AIDS, Rhetoric, and Medical Knowledge

ALEX PREDA

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Introduction

AIDS and Scientific Knowledge

Brightly colored condoms, arranged in the shape of bicycles, eyeglasses, or flowers: part of an extensive campaign against the AIDS risk, these have been a common sight on billboards in Germany for several years now. An advertising spot presented on the Arte television channel (which defines itself as the cultural television channel of Europe) calls on viewers to “fight together.” The spots on German television (distributed by both private and public channels) are about “not giving AIDS a chance.” At the beginning of December, the major television and radio stations, advertising companies, and the press reminded the public not only about Christmas and family values, but also about risks, being safe, and not giving viruses any chance to spread. Since December 1st was declared World AIDS Day, the AIDS risk has been featured regularly in the media in the pre-Christmas period. Not that this topic is completely absent from the media in the first eleven months of the year; in fact, the opposite is true. The activities around December 1st are simply an extra reminder to be vigilant, keep up the fight, and not give this deadly enemy any opportunity. And fight it the populace must because these risks seem now to be almost everywhere.

The media have alerted people to “contamination risk,” “occupational risk,” “technological risk,” and “Third World risk.” In the 1990s, cases of patient–physician or patient–dentist contamination (Stine 1993, p. 418), and blood bank and organ transplant contamination – to name only a few of the situations highlighted by the media in
Western Europe and the United States—gained prominence.1 The rapid spread of AIDS in underdeveloped and developing countries has also been a major topic. Issues such as “risk factors” and “risk behavior,” along with the latest epidemiological trends and “risk groups,” old and new, have received media attention. With the advent of a number of epidemiological models, there has been a globalization of “AIDS risk” as well (Mane and Aggleton 2001, p. 23; Maticka-Tyndale 2001); since the end of the 1980s, the AIDS risk topics featured in the press and on radio and television have multiplied and diversified. This public presence of AIDS has been amplified by its being made a subject for novels, plays, docu-fictions, Hollywood-style and French existentialist movies, television medical drama series, votive painting, and avant-garde artworks, among other things (Treichler 1993; Miller 1992).

Reports and articles about “risk behavior” and “factors” in various parts of the world are not a rarity. Tourists and travelers are warned about them when traveling to some region with a “risk pattern.” Host countries, when not adopting concrete legislation, are thinking aloud about screening the risks tourists might bring in with them. In 1994, when the organizers of the Tenth International AIDS Conference in Yokohama announced in their preliminary programs2 that nobody coming to Yokohama to discuss risk reduction (among other topics) would be denied a visa because of his seropositive status, they implicitly asserted that the exceptional character of the occasion legitimated an exceptional, temporary suspension of risk screening.3

Health institutions have been confronted with the topic of “AIDS risk” from the beginning: the reaction to this challenge has been to enact measures for preventing, screening, coping with, controlling, or minimizing risks. This implies, among other things, increasing the knowledge of various social groups about AIDS risk; inducing overall

1 Cases of dentist–patient contamination have been much publicized in the United States, whereas the theme of blood bank contamination seems to be a European one; the most prominent cases were recorded in France at the end of the 1980s and in Germany in 1993–4. Both events enjoyed a large amount of publicity and have been debated in courts of law.
2 See, for example, the Advance Program of the Conference, p. 41; also, www.aidsinfobs.org/periodicals/amin/1987/127.05. Downloaded on May 13, 2004.
3 According to reports in German newspapers (Tageszeitung, August 6, 1994, pp. 1, 3; Frankfurter Allgemeine Zeitung, August 6, 1994, p. 7) there were attempts on the part of the organizers to forbid seropositive conference participants from entering Japan.
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behavioral changes supposed to be risk-reductive; increasing the knowledge of public health institutions about individual and collective risks; systematically monitoring these risks in one form or another; preparing healthcare institutions to meet future challenges, according to knowledge about risk; and modifying other policies (concerning insurance and immigration, for example) according to the same knowledge. This broad spectrum of risk-reduction policies has been implemented in many countries.

