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Domestic AIDS Vaccine Trials:
Addressing the Potential for Social
Harm to the Subjects of
Human Experiments

Philip A. Leider†

In 1998, the FDA approved the first large-scale human trials of a candidate AIDS vaccine in our nation's history. While the legal issues raised by these trials are manifold, the academic literature has focused almost exclusively on the potential for mass tort liability and the resulting hesitancy of biotech and pharmaceutical firms to enter the field. This Comment argues that another issue of vital concern demands attention: the potential for social harm to the human subjects of AIDS vaccine trials. After providing an overview of the current epidemiology of HIV/AIDS and explaining why a safe, effective AIDS vaccine represents the best way to control the pandemic, this Comment analyzes the scientific and social obstacles to production of such a vaccine.

In order to know whether a candidate AIDS vaccine is truly effective, researchers will have to test the product in HIV-negative volunteers at high risk of infection. Since these volunteers may subsequently test positive for HIV on standard blood tests, they will be vulnerable to discrimination on that basis in such areas as employment, insurance, immigration, and

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incarceration. Moreover, by participating in vaccine trials, volunteers will be marking themselves as people at high risk of HIV infection, another basis for disparate treatment. Researchers have suggested that federal disability discrimination law may afford protection against research-related social harms. Through close analysis of the Americans with Disabilities Act of 1990 and the Supreme Court’s decision in Bragdon v. Abbott, this Comment demonstrates that optimistic reliance on federal disability law is misplaced. The unique issues raised by domestic AIDS vaccine trials must be addressed in their own right. The Comment accordingly concludes with a broad range of legislative and regulatory proposals to protect trial participants and advance the AIDS vaccine research agenda.

INTRODUCTION

In May of 1998, the U.S. Food and Drug Administration (FDA) granted permission for the first domestic efficacy trial of a preventive AIDS vaccine in our nation’s history. The vaccine, AIDSVAX, has already been tested for safety in over 1,200 human subjects. In Phase II trials, AIDSVAX proved safe, with no significant side effects, and it induced HIV antibody response in most vaccinated individuals. Five thousand HIV-negative volunteers in clinics across North America, Puerto Rico, and the Netherlands will now offer up their forearms to see if AIDSVAX is not only safe but also effective in preventing HIV infection.
Although it is a genuine cause for celebration in the battle against AIDS, the progress to large-scale testing of AIDS vaccines such as AIDSVAX also raises a host of vexing legal issues. What will happen if trial participants have adverse physical reactions to candidate vaccines? Who will pay for their medical care? Should insurance companies have to pay for injuries caused by experimental products? Or should biotechnology companies, like VaxGen, be liable for trial-related injuries? If the latter, are these smaller spin-offs of larger corporations sufficiently capitalized to compensate victims? If they are not, will courts pierce the corporate veil and make larger corporate shareholders, like Genentech, bear the risk? Or should the state subsidize the efforts of companies like VaxGen—those that take risks in the interest of public health—by shielding them from unlimited tort liability and paying victims out of some form of state fund? These and many other questions clustered around injury compensation and tort relief for biotechnology and pharmaceutical companies have dominated the burgeoning legal literature on AIDS vaccine research.\footnote{Eclipsed by the potential for mass tort liability, another issue of substantial concern has received scant legal attention thus far: The prospect of social harms facing participants in AIDS vaccine trials.\footnote{Although the legal community has been slow to awaken to it, researchers and AIDS activists have foreseen the problem of vaccine trial participants being subjected to social harms. See, e.g., Christine Grady, The Search for an AIDS Vaccine: Ethical Issues in the Development and Testing of a Preventive HIV Vaccine 141-42 (1995); Chris Collins, Sustaining Support for Domestic HIV Vaccine Research: Social Issues Over the Long Haul of Human Trials 9 (Center for AIDS Prevention Studies, Univ. of Cal. San Francisco, Monograph Series, Occasional}
trial serves as a good illustration: In Phase II trials of the product, 95.5% of vaccinated individuals generated HIV antibody responses. Because standard blood tests for HIV detect the presence of HIV antibodies, it is possible, if not probable, that many of these individuals will test positive for HIV even though they are not actually infected.

What will happen if these trial participants test positive for HIV when they apply for health insurance or for a job? How will employers, prison officials, customs officers, and others who rely on medical information regard a positive ELISA test? If they react negatively, should trial participants reveal that they are taking part in an AIDS vaccine trial? Since AIDS vaccine trials are widely publicized as enrolling only "high-risk individuals" (for example, gay men, prostitutes, and injection drug users), does revealing trial participation simply substitute one potential basis of disparate treatment for another? Finally, who would enroll in an AIDS vaccine trial given the evident physical and social risks, and how, as a result, will five thousand volunteers for the Phase III trial of each candidate vaccine be assembled?

This Comment addresses the concerns outlined above by advancing a four-fold thesis: (1) the world needs a safe, effective preventive vaccine to curb the global AIDS epidemic; (2) production of such a vaccine will

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8. See VaxGen, supra note 2.

9. The ELISA, or enzyme-linked immunosorbent assay, is a blood test that detects the presence of antibodies to HIV in the blood. See NIAID, HIV Vaccine Glossary (visited Apr. 3, 2000) (<http://www.niaid.nih.gov/factsheets/GLOSSARY.htm>) (defining "ELISA"). It is commonly used as an initial screening test for HIV because it is "relatively easy and inexpensive to perform." Id. Because the ELISA test often registers false positive results, identifying antibodies where there are none, it is usually confirmed by "a second, more specific test such as an HIV Western Blot." Id.

One problem in AIDS vaccine trials is that standard assays do not readily distinguish HIV antibodies from vaccine-induced antibodies. As NIAID explains,

[id] when people develop antibodies to HIV or an experimental HIV vaccine, they "seroconvert" from antibody-negative to antibody-positive. Vaccine-induced seroconversion does not represent an infection. Instead, vaccine-induced seroconversion is an expected response to vaccination that may disappear over time.

Id. (defining "seroconversion"). Thus, even though researchers "expect" vaccine-induced seroconversion, standard blood tests may not register a difference between infection and vaccination. As a result, researchers can anticipate and test for vaccine-induced antibodies using specialized tests, while the general public might easily confuse vaccination with HIV infection. For a helpful overview of currently available blood assays and their relative advantages and disadvantages, see Niel Constantine, HIV Antibody Testing (last modified Feb. 1998) (<http://hivinsite.ucsf.edu/akb/1997/02abtest/index.html>.

10. See discussion infra Part III.A.

11. Due to the double-blind structure of vaccine trials, discussed infra Part II.D, participants will not be informed whether they have received vaccine or placebo. As a result, they will not readily know whether they have actually seroconverted or are simply reacting to vaccine. As discussed infra Part III.A, researchers are offering on-site testing and participant ID cards as possible responses to this problem.
DOMESTIC AIDS VACCINE TRIALS

take place only if human subjects are reasonably safeguarded against physical and social harm; (3) extant legal and regulatory protections will not suffice to minimize trial-related social harms, and, correlatively, this will adversely affect trial enrollment in the long-term; and (4) appropriate federal measures should be taken to protect research subjects and enhance the prospects for a successful AIDS vaccine.

The Comment is divided into five Parts. Part I provides an overview of the current epidemiology of AIDS. After fleshing out why the treatment options currently available will not suffice to curb the global AIDS epidemic, Part I outlines the rationale behind preventive intervention and the considerable advantages a safe, effective AIDS vaccine could provide.

Part II discusses the basic scientific obstacles to AIDS vaccine research that make the participation of human subjects at high risk of infection essential to determining the efficacy of a candidate vaccine. Since individuals at high risk of infection are often also socially vulnerable, Part II links the scientific barriers facing AIDS vaccine research to the social impediments at the core of the Comment's analysis.

Part III focuses on the specter of social harms to human subjects in domestic AIDS vaccine trials. Since candidate vaccines may make volunteers test positive for HIV on standard assays,12 trial participants will be made vulnerable to the various forms of discrimination facing HIV-positive people today. In addition, AIDS vaccine trial participants will be marking themselves as members of a "high-risk group" since the entry criteria for vaccine trials will necessarily focus on risk behavior such as anal sex or injection drug use.13 Research subjects will accordingly face disparate treatment on at least two grounds: Their apparent HIV-positive serostatus and their membership in a group at high risk of exposure to the AIDS virus. Without some form of ex ante legal protection, potential human subjects will be exposed to considerable prospective harm and, as a result, will be discouraged from participating in clinical vaccine trials.

Part IV begins this Comment's analysis of potential sources of legal protection for AIDS vaccine trial participants by focusing on the Americans with Disabilities Act of 199014 (hereinafter "ADA" or "the Act"). Part IV evaluates whether AIDS vaccine trial participants who falsely test positive for HIV will be protected by the ADA, as researchers have optimistically suggested. It concludes that ADA jurisprudence is not sufficiently settled to answer the question definitively. It further illustrates that disability law is, at best, an awkward fit for trial subjects. Part IV therefore underscores the need for a legislative and regulatory regime uniquely tailored to large-scale vaccine trials.

12. See discussion infra Part III.A.
13. See discussion infra Part III.B.
Part V proposes a series of federal legislative and regulatory measures to address the problem of social harms to AIDS vaccine trial participants. These measures must take stock of the existing patchwork of regulations and procedural controls, such as the role of Institutional Review Boards in the design and oversight of study protocols, that could be enhanced. In addition, the proposed measures must be coordinated with related measures, such as limited tort relief for biotechnology and pharmaceutical companies and an indemnification fund for trial-related injuries, to give the U.S. a coherent, effective AIDS vaccine policy.

I

THE NEED FOR AN AIDS VACCINE

A. Current State of the AIDS Epidemic

It is customary to begin any discussion of AIDS vaccine research with what one report aptly calls "a recapitulation of the latest epidemiological horrors."\(^{15}\) The latest global statistics are indeed grim. Estimates by the Joint United Nations Programme on HIV/AIDS (UNAIDS) and the World Health Organization (WHO) indicate that 16.3 million people around the world have died of Acquired Immune Deficiency Syndrome (AIDS) since the beginning of the epidemic.\(^{16}\) By the end of 1998, over 33.6 million people were living with Human Immunodeficiency Virus (HIV), the virus that causes AIDS.\(^{17}\) In addition, an estimated 5.6 million people worldwide were newly infected with HIV in 1999.\(^{18}\)

In sharp contrast to the rest of the world, Western nations have experienced a rapid decline in AIDS cases.\(^{19}\) The dramatic turnaround can be attributed in large measure to remarkable advances in the therapeutic treatment of HIV. Most notable among these advances has been the advent

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17. See id.
18. See id.
of protease inhibitors and so-called "combination" or "triple cocktail" therapy. However, these treatment regimens are expensive and difficult to adhere to; the drugs often require refrigeration and they can cause toxic side-effects in HIV-infected individuals. As a result, developing countries, where the AIDS pandemic is spreading uncontrollably and the infrastructure to deliver medication is largely non-existent, have not appreciably benefited from the latest advances. The result is as predictable as it is lamentable: Western nations are generally seeing a steep decline in the rate of AIDS-related deaths (accompanied by a large increase in the number of people living with HIV), while in developing countries both infection and mortality rates continue to climb precipitously.

B. Recent Advances Do Not Eliminate the Need for Preventive Intervention

It is important to recapitulate the "latest epidemiological horrors," then, for at least four reasons. First, the media have loudly trumpeted recent advances in AIDS treatment. Anecdotal evidence suggests that people who hear of these advances are increasingly engaging in risky behavior, presumably because fear of disease is on the decline. In other


21. The situation in southern Africa is particularly stark. According to UNAIDS and WHO estimates, seven out of ten people who became infected with HIV in 1998 live in sub-Saharan Africa. Among children under fifteen years of age, the proportion is nine out of ten. Of all AIDS deaths since the epidemic began in 1981, 83% have occurred in sub-Saharan Africa. Only one-tenth of the world’s population, however, lives in Africa south of the Sahara. See UNAIDS, AIDS in Africa (last modified Nov. 30, 1998) <http://www.unaids.org/publications/documents/epidemiology/determinants/esaepap98.html>. Needless to say, antiretroviral therapy regimens have not yet had an impact in sub-Saharan Africa.

22. See MAP NETWORK, supra note 19, at 7.


24. "Barebacking," that is, engaging in anal sex without a condom, is reportedly on the rise among gay men. See, e.g., Robert B. Hays et al., High HIV Risk-Taking Among Young Gay Men, 4
words, as people come to regard HIV as a treatable disorder, their fear of infecting others or getting infected themselves may correspondingly decrease. Needless to say, current HIV prevention efforts will founder if safer-sex and clean-needle campaigns must constantly counterspin misleading media coverage of advances in AIDS treatment. The irony is patent: Advances in AIDS treatment may increase, to some degree, the incidence of HIV infection.

Second, while treatment efforts have shown remarkable success in industrialized countries, there has been little headway in developing countries. The success of new treatments has virtually blinded people in Western countries to the continuing ravages of the pandemic elsewhere in the world: Their frame of reference no longer includes the unchecked progression of disease still so prevalent "over there." Since the resources to combat the disease remain largely in Western hands, awareness of the epidemic's worldwide effects remains a vital component of AIDS advocacy and political lobbying efforts (for example, to increase availability of antiretroviral therapies in developing countries or sustain domestic research


25. See generally Jeffrey A. Kelly et al., Protease Inhibitor Combination Therapies and Perceptions of Gay Men Regarding AIDS Severity and the Need to Maintain Safer Sex, 12 AIDS F91 (1998) (finding that a substantial number of individuals believe AIDS is now nearly cured and concluding that it is necessary to temper the optimism regarding treatment advances); S. Murphy et al., Antiretroviral Drugs and Sexual Behavior in Gay and Bisexual Men: When Optimism Enhances Risk, 12 INT'L CONF. AIDS 209 (1998) (concluding that individuals engaging in risky sex are either rationalizing their behavior through optimism about the effectiveness of treatments or are actually less concerned about contracting or transmitting HIV); J. Toeppeich ct al., Does Safer Sex Survive the Therapeutic Optimism in the General Public?, 12 INT'L CONF. AIDS 947 (1998) (reviewing comparative historical data drawn from questions posed during the German national AIDS prevention campaign).


