Male Patient Perceptions of HIV Stigma in Health Care Contexts

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ABSTRACT

Individuals living with HIV may have a heightened sensitivity to the behaviors of others that may signal bias or discrimination. Identifying and avoiding these potentially problematic behaviors may be especially important for service providers, such as health care personnel, who regularly interact with HIV-positive clientele. This study examines the experiences of 50 male American military veterans living with HIV and their perceptions of HIV stigma within health care contexts. Participants described a variety of behaviors performed by health care personnel that they perceived to be indicative of HIV stigma, ranging from ambiguous nonverbal cues (e.g., minimal eye contact) to blatant discrimination (e.g., physical abuse of HIV-positive patients). These findings extend previous research on HIV stigma in health care settings by (1) focusing on health care personnel’s actual behaviors rather than their attitudes and beliefs about HIV-positive patients, (2) including patients’ perceptions regarding the behaviors of both clinical and nonclinical health care personnel, and (3) identifying behaviors patients perceive as stigmatizing that are unique to health care contexts. Combined, these findings provide health care personnel a tangible list of behaviors that should either be avoided or further explained to HIV-positive patients, as they may be interpreted as stigmatizing.

INTRODUCTION

RECENT ESTIMATES indicate that well over a million people are currently living with HIV in the United States. 1 Although the introduction of antiretroviral treatments has dramatically increased the life span of those infected by the virus, 2, 3 progress in addressing the widespread social stigma surrounding HIV has lagged behind these biomedical advances. 4, 5 Despite public health efforts to educate the general public about HIV, a host of studies indicate that misconceptions of how the virus is transmitted and disparaging attitudes toward those infected with the virus are still prevalent. 5, 6 Subsequently, people living with HIV often face ridicule, ostracism, and persecution in many forms, ranging from caustic social interactions to draconian legislative efforts designed to strip people living with HIV of their civil rights. 7–13 In its extreme form, hostile expressions of HIV stigma have even driven people from their homes and led to acts of physical violence against them. 8, 14–16 The effects of HIV stigma on those living with the disease are significant and include adverse mental and physical health outcomes, such as elevated stress, depression, immune

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suppression, and suicide.\textsuperscript{17–23} In addition, concern over being stigmatized can lead people to conceal their HIV status from others with whom they engage in risky behaviors, such as sexual activity or injection drug use.\textsuperscript{14,24–26} Similarly, to avoid discovery and potential stigmatization, people have even been found to forego their HIV medications in the presence of others, the consequences of which can be both viral resistance and clinical failure.\textsuperscript{27–29} Given its formidable range of social, psychological, and physical consequences, it is of no surprise that stigma as been labeled the most significant social and psychological challenge of the HIV experience.\textsuperscript{30–32}

One of the most troubling aspects of HIV stigma involves its prevalence and expression among health care personnel, who include all clinically and nonclinically employed individuals working in health care settings. Health care personnel have been instrumental in managing the HIV epidemic in the United States and abroad, serving both to treat people living with the virus and to educate the general public about HIV. However, health care staff are not immune to HIV stigma and some have, in fact, also been shown to openly report disparaging attitudes toward people living with the disease.\textsuperscript{33–35} Such attitudes have been documented among a variety of health care personnel, including practicing physicians, medical students, nurses, and psychologists.\textsuperscript{36–38} Although such negative attitudes are shown to diminish as health care personnel gain experience working with HIV-positive patients, they may still persist even among those who have worked with such clientele for many years.\textsuperscript{39,40} Such attitudes have been linked to reluctance in health care personnel to interact with those living with the disease, in some cases leading to their refusal to provide care for HIV-positive patients.\textsuperscript{41–44}

Beyond such blatant incidents as refusing to treat HIV-positive clientele, stigma may also be expressed by health care personnel through more subtle communication. Whether they are conscious of it or not, people frequently express their biased attitudes during social interaction, often through nonverbal gestures.\textsuperscript{45} In the context of HIV, relatively little research has examined expressions of stigma at such micro-social levels; however, the existing literature suggests that people living with HIV are attuned to subtle behaviors from others that alone may be explained away, but taken together cause people to feel stigmatized.\textsuperscript{46}

Perhaps because of their experiences with bias and discrimination, or simply because of their mindfulness that they may be devalued by others, people who belong to stigmatized groups are shown to develop a heightened sensitivity to behaviors that may indicate the presence of prejudice or discrimination.\textsuperscript{47} Some research has revealed this type of sensitivity among people living with HIV and has shown that it may be especially acute when dealing with individuals on whom people must depend, such as health care personnel.\textsuperscript{46} Stigmatized people are more prone to interpret ambiguous and subtle behaviors as indicative of social stigma rather than the result of other equally plausible explanations.\textsuperscript{48–50} Subsequently, patients’ heightened sensitivity to stigma may lead them to interpret ambiguous actions of health care personnel as stigmatizing, even when it is unclear that stigma is, in fact, being expressed. Regardless if these perceptions and interpretations of such behaviors are accurate, the impact on patients is no less significant. This suggests that even well-intentioned health care personnel may inadvertently exhibit behaviors that HIV-positive patients interpret negatively.\textsuperscript{51}