Many social studies of AIDS operate with and have a concept of “risk” at their core: they describe individual and collective risks, analyze their avoidance, or examine social and behavioral “risk factors.” “AIDS risk” has also become an important topic for health economics and for calculating the present and future costs of medical care, research, and drug development. Social security institutions, insurance firms, as well as courts of law, have been confronted with the relationship between AIDS risk on the one hand, and responsibility, care, partnership, and general human rights on the other.

At perhaps a deeper level, “AIDS risk” continues to be a topic for biomedical research. In its basic and applied aspects, research is oriented according to certain criteria of “risk persons,” “risk groups,” “behavior,” and the like. Drug design and clinical trials, as well as clinical and epidemiological studies, constantly operate with notions of risk: at their core is the effort to construct trial groups as homogeneously as possible according to risk criteria. Especially in the United States, this has generated much criticism from activist organizations; counter-trials have become part of an alternative expert culture (Arno and Feiden 1993; Epstein 1992, 1996).

AIDS risk is then a topic for (1) clinical and epidemiological research; (2) applied pharmaceutical research; (3) public and health policy; (4) politics, economics, ethics, and law; (5) the social sciences; (6) the media; and (7) the arts and entertainment industries. What these approaches have in common, in spite of their diversity, is the assumption that notions such as “AIDS risk,” “risk factors,” “risk behavior,” “risk groups,” and “populations at risk” can be understood because they are ultimately grounded in a body of expert medical knowledge about AIDS. In other words, this body of knowledge about the syndrome, its modes of transmission, and the nature of the infectious agent is taken as reliable ground for specifying other aspects and implications
of “risk.” “AIDS risk” as an issue for expert, scientific knowledge precedes particular (political, juridical, economic) redefinitions of risk. Scientific knowledge determines what “risk” is and what it is not, and how it can be assessed in its various aspects.

The relation of precedence is understood as a logical as well as an empirical–historical one. Its empirical–historical dimension is given by “AIDS risk” initially appearing as a medical issue. Its logical dimension is that “AIDS risk” as a medical topic is necessarily prior to its being a topic for health, insurance, or legal policies. It is hardly imaginable that “AIDS risk” would be referred to without appealing in some way to scientific knowledge. Even mid-1980s televangelists preaching that AIDS was the wrath of God visited upon sinners took care to legitimate their statements by constantly referring to this knowledge (Patton 1985; Treichler 1988b). References to expert knowledge and the experts’ presence are constant features of the media’s handling of the issue. The idea that this knowledge is a necessary condition (in both the logical and the empirical sense) for analyzing particular aspects of AIDS risk can also be found in historical accounts (e.g., Grmek 1990), as well as in many social studies. They all refer to expert knowledge not only as a source of authority and legitimation but also as the epistemic condition for “AIDS risk.”

Scientific Knowledge and Rhetoric

At the center of this book lies the relationship between rhetoric and scientific knowledge about AIDS. In this, I depart from the thesis of AIDS as a “full blown medical and cultural phenomenon” (Sturken 1997, p. 147), which implies that these two aspects are completely separate and brush against each other only at their fringes. I examine here their entanglement at the core of scientific knowledge. There are several social sites where scientific knowledge about AIDS is produced: research institutions, laboratories, clinics, operating theaters, and treatment centers. Moreover, as Steven Epstein (1996) has shown, social movements and alternative organizations are large, significant sites of knowledge production. The study will concentrate on only one such site, one which does not even appear in the previous enumeration; indeed, it does not appear to be a site at all: or, if it is one, then it is very, very flat. It seems to lack the richness, depth, and complexity of the
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lab, the clinic, and the operating theater, and the vigor, determination, and commitment of social movements. It consists of a thousand disparate pieces which circulate constantly, continuously appearing and disappearing in all sorts of places. This site consists of expert articles on AIDS in medical journals; they are what form the core of what is known as medical AIDS discourse. (That a text can be and is a social site is argued at length in the pages to come.)