27. The UNAIDS HIV Drug Access Initiative commenced in late 1997. It aims to make HIV-related drugs more accessible to broad sectors of populations in developing countries. The Initiative has begun pilot-testing of antiretroviral therapies in Chile, Vietnam, Uganda, and the Côte d'Ivoire. See UNAIDS, UNAIDS Launches Initiative to Help Bridge Gap in Access to HIV/AIDS-Related Drugs in Developing World: Multinational Pharmaceutical Companies Already Committed to Making Drugs More Affordable (last modified Nov. 5, 1997) <http://www.unaids.org/whatsnew/press/eng/pressare97/drug.html>. Organizations such as the International AIDS Vaccine Initiative (IAVI), the AIDS Vaccine Advocacy Coalition (AVAC), the American Foundation for AIDS Research (amfAR), and the Elizabeth Glaser Pediatric AIDS Foundation (PAF) are also striving to keep the global nature of the epidemic squarely before philanthropic and governmental bodies. See AVAC, EIGHT YEARS AND COUNTING: WHAT WILL SPEED DEVELOPMENT OF AN AIDS VACCINE? 47-51 (1999).
for an AIDS vaccine). Successful vaccine efforts in the past, such as those for polio and smallpox, have originated in Western countries; it is therefore crucial that the citizens of these countries maintain an undistorted understanding of the pandemic's scope so as to spur further research.

Third, although current treatment options may help to stave off progression to disease, they do not do so uniformly. Many people cannot adhere to the exacting regimen of pills and diet involved; others do not respond well to the medication; some have even developed drug-resistant strains of the virus. And even with perfect compliance, none of the current antiretroviral therapy regimens has proven to be completely effective against the disease. More significantly, the treatments are largely unavailable in developing countries. It remains to be seen whether treatments developed in the West will prove as effective against other "clades" or genetic subtypes of the virus more prevalent in other parts of the world. Moreover, given HIV's unfortunate ability to mutate, new strains of the virus are likely to emerge and familiar strains may themselves eventually become resistant to medication. In short, the long-term prospects for today's promising treatment regimens remain, as yet, uncertain.

Fourth and finally, although decreases in domestic mortality rates are a genuine cause for optimism, the great expense of caring for people living with HIV/AIDS vastly increases as more people survive for longer periods of time. While wealthy nations such as the U.S. may be able to sustain the

31. The side effects of the various antiretroviral drugs are meticulously catalogued by Drs. Paul Volberding and Steven G. Deeks in Antiretroviral Therapy for HIV Disease (last modified Sept. 1999) <http://www.hivinsite.ucsf.edu/akb/1997/04arvtx/index.html>.
33. See discussion supra Part I.A.
36. The drugs for the combination therapy cocktail alone cost $10,000 to $15,000 annually, and most HIV/AIDS patients require several other drugs as well. See Linda C. Fentiman, AIDS as a Chronic Illness: A Cautionary Tale for the End of the Twentieth Century, 61 Alb. L. Rev. 989, 1004
heavy economic burden imposed by chronic AIDS care, other economies cannot absorb the crushing expense. In countries where famine and other fatal diseases pose imminent threats to the population, paying the high price required for AIDS treatments is difficult to rationalize. Even in wealthier countries, AIDS threatens to overwhelm well-funded public health systems both logistically and financially. Briefly put, the massive expense of AIDS treatment militates for a viable preventive option.

In sum, therapeutic treatment of the AIDS virus, though promising, will not do the job alone. As AIDS treatments progress, people might be more likely to become infected with HIV due to a concomitant decrease in the fear of AIDS. In addition, current treatment regimens are difficult to comply with, and their long-term efficacy is still in question. Even if combination therapy were completely effective, it remains an option open only to industrialized nations at this point. Antiretroviral medications have toxic side-effects in some infected individuals, they are difficult to preserve in transport, and it is far from certain that they will prove effective against variant strains of the virus. They are also prohibitively expensive for most countries, the infrastructure is not in place to deliver them or monitor their effects, and there are cultural divides that make compliance with the rigorous regimen involved highly challenging. All of these factors limit the global impact of promising new AIDS therapies. Treatment of AIDS must be supplemented with effective strategies to prevent infection with HIV in the first instance.

(1998); see also NATIONAL AIDS STRATEGY, supra note 19, at 9 (“The average lifetime cost of medical care after an HIV diagnosis is $119,000.”).

37. In fiscal year 1998, total federal HIV/AIDS spending was estimated to be $8.7 billion. This includes amounts for programs specifically targeted to HIV/AIDS, as well as amounts designated by federal agencies as HIV/AIDS spending within more general activities. Compared to total federal spending, estimated to be $1.8 trillion, funding for HIV/AIDS represents approximately one half of one percent of total federal spending. See Scott Foster et al., Federal HIV/AIDS Spending: A Budget Chartbook (visited Mar. 26, 2000) <http://www.hivinsite.ucsf.edu/social/kaiser_family_found/2098.3d3f.html>.

38. See NATIONAL AIDS STRATEGY, supra note 19, at 29; Fentiman, supra note 36, at 1005; Naomi Freundlich et al., AIDS: Hope is Where the Money is, BUSINESS WEEK, July 22, 1996, at 34.

39. “Companies in hard-hit countries are losing trained staff at rates unheard of in the industrialized world. Extra staff are hired in anticipation of workforce losses to AIDS. Profits are eaten up by health care costs and work-hours lost to illness and attendance at funerals.” UNAIDS, UNAIDS Executive Director Warns of Unprecedented Emergency in Southern Africa (quoting Dr. Peter Piot, UNAIDS Executive Director) (last modified Nov. 30, 1998) <http://www.unaids.org/whatsnew/press/eng/pressarc98e.html>.

C. The Goal: A Safe and Effective AIDS Vaccine

As the dimensions of the AIDS pandemic continue to expand, a growing international cadre of scientific, political, and philanthropic organizations and leaders is calling for the production of a safe and effective preventive vaccine against the disease.\(^{41}\) Citing the global eradication of smallpox and poliomyelitis as precedents, they argue that an AIDS vaccine is the only comprehensive way to stem the epidemic worldwide.

A vaccine is "a substance used to teach the immune system how to defend itself against a pathogen, such as a disease-causing virus or other organism."\(^{42}\) Deliverable in various ways, a vaccine helps the body to resist infection or progression to disease: "If a vaccine is administered before the body's exposure to a pathogen, it can help the body to completely rid itself of the pathogen or to at least control the pathogen enough to prevent the development of clinical manifestations of disease and to hinder transmission."\(^{43}\) A safe and effective HIV vaccine ideally would "be inexpensive to manufacture, provide protection against all subtypes of HIV, require minimal if any boost, protect against all methods of spread of HIV for years, and be easily administered, stable to heat, and widely accessible."\(^{44}\) Even if the "ideal" AIDS vaccine cannot be created, though, a substantially protective vaccine will accomplish much.

Vaccines present considerable advantages over established modes of HIV prevention such as: (1) risk-reduction counseling, which seeks to help people alter their high-risk behaviors; (2) explicit safer-sex education programs; (3) condom distribution; and (4) provision of sterilized needles to injection drug users. Each of these prevention techniques has disadvantages that reduce its efficacy. Counseling and education programs rely on genuine behavioral change, which is notoriously difficult to achieve and sustain over time.\(^{45}\) Condoms break, and for reasons ranging

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43. Id. Researchers distinguish between preventive vaccines (those that help the body prevent or resist infection with HIV, the virus that causes AIDS) and therapeutic vaccines (those that inhibit progression to AIDS once an individual is already infected with the virus). See NIAID, supra note 9 (defining "preventive vaccine" and "therapeutic vaccine"). Throughout this Comment, I use the term "AIDS vaccine" to mean a vaccine aimed at preventing infection by HIV unless otherwise indicated.

44. Margaret I. Johnston & Sam Avrett, Developing HIV Vaccines and Other Interventions to Prevent AIDS Worldwide, in TEXTBOOK of AIDS MEDICINE 725, 728 (Thomas C. Merigan, Jr. et al. eds., 1998).

from physical discomfort to religious objection, people resist using them.\textsuperscript{46} Although sterilized needle campaigns have proven successful in reducing rates of HIV transmission among injection drug users, political and moral resistance to needle distribution has thrown up seemingly insurmountable roadblocks in this area.\textsuperscript{47} In short, current modes of HIV prevention, though assuredly a step in the right direction, have failed to contain the epidemic.\textsuperscript{48}

It is easy to see why a safe, effective vaccine presents an attractive alternative. Once inoculated, an individual would not have to completely alter long-engrained patterns of behavior to remain protected. Nor would lapses into risk behavior or barrier failures represent the threat they now do; an effective vaccine would offer a second layer of protection. A vaccine would also obviate, to some degree, making clean needles available to injection drug users, a campaign that has drawn vociferous opposition from those who see it as an encouragement to use drugs. Finally, many of the drawbacks to AIDS treatments, including prohibitive expense, compliance issues, toxicity, and viral resistance, would be eliminated for a large number of individuals who might otherwise become infected. Production and dissemination of a safe, effective vaccine against HIV would be the ideal preventive measure in the fight against AIDS.

An AIDS vaccine will not, however, magically spring from a test tube like Venus from the foam. The obstacles to discovery, production, and dissemination of a vaccine are formidable, if not insurmountable. Because the design of AIDS vaccine trials raises novel and important issues, I will briefly describe the basic scientific obstacles facing researchers before


\textsuperscript{48} Targeted intervention has helped decrease infection rates in the gay community, once considered the epicenter of the epidemic. The global trend towards heterosexual and IV drug transmission, though, has fast outstripped any gains in targeted communities. See discussion supra note 19 and accompanying text.
turning to the legal and political hurdles facing AIDS vaccine research more generally.

II

SCIENTIFIC OBSTACLES TO AIDS VACCINE RESEARCH AND THEIR EFFECT ON TRIAL DESIGN

A. The Unique Challenges of AIDS Research

Although vaccine science flowered in the twentieth century, HIV/AIDS poses unique challenges to vaccine researchers. In the early years of the AIDS epidemic, isolation of the causative agent and identification of the pathways to infection were the paramount issues for scientists.\textsuperscript{49} Public uncertainty and fear about the causes and contagiousness of the disease were widespread; rumors of a "gay pneumonia" hit the newspapers,\textsuperscript{50} and AIDS was treated as what one court has called "the modern day equivalent of leprosy."\textsuperscript{51} Since the underlying immunosuppressive disease was made visible only by the various opportunistic infections that set in—Kaposi’s Sarcoma (KS) lesions, candidiasis (thrush), and the rattling cough of pneumocystis carinii pneumonia (PCP)—people saw the lethal effects of infection and remained extremely cautious about even casual contact with those infected.\textsuperscript{52}

Once HIV was isolated as the etiologic agent that causes AIDS\textsuperscript{53} and the principal modes of communication of the virus were identified, many other advances followed. The first test for HIV was soon available,\textsuperscript{49} and the intellectual property rights to the discovery precipitated infamous internecine battles between Gallo and Montagnier.\textsuperscript{54}
making epidemiological tracking of the virus possible. HIV testing also enabled preventive intervention since individuals could find out whether they were infectious before progressing to full-blown AIDS. In addition, discovery of the pathways to infection aided preventive efforts because identifying people infected or at risk of infection allowed for targeted intervention, for example, condoms for those at sexual risk, or blood-product testing for those in need of transfusions.

Basic science also profited from isolation of the virus. Scientists were now able to begin natural history studies of AIDS to begin to understand the way HIV infection progresses to clinical AIDS and eventually to death. HIV was identified as a retrovirus, that is, a virus that uses the replicatory machinery of the cells it infects to reproduce itself. HIV was also identified as a lentivirus, that is, a virus that engenders acute symptoms in the blood upon infection and then remains clinically latent for a long period as it progresses clandestinely throughout the lymphatic system. These insights helped scientists devise antiretroviral therapies such as AZT and, later, protease inhibitors, which have shown great promise in slowing progression to AIDS after initial infection with HIV.

While advances in the understanding and therapeutic treatment of AIDS have been remarkable, several intractable issues have hampered vaccine research. Primary among these have been: (1) variability of the AIDS virus; (2) lack of an appropriate animal model for HIV/AIDS; and (3) uncertainty as to the correlates of protection from the virus. Since these three obstacles affect trial design and, consequently, the legal and ethical dimensions of vaccine trials, I will discuss each briefly.

57. See Donald P. Francis et al., Targeting AIDS Prevention and Treatment Toward HIV-1-Infected Persons: The Concept of Early Intervention, 262 JAMA 2572, 2572 (1989); GRAY ET AL., supra note 52, at 7; NATIONAL AIDS STRATEGY, supra note 19, at B-31; REPORT, supra note 40, at 83-91.
B. Variability of the AIDS Virus

Several different "clades" or genetic sub-types of HIV exist worldwide.64 Five to seven principal subgroups of the virus have been identified.65 Although these variations are generally divisible geographically, two or more viral clades can exist in a given region. In fact, more than one clade can infect a single individual.66 More commonly, variant sub-strains of HIV develop within individuals as the virus mutates irregularly in the body.67

One of the most formidable challenges to AIDS vaccine research, then, is to identify a vaccine or vaccines that can protect against all of the extant variations of the virus.68 Logistically speaking, of course, this means that vaccine trials cannot be conducted solely in one region of the world.69 The viral subtype predominant in Thailand, for example, may overcome vaccine-induced immune defenses specific to viral subtypes generally found in the U.S. and Europe. The genetic variation in the AIDS virus, therefore, puts a premium on international cooperation and coordinated research efforts on a global scale.70

Cultural and political differences among the nations can also raise vexing research issues. For instance, western-style informed consent may not "translate" in a country where the infection rate is high and the pressure to devise a viable preventive mechanism is strong.71 Moreover, the inherent danger in an AIDS vaccine trial—that subjects might believe they

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65. See Grady, supra note 7, at 99.
66. See Jon Cohen, Can One Type of HIV Protect Against Another Type?, 268 SCIENCE 1566, 1566 (1995); J. Travis, HIV-2 Offers Protection Against HIV-1, 147 SCI. NEWS 373, 373 (1995).
67. See NIAID, CHALLENGES IN DESIGNING HIV VACCINES 3 (1997).
68. See id.
69. In vitro assays might seem to be a logical alternative, but HIV antibodies in a test-tube (laboratory isolates) behave differently than HIV "out there" in the world (field isolates). "What is needed is a vaccine capable of inhibiting a spectrum of field isolates, not just laboratory strains of virus." Grady, supra note 7, at 100.
70. See discussion supra notes 27-29 and accompanying text. The ethical and legal issues presented by international drug trials are very complex and, as such, they lie beyond the scope of this Comment. Except for brief allusions, the focus here is restricted to domestic AIDS vaccine trials. For more extensive treatment of the ethical issues at stake in international trials, see UNAIDS, ETHICAL CONSIDERATIONS IN HIV PREVENTIVE VACCINE RESEARCH (2000). On some of the legal issues presented, see John P. Wilson, Limitation of Manufacturer Liability for Administration of an AIDS Vaccine Overseas, 30 INT'L LAW. 783 (1986).
are protected and engage in heightened risk behavior as a result—may prove to be particularly acute in cultures unfamiliar with Western forms of medicine. Finally, stigmatized groups, such as homosexuals in many countries, may encounter severe forms of persecution associated with trial participation. In sum, the genetic variation in the AIDS virus necessitates trial design on an international scale with all of the attendant cultural challenges.