Despite the obvious implications HIV stigma has for quality of care, surprisingly little research has been conducted in this area. The literature on HIV stigma includes only a handful of studies on HIV stigma in health care settings, with existing studies being limited in scope (e.g., focusing solely on one type of health care personnel, such as physicians or nurses) and emphasizing the antecedents of stigma (e.g., attitudes toward people living with HIV), rather than its behavioral manifestations.\textsuperscript{33,34,52,53} Exploration of patient’s lived experiences and perceptions of HIV stigma in health care contexts is the next step in this line of inquiry. Such exploration is not only of great academic interest and pragmatic value, but may hold considerable legal consequence, as well. Identifying and avoiding such problematic behaviors may help health care personnel more quickly
develop open and trusting relationships with HIV-positive patients. Avoidance of such behaviors may subsequently minimize the likelihood of malpractice lawsuits, because patients who like their care providers are less likely to sue than are those who experience troubled patient–provider relationships. Above all, avoidance of such behaviors will better fulfill the ethical obligations of health care personnel and improve clinical outcomes of their patients.

Exploration of patient’s experiences and perceptions of stigmatizing behaviors among health care personnel includes identification of the more blatant and extreme expressions of HIV stigma among health care personnel, which to date have not been documented beyond occasional refusals to provide care for HIV-positive clientele. So, too, must it include the identification of the more subtle behaviors to which HIV-positive patients are acutely attuned. Assessment of patients’ experiences encountering such behaviors must also include all types of health care personnel, not just clinical care providers. With these objectives in mind, the present study was conducted to explore patients’ reported perceptions of and experiences with HIV stigma in health care contexts.

MATERIALS AND METHODS

As part of a larger study on coping with HIV stigma, American military veterans living with HIV were interviewed regarding their experiences with stigma and discrimination when interacting with health care personnel. A convenience sample of 50 participants was recruited through the infectious disease units at three Veterans Administration (VA) hospitals in a large city in the Midwestern United States. Participant recruitment entailed passive solicitation via handouts distributed onsite by infectious disease staff. Fliers were placed in common areas of the infectious disease clinic areas and waiting rooms at the various facilities (staff at these centers routinely provide information about such opportunities to their patients and actively requested the right to do so). Participants interested in taking part in the study contacted the primary investigator, who then scheduled them for either a focus group or one-on-one interview, depending on participants’ preferences and availability. Participants received $25 in remuneration for their involvement in either the focus group or the one-on-one interviews.

This study utilized a grounded theory approach during data collection, which involved reformulating and refining research questions as the study progressed to pursue promising lines of inquiry. This exploratory and hypotheses-generating process consisted of two stages. Stage 1 involved two semistructured focus group interviews (n = 3 and 5, respectively) regarding participants’ experiences with stigma and discrimination by health care providers; stage 2 involved one-on-one interviews (n = 42) on the same topics. This two-tiered process was utilized to produce a data set that combined the synergy of focus groups with the depth of responses afforded by one-on-one interviews.

In stage 1, the lead author led two focus groups through a semistructured interview. The interview included questions regarding participants’ experiences with the forms and effects of stigma and discrimination resulting from their HIV-positive status, including stigmatizing experiences with health care personnel (a total of 18 stigma-related questions were asked during the focus groups, 3 of which dealt directly with participants’ perceptions of stigma among health care personnel). Probe and follow-up questions were included, when appropriate, to clarify issues and validate the interviewer’s interpretations of responses. These focus groups lasted 120 minutes each. Based upon the findings from these focus groups, a more refined set of questions was developed to explore the problems participants experienced with health care personnel. These questions were then used during stage 2 of the study, which involved the primary investigator conducting in-depth, one-on-one interviews with participants regarding their stigma experiences. These one-on-one interviews lasted between 45 and 230 minutes (median = 70). Digital recordings of the focus groups and interviews were transcribed verbatim and distributed to each member of the research team for coding.

Both the focus group and one-on-one interview transcripts were analyzed using latent content and constant comparative techniques. Coding included two stages of anal-
ysis, with the first stage following the focus group interviews and the second stage follow-
ing the one-on-one interviews. In stage 1 of analysis the primary investigator first reviewed
the focus group data to identify focal themes among the participants’ responses. This pro-
cess helped identify and refine interview top-
ics that required clarification or further explo-
ration during the one-on-one interviews. Stage
2 of the analysis occurred after the one-on-one
interviews were transcribed. The entire re-
search team independently coded both the fo-
cus group and one-on-one interviews for focal
themes, after which they convened to compare
and compile their findings. Through consen-
sus, the team constructed a detailed categori-
cal system for describing the forms of prob-
lematic encounters participants experienced
with health care personnel. Through this pro-
cess, the research team codified an inventory of
the ways in which participants experienced be-
haviors as stigmatizing and discriminatory
when interacting with care providers and other
personnel in health care contexts.