Seeing journal articles as knowledge-producing social sites may appear paradoxical; after all, a (scientific) text is ultimately merely a vehicle for expressing knowledge produced elsewhere, a means for transmitting knowledge, not an engine that constitutes it. In expressing knowledge, texts may rearrange and reconfigure it according to the logic of literary representation and the canons and conventions of scientific prose (e.g., Gross 1999; Prelli 1989; Knorr 1981). Textual resources, the nature of which is ultimately rhetorical (Fish 1989, pp. 472–3), can perhaps persuade (which is in itself bad enough) but cannot produce knowledge. In other words, a (scientific) text can (more or less successfully) convey its knowledge content to the reader by using rhetorical devices – i.e., it can persuade the reader that something is the case, but its task ends there. Instruments of persuasion may have different forms: coherence and rigor in textual organization and an apparent minimum of rhetoric (as is common in scientific texts) are only two examples of rhetorical strategies. However, such texts remain no more than instruments for transmitting something, or to put it more colloquially, for selling some knowledge content to the reader.

Moreover, isn’t rhetoric (that of scientific texts included) contingent upon the skills of the author and, therefore individual, fluctuating, and non-standardizable? Does it not, ultimately, belong to the realm of the literary critic, and exclusively so? To make matters even worse, what about the rhetoric of this text? Isn’t it proof of what Woolgar and Pawluch (1985) would call ontological gerrymandering, when a text claims to have something sociologically relevant to assert about the textual (i.e., rhetorical) production of knowledge by pretending not to have any rhetoric – or, if it has, that it is just an innocent means of conveying some external knowledge?

In setting myself the aim of looking more closely at “AIDS risk” in this book, I was confronted with the ways in which rhetoric appears to insinuate itself parasitically into scientific knowledge. For if rhetoric
is supposed to not have any place in scientific texts, yet invariably insinuates itself into them, how else can it be regarded than as a parasite that lives and feeds on the knowledge content it helps convey to readers? It may successfully persuade skeptical readers; the usual scientific rhetoric of clarity and rigorousness may help convey the message better, but it is still a parasite. Worse still, in this light, do (scientific) texts not actually start to look like parasites on the activities through which scientific knowledge is produced? Do they not live on the richness and complexity of the local production of (scientific) knowledge? If there is something to be said about this, then texts are not the place to look: they may say something about communicating, about transporting this knowledge, about making it available to the public – but not about its production. In the flatness of a (scientific) text, one is confronted with the rhetoric that lives and feeds on the knowledge content and therefore should be rigorously separated from it, but how?

I argue that:

1. Texts are not to be viewed as flat, thin conveyors of knowledge, but rather as social “dispositives” (Derrida 1972a, p. 339).
2. Rhetoric is not the (more or less sophisticated) form of the knowledge content, meant only to persuade the reader that something is the case, but a social practice producing knowledge.

Arguments contesting the parasitic position of rhetoric with respect to the authorial intention and to content are not new: they have almost become commonplace in the fields of literary studies (De Man 1983; Fish 1989), historiography (White 1985, 1987), anthropology (Geertz 1988), and economics (McCloskey 1998, 1990, 1994). Arguments about the conceptual primacy of writing and texts for the social constitution of meaning are also commonplace in so-called deconstructivist philosophy (e.g., Derrida 1972a,b, 1979; Sarup 1988; Norris 1990). In the field of sociology, the idea that texts should be viewed as social dispositives and rhetoric as a social practice is a matter of debate and dissension. More recently, Actor-Network Theory (ANT) has argued that texts act as “immutable mobiles” (Latour 1999), transporting knowledge across various contexts and disentangling it from local practices. The sociology of knowledge and science has shown the double (local and textual) embeddedness of scientific knowledge (Knorr 1981; Latour and Woolgar 1986), its reconfiguration according to the
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logic of literary representation (Woolgar 1988; Potter 1988), as well as
the role played by rhetoric in the constitution of scientific knowledge
(Prelli 1989; Gross 1996; Gragson and Selzer 1993; Berkenkotter and
Huckin 1995; Ceccarelli 2001; Fahnestock 1999; Halliday and Martin
1993; Montgomery 1996; Myers 1990; Swales 1990). The arguments
for rhetoric as a social practice are presented and detailed throughout
the study not in a purely theoretical fashion but by means of examin-
ing the concrete historical constitution of scientific knowledge about
AIDS.

