need more prospective studies that ask doctors and patients about how they make decisions to offer and to accept, respectively, particular treatments. The following vignette shows just how difficult it is to interpret “bias” in medical records without an accompanying narrative from the clinician:

Kathy A. is a nurse practitioner in a public health clinic near Washington, D.C. She treats many young African American women. As part of the routine gynecological exam she asks them whether they had a PAP smear within the last two years. Typically, they say yes and Kathy A. does not perform one. When she started looking through records systematically, Kathy A. realized that many of the women who said they had a PAP smear never actually did. Soon she realized that many of the patients had mistaken a genital swab for STD for a PAP smear and has since kept this in mind during her history-taking (not to mention intensified her ongoing plea to the clinic director for computerized record-keeping). (Johns Hopkins School of Medicine 2003)
Conceivably, the innocent—though avoidable—mistake that Kathy A. made goes on daily in many inner city clinics. On chart review, Kathy A. would appear to be a (white) clinician who was short-changing black patients by not offering a routine PAP smear. But to allege that her error was born out of “prejudice,” “bias,” or “discrimination” is misguided.

We must also consider the matter of rational inference. To be sure, physicians base their clinical decisions on experience and statistical norms. These are sometimes influenced by race (or sex or class, for that matter). Furthermore, judgments that appear to be made on race may actually be made on other variables that simply correlate with race. Level of education is such a variable. As recent work by Dana Goldman and James Smith (2002) of RAND shows, adherence to treatment regimens in patients with HIV and diabetes varied greatly with the level of patients’ schooling; compliance, in turn, had a meaningful impact on patients’ overall health status.

In practical terms, if a physician thinks that a patient will not comply with triple therapy for HIV, he might either forgo the medication, give the patient a compliance “trial” wherein he must at least keep a second appointment in order to receive medication, or prescribe it while monitoring the patient especially closely. To the extent that a physician does the former, he has acted unethically, in our view. In this respect, the IOM report serves a consciousness-raising function, prompting doctors to ask themselves whether they are giving every patient the opportunity to benefit from treatment. Journal clubs or weekly rounds are good venues for group discussion of the potential miscommunications, clinical discretion, and problematic inferences that arise within the complexity of the doctor-patient relationship (Balsa et al. 2003). But to suggest that making clinical generalizations could rise to the level of “disparate impact discrimination” under Title VI of the Civil Rights Act, as the IOM report does, seems a large leap (Perez 2003).

Audit Studies

Audit studies are highly controlled, labor-intensive investigations in which only one variable—race, in this case—is altered. There are very few such studies in the area of health disparities research. In 1999, Kevin Schulman and colleagues at Georgetown University School of Medicine published an audit study in the New England Journal of Medicine. Briefly, the team made videos of black and white men and women who were actors playing patients with chest pain. The actor-patients were dressed in hospital gowns and described identical symptoms, and had the same EKG findings and health insurance. About 700 physicians viewed these tapes and were asked whether they would refer the “patients” to catheterization.

Schulman himself erroneously stated to the press that African Americans were 40% less likely to undergo catheterization and explicitly attributed the discrepancy to bias. The 40% estimate appeared to have been a misstatement based on an odds ratio finding, as demonstrated by a recalculation of the Schulman data.
by a team at the White River Junction VA (Schwartz, Woloshin, and Welch 1999). More accurately, white men, white women, and black men were referred at the same rate of 90%. Black women were referred at a mean rate of 80%, largely due to the low referral rate for a single black actress-patient—probably more a reflection of her acting than anything else. In all, the probability of referral for all black actors in the Schulman study was 7% lower than for whites, not 40%. As the White River Junction team wrote in the New England Journal of Medicine several months after publication of the Schulman article: “These exaggerations [of 40%] serve only to fuel anger and undermine the trust between physicians and their patients” (Schwartz, Woloshin, and Welch 1999).

Though there ended up being little difference in referral rates, the Schulman study galvanized the press. Perhaps the most egregious report appeared on the ABC news program Nightline. Here is how Ted Koppel introduced the segment: “Last night we told you how the town of Jasper, Texas, is coming to terms with being the place where a black man was dragged to his death behind a truck by an avowed racist. Tonight we are going to focus on [doctors] . . . who would be shocked to learn that what they do routinely fits quite easily into the category of racist behavior” (Nightline 1999). Not surprising, the IOM report was also greeted with sensationalism: “Color-Blind Care Is Not What Minorities Are Getting,” declared Newsday (2002); “Fed Report Cites ‘Prejudice’ in White, Minority Health Care Gap,” the Boston Herald charged (Lasalandra 2002); “Separate and Unequal,” said the St. Louis Post-Dispatch (2002).

