Appropriateness of Disclosure of HIV Testing Information: The Perspective of PLWAs

Kathryn Greene and Julianne M. Serovich

ABSTRACT This paper explores perceptions of the appropriateness of disclosure of HIV testing information from the perspective of people living with HIV/AIDS (PLWAs). A systems framework, communication boundary management, is employed as a theoretical approach. Seventy-seven HIV-positive individuals responded to a questionnaire asking if they thought particular individuals (also labeled targets or recipients) should have access to results of HIV tests. Results indicated PLWAs created three grouping of recipients based on perceptions of who should have access to this information. As expected, participants reported most desire for access to results of HIV testing information to the immediate family subsystem (e.g., spouses, fathers), next most to the extended family (e.g., aunts, grandfathers), and least access to non-family (e.g., employers, teachers); actual disclosure patterns were similar. HIV-positive individuals also reported there should be less overall disclosure of HIV testing information than other sample groups. Implications and directions for future research are discussed.

The incidence of AIDS/HIV infection, accompanied by a rise in HIV testing, has increased dramatically in the past ten years. People who test HIV-positive, even those tested for HIV, must deal with issues surrounding disclosure of this sensitive information. As the ranks of those infected swell, there is growing controversy regarding the rights to privacy of HIV-positive individuals on the one hand and the perceived public good (disclosure) of this sensitive information on the other (see: Bayer & Toomey, 1992; Krupka & Vener, 1988; Lang, 1991; Swartz, 1990). As Silin (1987) noted, “because AIDS has been constructed as a disease of unknowns affecting the other, public debate has been most easily framed in terms of individual rights versus the public good” (p. 13). Most Americans see the control of HIV/AIDS as requiring some loss of individual privacy and possible restrictions on civil rights (Blendon & Donelan, 1988).

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An earlier version of this paper was presented at the meeting of the Speech Communication Association, Miami, 1993. Authors would like to thank Jerold L. Hale (University of Georgia) for his helpful comments on an earlier draft of this manuscript. Completion of this project was made possible by travel funding from the Deans Office, College of Arts and Sciences at East Carolina University.
Despite some literature on the perceived conflict of public and private interests regarding HIV/AIDS, there has been a marked absence of research on the consequences of disclosure of HIV test results for those most directly affected, those who have tested HIV-positive as well as people living with AIDS (PLWAs). This article investigates disclosure in the context of HIV in two ways: first, it explores PLWAs perceptions of who should have access to HIV testing information; second, it explores to whom PLWAs disclose their HIV infection.

**REVIEW OF LITERATURE**

Little is known about why, to whom, and with what effects people choose to disclose their serostatus. Even with this absence of understanding, policies regulating HIV/AIDS and disclosure continue to be made. For example, third party notification programs are in place in some states, but they may create ethical dilemmas for those involved (see: Bayer & Toomey, 1992; Kermani & Weiss, 1989; Melton, 1988; Swartz, 1990). Identification of people with HIV antibodies has been rationalized as a way to protect uninfected persons from possible transmission of the virus (Silin, 1987).

There is an available model to assist in understanding how HIV-positive individuals might want to regulate information about their serostatus. Using a systems approach, Petronio (1991) proposed a theory of boundary management to describe how people control potential risk to self when disclosing. Petronio described how individuals regulate disclosure of private information in relationships to attain a balance between disclosure and privacy, between intimacy and autonomy. To achieve this balance, individuals erect metaphoric boundaries to protect themselves and to reduce their chance of losing face. The risks associated with disclosing HIV infection are tremendous, perhaps leading to more rigid boundaries (little or no disclosure) where PLWAs might perceive any sharing of information about their disease status as a threat.