The first argument is this: what would appear to be nothing more
than strategies of argumentation actually played a constitutive role
with respect to the primary knowledge about the nature of the infec-
tious agent, its means of transmission, and its causal role in the Ac-
quired Immunodeficiency Syndrome. In other words, social representa-
tions of “risk” are intrinsic to this knowledge. This means that both the
conditions under which it becomes possible to speak about a new syn-
drome and the concrete forms taken by the scientific knowledge about
the syndrome, its causal agent, and its modes of transmission were gen-
erated by representations of risk. They played a central part in making
the Acquired Immunodeficiency Syndrome the Acquired Immunodeficiency
Syndrome – that is, a condition under which old, familiar dis-
eseases became new, complex, previously unseen diseases. Moreover, they
were central in shaping knowledge about the nature of the infectious
agent: something coming out of the environment, a behaviorally de-
termined agent, a gender- or genetically determined predisposition, or
a mixture of all of these. Later on, when it was debated whether the
French or the American retrovirus was the causal agent, these represen-
tations were at the core of the two sides’ arguments: both vigorously
contended that theirs was the etiological agent because it fit patterns of
risk. In shaping medical knowledge about the retrovirus, its effects, and
its means of transmission, risk representations also constituted an order
of knowledge from which they themselves emerged as secondary and
derived, and as feeding on the essential medical knowledge about the
syndrome. Risk representations emerged as dependent on whether the
causal agent is environmentally or sexually transmitted, spatial loca-
tion, gender particularities, and membership in certain population seg-
ments – i.e., on factors derived from knowledge about the causal agent
and how it is transmitted, which, in turn, were constituted by “risk.”
Scientific Knowledge and the World Risk Society

Scientific representations of risk become fully relevant only if we consider them against the broader picture of the world risk society. In the past decade, the notion of risk society has attained a visibility comparable to that attained by the concept of “postmodern society” in the 1980s; intellectual fashions aside, this notion helps us better understand the broader significance and consequences of scientific knowledge about risk.

The sociological concept of risk is usually understood in opposition to the notions of uncertainty and danger. Whereas uncertainty designates lack of valid knowledge about a present or future event, risk implies a set of procedures and techniques through which valid, albeit probabilistic knowledge about the event in question is obtained. Risk emerges when social actors are able to compute the probability of a (natural or social) event, as, for example, when social organizations compute the probability of a technological failure and forecast its consequences (as in the case of electricity grid failures) or compute the rate of spread of infectious diseases (SARS is a good example here).

Analogously, at a basic level the notion of danger presupposes an undesirable (natural or social) event occurring with a lack of social knowledge about its causes, concrete shape, and consequences. By contrast, risk implies a set of tools and procedures through which knowledge about the causes, shapes, consequences, and means of prevention of undesirable events is gained. In both pairs (risk/uncertainty and risk/danger), the concept of risk is grounded in tools and procedures through which unknown events are made into an object of analysis and valid expert knowledge is gained. This body of knowledge enables social actors and institutions to devise paths of action, maintain trust, make decisions, and prevent or reduce the consequences of undesirable events.

It follows, then, that expert scientific knowledge plays a central role with respect to risk. At the macro-social level, however, the picture becomes more complicated. Roughly speaking, we encounter two main theories about how risk works at this level: a systemic approach promoted mainly by Ulrich Beck (1992), Scott Lash (2000), and Niklas Luhmann (1990), and an anthropological one promoted by
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Mary Douglas (1992a, 1985) and Aaron Wildawksy (Douglas and Wildawksy 1982).

Beck’s argument is that processes of social modernization (individualization, industrialization, the penetration of technology into all spheres of social life, and the expansion of capitalist exchanges) bring with them not only benefits, but also undesirable effects (e.g., technological failures, epidemics, economic recession, and environmental destruction). Once these are recognized, science is called upon to analyze them and devise countermeasures. Scientific knowledge lies at the core of modernization processes, and the solutions it provides are inescapably scientific: analysis and knowledge will be used to counteract the undesirable effects of modernization. But there is no guarantee that these measures designed to counteract risks will not, in turn, have undesirable side effects. This, in fact, happens in many cases. The social consequences, argues Beck, are manifold: late modern societies learn that total indemnity from risks is impossible. They have to reflect constantly upon the social consequences of the decisions taken at the collective, institutional, and individual level; risk society implies then a stage of advanced modernization, where society “disenchants and then dissolves its own taken-for-granted premises” (Beck, Bonss, and Lau 2003, p. 3).