C. The Lack of an Appropriate Animal Model for HIV/AIDS

Clinical research involving experimental substances is usually preceded by testing in animals. Generally speaking, before a candidate vaccine is approved for testing in humans, it must be demonstrated to be safe, immunogenic, and effective in animals. Scientists usually evaluate a new vaccine by testing its ability to generate antibodies in animals. They then test the effectiveness of these antibodies in neutralizing the virus through in vitro laboratory tests. With some diseases, however, animal infection does not closely mimic the characteristics of infection and immune response in humans. In such cases, the analogy between animal disease and human disease breaks down, making it difficult to draw reliable inferences from the animal data. Initial testing, therefore, is often a "guessing game until the human model is used."

To date, no suitable animal models for HIV and no reliable in vitro markers for efficacy have been identified. The closest candidate is Simian Immunodeficiency Virus (SIV). SIV infects several subspecies of monkey, and macaques have shown varying degrees of protection against SIV challenge after vaccination with a variety of SIV vaccines. In the final

72. See discussion supra notes 24-26 and accompanying text.
74. "[T]he vaccine . . . must produce protective immunity with only minimal side effects (such as redness and soreness at the vaccination site) for the overwhelming majority of those who receive it." NIAID, supra note 1, at 4.
75. "The vaccine . . . must cause a strong and measurable immune response." Id. at 6.
76. The vaccine must "produce a desired clinical effect, such as protection against a specific disease, at the optimal dosage and schedule in a given population." Id. at 30.
77. See Grady, supra note 7, at 25.
78. Id.
79. Id. at 96.
analysis, though, "SIV is a surrogate for HIV, and it is not known whether successful SIV strategies can be translated to HIV."\(^8\)

Chimpanzees are the only animals that researchers have successfully infected with HIV.\(^8\) Although they exhibit HIV-like symptoms upon infection, chimps do not develop T-cell deficiencies or other disease symptoms similar to AIDS in humans.\(^8\) Chimps are a relatively rare species requiring extremely expensive care, which further inhibits experimentation.\(^8\) If researchers could pinpoint why chimpanzees do not progress to disease, it might be a significant advance in the treatment of AIDS. Currently, though, chimpanzees represent an expensive and incomplete analogue for human vaccine research.\(^8\)

The lack of an appropriate animal model raises a thorny question in much of the HIV/AIDS research to date: Should researchers be testing experimental substances in humans before they are known to be safe, immunogenic, and protective in animals?\(^8\) Since a preventive vaccine must, by definition, be tested in HIV-negative human subjects, the stakes are quite high: AIDS is currently a fatal disease with no known cure. Without reliable animal data, researchers take the risk that some individuals will expose themselves to HIV based on inaccurate assumptions about vaccine

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81. Grady, supra note 7, at 97.
83. See Grady, supra note 7, at 97.
84. At approximately $70,000 per chimp, animal tests exact a prohibitive toll for many research efforts. See id. at 99. Grady further notes that animal research has been limited by "sociopolitical concerns about the use of animals in research." Id. at 25.
85. This is not to say that primate research represents a dead-end in the search for an AIDS vaccine. As NIAID has recognized, important information has been obtained from both monkeys and chimpanzees. Experiments in both species have demonstrated the feasibility of developing a protective vaccine. Moreover, a new animal model—infection of macaques with a chimeric virus (SHIV) based on SIV but including the HIV envelope, with subsequent development of disease—may become extremely valuable for evaluating candidate HIV vaccines.
NIAID, supra note 67, at 6.
86. As Hans Jonas presciently argued, the transition from animal research to human research always involves a risk of this sort: "Up to a point, animals may fulfill the proxy role of the classical physical experiment. But in the end man himself must furnish knowledge about himself, and the comfortable separation of noncommital experiment and definitive action vanishes." Hans Jonas, Philosophical Reflections on Experimenting with Human Subjects, in LAW, SCIENCE AND MEDICINE 987, 988 (Judith Arcen et al. eds., 1996). The structural necessity to cross the threshold from animal to human experimentation, however, does not mean that ethical categories are thereby voided. On the contrary, Jonas asserts that "[h]uman experimentation for whatever purpose is always also a responsible, nonexperimental, definitive dealing with the subject himself. And not even the noblest purpose abrogates the obligations this involves." Id.
efficacy. On the other hand, the epidemic is spreading at such an advanced rate that a genuinely effective vaccine could save millions of lives. The societal benefits from vaccine research arguably outweigh the individual risks voluntarily assumed by trial participants.\textsuperscript{87} Such a risk/benefit analysis now supplies an \textit{ex post} rationalization for the AIDS vaccine effort. As Grady notes: "Because of the urgency of the public health need and public attention and political pressure, human Phase I trials of HIV candidate vaccines began before efficacy was demonstrated in chimpanzees or any other animal model."\textsuperscript{88}

\textbf{D. Uncertainty as to the Correlates of Protection}

Once testing begins in humans, researchers face another important question: How will they know if a candidate vaccine has actually resulted in some level of protection to an individual? Researchers do not currently know what a protected individual looks like because humans naturally immune to HIV infection have not been identified.\textsuperscript{89} To know whether a vaccine is protective, in other words, researchers must first identify the correlates of protection against HIV infection.\textsuperscript{90}

With many diseases, the presence of antibodies in the blood functions as a sign that the individual has either been infected with and overcome the disease or, in the alternative, has been successfully inoculated.\textsuperscript{91} This is not the case with HIV. HIV-infected persons register high levels of antibodies, including antibodies capable of neutralizing the virus in vitro, but they are still not protected against disease progression.\textsuperscript{92} Although antibody presence does not inhibit \textit{progression} to AIDS, however, this does not mean that antibodies afford no protection against HIV \textit{infection}: "Most of the vaccine experiments performed to date are banking on this hypothesis."\textsuperscript{93}

Grady’s choice of words—"banking on"—is curiously apt. At this point, there is no way to know whether HIV antibodies afford any
protection against HIV infection. AIDS vaccine trials are designed, in part, to identify the correlates of protection contemporaneously with the generation of protection. In plain terms, the only way to know if an AIDS vaccine is working is to test it in HIV-negative people who are likely to be exposed to the virus; researchers cannot ethically “challenge” subjects directly with a deadly virus to test vaccine efficacy. If high-risk individuals in the vaccine arm of a research study demonstrate a statistically significant difference in infection rates from similarly at-risk individuals in the placebo or control arm, then it is logical to assume that the vaccine has afforded some degree of protection. Researchers can then work backwards, in a sense, to identify the correlates of protection based on the empirical verification of efficacy.

To summarize, trial design in AIDS vaccine research is anomalous in at least three ways: (1) viral variation makes the target both multiple and self-modifying; (2) research has been initiated in humans before efficacy has been demonstrated in animals; and (3) the correlates of protection will theoretically emerge, ex post facto, from the comparative infection rates of high-risk individuals injected with vaccine versus those injected with placebo. Part III explores how these necessary anomalies in trial design affect the often vulnerable individuals and communities from which research subjects must be drawn in domestic AIDS vaccine research. In other words, Part III moves from the scientific challenges to the intimately related sociopolitical and legal challenges facing the domestic AIDS vaccine effort.

III

SOCIAL HARM AND AIDS VACCINE RESEARCH

In 1994, the AIDS Vaccine Evaluation Group (AVEG) conducted a confidential survey of 247 participants at high risk of HIV infection in 1994. Obviously it will take a large number of participants to power such a study. A statistically significant disparity can be generalized only if it occurs regularly across a significant population. “HIV-vaccine research and development differs from the development of treatments in that testing of efficacy cannot be targeted at specific individuals but must be done in large-scale trials to assess the effectiveness of the vaccine against a ‘natural’ rate of infection.”

ENCYCLOPEDIA OF AIDS, supra note 42, at 435.

95. The AVEG is a multi-centered organization established by the NIAID Division of AIDS (DAIDS) as part of its program to develop AIDS vaccines. See MICHAEL LANGAN & CHRIS COLLINS, PAVING THE ROAD TO AN HIV VACCINE: EMPLOYING TOOLS OF PUBLIC POLICY TO OVERCOME SCIENTIFIC, ECONOMIC, SOCIAL AND ETHICAL OBSTACLES 5-8 (Center for AIDS Prevention Studies, Univ. of Cal. San Francisco, Monograph Series, 1998) (providing a helpful overview of the U.S. government’s involvement in AIDS vaccine research). “The AVEG has historically focused on early safety and immunogenicity trials, and correlative laboratory studies of candidate preventive vaccines.” Id. at 7. DAIDS has also established the HIV Network for Prevention Trials (HIVNET), which “has a broader agenda that includes trials of vaccines, topical microbicides, STD treatment, behavioral interventions, and approaches to prevent mother-to-infant transmission. [HIVNET] has focused primarily on efficacy trials in an international multi-center network.” Id. NIAID is currently
Phase II trials of gp120. The survey was designed to support a systematic assessment of the social risk of HIV vaccine trial participation. A full eighteen percent of those surveyed reported trial-related adverse social events. Many of these adverse events were admittedly de minimis or beyond the scope of legal intervention (for example, "I told a friend I might test ELISA-positive on a test and he started being careful about sharing food with me."). Others, however, give serious reason for pause: A "volunteer reported that he tested positive during a hospitalization, and that the insurance company was notified and in turn notified the employer, who fired the volunteer. This volunteer reported losing health insurance when the job was lost.")

This nameless, faceless "volunteer" stands as an emblem of the social risks faced by participants in AIDS vaccine trials. The participant lost his job and his health insurance, but it is not altogether clear how this happened. Did the disclosure of his apparent HIV-positive serostatus (hospital to insurance company to employer) take place without his knowledge and without a chance to explain the test result? Or did the volunteer try to explain the result, with the insurance company, the employer, or both drawing inferences from his participation in the trial? Either way, this brief and anonymous anecdote serves as a cautionary tale to trial participants and researchers alike.

The following subsections examine two possible bases for trial-related discrimination: An antibody-positive HIV test, and disclosure of AIDS vaccine trial participation. Although the focus will be on potential harm consolidating and redistributing its vaccine research programs to develop an HIV Vaccine Trials Network (HVTN) and HIV Prevention Trials Network (HPTN). See AVAC, supra note 27, at 32-33. On NIAID's involvement in AIDS vaccine research generally, see NIAID, Clinical Research on AIDS Vaccines (visited Apr. 3, 2000) <http://www.niaid.nih.gov/factsheets/cclinrch.htm>.

96. See Sheon et al., supra note 7, at 520.
97. See id. at 523.
98. See id. at 524.
99. Id. at 522.
100. Id.
101. The use of the term "volunteer" to describe this participant begs the question: Was his participation truly voluntary? Did he know what he might get into? Hans Jonas's broad concept of voluntariness in human subject experimentation rings oddly in this context: "Only genuine authenticity of volunteering can possibly redeem the condition of thinghood to which the subject submits." Jonas, supra note 86, at 989. Jonas distinguishes genuine volunteering from consent narrowly defined: "Mere 'consent' (mostly amounting to no more than permission) does not right this reification." Id.
102. A third possibility is that the participant actually seroconverted. Due to an error in the design of the trial, however, "it was not possible to determine whether the reported positive test results were due to actual infections from high risk behavior or from vaccine-induced antibody reactions." Sheon et al., supra note 7, at 522.
103. It is worth emphasizing that discrimination against people who test positive for HIV, whether they actually have the virus or have only generated antibodies, derives from discriminatory attitudes towards HIV-positive people generally. In other words, the need to protect participants in AIDS vaccine trials (who are HIV-negative) derives, in large measure, from the stigmatization of HIV-positive people as such. Aside from its general desirability, then, reducing the stigma associated with
to participants, this harm can also impede AIDS vaccine research as a whole. If social harms in early trials are not managed effectively, publicized harms may dry up voluntary enrollment. Potential discrimination, then, will be analyzed both as an issue of individual harm to research subjects and as a more general threat to the vaccine effort.

A. **Discrimination Based on Vaccine-Induced Seropositivity**

The HIV Network for Prevention Trials (HIVNET)\(^{104}\) recently conducted a Vaccine Preparedness Study among HIV-negative homosexual men, male and female injection-drug users, and non-injecting women at heterosexual risk.\(^{105}\) Participants ranked false-positive HIV tests as their number one concern in the realm of social risks.\(^{106}\)

A major concern across all study populations was vaccine-induced seropositivity. Study sites will need to emphasize to participants that testing to discriminate actual HIV-1 infection from vaccine-induced seropositivity will be available at the study site, as needed. However, participants need to understand that if they test outside of the study site, HIV-1 antibody-positive results found by conventional serologic tests may not be able to distinguish infection from vaccine-induced seropositivity, and these results may lead to possible discrimination or other social harms.\(^{107}\)

As this quotation suggests, researchers expect some version of informed consent\(^{108}\) to minimize potential harm based on false seropositivity ("Study sites will need to emphasize" and "participants need to understand" being the operative phrases). They also emphasize that specialized tests available at the study site will ensure that vaccine-induced antibody response will be distinguishable from actual seroconversion.\(^{109}\) Ultimately, the goal is to

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HIV protects human subjects from potential social harm and enhances the likelihood that a successful AIDS vaccine will be produced for all.