To the extent that this kind of media coverage—and, indeed, the language of the IOM report itself—suggests that we know that important discrepancies exist and, further, that we know that the causal mechanism is racism, further research in this area will be stunted. The dearth of sophisticated research in this area will necessarily limit the ability of health care professionals to provide all of their patients with optimal treatment, and public health policy will also suffer.

Race Comparison Between Doctors

A third genre of study valuable for understanding race-related factors in treatment are ones comparing care provided by white and black doctors to white and black patients. For example, evidence that doctors of both races treat black patients similarly, say, in terms of rate of referral for catheterization—even if both refer black patients less often than they do white patients—would cause us to question a charge of bias. We are aware of only one study that has analyzed data with this question in mind.

Jersey Chen and colleagues at Yale University analyzed data from the Cooperative Cardiovascular Project (Chen et al. 2001). They evaluated 40,000 Medicare beneficiaries hospitalized for acute myocardial infarction in 1994 and 1995, to determine whether differences between black patients and white patients in the use of cardiac catheterization within 60 days after acute myocardial infarction varied according to the race of their attending physician. Black patients had sig-
Significantly lower rates of cardiac catheterization than white patients, regardless of whether their attending physician was white (rate of catheterization, 38.4% versus 45.7%) or black (38.2% versus 49.6%).

There was no significant interaction between the race of the patients and the race of the physicians in the use of cardiac catheterization, strongly suggesting that racial bias is not at issue. Critics of the Chen study have suggested that white cardiologists exhibited racial bias in their referrals for cardiac catheterization in regard to patients of the black attending physicians, but then, as Chen and colleagues reply, it would follow that black attending physicians concurred with and supported these racially biased decisions—a scenario they believe unlikely (Barr 2001; Chen, Rathore, and Krumholz. 2001). Moreover, the adjusted mortality rate among black patients was lower than or similar to that among white patients for up to three years after the infarction.

The mortality outcome in the Chen study raises an often-overlooked and perhaps counterintuitive point: differences in care do not inevitably translate into differences in outcome. That is, they do not invariably mean worse care. Granted, lower mortality may not reflect lower morbidity—another outcome variable that must be examined—but it adds an important, practical dimension to research in treatment disparity. The results of the RAND Health Insurance Experiment, conducted between 1974 and 1982, are instructive here. By randomly assigning subjects to different insurance arrangements, the researchers were able to induce different levels of care and expenditures unrelated to the subjects’ underlying health characteristics. By and large, the RAND research suggests that, in many contexts, increased treatment and expenditure levels do not translate into systematically better health (Newhouse 1993).

Chen’s finding of equal mortality between blacks and whites (including more favorable mortality rates for blacks) is by no means unique. In fact, according to a Kaiser Family Foundation (2002) review, the overwhelming majority of cardiac care studies found no mortality differences between races, despite lower rates of procedures for blacks. One possible explanation is that catheterization may be overused in white patients, meaning that the procedure is performed even when it probably will not benefit patients (Schneider et al. 2001).

One reason for this may be so-called “defensive medicine.” In a 1996 study, economists Daniel Kessler and Mark McClellan examined all elderly Medicare beneficiaries treated for heart disease in 1984, 1987, and 1990, to determine whether the liability exposure a doctor faces (as indicated by various medical malpractice reforms) affects treatment decisions. They found that while patients treated in states with lower liability exposure received almost 10% less in terms of medical expenditures, there was no statistically significant difference in mortality or medical complications as a result of these differences.

If doctors believe that white patients are systematically more likely to sue, or perhaps if the damages awarded or settlements reached in medical malpractice cases are significantly higher for white patients than minority patients in the
event of an adverse outcome (this might result from higher average incomes of the white population, which would translate into higher judgments for lost wages), doctors will have the incentive to engage in defensive medicine by expending more resources on white patients. If Kessler and McClellan’s results can be generalized, we would expect that white patients do not benefit, in terms of net health, from these additional health expenditures.

In any event, the important clinical issue suggested by comparable mortality findings is that we may be placing too much emphasis on tracking relative rates of specific forms of care when the critical issue is whether patients are getting relevant care.

Our Proposal for a Novel Study

What research design, then, would be needed to isolate discrimination in medicine? First, we must keep in mind that while demonstrating that doctors treat minority patients differently might be a necessary condition for proving the existence of systematic discrimination, it is far from sufficient. Sufficiency requires at least two additional findings. We must show that white doctors and minority doctors make systematically different decisions based on the patient’s race and, for those differences to be significant, we must show that this differential care results in worse health outcomes for the minority patients.