RESULTS

A total of 50 participants took part in either
the focus groups or one-on-one interviews,
ranging in age from 24 to 70, with a mean age
of 50 (standard deviation [SD] = 8.9). Twenty-
six (52%) participants identified as African
American, 17 (34%) as Caucasian, 4 (8%) as
Latino, 1 (2%) as Native American, 1 (2%) as
other, and 1 (2%) who chose not to self-iden-
tify by race. Thirty-four (68%) identified as het-
erosexual and 16 (32%) as gay or bisexual. Dur-
ing the time of this study, no female veterans
were receiving HIV care through the VA cen-
ters at which the study was advertised; subse-
quently, all study participants were male. Par-
ticipants possessed varying degrees of formal
education, with 6 (12%) having not completed
high school, 14 (28%) having completed high
school, 20 (40%) having some college or tech-
nical/trade schooling, 6 (12%) having a bache-
lors degree, and 4 (8%) having either master’s
or doctoral degrees. Participants also reported
a variety of occupational employments, with 12
(24%) fully employed, 6 (12%) unemployed, 26
(52%) on disability, 5 (10%) retired, and 1 (2%)
a full-time student. Time since diagnosis var-
ied from one to 23 years, with a mean time of
12.8 years (SD = 6 years). Twenty-eight (56%)
of the participants had previously received an
AIDS clinical diagnosis, although many had re-
sponded well to antiretroviral therapies and
currently maintained CD4 counts well above
200.

Although several participants reported only
positive experiences with health care pro-
viders, in which HIV stigma was noticeably ab-
sent, others reported encountering suspicious
or explicitly stigmatizing behaviors in a vari-
ety of health care settings and across a broad
range of health care personnel. Any behavior
patients interpreted as conveying discomfort,
ignorance, fear, contempt, or exclusion were
coded as problematic and potentially stigma-
tizing. These behaviors are organized and pre-
sented below, which involve issues of de-
meanor, the provision of care, or combinations
thereof. During events in which only stigma-
tizing demeanor was exhibited by health care
personnel, treatment and other services were
still perceived by patients to be fully and ade-
quately provided. In contrast, stigmatizing
events involving the provision of care involved
denial or disruption of appropriate services for
patients. In many instances, descriptions of
problematic care provision also entailed stig-
mating demeanor on the part of health care
personnel.

Demeanor

Participants identified a variety of encour-
ters in which nonverbal cues from health care
personnel sent ambiguous or explicitly prob-
lematic signals to patients. In particular, par-
ticipants reported eye contact, proxemics (dis-
tance maintained between conversational
partners), and paralanguage (vocal tone and in-
flection) as important indicators for interpret-
ing others’ feelings toward patients with HIV.
The convergence of these nonverbals commu-
nicated what participants identified as various
forms of negative affect, including irritation or
anger, nervousness, or fear at having to work
with HIV-positive clientele.

Eye contact. Participants described being
mindful of the amount of eye contact provided

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them during health care interactions. In particular, a lack of eye contact was interpreted as possible discomfort with or dislike of HIV-positive patients. Evan describes such behavior during an encounter with a neurologist, who, he explained, “never looked me in the eye. I still don’t know what his face looks like, which is not a crime, [but it] sends unfriendly signals. Or at least, cold signals.” Although participants could not quantify how much eye contact was appropriate and sufficient to alleviate their perceptions of stigma, those who described such events felt confident the eye contact given by the health care personnel in question fell far below conventional expectations.

Paralanguage. Participants also described a heightened sensitivity to the vocal tones with which health care personnel spoke to patients. Clipped, flat, or brusque tones, in particular, where identified as indicating potential unease, dislike, or disdain of the HIV-positive patient. For instance, Kobe described how the tone of voice with which a nurse delivered his AIDS diagnosis left him feeling alienated and dismissed. He explained,

She made me feel low. It was just so callous and cold the way she said it. I think my viral load or CD4, one of ’em went below 200 and she said, “You have AIDS.” And I said, “What?” The way she said, “Whenever you go below 200 you got AIDS.” It was just the coldness in it—there wasn’t no feeling. It’s almost like a stone-faced warden or something. No concern, you know? I was like, “Wow.” I felt awful.

Proxemics. Similarly, participants also described using the distance care providers placed between themselves and patients, or proxemics, as an indicator of providers’ comfort with HIV clientele. Standing or positioning one’s self close to the patient, such as within arm’s reach, was considered a reassuring sign that the provider was unafraid of a person with HIV. Conversely, participants interpreted standing further away while interacting with a patient as an indication of fear or dislike on the part of the care provider. Participants reported encounters in which health care personnel maximized the space between themselves and their patients by doing such things as addressing them from across a room rather than approaching their bedsides. Gabriel described one such encounter in which, “The doctor was young, foreign. Maybe he was a med student. He went to the other side of the other patient’s bed and talked loudly to me about having AIDS.” Similar encounters involved care providers conversing with patients from hallways rather than coming into close proximity, such as an experience described by Max, in which, “[the doctor] wouldn’t even come into the room.”