104. On HIVNET, see discussion supra note 95.

105. See Koblin et al., supra note 7, at 786.

106. See id. at 788-90.

107. Id. at 791-92.

108. Federal regulations promulgated by the Department of Health and Human Services (HHS) require researchers to disclose "any reasonably foreseeable risks or discomforts to the subject." 45 CFR § 46.116(a)(2) (1998). Full disclosure, however, may not always be in the interest of researchers: "There are, of course, countervailing pressures that work against complete honesty. For instance, if all possible harmful effects were discussed in detail, it is possible that few people would be willing to participate." Gray et al., supra note 52, at 42. For further discussion of informed consent in AIDS vaccine trials, see infra Part V.C.2.

109. VaxGen has issued a letter to participants assuring them that they will have access to tests that distinguish HIV infection from seroconversion due to vaccination. See Letter from Donald P. Francis, President of VaxGen, Inc. to volunteers (on file with author) [hereinafter VaxGen Letter]. While access to such a test may help to explain why a participant has tested positive for antibodies on a standard blood test, it does nothing to prevent misdiagnosis and ensuing discrimination in the first place. Nor are insurers or others who rely on blood testing data obliged to consult VaxGen. While antibodies to the AIDSVAX gp120 product are relatively easy to distinguish from HIV antibodies...
shield participants from harm by limiting disclosure of their blood-test results. Given the prevalence of blood testing in our society, though, it is unlikely that participants will be able to confine testing to the research site, even if they might otherwise choose to do so. "In most states, insurance carriers are permitted to require HIV testing of insurance applicants, and the U.S. federal government requires HIV testing for military recruits, foreign service officers, Job Corps participants, prison inmates, and persons applying to immigrate or become naturalized citizens." During a participant's lifetime, it is highly improbable that every blood test can take place at the study site. Many, if not all, potential study participants will therefore have to face squarely the possibility of discrimination based on a false-positive blood test.

Why should we assume that employers, prison officials, immigration officers, and other similarly situated decision makers might discriminate against people who seem to be HIV-infected? Unfortunately, attitudes towards people living with HIV/AIDS have not progressed as quickly as medical treatment of the disease:

About half of American adults say that they would avoid a grocery store if the owner had AIDS, and nearly one-third advocate the quarantine and publication of the names of people with the disease. Almost half continue to believe that some forms of casual contagion (e.g., a kiss on the cheek) or airborne transmission (e.g., sneezing) can result in HIV infection.

using specialized assays, moreover, "[m]ore complex vaccines that may be tested in the future could be more likely to react with commercially available tests." Sheon, supra note 7, at 524. Among the priorities established for the National Institutes of Health by President Clinton's National AIDS Strategy Taskforce is production of more sensitive blood assays to distinguish vaccine antibodies from natural infection. See NATIONAL AIDS STRATEGY, supra note 19, at B-29. See generally David H. Schwartz et al., Extensive Evaluation of a Seronegative Participant in an HIV-1 Vaccine Trial as a Result of False-Positive PCR, 350 LANCET 256 (1997) (concluding that the design of efficacy trials for new vaccines should be in parallel with the development of diagnostic tests capable of distinguishing vaccine-induced antibodies from true HIV infection); Jonathan Weber, Distinguishing Between Response to HIV Vaccine and Response to HIV, 350 LANCET 230 (1997) (predicting that newer vaccines will produce antibody responses which will be practically indistinguishable from true infection using standard ELISA/Western Blot assays).

110. Sheon et al., supra note 7, at 523.
111. VaxGen is offering participants a study ID card to document participation in the AIDSVAX trial. See VaxGen Letter, supra note 109. The use of study ID cards, however, has backfired in some instances in Phase II trials of gp120:

Inadvertent disclosure of trial participation was very distressing in a small number of cases. Identity cards confirming the volunteers' participation were used deliberately by only 3 volunteers but were used against 2 others. Prospective volunteers should thus be asked to weigh the risks of someone's finding the card against the potential benefits that the card could offer in the event of a false-positive HIV test result.

Sheon, supra note 7, at 524. By carrying a study ID card, in other words, participants may inadvertently jump from the "false seropositive" frying pan into the "trial participant = high risk individual" fire discussed in Part III.B infra.

112. GRAY ET AL., supra note 52, at 164 (references omitted).
With paranoia about contagiousness still rampant, it is no wonder that Americans treat people living with HIV/AIDS as social lepers.

Judge Broderick of the U.S. District Court for the Eastern District of Pennsylvania, under the palpable influence of Susan Sontag's *AIDS and its Metaphors*, has written eloquently about American society's attitudes towards perceived "HIV carriers":

[S]ince first identified in the early 1980s as a distinct medical condition, AIDS has engendered such prejudice and apprehension that its diagnosis typically signifies a social death as concrete as the physical one which follows . . . . The particular associations AIDS shares with sexual fault, drug use, social disorder, and with racial minorities, the poor, and other historically disenfranchised groups accentuates the tendency to visit condemnation upon its victims. AIDS mythology has fomented not only private judgments about carriers of the virus. It has spawned calls for punitive, oppressive official action against them . . . . Vast segments of the American populace favor the forced quarantine of persons with AIDS, tattooing HIV-positive persons for ready identification, and banishing HIV carriers from the workplace and school. Thus, to conclude that persons with AIDS are stigmatized is an understatement; they are widely stereotyped as indelibly miasmic, untouchable, physically and morally polluted.

If Judge Broderick's rhetoric seems somewhat melodramatic a decade later, consider the remarks of some our country's most powerful legislators opposing the Americans with Disabilities Act of 1990. The ubiquitous Senator Jesse Helms of North Carolina opined: "I do not understand why . . . you went down the road of including in your definitions people who are HIV-positive, because 85 percent or more of the HIV positive people in this country are known to be drug users or homosexual or both." Not to be outdone, Representative Dan Burton of Indiana tersely stated: "The ADA is the last ditch attempt of the remorseless sodomy lobby to achieve its national agenda before the impending decimation of AIDS destroys its political clout." Heated rhetoric aside, the point need not be belabored ad nauseam: Discrimination against people perceived to have HIV or AIDS is an American reality.

B. Discrimination Based on Trial Participation
   as an Indicator of Risk-Group Status

As the remarks of Senator Helms and Representative Burton attest, the association between AIDS and homosexuality or drug use is commonplace. The association works in both directions: Injection drug users and homosexuals are often presumed to be HIV-positive, and those who are HIV-positive are often presumed to be homosexuals or injection drug users.

Given this reflexive association, the effort to enroll participants in an AIDS vaccine trial can be tricky business. On the one hand, researchers will need to reach out to affected communities to assemble trial cohorts that can tell them something about a candidate vaccine’s efficacy. As discussed earlier, researchers will need to test vaccines in HIV-negative subjects at high risk of infection. On the other hand, by reaching out to those who take part in high-risk behaviors, researchers will be relying upon and reinforcing some of the very associations that stigmatize affected populations. Future efficacy trials will have to assemble in the neighborhood of 5000 participants for each Phase III trial. Each time a flyer, billboard, newspaper, radio, television, or Internet advertisement publicizes the risk-groups needed for AIDS vaccine trials for recruitment purposes, it also informs those who would discriminate that participation correlates with high-risk behavior.

C. Trial-Related Discrimination as a Disincentive to Participate

The two forms of discrimination discussed above—discrimination based on vaccine seropositivity or on trial participation itself—could have adverse consequences for the AIDS vaccine effort. First, individual participants may suffer the consequences of trial-related discrimination. HIV-negative research volunteers, many of whom decide to participate based on altruism, would be exposed to social harms. Since these volunteers place

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117. See supra Part II.D.
118. The HIVNET Vaccine Preparedness Study, for example, recruited “HIV-1 negative homosexual men, male and female injecting drug users and non-injecting women at heterosexual risk,” Koblin et al., supra note 7, at 785, clinical-speak for gays, addicts, and prostitutes. Sheon and her colleagues made a good-faith, if half-hearted, stab at addressing this problem: “[I]t seems prudent to consider including both high-risk and low-risk volunteers in future efficacy trials. If trial sponsors widely publicize the fact that trials include both high-risk and low-risk volunteers, stigmatization of volunteers may be reduced.” Sheon et al., supra note 7, at 525. Such a remedial measure is questionable on at least three grounds. First, it entirely neglects the distorting effects enrolling low-risk individuals would have on trial data. Second, it also neglects the considerable expense involved in screening, consenting, enrolling, and processing individuals who would contribute nothing to the actual findings of the experiment. Third, it naïvely assumes that private sponsors will expend their resources on media counter-spin campaigns that have, at best, little chance of reducing stigma. As suggested infra Part V, legislative and regulatory measures aimed at protecting participants offer a more robust alternative.
themselves in a vulnerable position in the interest of public health, it is imperative as a matter of individual justice that we protect them. A second consideration is potential harm to the vaccine effort itself. If trial participants are exposed to unnecessary harm and their experience is communicated to their constituent communities, the high-risk volunteers so essential to the success of efficacy trials might quickly become averse to participation. As Mark Harrington has observed, “vaccine trials will be subjected to unprecedented worldwide scrutiny.” One adverse event, with enough publicity, could radically impede recruitment efforts and trials generally.

Although researchers recognized the potential for social harms long ago, the remedial measures they have devised are, at best, weak. Providing for HIV testing at the study site is helpful, as far as it goes, but participants will often be required to undergo HIV tests off-site. Vaccine-induced seropositive results are bound to occur, especially as vaccines advance and antibody responses grow increasingly difficult to distinguish from natural infection. Specialized assays to distinguish HIV antibodies from vaccine-induced antibodies will be a welcome innovation, but they must be widely produced, disseminated, and employed to help in any meaningful way. In addition, specialized assays may become a logistical nightmare as several different vaccine candidates reach Phase III trials and therefore turn up more frequently in the blood of the general population. Study ID cards and phone calls from researchers assuring individuals that trial participation is the source of false-positive HIV tests also provide little solace. Since study participation stands as a virtual proxy for risky behavior such as sodomy, drug use, and promiscuity, disclosure of trial participation simply replaces one potential basis of discrimination with another. Finally, traditional informed consent procedures do not appear to square well with the unique elements of the AIDS vaccine effort. It is hardly possible to anticipate all

119. As Gary Ellis, director of the Office for Protection from Research Risks (OPPR), has stated: “Respecting the rights of research subjects and providing for their welfare is to honor a deep obligation to those individuals who make a remarkable contribution to the common good by volunteering to serve as research subjects.” Gary Ellis, Keeping Research Subjects Out of Harm’s Way, 282 JAMA 1963, 1964 (1999).

120. In African American communities, for instance, suspicion of medical experimentation runs deep ever since word of the shockingly exploitative Tuskegee syphilis experiments became public. Although unquestionably justified by the historical record, such suspicion can also do considerable damage. Now that effective treatments for AIDS exist, for instance, it is becoming increasingly clear that African Americans are not benefiting from antiretrovirals as much as whites. See JAMES H. JONES, BAD BLOOD: THE TUSKEGEE SYPHILIS EXPERIMENT 220-42 (2d ed. 1993). For other trenchant discussions of the shadow Tuskegee casts over AIDS research, see Allan M. Brandt, Racism and Research: The Case of the Tuskegee Syphilis Study, 8 HASTINGS CTR. REP. 21 (1978); David L. Kirp, Blood, Sweat, and Tears: The Tuskegee Experiment and the Era of AIDS, 10 Tikkun 50 (1995); Deborah L. Shelton, Legacy of Tuskegee, AM. MED. NEWS, June 3, 1996, at 11.

potential "risks or discomforts"¹²² that may arise socially as a result of trial participation or false-positive HIV tests. An attempt to catalogue exhaustively the potential social risks might simply produce debilitating anxiety in the vulnerable participants essential to AIDS vaccine research.

In conclusion, the potential for social harms, when combined with the physical risks of vaccine research, may prove an overwhelming disincentive to participate in AIDS vaccine trials. Given the weak remedies that researchers have proposed thus far, participants in efficacy trials may well be exposed to considerable social harm. If participants are harmed, this will be problematic both as a matter of individual justice and as a potential threat to recruitment for future trials. Since it is likely that several Phase III trials will be necessary to identify an effective product, and since the need for an effective AIDS vaccine is so acute, more robust ways must be found to protect trial participants. Part IV therefore turns to federal disability discrimination law to examine one viable, but ultimately uncertain, avenue of protection.

IV

DOES THE ADA PROTECT AIDS VACCINE RECIPIENTS?

Anticipating trial-related social harms, researchers naturally turned to extant legal protections to see if the fears of participants might be reasonably allayed. One potential source of protection seized upon early by legal analysts was the Americans with Disabilities Act of 1990.¹²³ The following Part evaluates whether the ADA offers the kind of protection against trial-related discrimination that AIDS vaccine researchers and their legal advisors have envisioned. In light of recent Supreme Court decisions construing the Act, this Part concludes that ADA protection is, at best, unreliable in this context.

A. HIV/AIDS and the Americans with Disabilities Act of 1990

The ADA was promulgated "to address the major areas of discrimination faced day-to-day by people with disabilities."¹²⁴ Title I of the Act applies to discrimination in employment,¹²⁵ Title II to public services.¹²⁶

¹²². 45 C.F.R. § 46.116(a)(2) (1998) (providing that a description of foreseeable risks and discomforts involved in participation be given to participants as part of the informed consent process). For suggestions as to how informed consent might be tailored to AIDS vaccine trials, see discussion infra Part V.C.2.

¹²³. See discussion infra Part V.A.

¹²⁴. 42 U.S.C. § 12101(b)(4) (1994). The Act mentions "discrimination against individuals with disabilities . . . in such critical areas as employment, housing, public accommodations, education, transportation, communication, recreation, institutionalization, health services, voting, and access to public services" in legislative findings. Id. § 12101(a)(3).

¹²⁵. See id. §§ 12111-12117.