To investigate the issue of differences between white and minority doctors, we propose audit studies along the lines of the Schulman study mentioned above but with an important modification. The sample of doctors on whom the audit is run should be composed of pairs of white and minority doctors who are matched by as many non-race characteristics (years of experience, location, etc.) as is possible. Once the matching is done, the audit participants can (and should) be assured of complete anonymity, with the researcher only being able to tell whether the participant is a white or a minority doctor. The matching allows the researchers to define a presumptively non-discriminatory control group from the minority doctors.

This control group is important to net out the possibility that there may be medical reasons why a conscientious doctor would choose to treat white patients and minority patients differently. Although the most sophisticated audit studies make efforts to normalize the observable characteristics (except for race, in this case) of the test cases, competent doctors will necessarily draw probabilistic inferences about the patients’ unobservable characteristics based upon their own practice experience (see Heckman 1998). Assuming that white and minority doctors have similar estimates of the underlying distribution of unobservable characteristics, any differential treatment of white and minority patients found within the minority doctor control group should be treated as non-discriminatory. If the differential for white doctors exceeds that found in the control group, our confidence in drawing the inference of discrimination would increase.
However, even with such a finding, it might be premature to infer that significant discrimination exists. Despite a researcher’s best efforts, it will not be possible to match white and minority doctors perfectly. To mitigate the error induced by imperfect matching, replication of the audit results is important. Also, to answer the question of whether differential treatment translates into worse health outcomes for minority patients, researchers should use the audit results to inform the construction of prospective studies that focus on health outputs rather than health inputs. If both the audit studies and the prospective trials indicate that minority patients fare worse than their white counterparts, the inference of discrimination would be unavoidable. In the absence of such a rigorous research design, we are left with mere conjecture that does little to inform public health policy.

Thus, in order to speak in a more informed way about clinical uncertainty we require more of the kinds of studies just described—detailed prospective studies, audits, black-white doctor comparisons, and outcome analyses. Currently, there are very few. Yet many medical schools, health philanthropies, policymakers, and politicians are proceeding as if “bias” were an established fact. In other words, they consider part of the solution to the disparities problem to be located in the arena of race politics. This has led to some highly questionable developments—developments that the IOM report endorses.

For example, there is now a veritable “cultural competence training” industry that, among other activities, has been known to conduct patronizing racial sensitivity training for doctors (AAMC 2003). This is not to be confused with the need to learn local anthropology of unacculturated populations, a vital necessity for physicians who work with cultural minorities. In addition, medical schools are forthright in pursuing racial preferences wherein academic standards and performance are clearly lowered in the service of building a more racially diverse workforce (Cohen 2003; Satel 2000, 183–89). The main rationale for racial preferences is to funnel minority doctors into minority areas (based on the unfounded premise that minority patients overwhelmingly prefer a same-race doctor; Stinson and Thurston 2002). Yet a more fair and clinically responsible way to get good doctors into poor neighborhoods is to offer financial incentives. A third alternative for solving the disparity problem is the identification of civil rights violations under Title VI. In some cases, this activity might include bureaucrats parsing clinical decisions and second-guessing physicians (U.S. Commission on Civil Rights 1999, 14).

In our view, these race-based remedies pose a divisive distraction from more constructive solutions—solutions that the IOM promotes as well. These include the expansion of community health clinics (with evening hours and a responsive hotline for patients to call with acute medical questions) as well as grassroots outreach efforts to minority communities (Shin, Jones, and Rosenbaum 2003). These efforts, underway across the country, are designed to enhance knowledge...
about common health problems and healthy lifestyle; perform screenings (e.g., for hypertension, breast cancer); and educate about organ donation. Community-based interventions draw on local members as educators, translators, and nursing/physician assistants. To the extent that many minority patients are distrustful of the more conventional medical establishment—a claim frequently made by the proponents of racial preferences in medical schools—receiving care from and interacting with culturally compatible staff should be conducive to compliance with care. Doubtless, for patients who cannot easily travel to major hospital centers (a definite problem for people with disabilities and inflexible working hours), local clinics will improve access significantly.

Understanding health disparities as an economic problem tied to issues of access to quality care and health literacy, rather than a civil rights problem borne of overt or unconscious bias on the part of physicians, is a more efficient and rational way to address the problem of differential health outcomes.

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