Irritation/anger. Participants also identified specific affective states among health care personnel that HIV-positive patients may be prone to attribute to HIV stigma. For instance, when health care personnel expressed anger or irritation with or around HIV positive patients, the participants noted that they questioned if these negative emotions were due to the health care personnel having to deal directly (and unwillingly) with HIV-positive clientele. For instance, Tyler described having dealt with clerical staff who spoke to him in ways that he described as “standoffish.” He further explained, “you get that feeling, that standoffish feeling, from the support services—from the clerks,” which he attributed to his HIV-positive status. These problematic demeanors sometimes involved health care personnel expressing more than mere irritation, including what patients interpreted as full-fledged anger. Jacob’s experience with an ambulance driver provides one such example, in which the participant attributed the health care provider’s negative emotions to her having to work with an HIV-positive patient,

One time, my brother was washing dishes in the house. He had this big ceramic bowl and he dropped it. It bounced back and cut his wrists in a perfect football shape—took the skin right off. He was bleeding profusely. He called 911 and the ambulance came, but he forgot to tell them right away that he had HIV. When he remembered to tell them, this ambulance driver was so pissed! It just blew me away. She took her gloves off, which were extremely bloody, and just threw them out onto the street. Left them there. And she wasn’t exactly nice to him after that.

Nervousness/fear. Anger was not the only affective response participants described encountering amongst health care personnel that
was attributed to HIV stigma. Participants noted such things as awkward glances, fidgeting, and agitated speech on the part of health care personnel as signs of nervousness around patients with HIV. Nervousness, though possibly attributable to a variety of factors, was credited to unreasonable fear of contagion when working with HIV-positive patients. For instance, Jerome described sensing a “nervous vibe” when seeking care from a dentist. He explains,

The dentist that was actually going to work on me, I felt like, the vibe that I got from him, the energy that I got from him, or at least, the demeanor that I got from him, was that he really didn’t want to work on me or he wasn’t comfortable working on me. And that was real disconcerting because I thought, “Where am I going to go to be able to get this done?”

Panic. Participants described inferring ignorance on the part of health care personnel when such events occurred. In particular, patients suspected that irrational fear of contagion and/or incomplete understanding of HIV’s transmission routes were to blame for such behaviors. Sometimes this translated into care providers panicking when faced with the task of working with an HIV-positive patient. For instance, Roosevelt recounts an appointment with his dentist, who was informed of his patient’s HIV status, but whose assistant was not. The assistant’s reaction upon learning this information left a lasting impression with Roosevelt, who explains,

As I’m sitting there and we’re talking, and [the assistant] comes running over. I could tell it was a big emergency. He says, “wait a minute!” and grabs the manila folder. Because he’s writing it so large, I can tell what he’s writing across the front of this manila folder on the side that I can’t see, and he’s writing the word “AIDS!”

Demeanor shifts. Such nonverbal and affective cues were particularly salient to patients when they were preceded by warmer, friendlier interactions. Participants explained that such noticeable shifts in demeanor often occurred after the patients’ HIV status was revealed. Ahmad described one such example of inconsistent nonverbal behavior, stating, “I went to this office and the lady asked what my disability was. When I told her, it seemed like her whole attitude changed.” Carlos explained his perception of being judged during such encounters and his resulting frustration, stating,

It usually starts out with the front desk and the support staff, who are not educated. I mean, they should not care what I have, whether it’s cancer, AIDS, or whatever. They should just be concerned that I get the best care I possibly can, you know? They have no right to make a judgment on me!

In several accounts of health care personnel’s anger when dealing with HIV-positive patients, the patients failed at the onset of the encounter to inform the provider of their HIV status. However, patients during these encounters were often experiencing some form of duress, such as having been injured or suffering from another acute illness. Under these circumstances, Xavier explained how disclosing one’s HIV status may not be on the patient’s mind,

When you’re in pain, that’s not the first thing you’re thinking about! That’s not the first thing. When they picked me up, I was in pain and that was not the first thing I thought to tell them. It came to me a little bit later in the ambulance. But when you’re in intense pain, HIV is not the first thing to come out of your mouth. They tell you it should be, but I was too focused on telling them where it hurt.

Excessive precautions. Perhaps as a result of these experiences of fearfulness among health care personnel, some participants described a heightened awareness of the safety precautions clinicians used when working with HIV-positive patients. In particular, they described being critical of what appeared to be excessive precautions or use of protective gear. For instance, Carlos described his trepidation over the protective gear his surgeons wore when they came to talk with him well in advance of his actual surgery. He recalled wondering the following questions,

Are the doctors dressed the way they’re dressed because they’re afraid they’re going to get infected? They looked like they were ready for war. I mean, they had shields on—it was like looking at Roman gladiators! They were completely hermetically sealed.
Jerome further explained the confusion HIV-positive patients feel when faced with such elaborate safety measures, by asking aloud, “Is this typical? Do they do this for everyone?” Unsure as to what constitutes standard procedure and what constitutes paranoid overprecaution, in the absence of explanation from care providers the patients were inclined to infer the latter.