¹²⁶. See id. §§ 12131-12165.
and Title III to public accommodations and services operated by private entities.\textsuperscript{127} The Act covers private employers with fifteen or more employees\textsuperscript{128} as well as state and local governments.\textsuperscript{129} The ADA was not the first federal legislation to address discrimination against disabled people. Congress had earlier passed the Rehabilitation Act of 1973,\textsuperscript{130} which applies to federal agencies\textsuperscript{131} and private entities receiving federal assistance.\textsuperscript{132}

The ADA sets forth a three-pronged definition of disability. It defines "disability" with respect to an individual as:

\begin{enumerate}
  \item (A) a physical or mental impairment that substantially limits one or more of the major life activities of such individual [the "actual disability" prong];
  \item (B) a record of such an impairment [the "record of" prong];
  \item (C) being regarded as having such an impairment [the "regarded as" prong].\textsuperscript{133}
\end{enumerate}

Individuals who fit into any of these three prongs of the definition of disability are protected from discrimination if they are "otherwise qualified."\textsuperscript{134} In the employment context, individuals are otherwise qualified if they can perform the essential functions of the position with or without reasonable accommodation.\textsuperscript{135} Individuals are not otherwise qualified if they pose a direct threat to the health or safety of others.\textsuperscript{136}

The question that might arise in future ADA litigation involving AIDS vaccine recipients can be concisely stated: Does an individual who

\textsuperscript{127} See id. §§ 12181-12189.
\textsuperscript{128} See id. § 12111(5).
\textsuperscript{129} See id. § 12202.
\textsuperscript{130} 29 U.S.C. §§ 701-797(b) (1994).
\textsuperscript{131} See id. § 791(b).
\textsuperscript{132} See id. §§ 793-794. The ADA expressly incorporates Rehabilitation Act standards as a "floor" for purposes of the Act. See 42 U.S.C. § 12201(a)(1994). In addition, Congress directed that enforcement agencies develop procedures to ensure that administrative complaints filed under the ADA and under the Rehabilitation Act are dealt with in a manner that "avoids duplication of effort and prevents imposition of inconsistent or conflicting standards." Id. § 12117(b). The Equal Employment Opportunity Commission (EEOC), responsible for enforcement of Title I of the ADA, made Rehabilitation Act regulations and case law a cornerstone of its regulatory drafting for the ADA:

   The format of part 1630 reflects congressional intent, as expressed in the legislative history, that the regulations implementing the employment provisions of the ADA be modeled on the regulations implementing section 504 of the Rehabilitation Act of 1973, as amended, 34 CFR part 104. Accordingly, in developing part 1630, the Commission has been guided by the section 504 regulations and the case law interpreting those regulations. 56 Fed. Reg. 35,726 (1991). Since Rehabilitation Act regulations and precedent have been widely held to apply in ADA cases, the following discussion assumes this without further analysis.
\textsuperscript{133} See id. § 12111(8).
\textsuperscript{134} The term "reasonable accommodation" (defined at id. § 12111(9)) has been the source of extensive litigation. See generally Mary L. Topliff, Remedies Available under Americans with Disabilities Act (42 U.S.C.A. § 12101 et seq.), 136 A.L.R. Fed. 63 (1997). Because reasonable accommodation is not, as a general matter, required by people who test false-positive for HIV, this issue will not be treated further.
\textsuperscript{135} See 42 U.S.C. § 12113(b).
tests false-positive for HIV potentially qualify for protection as "disabled" under the Act? The first prong of the Act's disability definition clearly seems inapposite: A false-positive HIV test probably does not constitute "a physical or mental impairment." The second and third prongs, however, may plausibly be construed to cover the scenario. A false-positive HIV test arguably constitutes a "record of" impairment. Moreover, a complainant alleging discrimination based on a false-positive result could logically claim she was "regarded as" impaired by an employer or other relevant decision maker.

Before turning to the relevant case law, it is helpful to explain why the ADA protects individuals as "disabled" who do not actually suffer any mental or physical impairment in the traditional sense. In other words, what is the rationale behind the "record of" and "regarded as" prongs of the ADA's definition of disability? Why should an entirely healthy individual, such as an AIDS vaccine recipient who suffers no actual physical harm from the vaccine but who tests HIV-positive nevertheless, qualify as "disabled"? Justice Brennan eloquently stated the rationale behind the protection of perceived disabilities in School Board of Nassau County v. Arline: 3

[S]ociety's accumulated myths and fears about disability and disease are as handicapping as are the physical limitations that flow from actual impairment. Few aspects of a handicap give rise to the same level of public fear and misapprehension as contagiousness. Even those who suffer or have recovered from such noninfectious diseases as epilepsy or cancer have faced discrimination based on the irrational fear that they might be contagious. The [Rehabilitation] Act is carefully structured to replace such reflexive

137. The EEOC and Department of Justice (DOJ) regulations attempt to capture the full range of impairments that qualify under the Act:

Physical or mental impairment means:

(1) Any physiological disorder, or condition, cosmetic disfigurement, or anatomical loss affecting one or more of the following body systems: neurological, musculoskeletal, special sense organs, respiratory (including speech organs), cardiovascular, reproductive, digestive, genito-urinary, hemic and lymphatic, skin, and endocrine; or
(2) Any mental or psychological disorder, such as mental retardation, organic brain syndrome, emotional or mental illness, and specific learning disabilities.


138. 480 U.S. 273 (1987). Arline was a Rehabilitation Act case in which plaintiff Arline, an elementary school teacher suffering from chronic tuberculosis, alleged she was fired out of fear of contagion. The Supreme Court held that Arline could be considered a "handicapped individual" within the meaning of § 504 of the Rehabilitation Act because her hospitalization for tuberculosis sufficed to establish a record of impairment. See id. at 281. The Court remanded on the question whether Arline was otherwise qualified, that is, whether she posed "significant health or safety risks" to others. Id. at 287. Arline played a seminal role in the drafting of the ADA and its implementing regulations and interpretive guidelines. See, e.g., 29 C.F.R. § 1630.2(l) (app. 352) (1999) ("The rationale for the "regarded as" part of the definition of disability was articulated by the Supreme Court in the context of the Rehabilitation Act of 1973 in School Board of Nassau County v. Arline.").
reactions to actual or perceived handicaps with actions based on reasoned and medically sound judgments.139

According to Justice Brennan's logic, perceived impairments can have handicapping effects just as severe as those caused by actual physical impairments. The locus of limitation is immaterial: Whether the limitation arises from an individual's own body or from "society's accumulated myths and fears about disability and disease," it debilitates just the same. Accordingly, the first prong of the ADA's disability definition addresses actual physical and mental impairments whereas the second and third prongs focus on perceived disabilities, whether actual or "mythical."

Although the actual impairment prong of the definition is analytically distinct from the other two, the "record of" and "regarded as" prongs relate back to the first prong in their reference to "such an impairment" (that is, one that substantially limits a major life activity).140 To know whether an individual has a "record of" or is "regarded as" having "such an impairment," we must first determine what "such an impairment" would mean for an AIDS vaccine recipient. A false-positive HIV blood test, although incorrect, may arguably constitute a record of infection with HIV. As the EEOC regulations construing the definition state: "Has a record of such impairment means has a history of, or has been misclassified as having, a mental or physical impairment that substantially limits one or more major life activities."141 Assuming arguendo that a false-positive test "misclassifies" an individual as HIV-positive, the question then becomes: Is HIV infection a physical impairment that substantially limits one or more major life activities?

The analysis follows a similar trajectory under the "regarded as" prong. Assuming that an employer or other decision maker has learned about a plaintiff's apparent HIV-positive status, and assuming that the employer or decision maker "treats"42 the plaintiff adversely based on the impression that he or she is actually infected with HIV, the analysis again turns on the definition of "such an impairment." Did the employer or

139. Arline, 480 U.S. at 284-85. Although the language cited has obvious implications for people with AIDS, the Court carefully limited its holding:

This case does not present, and we therefore do not reach, the question whether a carrier of a contagious disease such as AIDS could be considered to have a physical impairment, or whether such a person could be considered, solely on the basis of contagiousness, a handicapped person as defined by the [Rehabilitation] Act.

Id. at 283 n.7. As discussed infra, the question identified but left unanswered in Arline was addressed in Bragdon v. Abbott, 524 U.S. 624 (1998) (holding, inter alia, that HIV is an impairment from the moment of infection).

140. See Bragdon, 524 U.S. at 658 n.1 (Rehnquist, C.J., concurring in part and dissenting in part).


142. The EEOC regulations offer three alternative readings of the "regarded as" prong. The third is relevant in this context: "Is regarded as having such an impairment means...[It]as none of the impairments defined...but is treated by a covered entity as having a substantially limiting impairment." 29 C.F.R. § 1630.2(1)(3).
relevant decision maker regard the plaintiff as having an impairment that substantially limits a major life function? The outcome depends on whether HIV infection substantially limits a major life function. As discussed in the next Section, this was precisely the question presented in Bragdon.

B. Bragdon v. Abbott and Asymptomatic HIV Infection

In Bragdon, the U.S. Supreme Court granted certiorari on the following question: "[W]hether HIV infection is a disability under the ADA when the infection has not yet progressed to the so-called symptomatic phase." The facts of the case were simple. Plaintiff Abbott went to dentist Bragdon for an appointment in 1994. She had been infected with HIV since 1986, and she disclosed her HIV status on the dentist’s patient registration form. When Bragdon discovered a cavity in the course of his examination of Abbott, he informed her of his policy against filling cavities of HIV-infected patients in his dental office. Bragdon offered to perform the work at a hospital with no added fee for services. Abbott, however, would have to foot the bill for the hospital’s facilities, he explained. Abbott declined and sued Bragdon under state law and the ADA for discrimination in public accommodations. The U.S. District Court for the District of Maine granted summary judgment for plaintiff Abbott, and the First Circuit affirmed. On appeal to the U.S. Supreme Court, Justice Kennedy, writing for a 5-4 majority, held that: (1) HIV is a physiological impairment

143. The EEOC’s Interpretive Guidance on Title I of the Americans with Disabilities Act, appended to the implementing regulations, offers an example of this situation that is telling for our purposes:

An individual satisfies . . . the “regarded as” definition of “disability” if the employer or other covered entity erroneously believes the individual has a substantially limiting impairment that the individual actually does not have. This situation could occur, for example, if an employer discharged an employee in response to a rumor that the employee is infected with Human Immunodeficiency Virus (HIV). Even though the rumor is totally unfounded and the individual has no impairment at all, the individual is considered an individual with a disability because the employer perceived of this individual as being disabled. Thus, in this example, the employer, by discharging this employee, is discriminating on the basis of disability.


145. Id. at 628.
146. See id. at 629.
147. See id. "No individual shall be discriminated against on the basis of disability in the full and equal enjoyment of the goods, services, facilities, privileges, advantages, or accommodations of any place of public accommodation by any person who owns, leases (or leases to), or operates a place of public accommodation." 42 U.S.C. § 12182(a) (1994). The term “public accommodations” is defined by the ADA to include the “professional office of a health care provider.” Id. §12181(7)(F).
149. See Abbott v. Bragdon, 107 F.3d 934 (1st Cir. 1997).
of the hemic and lymphatic system "from the moment of infection;" and (2) reproduction is a major life activity contemplated by the ADA; and (3) Abbott's HIV infection substantially limited her ability to reproduce. Accordingly, the Court concluded that Abbott was actually disabled under the ADA. It remanded the case to the First Circuit for further development of the record on the question whether Abbott's HIV infection posed a "direct threat" to the safety of her dentist Bragdon.

The majority opinion in Bragdon is as noteworthy for what it declined to answer as for what it addressed. Because the Court found Abbott actually disabled, it declined to consider whether she would qualify for protection under the "record of" or "regarded as" prongs of the ADA's definition of disability. For the same reason, the Court also declined to answer the second question presented on appeal, that is "whether HIV infection is a per se disability under the ADA." Similarly, the Court refused to address whether analysis of the limitations posed by an individual's impairment should proceed from the individual's status with or without mitigating measures such as medications that keep symptoms in check. Finally, the Court's tight focus on the major life activity of reproduction left open the

151. See id. at 638.
152. See id. at 639. The Court developed two independent paths to this conclusion. The first analyzed potential transmission of HIV to Abbott's sexual partner as a limitation on reproduction. See id. The second focused on the risk of perinatal (mother-to-infant) transmission as such a limitation. See id. at 640.
153. See id. at 631.
154. Id. at 655.
155. "We hold respondent's HIV infection was a disability under subsection (A) of the definitional section of the statute. In light of this conclusion, we need not consider the applicability of subsections (B) or (C)." Id. at 631.
156. "Respondent's HIV infection is a physical impairment which substantially limits a major life activity, as the ADA defines it. In view of our holding, we need not address the second question presented, i.e., whether HIV infection is a per se disability under the ADA." Id. at 641-42.
157. Given the procedural posture of the case, that is, summary judgment in favor of plaintiff Abbott below, the Court characterized the issue as a matter of law it did not need to resolve in order to decide the instant case:

The Solicitor General questions the relevance of the 8% figure [of perinatal transmission of HIV when the mother is treated with AZT], pointing to regulatory language requiring the substantiality of a limitation to be assessed without regard to available mitigating measures .... We need not resolve this dispute in order to decide this case, however. It cannot be said as a matter of law that an 8% risk of transmitting a dread and fatal disease to one's child does not represent a substantial limitation on reproduction. Id. at 640-41. The Court has recently settled this question, flatly contradicting the regulatory language mentioned by Justice Kennedy, establishing that "disability under the Act is to be determined with reference to corrective measures." Sutton v. United Air Lines, Inc., 119 S.Ct. 2139, 2149 (1999) (holding that severely myopic job applicants must be evaluated with corrective lenses to decide whether they are disabled under the ADA); see also Murphy v. United Parcel Service, Inc., 119 S.Ct. 2133 (1999) (holding that an employee with severe hypertension must be evaluated for disability in medicated condition). See generally The Supreme Court, 1998 Term-Leading Cases, 113 Harv. L. Rev. 326, 337-49 (1999) (analyzing the trilogy of ADA cases, including Murphy and Sutton, decided by the Court in its 1998 term).
question of other major life activities that might be substantially limited by HIV infection.158

C. Questions Remaining after Bragdon: Are Individuals with a “Record of” HIV Infection or Who are “Regarded as” HIV-Infected Protected by the ADA?