Another consequence is that risk groups occupy a prominent place in the social fabric: they are defined by their exposure to undesirable events and by their means for reducing exposure (Scott 2000, p. 35). This is evident in the process of biomedicalization, among others, where the health state of individuals is comprehensively monitored on a mass level with the help of standardized risk-assessment tools (Clarke et al. 2003, p. 172).

Yet another consequence is that, due to globalization processes, risk society becomes world risk society: undesirable events can no longer be geographically contained but rather unfold on a planetary scale. Epidemics such as SARS (which surfaced simultaneously in several cities on two continents) and AIDS are cases in point.

There are, however, still more implications: developed societies learn that the total management of undesirable effects is impossible, but in this process they are confronted with the fears and anxieties of their citizens. A major social institution that should alleviate fears and
restore trust is science itself, because undesirable effects cannot be managed without scientific expertise. The increased need for expertise in all domains of social life gives rise to a class of “professionals of representation, simultaneously oriented towards their constituency (social reality, the citizenry) and their professional rivals (fellow scientists and politicians)” (Pels 2000, p. 7). Several levels of dialogue have to be maintained in the social management of risks: a dialogue among experts/scientists, as well as dialogues between the general public and scientists, and between policy makers and scientists. In many cases, group interests intervene in this dialogue and can shape it in decisive ways (Brint 1994, p. 18).

The maintenance of social order also requires trust in social institutions, which in turn requires the ability of these institutions to account for events. This implies, among other things, that responsibility is assumed and blame is ascribed. The notion of risk intervenes in this process: Niklas Luhmann (1990, pp. 10, 23; see also Nelkin and Gilman 1988) argues that causes of undesirable events can be attributed either to one’s own social institutions (and they become risks) or to external entities (natural and supra-natural forces, external enemies, and radically different societies), in which case they become dangers. “Risk” is not only a tool for assessing the probability of undesirable events, but also a device for attributing responsibility, maintaining trust, and ensuring social order.

In a similar line of argumentation, Mary Douglas (1967) sees risk as a cultural component of social order: social cohesion, she argues, is determined by the degree of internal and external cohesion of social groups, among other things, as well as by the categories with which these groups operate. In making use of categories such as pure/impure, safe/unsafe, social groups establish paths of individual and collective action and, at the same time, trace the boundaries of their social world. From this perspective, risk appears as one of the categories with the help of which social actors make sense of their world: it is used for defining responsibility, placing blame, establishing accountability, and maintaining trust. At the same time, risk is a device with the help of which fundamental distinctions between society and nature are established: we talk about risks generated in our own society, but we talk about dangers coming from nature or from other societies perceived as radically different (e.g., in the case of terrorism).
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Ultimately, risk appears as irreducible to a set of technical procedures for estimating the probability and harm degree of events: “it is cultural perception and definition that constitute risk” (Beck 2000, p. 213; emphasis in original).

There are several important implications here: the first is the distinction between scientific knowledge and cultural definitions of risk. According to this distinction, scientific knowledge is influenced in its interests, but not in its substance, by cultural perceptions of risk. These may orient the focus of research, whereas the content of scientific knowledge is determined by other factors.

The second implication derives from the first: because society is constrained to reflect upon the risks it generates and scientific knowledge is distinct and separated from broader cultural perceptions, experts must enter into a dialogue with a concerned public to find effective ways of preventing and/or avoiding risks. This dialogue is an intrinsic feature of reflexive modernization: examples here are the dialogue between AIDS experts and alternative AIDS organizations (Epstein 1996), between experts and environmental groups, and between nuclear scientists and concerned farmers (Wynne 1996). Such dialogue requires a “public understanding of science” (see, e.g., Locke 2002), that is, social groups that acquire a relevant amount of expert knowledge and efficiently translate their own viewpoints into the language of science.