**Differential precautions.** Another perceived problem regarding the use of protective gear involved clinicians using one set of precautions when working with some patients, then switching to other sets when working with HIV-positive patients. Although participants who witnessed such events applauded appropriate safety measures, they also felt the application of universal precautions when working with all types of patients would avoid the conspicuous altering of procedures that provokes suspicion and resentment from HIV-positive patients. Isaiah recounted one such situation when having his labs drawn by a phlebotomist,

> Even today, there was a girl down there who took my blood. There was a guy in there before me whose blood she took—she didn’t put on gloves. I never would have thought about that before, except I had somebody else there who had AIDS, who was saying, “Oh that’s discrimination, because then they should do it [use consistent precautions] with everybody.”

**Labeling.** Another set of problematic encounters some participants linked to fear and ignorance involved how health care personnel labeled HIV-positive patients. Although disconcerting to patients, some labeling experiences described by participants may have been part of a standard procedure. Terrell described one such situation,

> I was in the emergency room and they were taking some blood and I said, “You know, you need to put some gloves on, ‘cause I’m HIV positive.” The thing was, after that I was admitted to a room and I had signs on my door. ‘Biological fluid,’ ‘ah, ‘caution,’ you know? It was terrible. It made me feel really dirty.

Other labeling events, however, appeared more derogatory in nature and designed to belittle HIV-positive patients. Some participants described health care personnel who insisted on labeling patients according to a particular high-risk behavior group, even when the patient is adamant that he doesn’t belong to one. Malik described dealing with one such nurse practitioner, stating,

> She came in and she’s like, “well, how’d you get it?” I said, “I don’t know.” She asked me about being in the risk categories and I said, “well, I wasn’t.” And then she said, “no, you’re just in denial. You had to have been. You don’t ‘just get it.’ ” I’m like, “well, I . . .” I didn’t know. I was just thinking, going by those categories, I didn’t fit in any of them. It’s like, “no, I’m not in any of them.” And she’s like, “well, yes you are.” And I’m like, “no, I’m not.” And it became real big confrontation.

**Scaring patients.** Another set of stigma-related behaviors patients attributed to ignorance or, potentially, cruelty on the part of health care personnel involved scaring patients with issues of mortality. Participants reported a variety of encounters with assorted clinicians in which patients were told their lives were over or that they were lost causes because they were HIV-positive. Noah, for example, recounted how one physician responded in such a way immediately after his diagnosis. He stated, “I was like, ‘okay, what’s the next step?’ And the immunologist just went, ‘you’re gonna die.’ ” DeShawn described a similar encounter with a physician who sought him out one evening in his hospital room to seemingly taunt him for his diagnosis. He explained,

> This little, goofy, bearded guy, says, “You know you gonna die.” And he was a doctor. He said, “Do you know what kinda AIDS you got?” I said, “Yeah. I got the AIDS that you can’t cure.” He said, “Well, you know, you don’t got long to live.”

**Mocking patients.** In addition to health care personnel scaring patients, participants also described being the targets of outright contempt and ridicule. Participants described these various forms of mockery and contempt as explicit expressions of HIV stigma, which were performed by a host of health care personnel ranging from desk receptionists to attending physicians. Evan provided one such example, in which he explained,

> I had a negative thing from a dentist. He was gonna pull a tooth and he gave this sort of smug, superi-
Blaming patients. Coupled with these contemptuous episodes, participants described a host of encounters in which health care personnel openly blamed patients for their HIV. Participants described feeling shamed during these episodes, even if they had contracted HIV simply while "living their lives." Cedric described one such encounter with a phlebotomist as follows,

I had one blood draw—she had a hard time drawing my blood. She got really angry and agitated about it. She kept poking me and I told her it hurt, "It’s hurting," you know? And she said, "if you hadn’t of done this to yourself, we wouldn’t have to be going through this!"

In summary, participants reported being sensitive to health care personnel’s demeanor, which patients used to determine other’s comfort and attitude toward HIV-positive patients. Nonverbal cues such as proxemics, vocal tones, and eye contact, in particular, were used to infer irritation, anger, nervousness, or discomfort among health care personnel when working with HIV-positive clientele. Participants were especially mindful of shifts in health care personnel’s attitudes after their HIV-status was revealed. Similarly, labeling patients and inconsistent or excessive use of safety precautions when working with HIV-positive clientele were each interpreted as indicative of HIV stigma. Although participants reported feeling resentful and unsatisfied after such encounters with health care personnel, they noted that such behaviors did not necessarily impede the delivery of sufficient and effective health care. In other words, they often felt their health care was not compromised, despite the misgivings and stigmatizing demeanors of health care personnel during these encounters. Unfortunately, stigmatizing behaviors exhibited by health care personnel did not always have such benign results, as demonstrated in the following section.