The Bragdon decision resolved a notorious split in the circuits. In Runnebaum v. NationsBank of Maryland,159 the Fourth Circuit had held that the “plain meaning” of the term impairment, as used in the ADA, “suggests that asymptomatic HIV infection will never qualify as an impairment.”160 Equating the misnomer “asymptomatic” with a complete absence of physical effects, the Court reasoned that no limitation could reasonably be found: “[W]ithout symptoms, there are no diminishing effects on the individual.”161 The First Circuit, on the other hand, took the opposite tack in Abbott v. Bragdon:162 “We hold unhesitatingly that HIV-positive status, simpliciter, whether symptomatic or asymptomatic, comprises a physical impairment under the ADA.”163

In Bragdon, the Supreme Court expressly overruled the interpretation of impairment articulated in Runnebaum: “HIV infection must be regarded as a physiological disorder with a constant and detrimental effect on the infected person’s hemic and lymphatic systems from the moment of infection. HIV infection satisfies the statutory and regulatory definition of a physical impairment during every stage of the disease.”164 By holding that infection with HIV is a per se impairment, however, the Court did not hold that HIV is a per se disability. In refusing to address whether HIV is a per se disability, the Court implicitly suggested that the inquiry as to substantial limitation of a major life activity should take place on an individualized basis.165 The resulting indeterminacy, arising from the fact-specific nature

158. “[I]t may seem legalistic to circumscribe our discussion to the activity of reproduction. We have little doubt that had different parties brought the suit they would have maintained that an HIV infection imposes substantial limitations on other major life activities.” Bragdon, 524 U.S. at 637. See generally Michelle T. Friedland, Note, Not Disabled Enough: The ADA’s “Major Life Activity” Definition of Disability, 52 Stan. L. Rev. 171 (1999) (analyzing the disjunction between the ADA’s dual goals of encouraging reasonable accommodation and ferreting out discrimination with respect to the major life activities requirement).

159. 123 F.3d 156 (4th Cir. 1997).

160. Id. at 169.

161. Id. at 168 (citing Webster’s Dictionary definitions of “impairment”).

162. 107 F.3d 934 (1st Cir. 1997), vacated and remanded, 524 U.S. 624 (1998).

163. Id. at 939. The First Circuit joined the Ninth and Eleventh Circuits in so holding. See Gates v. Rowland, 39 F.3d 1439, 1446 (9th Cir. 1994); Doe v. Garrett, 903 F.2d 1455, 1459 (11th Cir. 1990), cert. denied, 499 U.S. 904 (1991).

164. Bragdon, 524 U.S. at 637.

165. Justice O'Connor stressed this point in her brief opinion: “[R]espondent’s claim of disability should be evaluated on an individualized basis and... she has not proven that her asymptomatic HIV status substantially limited one or more of her major life activities.” Id. at 664 (O’Connor, J., concurring in part and dissenting in part). She has since clarified her reasoning on this point in Sutton v.
of the substantial limitation inquiry, effectively undercuts much of the precedential value of Bragdon.

To illustrate this point, we turn back to our hypothetical AIDS vaccine recipient. If we posit a menopausal female vaccinee, for example, it is not at all certain that she will be protected by the ADA after Bragdon if she tests positive for HIV antibodies. While the Court focused solely on reproduction as a major life activity in Bragdon, an individualized, fact-specific inquiry may reveal that the hypothetical discriminating decision maker did not "regard" the plaintiff as substantially limited in the major life activity of reproduction. Since the decision maker may regard the plaintiff as infertile, and therefore may regard reproduction as an insignificant life activity for the plaintiff, rightly or wrongly, the decision maker may not regard the plaintiff as substantially limited in a major life activity. A similar hitch upsets the "record of" analysis under this scenario. If a plaintiff must show that a false-positive blood test constitutes a

United Air Lines, 119 S.Ct. 2139, 2147 (1999) (insisting on individualized inquiry under the ADA and rejecting an approach that "would often require courts and employers to speculate about a person's condition and would, in many cases, force them to make a disability determination based on general information about how an uncorrected impairment usually affects individuals, rather than on the individual's actual condition.")

166. In Abbott v. Bragdon the First Circuit had held that "HIV-positive status is a physical impairment that substantially limits a fecund woman's major life activity of reproduction." Abbott, 107 F.3d at 942 (emphasis added). This awkward holding has led to speculation about its applicability to non-fecund women or fecund and non-fecund men. Compare Wendy E. Parmet & Daniel J. Jackson, No Longer Disabled: The Legal Impact of the New Social Construction of HIV, 23 AM. J. L. & MED. 7, 35-36 (1997) (arguing that "the court's conclusion that Ms. Abbott was disabled was based in large part on the fortuity of her own fertility"), with Theresa A. Schneider, Note, Stretching the Limits of the ADA: Asymptomatic HIV-Positive Status as a Disability in Bragdon v. Abbott, 118 S.Ct. 2196 (1998), 77 Neb. L. Rev. 206, 221 (1998) (contending that individualized inquiry is necessary to prevent "absurdities" such as gay men or menopausal women basing ADA claims on the major life activity of reproduction).

167. The issue discussed here was nicely captured in a recent district court case involving a gay male, HIV-positive plaintiff:

Plaintiff asserts that he has wanted to have children since he was nineteen years old . . . . On the other hand, defendant argues that since plaintiff claims he wanted to father a child since he was nineteen, but did not discover he was HIV-positive until he was 33, and plaintiff did not take any steps to consummate his alleged desire to father a child, this demonstrates that his subsequent diagnosis of HIV-positive did not substantially limit this activity. In addition, defendant contends that plaintiff cannot demonstrate that his HIV condition substantially limited his ability to reproduce because plaintiff is a sexually active homosexual male, who during the relevant time period, was involved in an exclusively homosexual relationship. Contrary to defendant's arguments, plaintiff's homosexual relationship would not preclude him from fathering a child.

Hernandez v. Prudential Ins. Co. of America, 977 F. Supp. 1160, 1164 (M.D. Fla. 1997). Although plaintiff may be able to father a child, as the court observes, the question here is whether reproduction is a major life activity of plaintiff. As Chief Justice Rehnquist has pointed out, the ADA provides that "the 'major life activities' allegedly limited by an impairment must be those 'of such individual.'" Bragdon, 524 U.S. at 657 (Rehnquist, C.J., concurring in part and dissenting in part) (citing the ADA at § 12102(3)(A)) (emphasis added). Furthermore "the ADA's definition of a disability is met only if the alleged impairment substantially 'limits' (present tense) a major life activity." Id. at 661. Hypothetical attempts to reproduce, hence, are not necessarily pertinent to the analysis.
record of impairment that substantially limits a major life activity, then, once again, the plaintiff will have to show on an individualized basis that the life activity alleged is "major" for him or her. In short, the Court's self-described "legalistic" circumscription of the major life activity analysis in Bragdon leaves many key questions unsettled for subsequent suits.

The Court is not solely responsible for this continuing ambiguity. The problem also stems from the drafting of the "major life activities" definition in the ADA's implementing regulations: "Major life activities means functions such as caring for one's self, performing manual tasks, walking, seeing, hearing, speaking, breathing, learning, and working." As the Court correctly states, the use of the term "such as" clearly demonstrates that the list is intended to be "illustrative, not exhaustive." This is why the Court is free to consider whether reproduction qualifies as a major life activity even though it does not appear on the list. The trouble with an "illustrative" list, however, is that there is no articulated principle allowing lower courts to determine which activities qualify as major life activities. On the one hand, the illustrative list allows room to construe the term "major life activities" liberally. On the other hand, uncertainty in construction reigns until a new major life activity has been anointed by the Supreme Court.

The upshot of the Court's constrained holding in Bragdon is readily apparent. An HIV-positive "fecund woman," as the First Circuit clinically put it, is now clearly protected by the ADA as actually disabled. Is there any legitimate reason why HIV-infected infertile women or non-procreating men should not be equally certain of their protection by federal disability law? Is one any less disabled by HIV simply because one does not presently reproduce? By treating all sexual activity under the rubric of reproduction, as if all sexual activity capable of transmitting HIV were procreative in nature, the Court draws an artificial line. On one side of this line fall fecund women, and on the other side falls everyone else. In explaining why reproduction qualifies as a major life activity, Justice Kennedy writes: "Reproduction and the sexual dynamics surrounding it are central to the life process itself." Although a subtle parochialism, this strong association of sex with reproduction narrows the field of sexuality considerably. For instance, anal and oral sex are not necessarily "sexual

168. 28 C.F.R. § 41.31(b)(2) (1998); 29 C.F.R. § 1630.2(i) (1999).
170. The EEOC's interpretive guidance does not help significantly:
'Major life activities' are those basic activities that the average person in the general population can perform with little or no difficulty. Major life activities include caring for oneself, performing manual tasks, walking, seeing, hearing, speaking, breathing, learning, and working. This list is not exhaustive. For example, other major life activities include, but are not limited to, sitting, standing, lifting, reaching.
171. Bragdon, 524 U.S. at 638.
dynamics surrounding [reproduction].” Justice Kennedy’s analysis thus
constrains the activity of sex to reproductive sex without ever asking
whether non-procreative sex constitutes a major life activity.

The bright line drawn by the Court, moreover, does not appear so
bright when it comes to the “record of” and “regarded as” prongs. By de-
clining to reach Abbott’s claims under these theories, the Court left the
analysis of perceived disabilities, as described by Arline, entirely vague.
Although we now know that HIV, whether symptomatic or asymptomatic,
qualifies as a physical impairment, we do not know how the perception of
such an impairment by a decision maker should be treated. Is it enough that
the defendant regards the plaintiff as HIV-positive? Or must the defendant
regard the plaintiff as HIV-positive and therefore impaired in her ability to
reproduce?172 If the plaintiff cannot demonstrate by record evidence that
reproduction is a present major life activity for her, does this preclude a
finding of disability after Bragdon?173 Or is the relevant decision maker’s
perception of the significance of the life activity to the plaintiff dispo-
sitive? Given the conflict between the plain language of the ADA, the inter-
prediction offered by the agencies charged with implementing it, and
the Supreme Court’s “miserly construction”174 of the Act, such questions
can be multiplied indefinitely.

V
STATUTORY AND REGULATORY RESPONSES: MINIMIZING SOCIAL HARMs
TO AIDS VACCINE RECIPIENTS

A. Disability Discrimination Law as a Poor Fit for Research Subjects

As discussed in Part IV, the Supreme Court’s holding in Bragdon
does not sufficiently settle the construction of the ADA’s definition of dis-
ability to grant AIDS vaccinees any reasonable measure of security in the
face of prospective social harms.175 A false-positive HIV test may or may

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172. Chief Justice Rehnquist apparently views this as the correct inquiry: “Respondent has
offered no evidence to support the assertion that petitioner regarded her as having an impairment that
substantially limited her ability to reproduce, as opposed to viewing her simply as impaired.” Id. at 658
n.1 (Rehnquist, C.J., concurring in part and dissenting in part).

173. Again, this seems to be the Chief Justice’s opinion: “[T]here is not a shred of record
evidence indicating that, prior to becoming infected with HIV, respondent’s major life activities
included reproduction.” Id. at 658.

Colker has similarly criticized the courts for treating the ADA parsimoniously: “[D]efendants prevail
in more than ninety-three percent of reported ADA employment discrimination cases decided on the
merits at the trial court level. Of those cases that are appealed, defendants prevail in eighty-four percent
of reported cases.” Ruth Colker, The Americans with Disabilities Act: A Windfall for Defendants, 34
Harv. C.R.-C.L. L. Rev. 99, 100 (1999) (citations omitted). Colker further argues that courts have
abused the summary judgment device in ADA litigation. See id. at 101.

175. Even if we assume that the “regarded as” and “record of” analysis would conclude favorably
in those jurisdictions where HIV infection has been deemed a per se disability for ADA purposes (that
is, in the First, Ninth, and Eleventh Circuits), the lack of uniformity across the circuits remains a cause
not lead to liability if used against a given plaintiff. Without a stronger sense of the scope of protection afforded by the ADA, AIDS vaccine trial participants cannot rely on the Act to shield them from discrimination based on vaccine seropositivity. More significantly, disability law does not extend to discrimination based on AIDS vaccine trial participation. Many of the high-risk behaviors required for participation in vaccine trials, such as prostitution, illicit drug use, and sodomy, are themselves illegal in many states. It would be anomalous indeed if federal disability law prevented "discrimination" against individuals for behavior that would otherwise subject them to criminal sanction. In short, disability law seems poorly suited to cover the myriad adverse situations vaccinees may face as a result of their participation in experimental trials.

If this Comment has focused seemingly inordinate attention on the inadequacy of federal disability law as a potential source of protection for AIDS vaccine trial participants, it is because much of the extant literature uncritically assumes that ADA protection extends to asymptomatic HIV infection. One scholar writes: "As a result of the Bragdon decision, individuals may feel less hesitant about submitting to testing for the HIV virus because now they can feel confident that, if they are HIV-positive, they are protected from discrimination by the ADA." Another statement carries the same import and extends it to perceived infection: "[D]iscrimination on the ground of HIV seropositivity—or even suspicion that an individual is HIV-seropositive—is now illegal in many circumstances." Even President Clinton's National AIDS Strategy statement wishfully overstates the scope of federal disability law, mentioning among the White House's...
accomplishments "[e]nactment of the Americans with Disabilities Act of 1990 (ADA), which protects individuals living with HIV and AIDS and people perceived to be at risk for HIV from discrimination in housing, employment, and public accommodation." 179

Why do such overstatements of the protections afforded by federal disability law really matter in the context of AIDS vaccine research? The answer lies in the risk/benefit calculus a potential vaccine trial subject will perform in deciding whether or not to participate. Scott Burris has advanced a useful notion of "reliance" in the context of HIV testing that is helpful when thinking about vaccine trials:

Theories about compliance might explain why the ADA prevents a dentist from discriminating against a patient with HIV, and disputing theory will explain why most victims of this sort of discrimination do not sue, but neither addresses the question of why and how a person who is protected by law and who wishes to avoid an injury will integrate the law into a prospective decision to run the social risk of testing. For this, I suggest, we need a new concept, which I am calling "reliance." 180

Like someone deciding to get tested for HIV, a potential AIDS vaccine trial participant might make the "prospective decision to run the risk" of trial participation by integrating misstatements of the law into their decisionmaking process. "Reliance," in this case, leads to unfounded security. When researchers, who are not legal experts, incorporate broad statements about protection against discrimination such as those cited above into their consent procedures, trial participants might underestimate their risk of social harm. In this way, informed consent potentially shades into misinformed consent. 181 "Reliance" on unsettled law may lead, unsettlingly, to questionable voluntarism.