A third and even larger implication concerns democracy itself: if scientific expertise plays such a prominent role in all domains of social life, to what extent is the democratic decision-making process influenced by it? Several authors have recently argued that “technical democracy” (Callon, Lascoumes, and Barthe 2001) or “expert democracy” (Turner 2003), with scientific expertise at its core, raise important problems with regard to transparency, dialogue, civil society, and participation in policy-making.

With respect to the topic examined here, these implications can be specified as follows:

1. Can we maintain a sharp distinction between scientific knowledge about AIDS and cultural representations of risk?
2. To what extent is this knowledge influenced in its very substance by cultural representations of risk?
3. How do such representations work and what is their effect?
4. What are the practical consequences of (2) and (3) for the organization of AIDS research, prevention, and treatment policies?

5. What are the challenges posed to the “expert democracy” by scientific knowledge of AIDS?

Seen in this perspective, an examination of the ties between scientific knowledge and “AIDS risk” has deep implications, addressing the possibility of an informed dialogue, the participation of the public in policy-making, and the nature of the “knowledge society” itself. In Chapter 7, I discuss these implications in more detail. For now, I turn to how “AIDS risk” works with regard to scientific knowledge.

What Is “AIDS Risk?”

(1) At the first, basic level, “risk” can be regarded as a rhetorical device aimed at enhancing authors’ illocutionary force (Austin 1970, pp. 235–52). This is what emerges if we look at the usual opening or closing sequences of a medical paper on AIDS. Many opening sentences say something like, “In this paper, we study the risk of transmission . . . ,” or “We report [the occurrence of x] in a risk population . . . .” Closing sequences repeat the pattern in a somewhat changed form: “The findings support the view that risk of transmission . . . ,” or “The study of this risk population shows that . . . .” In these cases, the illocutionary force of “reporting x” or “studying y” is enhanced by “risk”: one reports or studies this or that not for its own sake but because of risk. In other words, “risk” is a tool or device by which a text formulates claims about its epistemic intentions and assertions, and about its position with respect to other texts.

(2) At a further level, “risk” can be seen as a classifying device: it establishes limits (i.e., categories) within which a certain form of pneumonia or skin cancer is to be seen as “normal” or “usual.” It also establishes by whom a retrovirus can be sexually transmitted, and how. One and the same form of pneumonia or skin cancer can be classified with the help of “risk” as unusual, problematic, previously unseen, or as seen in a category where it is not possible for it to be seen otherwise. Risk defines the domain of the possible, traces its limits, and shapes a pattern of knowledge. As such, “risk” produces categories of everyday medical practice and of everyday life. These categories are
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constitutive for the patients’ identities, how they account for infection, and the physicians’ management of the syndrome. One example is the classification of risk subjects through medical interview practices, in which the interviewees ascribe themselves to a category that is taken for granted by virtue of the operation of ascribing. Another example is that of AIDS patients’ self-classification in everyday life, as belonging to a clear-cut risk category, and their continuous identification with that category, even if their personal circumstances are much more complex (Carricaburu and Pierret 1992). Another example is that patients classified as belonging to a risk category are more ready to accept (and in some cases even expect) a diagnosis of HIV infection. Patients with similar symptoms who perceive themselves as non-risk are much more reluctant to accept such a diagnosis.

More generally, “risk” is a device that classifies and reclassifies diseases as seen/unseen, usual/unusual. It is generated by the work of ascribing different meanings to these diseases according to the social categories to which they are assigned. The figure of “risk” plays an instrumental role in the construction of AIDS as a phenomenon in its own right and acts as a negotiating device with respect to its definition. Moreover, the syndrome has varying meanings depending on the risk categories to which it is ascribed. Because “risk” is a device for defining the disease and classifying its forms, it can be seen as a set of classificatory operations and their outcomes.