Care provision

Dealing with fearful and/or demeaning behavior from health care personnel were unpleasant and upsetting experiences for those who endured them; however, participants also described encountering a variety of stigmatizing behaviors that far exceeded funny looks or verbal ridicule and included actions that impaired or even prevented patients’ utilization of health care. These encounters ranged from health care personnel ignoring patients or denying services, up to and including the physical abuse of HIV-positive patients. Unlike facets of demeanor, which participants admitted were sometimes ambiguous and potentially attributable to issues other than HIV stigma, events in which health care provision was impaired left little doubt in the minds of those affected that such experiences were rooted in bias and discrimination.

Ignoring patients. One behavior patients felt was rooted in HIV stigma and impaired their health care involved health care personnel ignoring HIV patients, even when patients were in great need. Such events left patients not only in distress, but also angry and resentful of the care they received. Gabriel described one such encounter, in which a physician refused to respond to him, even when being directly addressed.

I was here for a week and there was a doctor who was attending. He came in my room—he didn’t say a word to me. When I caught his attention, it was as if I hadn’t said anything! Yeah, I felt that was derogatory about my [HIV] status. I recall that because it actually prompted me to make a formal complaint. I went to the patient advocate and explained what had happened because he was the doctor, he was outside my door, I needed something—I was trying to call his attention and he didn’t respond!

Substandard care. Patients also recounted instances in which they received other forms of substandard care, ranging from health care personnel spending inadequate time on a patient’s needs to leaving patients in pain to wait upon the care providers’ convenience. Jack, who reported a number of such events when seeking dental care, described one such circumstance in which he had consistently been provided inadequate care, despite the fact that he was paying full price for such services. He explained,
I tell [the hygienist] I’m having problems with one tooth causing pain. The hygienist says, “ok” and 2 months later when I come back I said, “you know, the pain—it’s still there and, if anything, a little worse.” She calls the dentist in and the dentist says, “well, you’re so hard to anesthetize that we don’t, we can’t do a thorough cleaning on you. And,” he says, “in your condition . . .” You know? And I just want to turn and slug him! I’m paying you your regular prices every three months to make sure I don’t get cavities and you’re telling me you aren’t going to anesthetize me because I’m too hard and it really doesn’t matter because of my condition?

Not only did Jack experience these events, he saw others deal with similar treatment. As a result, had had come to view HIV stigma and poor quality of care as essentially indistinguishable. He explained,

It’s gotten to the point that the last time I was at the clinic, which I was in this week, they had a patient they wanted to pull a tooth for and couldn’t get the tooth all the way out. Instead of immediately calling Big University Hospital and getting him in, they told the patient to go home and to come back at 9:00 AM [the next day] as an emergency patient when the clinic opened. So there is a patient with a tooth half-pulled, in pain, and this is an HIV clinic treating an HIV patient. So is there still a stigma? I consider that a stigma. I try to differentiate stigma and the quality of the medical care that is being given to [us], but I can’t differentiate the two anymore. They are so thoroughly entwined that everyone I know kind of expects that, as an HIV person, we’re going to get lower quality health care in all areas, but in dental it’s extreme.

Denied care. At times, participants reported being denied even substandard care because of their HIV-positive status. For instance, some reported encounters in which health care personnel, apparently uncomfortable with HIV-positive clientele, attempted to redirect such patients to alternative sites of care rather than provide any services themselves. For instance, Ahmad explained that after learning he had come in search of HIV services, the care provider with whom he was speaking tried to send him away to clinics in other parts of the city. In particular, he explained, “What she did was start telling me, steering me toward other places to go and ask for the same help that they give, you know?”

Some provision of care or redirection to other helpful services is, at least, better than being fully rejected outright and denied services altogether. For some participants, such as Jack, this occurred due to the inability to find providers willing to take on HIV-positive clients. He explained, “There were years when even my private doctor didn’t know of a doctor in the entire city that would treat an [individual with] HIV.” Similar accounts were provided by participants, who described both clinical and non-clinical health care personnel refusing to provide services to patients living with HIV. In clinical contexts, for instance, Eric mused, “I’ve had a couple of incidents where a nurse or a phelmbologist didn’t want to draw blood because it was clear to her that she was drawing for a CD4 count.” Similarly, he explained,

One time I went to a county hospital to have a tooth pulled. I guess the dentist was a student or an intern or whatever. He refused to do it once he realized I was HIV-positive. They had to send for another doctor.

Clinicians, however, were not the only health care personnel described as refusing to provide services to HIV-positive clientele. Such behaviors also were portrayed by a variety of non-clinicians, such as receptionists, administrative staff, and custodial staff. Evan described one such case, explaining,

I was an inpatient at Big University Hospital and I don’t know what they put on the door of the room, but a woman who would normally come in and mop up and sweep, wouldn’t. It was really wild, with her shrieking, “I ain’t goin’ in there!”

Abusive treatment. Perhaps the most blatant and potentially dangerous experiences with stigma reported by participants involved some form of abuse at the hands of health care personnel. Certain abusive events involved malignment, in which health care personnel strategically defamed HIV-positive patients to other health care personnel, potentially ensuring these patients received less favorable treatment from other health care personnel. One such instance was described by David, who explained, I was treated like garbage. I don’t know the lady’s name, I could only go and look for her and see her and say, “That’s her,” but I couldn’t say her name because I was so drugged out. She treated me like crap. She called the police on me and said I was abusive. I couldn’t even move, but she said I was
abusive. And she called the floor that I was gonna be admitted to and said that I was abusive, that they’d better watch themselves because I’m being abusive and she doesn’t want to get next to me and get scratched, because she may get “the same thing.” She didn’t think I was listening and she didn’t think I understood what she what she meant by that, but I did.