179. NATIONAL AIDS STRATEGY, supra note 19, at 27 (emphasis added).
181. As part of the informed consent process, for example, a number of research entities in the San Francisco Bay Area distribute to potential AIDS vaccine trial participants a brochure prepared by the AIDS Legal Referral Panel (ALRP) entitled The Rights of HIV Vaccine Trial Participants. The brochure contains the following advice concerning potential employment discrimination:

As a person who is vaccine positive but not HIV infected, it is unlikely that you will be considered "disabled" under [the ADA], where it is defined as a physical or mental impairment that substantially limits one or more major life activities. But [the ADA] also protect[s] a person who is "regarded as having such an impairment." In other words, if you are discriminated against because your employer presumes you to be HIV infected, you would be entitled to the full protection of [the ADA] as if you were actually HIV infected.

ALRP, THE RIGHTS OF HIV VACCINE TRIAL PARTICIPANTS 4 (1997) (citations omitted). This misleadingly optimistic characterization of the law was picked up by another recent handbook: "The San Francisco-based AIDS Legal Referral Panel has noted that U.S. law may protect individuals from discrimination based on perceived HIV status resulting from HIV vaccine trial participation. This protection comes, in part, under the Americans with Disabilities Act." AIDS VACCINE ADVOCACY
I do not mean to suggest that researchers and study recruiters are intentionally disseminating false or misleading statements of the law to dupe prospective participants. I am, however, arguing that the legal issues raised by AIDS vaccine trial participation are complex and easily susceptible to misstatement. In addition, I am arguing that optimistic reliance on legal protections afforded by disability law can lead to uninformed decision-making when it comes to AIDS vaccine trial participation. In short, the social harm issues posed by AIDS vaccine trials go far beyond the bounds of disability discrimination law. They implicate the very foundational notions of human experimentation itself: Respect for the autonomy of persons, beneficence, and justice. These overarching principles must be taken into account in addressing the unique social challenges of AIDS vaccine research.

B. The Inadequacy of State-Initiated Protection Schemes

Apart from federal disability discrimination law, what other sources of legal protection are available to trial participants? Very few state governments have addressed the issue. California has taken the national lead in promulgating laws to accelerate production of a safe, effective AIDS vaccine. In 1983, the Golden State initiated a comprehensive legislative effort to address the AIDS crisis. As part of this scheme, which includes provisions for an AIDS vaccine victims compensation fund, an AIDS COALITION (AVAC), HIV VACCINE HANDBOOK: COMMUNITY PERSPECTIVES ON PARTICIPATING IN RESEARCH, ADVOCACY, AND PROGRESS (Bill Snow ed., 1999) [hereinafter HIV VACCINE HANDBOOK].

182. These three basic ethical principles were articulated by the National Commission for the Protection of Human Subjects of Biomedical and Behavioral Research in the famous Belmont Report of 1976. See DENNIS M. MALONEY, PROTECTION OF HUMAN RESEARCH SUBJECTS: A PRACTICAL GUIDE TO FEDERAL LAWS AND REGULATIONS 31-40 [hereinafter BELMONT REPORT]. Dr. Harold Varmus, Director of the National Institutes of Health (NIH), has called these principles "the three quintessential requirements for the ethical conduct of research involving human subjects." Testimony of Harold E. Varmus, M.D., Director, National Institutes of Health, Before the Subcommittee on Human Resources and Intergovernmental Relations Committee on Government Reform and Oversight, U.S. House of Representatives, in II BIOLAW: UPDATES AND SPECIAL SECTIONS S:343 (October 1997). My thinking about the ethics of human research has been guided by the seminal text on the topic, JAY KATZ, EXPERIMENTATION WITH HUMAN BEINGS (1972).

183. Connecticut has passed legislation designed to foster research into AIDS vaccines. See CONN. GEN. STAT. §§ 19a-590—19a-591c (1991). Connecticut’s approach is to limit research liability to cases of “gross negligence or reckless, willful or wanton misconduct,” id. § 19a-591b, provided that the research subject has been given a “written explanation” of the limitation and “informed consent” is secured before administration of an AIDS vaccine, id. § 19a-591a. Connecticut does not indicate whether foreseeable trial-related social harms must be disclosed as part of the informed consent process.

184. See CAL. HEALTH & SAFETY CODE §§ 121200-121280 (West 1996). The history and relevant background of the California AIDS Initiative is helpfully recounted in Margaret Salmon Rivas, The California AIDS Initiative and the Food and Drug Administration: Working at Odds with Each Other?, 46 FOOD DRUG & COSM. L.J. 107 (1991); see also Wilson, supra note 6, at 556-59.

vaccine guaranteed purchase fund, and an AIDS Vaccine Research and Development Advisory Committee. California law also offers limited protection against research-related social harms.

California’s legislation, however, is severely limited in key respects. The California Health and Safety Code provides in relevant part: “No health care service plan, disability insurer, nonprofit hospital service plan, self-insured employee welfare benefit plan, or life insurer may withhold any settlement or coverage of an individual solely because of his or her participation in an AIDS/HIV vaccine clinical trial.” This language is deceptively reassuring for several reasons. First, although it appears to confer broad protection against trial-related discrimination in the insurance domain, it eliminates coverage for mixed-motive discrimination by including the words “solely because of.” For example, an insurer could easily claim that its choice not to cover a vaccinee was not “solely because of” trial participation; instead, the reason proffered might be that trial participants are, by definition, at heightened risk of HIV infection. Since trial participation correlates significantly with risky behavior such as injection drug use, a rational insurer would be remiss if she did not take this evidence of risk into account. Second, large employers, who are generally self-insured, are substantially exempted from state regulation of their insurance plans by the preemptive sweep of the Employee Retirement Income Security Act of 1974 (ERISA). Third, protections afforded to trial participants in California are largely unavailable in other states. This means that trial subjects injected in California remain unprotected when they leave the state. It also means that trial subjects injected in states other than California remain unprotected everywhere but in California. Finally, the California statute is confined to the realm of insurance; it does not address the myriad other contexts in which trial-related disparate treatment might arise. The upshot is that state-initiated measures, though well-meaning, substantially fail to advance AIDS vaccine research. The inadequacies of state measures, coupled with the shifting demographic patterns of the disease, highlight the need for a decisive federal response.

186. See id. § 121275.
187. See id. § 121260(f).
188. Id. § 121280(4)(b).
190. John Wilson has also underscored the inadequacy of state schemes to address the AIDS vaccine problem: “While in many instances the piecemeal development of law in the separate states permits each state to craft its own solutions to particular needs, uniform national legislation is still the
C. Crafting a Viable Federal Plan

1. AIDS Vaccine Research Legislation

In 1988, Congress declared AIDS a national emergency and authorized funds for research and public information efforts. The epidemic affects every state, and the search for a safe, effective vaccine surely qualifies as a national interest. I therefore concur with John Wilson's conclusion that "a way must be found to limit manufacturer concerns about liability yet provide timely and adequate compensation to injured vaccinees. Federal legislation is the best way to achieve this goal." Accordingly, the first step in addressing the AIDS vaccine effort should be to map out a coordinated national agenda.

Much ink has already been spilled in devising injury compensation schemes and federal tort relief for biotechnology and pharmaceutical companies. Virtually none of the literature addresses social harm to trial participants; rather, the focus has been almost exclusively on protecting pharmaceutical and biotechnology companies from crushing liability so as to accelerate research and development of candidate vaccines. As an initial matter, then, the federal government should develop a balanced legislative program akin to the one incipiently spelled out in California, that is, a program that addresses the needs of companies as well as the needs of those who participate in research as subjects.

Representative Nancy Pelosi and Senator John Kerry recently introduced a bill, entitled "The Lifesaving Vaccine Technology Act of 1999," that would provide a thirty percent tax credit to biotechnology and pharmaceutical companies for qualified research and development expenditures on vaccines for malaria, tuberculosis, HIV, and other diseases that kill one million or more people annually. This is a laudable first step, and

preferred route when dealing with a crisis of national—indeed, international—dimensions." Wilson, supra note 6, at 562.


192. Wilson, supra note 6, at 569.

193. See discussion supra note 6 and accompanying text.

194. As discussed supra note 6, tort relief and injury compensation schemes have almost uniformly disregarded social harms as a factor in AIDS vaccine research. For example, in a subsection entitled "What kind of injury should be compensated?" Catherine Polizzi restricts consideration to "serious, acute, systemic injury, such as anaphylactic shock or neurological disorders." Polizzi, supra note 6, at 20. Although she considers extending coverage to "development of AIDS after vaccination," Polizzi quickly decides against it because "this potentially broad coverage could drastically increase the cost of the program due to the extensive and prolonged medical care required by AIDS." Id. at 21. Polizzi's proposal is representative in this respect. Social harms rarely even appear on the radar screen of potentially compensable harms.

195. See H.R. 1274, 106th Cong. (1999); S. 1718, 106th Cong. (1999); see also Tom Abate, Should Vaccine Research for HIV Merit a Corporate Tax Break? Pelosi-Sponsored Bill May Wind Up Being a Pandora's Pork Barrel, S.F. CHRON., Apr. 5, 1999, at E1; Judy Holland, Pelosi Seeks
political leaders in both parties should take up the cause. Although the proper size of the credit can be debated, it should be beyond partisan cavil that "achieving effective and affordable vaccines for HIV, malaria, and tuberculosis will yield public benefits beyond those benefits captured by the manufacturer."196

While I agree with the general strategy incentivizing private sector research and development of vaccines, this can only be a first step in the process. Companion legislation must be drafted that will broadly outline the responsibilities of the research entities receiving these substantial tax credits. Since the trial of AIDS vaccines raises unique challenges, these anomalies must be addressed. At a minimum, participating entities should be required to do the following:

- Voluntarily submit to regulation and oversight by an expanded Office for Prevention of Research Risks (OPRR);197
- Guarantee that any successful vaccine product will be made freely available to those who have participated in clinical trials;
- Make blood assays capable of distinguishing HIV infection from vaccine-induced antibody response available to participants for life;
- Provide for risk-reduction counselling to participants throughout the trials, emphasizing that experimental vaccines will not confer protective immunity against HIV infection;
- Implement an agency-approved system of confidentiality protections to prevent disclosure of the fact of trial participation and trial data, especially blood testing results;198

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196. S. 2132 § 2(11). President Clinton recently signaled his approval of the Pelosi/Kerry approach in his state of the union address. See President William J. Clinton, State of the Union Address, January 27, 2000 (visited Feb. 2, 2000) <http://www.whitehouse.gov/WH/SOTU00/sotu-text.html> ("I propose a tax credit to speed the development of vaccines for diseases like malaria, TB and AIDS. I ask the private sector and our partners around the world to join us in embracing this cause. We can save millions of lives together, and we ought to do it.").

197. OPRR is currently located in the National Institutes of Health, but HHS Secretary Donna Shalala has directed that OPRR be relocated to the Office of Public Health and Science within the Office of the Secretary of HHS in order to "elevate its stature and effectiveness." HHS, Protecting Research Subjects (last modified Nov. 4, 1999) <http://www.hhs.gov/news/press/1999pres/991104c.html>. Currently, research funded by states and private entities may be conducted without regulation by federal agencies, a proposition that has drawn intensifying criticism. See, e.g., Jonathan Moreno et al., Updating Protections for Human Subjects Involved in Research, 280 JAMA 1951, 1954 (1998) (arguing, inter alia, that it is "striking and unacceptable that there is no universal statutory requirement for informed consent to research irrespective of funding source").

198. HHS recently promulgated proposed regulations to enhance the confidentiality of electronic patient records that could serve as a useful model for experimental research. See Standards for Privacy of Individually Identifiable Health Information, 64 Fed. Reg. 59,918-60,065 (to be codified at 45 C.F.R. pts. 160-164) (proposed Nov. 3, 1999).
• Recognize an express right to compensation for unconsented disclosure of confidential trial information (including the fact of trial participation, trial-related communications, and test results) based solely on the fact of disclosure itself—unlike a "regarded as" claim under the ADA, the claimant would be required to prove neither adverse action taken by a third party based on the disclosure nor the state of mind of the disclosing party;

• Set up a compensation fund or secure liability coverage sufficient to compensate for injuries fairly traceable to the product or to the negligence, recklessness, or intentional malfeasance of clinicians administering them;

• Agree to submit all claims for compensation to binding arbitration or agency adjudication, with provision for such a mode of dispute resolution clearly spelled out in the informed consent process.

Once these minima are established by statute, ongoing regulation and oversight should be supplied by the agency charged with implementation, most likely OPRR within HHS, as discussed below.

Supplementing this entity-regulating scheme, additional legislation must be introduced to reduce the risk of social harms to vaccine recipients more generally. This legislation could draw on various legislative precedents, but it should:

• Expand the role of OPRR, granting it the power to promulgate regulations and guidelines governing AIDS vaccine research after notice and comment from the agencies involved, affected industries, researchers, community based organizations, targeted populations, and other interested parties;

• Create an ombudsman in OPRR specifically charged with responsibility for advocating on behalf of vaccinees, providing them with legal advice as to their rights, litigating select cases on their behalf, filing amici curiae on relevant matters, and providing legislative policy

199. John Wilson has suggested, as an alternative, a liability scheme based on the National Childhood Vaccine Injury Act, 42 U.S.C. §§ 300a-1 to-34 (1988):

A victim would be required first to make a claim for compensation from a fund established by the federal government... and an award from this fund would be based upon the fact of injury and not the manufacturer's behavior. Only after the receipt of compensation, or the denial thereof, would a state claim be permitted against a manufacturer, and an election at that point (to accept the judgment or file a civil action) would be required. In the application for compensation from the government, an injured vaccinee would have to prove injury and, presumably, causation by a preponderance of the evidence and, as is the case under the National Childhood Vaccine Injury Act, that there is not a preponderance of evidence that the illness or disability is unrelated to the administration of the vaccine.