(3) At a deeper level, “risk” acts as a device for producing causality from and through agency. This may seem paradoxical, because causality and agency are mutually exclusive: the retrovirus entering the bloodstream and attaching itself to the surface of CD4+ cells, reproducing itself in these cells and exhausting them, and so forth, cannot be represented as having purposeful agency. But it is agency, presented in terms of risk, that makes possible the construction of various forms of natural causality: descriptions of natural events leading to infection and to the syndrome (even when given in the language of protein strings and biochemical reactions) are embedded in discourses about agency. The natural history of the causal agent is produced from the social history of the patient. For example, the (biochemical) description of the way in which amyl nitrites may affect the immune system and lead to immunodeficiency is made possible by, and grounded in, a discourse about the risk agency of people belonging to some urban
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subcultures – people who sleep little, spend a good deal of time in discotheques, and have excessive amounts of sex, exhausting their bodies to the point where the amyl nitrates consumed interact with parts of a weakened immune system whose cells have been partially depleted. Another example: the (physiochemical) description of how the HI-virus is passed from women to men (a medical mystery for a very long time) is tied to narratives about uncircumcised African men, whose long foreskins covering the penile shaft oversensitize the penile glans and are a medium for infections. These narratives are complemented by those about tribal traditions forbidding circumcision, which are encountered in exactly those places where infection with HIV is at its highest. Representations of social agency frame the physiological discourse about the HI-virus entering the body; the latter, in turn, confirms and legitimates the social risk.

(4) “Risk” is a device that accounts for the order it produces and for the construction of natural causality through agency. In an order of knowledge with heterogeneous categories (homosexuals, Haitians, African men, prostitutes, female sexual partners, drug users, infants, blood parts recipients), each risk category defines itself via difference (homosexuals are non-Haitians, non-Africans, non-infants) and by reference to the classification system.

Another device is provided by the narratives on how the infectious agent was transmitted from one risk category to another: from primates carrying the virus to Africans, then from the latter to Haitians working in Zaire, from Haitians to homosexuals, from them to drug users, then to female sexual partners, to infants, prostitutes, and so on.

A third mechanism is that of constructing a past for the present – by showing, for instance, that risk had already been there for a long time. This is illustrated by the post hoc (and ad hoc) proofs of antibodies to varieties of HIV in blood probes from various risk categories, collected well before the first reports on the syndrome. Another illustration for the case in point is the reinterpretation of clinical files of persons deceased in the 1960s and the 1970s as being actually indicative of an AIDS diagnosis.

A fourth device is the reconstruction of social agency from relations of natural causality: of the heterosexual male risk as derived from the retrovirus entering the body through the oversensitized penile glans, or of the homosexual male risk as derived from the single-cell lining of the
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rectum. From this perspective, “risk” appears not as a result of etiologic and epidemiologic models of disease, but rather as a device that plays a role in the construction of these models, enabling the representation of (1) disease origins and (2) etiologic agents.

(5) “Risk” also appears as a device for producing the future from present orders of knowledge and the corresponding relations of natural causality. The common acceptance of risk as the computable probability of something occurring in the future occurs at this level. In the present context, however, the question is a more complex one. It can be formulated as follows: how does it become possible to produce computable probabilities from heterogeneous social categories? Under which conditions are these categories invested with forecasting power? For example, how does it become possible to compare “quantities” of risk of Kaposi’s sarcoma in homosexuals and in the general population, starting from the premise that Kaposi’s sarcoma is so rare, so problematic, that it is not even seen in the general population? How does it become possible to compute the “quantity” of risk of AIDS in the general population, under the premise that AIDS risk is actually category-specific? The answer requires taking into account the devices by which “quantities” of risk are produced from qualitatively different risk categories, as well as the ways in which these “quantities” of risk (re)produce qualities of risk. The construction of risk-in-the-future implies several transformations of distinct risk qualities into “quantities” and the reworking of “quantities” into distinct qualities. It implies, moreover, a “normal” risk – expressed in the statistical figures showing how many sexual contacts, what age, what geographical location, and which gender constitute the norm of being at risk of getting the HI-virus. It implies wiping out accidents, individual idiosyncrasies, and so forth, in favor of figures showing what it means to be a person normally at-risk, with which everyone can be compared. The future can be produced from the present because of the work done by “risk.”

The representation of AIDS risk as a computable probability rests on this classificatory system, which allows the transformation of heterogeneous categories into “quantities.” Conversely, risk as a quantity reinforces the categories of the system. Consequently, “risk” should be regarded neither as a natural fact mirrored by the expert discourse, nor as a simple corollary of medical knowledge about the infectious agent. It is, rather, a complex, multilayered result of classification
operations, a device for producing classifications, a strategy for setting up etiologic models, a device for providing the syndrome with a cluster of meanings, and a concrete quality resulting from quantifying and amalgamating various other qualities.