Finally, patients reported acts of physical abuse perpetrated against them by various health care personnel, attributing each of these events to their HIV status. This included patients being pummeled or put into situations in which they were likely to be hurt. David, who had endured malignation by a nurse in a previous encounter, also described being physically mistreated by paramedics. In his own words, he explained,

Being rushed to the hospital a few times I’ve dealt with paramedics. I’m epileptic, I have seizures. When the paramedics do pick me up, they man-handle me. In other words, they treat me bad . . . Once they ask whoever’s called the ambulance that knows my history, who tells ‘em I’m HIV-positive, they start thrashing me around. The only part they’re careful of is putting in a line in my vein because they don’t want to get sick. I mean, the paramedics are cruel and I prefer not to be handled by them at all. Just let me die. If I’m gonna die, let me die, but don’t, don’t call the ambulance. That’s how bad they are.

In summary, participants described a number of stigmatizing experiences at the hands of health care personnel that compromised the quality and effectiveness of the health care patients received. This included ignoring patient needs, providing insufficient and substandard health care to HIV-positive patients, denying and refusing to provide health care services to these patients, and abusing HIV-positive patients, which ranged from malignation to physical abuse. Participants explained that stigmatizing events that impaired the provision of health care could also entail problematic demeanor, such as mocking patients while providing them insufficient services.

**DISCUSSION**

In this study, participants reported being both mindful of health care personnel’s behaviors and sensitive to anything that may indicate bias or stigma toward HIV-positive patients. Participants reported encountering what they perceived to be expressions of HIV stigma in a variety of health care environments and performed by a broad range of health care personnel. Such encounters took place in ambulances, doctor’s offices, dental care facilities, inpatient hospital rooms, and the common areas of hospitals, such as hallways and reception desks. Problematic behaviors identified as stigmatizing by participants included such things as awkward or nervous nonverbal behaviors, excessive safety precautions, avoidance, refusal to provide care, as well as anger toward and even abuse of HIV-positive patients. These findings extend previous research on HIV stigma in health care settings by (1) focusing on health care personnel’s actual behaviors rather than their attitudes and beliefs about HIV-positive patients, (2) including patients’ perceptions regarding the behaviors of both clinical and non-clinical health care personnel, and (3) identifying behaviors patients perceive as stigmatizing that are unique to health care contexts.

These results complement and expand upon earlier findings that reveal some health care personnel are reluctant to interact with HIV-positive patients.34,46 In some instances, participants reported how this translated into health care personnel’s refusal to provide services to HIV-positive patients. This apparent aversion to interacting with individuals who have HIV is especially troubling in light of evidence that being touched and being checked upon signals high quality of care to patients.57,58 In fact, egalitarian treatment of patients, regardless of HIV status, has been linked to patients’ perceptions of higher quality of care.53 Because individuals with HIV wish to be touched, cared for, and treated like any other patients, health care personnel’s apprehensiveness could affect patients’ confidence in and evaluations of a care provider’s competence.

Beyond fearful and aversive behaviors, participants also described feeling disdain, contempt, and even anger directed toward them from various health care personnel. Sensing dislike from care providers has special significance for patients’ perceptions of the quality of care they receive, as well as their likeliness to trust or return to their care providers.59–61
However, patients reported experiencing far more than covert affect, which included events in which health care personnel openly mocked or blamed them for their HIV status, unfairly labeled them or maligned them to other health care personnel, and even physically abused HIV-positive patients. Although violence directed toward people living with HIV has been reported in earlier HIV research, it has almost exclusively been discussed in the context of romantic relationships.17,22 These findings reveal a frightening dilemma, in which seeking health care may, in some cases, actually place HIV-positive patients’ health and safety at risk. Evidence of events in which health care personnel intentionally jeopardize the well-being of their patients should concern all members of the allied health professions and the appointed stewards of the justice system.

Although such things as nervous nonverbal cues or even physical violence toward people living with HIV may also occur in other social contexts, these behaviors have special significance coming from those upon whom one must depend for health care. Additionally, some of these behaviors are unique to health care contexts, such as care providers using differential or excessive protective gear when working with HIV-positive clientele. Although it may be tempting to dismiss such things as problems of the past, claiming that advances in education on HIV have eradicated these issues, it is noteworthy that participants described such events occurring within weeks or even days of their interviews, making these findings both timely and relevant.

Clinicians such as physicians and nurses should now understand the true risks involved in working with HIV-positive patients and not succumb to irrational fears of contagion; however, participants recounted problematic events occurring with both sets of clinicians within the year preceding the interviews. Although other care providers, such as laboratory technicians, emergency medical technicians, and dentists may not have received sufficient training and education regarding HIV earlier in the epidemic, this should no longer be the case. Yet, participants reported recent problems with these care providers, as well.