Wilson, supra note 6, at 564. While I am comfortable entertaining the notion of a federally administered compensation system, such as the one Wilson describes, I believe that research entities should provide compensation through a private fund or liability insurance in the absence of such a program.
support to aid Congress in adjusting the statute, where needed, as trial-related issues arise;

- Incentivize research and development of more sophisticated blood screening mechanisms to distinguish natural infection from vaccine response in parallel with pre-clinical and clinical evaluation of candidate vaccines;

- Mandate that entities performing blood tests, in particular hospitals, laboratories, and insurance companies, shall not disclose the fact of vaccine seropositivity to employers, insurers, or other entities, instead reporting only that a test is either positive or negative for HIV;

- Craft a system whereby trial participants can self-identify as such solely for the purpose of diagnostic blood tests, so that specialized assays can be employed in a cost-effective manner;

- Prohibit use of the fact of vaccine positivity or participation in an AIDS vaccine trial as a basis for disparate treatment in employment, in any federally funded or operated program, or as a reason for denial or limitation of insurance coverage;

- Make evidence of vaccine trial participation inadmissible at trial for criminal acts, as a basis for denial of a visa or immigration, or as a basis for adverse employment action, including failure to hire or promote, in any federal or state agency or department;

- Redouble efforts to educate the American public about the prevalence of AIDS worldwide, the principle modes of transmission, the limitations of antiretroviral therapies, the need for continued prevention, and the myths about people living with HIV/AIDS and contagion.

These and many other bold legislative solutions to the dilemma of social harms must be devised, debated, and implemented. Protection of human subjects must keep pace with accelerated research. This is especially true as the federal government wields its taxing power to incentivize research initiatives in the private sector because private money then becomes indistinguishable from public subsidies.

2. Enhanced Regulatory Oversight of AIDS Vaccine Research

While legislators should paint with a broad brush, the details of implementation should be left to the administrative bodies responsible for enforcement since they have scientific expertise and are more closely connected to our nation’s research institutions. The current system for administrative oversight of human experimentation is based on a network of local Institutional Review Boards (IRBs) concurrently regulated by the FDA and

200. Vaccine seropositivity might be treated in the same way that genetic information has been treated for insurance purposes. See, e.g., 29 U.S.C. § 1181(b)(1)(B) (1999) (restricting the use of genetic information as a preexisting condition exclusion “in the absence of a diagnosis of the condition related to such information”).
HHS regulations apply only to human subjects research "conducted, supported or otherwise subject to regulation" by federal departments or agencies that have adopted the so-called "common rule." Institutions subject to these regulations must sign prospective "assurances" of compliance with the requirements set forth in the regulations, including the establishment of an IRB to review and monitor research protocols. The FDA, by contrast, employs a system of retrospective audits and inspections to ensure that all research involving products regulated by the FDA has complied with its requirements, including in most cases IRB review and monitoring.

The IRB system jointly overseen by HHS and the FDA has recently received increasingly harsh criticism. Six areas of particular concern have been identified: (1) IRBs face major changes in the research environment; (2) IRBs conduct minimal continuing review of approved research; (3) IRBs review too much, too quickly, and with too little expertise; (4) neither IRBs nor the agencies overseeing them devote sufficient emphasis to evaluating IRB effectiveness; (5) IRBs face conflicts that threaten their independence; and (6) IRBs and their institutions provide little training for investigators and board members. Other commentators have emphasized that IRBs need to establish stronger ties with their experimental populations so as to understand and address their most vital concerns.

IRB underperformance clearly lies beyond the scope of this Comment; however, the weaknesses inherent in the current IRB system are likely to be magnified by the unique scientific and social demands of AIDS vaccine research discussed above. Enhanced administrative oversight will play a crucial role in the evolution of a safe, effective AIDS vaccine. When

201. See 21 C.F.R. pt. 56 (FDA regulations governing IRBs); 45 C.F.R. pt. 46 (HHS regulations governing IRBs). The IRB regulations provide that IRBs shall have "at least five members of varying background," 45 C.F.R. § 46.108(a). The IRB must have at least one member whose "primary concerns are in scientific areas and at least one member whose concerns are in non-scientific areas," Id. § 46.108(c). In addition, at least one member "must not be affiliated with the institution." Id. § 46.108(d). IRB members are usually "doctors, scientists, patient representatives and others who are charged with regularly monitoring the design, development and progress of the research projects being conducted at the institution." HHS, supra note 197.


204. See 46 C.F.R. § 46.103.

205. See 21 C.F.R. § 56.103.


207. See id.

208. See, e.g., Marjorie M. Shultz, Legal and Ethical Consideration for Securing Consent to Epidemiologic Research in the United States, in ETHICS AND EPIDEMIOLOGY 107 (Steven S. Coughlin & Tom L. Beauchamp eds., 1996) (arguing that IRBs need "to get a broad and deep, rather than token, response about risk and consent from the types of people who are likely to populate research studies").
combined with truly comprehensive AIDS vaccine legislation and supporting regulations, such oversight will protect the physical and social well-being of research participants. Specifically, a newly expanded OPRR should undertake the following:

- Promulgate regulations and guidelines governing appropriate recruitment and screening techniques for AIDS vaccine trial cohorts, informed consent procedures and documents, participant confidentiality, provision of risk-reduction counselling, access to product-specific blood assays, and researcher responsibilities upon termination of trials;
- Mandate that IRBs considering the recruitment, trial design, protocol drafting, informed consent procedures, implementation, and oversight of AIDS vaccine trials include an individual who will be vaccinated;
- Require that Community Advisory Boards (CABs) be established with trial participants, members of targeted communities, and representatives from community based organizations to provide community perspectives on all trial-related matters to researchers, IRB members, trial participants, and involved agencies;
- Assemble and disseminate educational materials and provide on-site trainings for IRBs concerning the physical and social risks inherent in AIDS vaccine trials—such materials and presentations should emphasize that potential social harms are to be factored into the calculus of risks and benefits for approval and oversight of study protocols.

209. IRB regulations presently include special provisions for research in vulnerable populations such as pregnant women, 45 C.F.R. § 46.207, children, id. § 46.408, and prisoners, id. § 46.305. IRBs are required to include a prisoner in reviewing any human subjects research involving prisoners. See id. § 46.304.

210. Although still a germinal concept, CABs should play an increasingly prominent role in the conduct of AIDS vaccine trials from trial design to vaccination and beyond. As Bill Snow observes, the Division of AIDS (DAIDS) already has seven fully operational CABs participating actively in clinical trials of AIDS treatments, prevention, vaccines, and in natural history studies of HIV. See Bill Snow, Community Advisory Boards, in HIV VACCINE HANDBOOK, supra note 181, at 149-53. These local CABs have elected a National CAB to coordinate conference calls, address issues of concern common to the study sites, and advocate for change through contact with government officials and principal investigators:

Hundreds of times CABs have identified potential problems for investigators . . . or helped make trials more ethical, attractive, and feasible. In addition to their advisory and watchdog function, CABs have helped their communities by disseminating information, arguing the case for volunteering, and reaching people with AIDS and the stigmatized communities that bear the brunt of the epidemic.

Id. at 150. CABs, and other mechanisms to insure that the input of affected communities is taken into account at every stage of the research process, can provide particularly helpful guidance where social harms issues may arise. They should therefore be mandatory, regardless of whether the study involved is conducted in the private sector, by government-funded entities, or some combination of the two.

211. As the Belmont Report states,

[i]the requirement that research be justified on the basis of a favorable risk/benefit assessment bears a close relation to the principle of beneficence . . . Many kinds of possible harms and benefits need be taken into account. There are, for example, risks of psychological harm, physical harm, legal harm, social harm and economic harm and the corresponding benefits.
Separate discussion of informed consent is in order since it is the mainstay of the current regulatory protection of research subjects. As Margaret Johnston and Sam Avrett explain:

The informed consent process should explain the goals of the trial, the reason that the trial would benefit by the participation of the individual, what participation means, the potential risks and benefits of participation to the volunteer and others, alternatives to participation, issues of confidentiality, any compensation for trial participation and for any costs or injuries resulting from trial participation, and the right to end participation at any time.

In addition to these generalized requirements of informed consent, I would stress:

While the most likely types of harms to research subjects are those of psychological or physical pain or injury, other possible kinds should not be overlooked. BELMONT REPORT, supra note 182, at 38. On the spectrum of potential harms that need to be assessed, see KATZ, supra note 182, at 323-69, 435-62.

212. HHS regulations provide that “no investigator may involve a human being as a subject in research covered by this policy unless the investigator has obtained the legally effective informed consent of the subject or the subject’s legally authorized representative.” 45 C.F.R. § 46.116. In her foundational book The Search for an AIDS Vaccine, Christine Grady has suggested that “community informed consent,” not simply individual consent, should be secured for research trials:

Conceiving of individuals as inextricably related to and deriving identity from their community gives the individual a stake in research that is good for the community. For vaccine research especially, a system whereby consent to conduct the research was obtained from the “community” and input was obtained from the community about how to conduct it, followed by individual consent from the each of the participant-subjects, should be the ideal. Such a system, in which community members would deliberate together about the ends and methods of research, would give the individual community members some responsibility in supporting the research. Such active participation in deciding the ends and the means promotes the interests of the individual and the community.

Grady, supra note 7, at 84-85. Grady envisions this “community consent” playing a role at all stages of the trial, starting with preliminary meetings between community leaders and scientists; leading to “town meetings” with researchers presenting the aims and methods of the research to community members to educate and inform them while also seeking their input; and, finally, the community would actually vote on whether the proposed trial should be conducted. Id. at 145. Once a trial is approved, “[a]n advisory board or working group representative of the community should be created to discuss and negotiate the specifics of protocol design and methods.” Id.

Chris Collins, however, has posed trenchant questions regarding Grady’s proposal. After praising Grady for her “laudable contribution,” Collins notes that embedded in her thinking are “major assumptions about communities,” for example,

that their membership is definable and relatively cohesive; that divisions within communities do not prohibit reaching credible decision for all members; that self-perceived membership by trial volunteers in multiple communities will not render decisions by one community illegitimate; and, finally, that communities are comfortable choosing representatives or will be satisfied with a majority vote at a meeting as a valid decision making process for the whole group.

Collins, supra note 7, at 8. Collins ultimately concludes that, although “communities may be the ultimate beneficiaries of vaccine research, individual trial volunteers remain the ultimate arbiters of the merits of a trial.” Id. at 9. Although I am generally sympathetic to the vision Grady advances, I agree with Collins that informed consent properly should remain a matter of individual conscience. As the rest of my proposal should make clear, though, I believe that all of the communities affected by AIDS vaccine research should play an active role in the decision making regarding trials.

That the potential risks and benefits must include social as well as clinical risks and benefits;
• That legal and administrative remedies are available should adverse events occur;
• That a clear, concise explanation of the process for filing research-related claims, together with the contact information for the OPRR ombudsman, should be provided;
• That study-related documents, videotapes and other recruitment and consent instruments should be translated into the language of all prospective participants to ensure genuine voluntarism.

Simply put, the “informed consent process should ensure that the potential participant understands the information and that the person’s choice to participate or not is voluntary.”

CONCLUSION

Unfortunately, most discussions of vaccine research schemes have either overlooked social and related harms or consigned them to vague treatment. President Bush’s Commission for the Study of Ethical Problems in Medicine and Biomedical and Behavioral Research, for example, repeatedly stressed the narrow scope of harms that should be addressed by national compensation programs: “Benefits should be provided on a nonfault basis to subjects sustaining nontrivial bodily injuries or death as a result of their participation in covered research.” The Commission briefly considered social injuries, but rejected suggestions that they should be compensable:

The Commission understands the term “social injury” to refer to injuries to reputation, personal relationships, or legal status resulting from the disclosure of identifiable personal information gathered in the course of research. The Commission believes that the most serious problems likely to arise in this area (e.g., unauthorized disclosures of sensitive private information) are better handled by a combination of careful prior review of the design of research projects posing such risks and, where necessary, pursuit of legal remedies through the courts.

Social harms, as I have argued, should be factored into the calculus of risks and benefits at every stage of the AIDS vaccine research agenda, not just in trial design and eventual litigation. Although compensation programs for

214. Id.
215. See discussion supra notes 6 & 194 and accompanying text.
216. President’s Commission for the Study of Ethical Problems in Medicine and Biomedical and Behavioral Research, Compensating for Research Injuries A Report on the Ethical and Legal Implications of Programs to Redress Injuries Caused by Biomedical and Behavioral Research 129 (1982).
217. Id. at 137.
trial-related social injuries may present administrative and budgetary chal-

lenges, they should not be dismissed without due consideration. Indeed,

if social harms reach a level sufficient to threaten enrollment in Phase III

trials, compensation guarantees may present one of the few viable options
to facilitate testing of promising candidate AIDS vaccines.

The transition to large-scale domestic AIDS vaccine trials represents a
daunting challenge and a prime opportunity to bring our nation’s experimental research structures into the twenty-first century. As the director of the Office for Protection from Research Risks has stated: “This is precisely the time to take constructive account of the notes of caution being sounded and to reform, correct, revise, and improve the dynamic and evolving system that keeps those who are enrolled in research out of harm’s way.”

A coordinated effort—including federal legislation authorizing limited tort relief for biotechnology and pharmaceutical companies engaged in AIDS vaccine research, a scheme for compensation of trial-related injuries, and enhanced regulatory oversight of the research process addressing not only physical but also social risks to participants in the experimental process—will be a signal achievement and necessary step on the road to a lasting cure for the AIDS pandemic. President Clinton has articulated the vision: “My fellow Americans, if the 21st century is to be the century of biology, let us make an AIDS vaccine its first great triumph.”

It remains to be seen how this triumph will be accomplished. One thing, however, is certain: The biomedical breakthroughs that President Clinton envisions will only be achieved if those who put their lives on the line in the fight against AIDS are adequately protected against physical and social harms.

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218. I agree with the UNAIDS guidance on this point: “In addition to compensation for biological/medical injuries, appropriate consideration should be given to compensation for social or economic harms, e.g., job loss as a result of testig positive following vaccine administration.” UNAIDS, supra note 70, at 12.