These different dimensions might give the impression that rhetorical practices of risk are graded from the simplest to the most complex, and that, accordingly, the simplest would matter less than the more sophisticated ones. In this perspective, risk as an illocutionary force in asserting epistemic claims is less important than the device producing causality through agency. This might also give the impression that only at the simplest level does risk act as a rhetorical device, whereas at more complex levels it is not rhetoric anymore. My argument is that all these dimensions of “AIDS risk” are imbricated, reciprocally reinforcing and (re)producing each other. In textual practices, they can never be regarded as distinct from one another.

One might think that “AIDS risk” is nothing but another piece of fiction or a fantasy, something that exists at best in the flat world of texts; given the death toll from AIDS, this view seems curious at the very least. My argument is that “AIDS risk” is a rhetorical (and therefore social) practice, and as such it is neither a product of authors’ imaginations nor an ideological instrument; it is something very real, and it has consequences, but its order of reality is not constituted according to a clear-cut distinction between soft and hard worlds that never mingle. It is the rhetorical practice of “AIDS risk” that constitutes the system of knowledge we have about the syndrome, its etiological agent, and its modes of transmission. It is this practice that generates the concrete, lived definitions of risk subject, the self, her means of protection, and her relationships to other subjects and to animate and inanimate objects. But in constituting the system of medical knowledge, this practice seems to withdraw to a marginal position, appearing as something derived from hard-won scientific concepts. Showing how it unfolds therefore means showing the moments through which it both co-constitutes the system of medical knowledge and withdraws to its present position.

In this case one could ask: where is this practice to be retrieved from? Is it to be recovered from the history of medical concepts about the Acquired Immunodeficiency Syndrome, from gradual progress in this area? Or is it to be recovered from the passage from a “primitive”
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to an “enlightened” stage in the mid-1980s, as historians of medicine argue? Showing how this textual practice unfolds presupposes examining the genealogy (Foucault 1966, 1989) of scientific knowledge about AIDS, with which it is coextensive. This examination is based not on the assumption of something being produced by external forces, but on that of collocation of producer and products. At the same time, it presupposes examining the body of scientific knowledge by questioning its claims of unity and homogeneity, and its inconsistencies, contradictions, and fragmentation. It presupposes inquiring how the rhetorical practices of “risk” are reproduced in various discourses which run parallel or in opposite directions, intersect each other, or stand in mutual contradiction. In other words, this practice reproduces itself not as the simple repetition of the same statements; rather, it unfolds in a variety of discourses, simultaneously performing different movements.

The expression “medical AIDS discourse” presupposes that there is a unitary body of medical knowledge about the syndrome, knowledge that has evolved from the simpler hypotheses (or the astonishment) of the beginning, along more or less straight paths, up to today’s sophisticated standpoint. This is indeed the position adopted by historians of AIDS, as well as by many social scientists (Treichler 1988a,b, 1992; Patton 1990; Seidel 1992): as more became known about the retrovirus and its means of transmission, the risks of different categories became better known, so that we can tell today what puts a woman at risk, or a heterosexual man, a man from Kinshasa, or an infant in Milan. There is indeed little doubt that considerable progress has been made in medical research on AIDS in the past twenty years. But take the risk category of women: a closer look reveals that in the 1980s there were several discourses on “female sexual partners,” “spouses,” “mothers,” “prostitutes,” and “African women,” which ran in parallel and sustained different, indeed conflicting epistemic claims about the retrovirus and the ways it was transmitted. Each of these discourses actually contradicts the others: a spouse having sexual intercourse only with her husband (and getting the retrovirus from him) cannot transmit it further. Hence, transmission from female to male is not possible. Or, if she can transmit it, this happens through “household contacts,” in which case the nature of the retrovirus must be revised. Prostitutes are “reservoirs” of the retrovirus: males get it through direct contact with the semen of other males, deposited in the vagina. Therefore, transmission