Although it may be surprising to find such stigmatizing behaviors among clinicians, perhaps it should not be that participants reported such encounters with non-clinical personnel, such as custodians and administrative staff. These individuals receive little (if any) training on the biomedical aspects of HIV and may rely on knowledge they have acquired about the disease during their daily lives. Given the rampant misinformation and stereotyping of HIV still prevalent in the lay population, encounters with nonclinical personnel who are reluctant to interact with or even clean the rooms of HIV patients may be a more common problem than is currently realized. These findings suggest that additional outreach and continuing education is required among health care providers and other personnel not only regarding HIV, but also regarding the impact of their behaviors on patients. Behaviors identified in this study as problematic from patients’ perspectives should be avoided or, when a necessary component of care, further explained by health care personnel, as they may be interpreted as discriminatory or stigmatizing by patients with HIV. These dialogues can help foster the image that health care personnel are knowledgeable about the disease and want to provide patients with the best care possible. Failing to do so may not only impair provider-patient relations, but could conceivably even lead patients to avoid health care in the future. Without modifying these behaviors, the likelihood of legal action taken against care providers and the facilities in which they are employed may increase, given the correlation between law suits and patients who are dissatisfied with or take offense to their care providers.54 This is especially important given how HIV-positive patients may sometimes identify innocent or unintentional actions as stigmatizing, given their hypersensitivity to social stigma.46,47 Such sensitivity may be particularly acute following physical changes endemic to prolonged use of antiretroviral therapies, which combat the virus but mar the body (e.g., lipodystrophy).62,63

This study raises important questions about both the perception and prevalence of stigmatizing behaviors in health care contexts in general, but in certain contexts, in particular. For instance, HIV stigma in the practice of dentistry has received little attention in the research literature, but was the source of considerable frustration and anxiety for a variety of participants in this study. Understanding of this prob-
lem can be enhanced by research that explores dentists’ perceptions of HIV, its etiology, and transmission. Also, additional research must focus on these problems among other health care personnel that have, to date, received little to no attention in the HIV stigma literature. Such research could also couple attitudes or behavioral intentions of health care personnel with their actual performance, rather than just concentrating solely on the antecedents to stigmatizing behaviors or the behaviors, themselves. Such research may even explore distinctions between provider perceptions of encounters and the perceptions of their patients, in order to identify discrepancies in what providers and patients consider appropriate or stigmatizing.

Although identification of problematic behaviors is an important step in disarming the heightened sensitivity and anxiety patients may have about being stigmatized, knowing what behaviors send positive, reassuring signals to HIV-positive patients is of equal significance. Identification and performance of these more positive behaviors has the potential to further enhance care providers’ work and inspire patients’ confidence in the care they receive. Therefore, future research should explore forms of proactive behaviors health care personnel can employ to generate this amity. Finally, this study raises questions regarding the role of stigma in the health care provision of other illnesses. All illness is stigmatized, though the reasons for and degree of stigmatization differ depending upon illness context. As such, research efforts should focus on better identifying the forms and functions of stigmatizing behaviors among health care personnel attending to people living with other highly stigmatized illnesses.

In evaluating our findings, several limitations to our study should be acknowledged. To begin, the methods used in this study are excellent for identifying the forms and functions of stigma experienced by HIV-positive patients in health care settings, but are unable to explain the frequency of these experiences or the magnitude of their effects on patient outcomes, such as health care satisfaction and likelihood of future health care utilization. In addition, this study was based on interviews with a sample of only 50 participants, all of whom were male and currently seeking treatment through VA hospitals. The fact that the sample was entirely male is particularly important, as men may be subjected to some forms of stigma that differ from those experienced by women living with HIV. The fact the participants were all veterans, most of whom are older and have dealt with HIV/AIDS for many years, also has implications for our results. Younger people with HIV or nonveterans may encounter different experiences with stigma or access different health care facilities than those described by the current study participants. Of note, however, is the fact that participants’ health care histories did include a wide variety of health care facilities outside of the VA, which strengthens the significance and applicability of these findings for nongovernmental health care organizations. In order to address these limitations and to provide a more accurate assessment of the forms, frequencies, and effects of patient perceived stigma among health care personnel, larger studies are currently being conducted utilizing structured interviews with significant numbers of participants spread across multiple states.

CONCLUSION

Participants described a variety of behaviors performed by health care personnel that they perceived to be indicative of HIV stigma, ranging from ambiguous nonverbal cues, such as minimal eye contact, to blatant discrimination, such as physical abuse of HIV-positive patients. These findings offer health care personnel a tangible list of behaviors that should be either avoided or further explained to HIV-positive patients, as they may be interpreted as stigmatizing. Also, this study reveals that patients are sensitive to such behaviors being performed by a variety of health care personnel, indicating the need for all such personnel to be mindful of their actions toward these patients. Further research is needed to identify the prevalence of such behaviors and their effects on HIV-positive patients.

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