One of the strengths of Petronio’s model is that it is transactional, focusing in part on why people make decisions to disclose. People use rules or criteria to control boundaries and regulate the flow of private information (Petronio, 1991), yet much work remains to establish how these rules function. In the context of HIV, the most useful information to consider is the factors that affect discloser decisions. Petronio argues there are three discloser factors: expectations of the discloser, message strategy, and message content. It is the expectations of the discloser that are likely to vary by relational type and intimacy, characteristics associated with perceptions of the potential disclosee. In this sense, disclosure is strategic.

Yep (1993) applied boundary management theory specifically to the potential vulnerability of disclosure of HIV infection. Yep elaborated on what factors might affect decisions to disclose, especially the balance between a need to receive social support yet avoid the emotional distress and stigma associated with HIV. Although Yep’s application of boundary management theory to the context of HIV is intriguing, it has not been fully tested to this point. Greene and Serovich (1995) used boundary management theory to explore disclosure of HIV, and they reported relational quality and anticipated response were the best predictors (in a non HIV-positive sample) of willingness to disclose information about HIV. The present investigation assessed the utility of boundary management theory in the context of disclosure of HIV infection by focusing on target or role of discloser.
Research on Privacy and HIV Testing

Disclosure is a complex process, and there is a need to adapt what is known about disclosure to the specific context of HIV/AIDS. Research on the process of disclosure also may prove beneficial in understanding how perceptions of privacy function in the context of HIV testing.

Greene, Parrott and Serovich (1993) compared college students' and parents' perspectives of privacy and appropriateness of disclosure of HIV testing information. Not surprisingly, they found people reported more overall support for access to results of HIV testing information than access to information that someone has been tested for HIV. Overall, findings indicated that attitudes toward disclosure of HIV testing information can be predicted by individual difference variables (e.g., perceptions of privacy, religiosity, and attitude toward homosexuality). In general, people with more accepting attitudes toward homosexuality and more liberal religious beliefs report that others should have less access to information about results of HIV tests.

In addition to research on predictors of individual perceptions of disclosure of information about HIV, research has examined if who the recipient is affects perceptions of appropriateness of disclosure. Researchers have established that the target or recipient of disclosure constrains how individuals reveal information about themselves (Stokes, Fuehrer, & Childs, 1980; Tardy, Hosman, & Bradac, 1981). Individuals delineate clear boundaries among (a) marital, community, and general public groups, and among (b) marital, nuclear, and extended family subsystems in perceptions of appropriateness of disclosure of HIV testing information (Serovich & Greene, 1993; Serovich, Greene, & Parrott, 1992). Specifically, those closer or more intimate are perceived as more appropriate for access to information about HIV, and those with less intimate relationships are perceived as less appropriate for access to this information. This process can be described as an attempt by individuals to create informational boundaries, a process consistent with that described by Petronio's boundary management theory. Some people are simply not viewed as appropriate for having access to information about results of HIV tests. As a result, a semi-permeable boundary may be necessary; such a boundary allows adequate input in the form of education, medical treatment, and social support—input which may be provided without negative consequences such as loss of employment or housing. It is necessary to engage in some self-disclosure to create a social support network. Thus, there may be a need for both privacy and disclosure for adequate functioning.

Using this boundary approach, Serovich et al. (1992) looked at the potential recipient of HIV testing information. It was expected that people viewed as most directly affected by the diagnosis (e.g., spouse, lover) would be seen as more appropriate recipients than those less directly affected (e.g., teachers, employers). Serovich et al. manipulated the target/recipient of the information, for example "Employers should have access to results of employees AIDS tests," using eleven recipients in Likert-type format. They reported a three-factor solution accounting for 73% of the variance. The "community" factor included employer, potential employer, co-worker, and teacher targets. The "general public" factor included newspaper, general public, and community leader targets. The "marital" factor included two targets, lovers and spouses. Two items in these analyses loaded on two factors, classmates and family members. Overall, individuals reported most
desire to restrict access to HIV testing information to the general public, less to the community, and least to the marital subsystem.

Serovich and Greene (1993) expanded on this work by looking at potential recipients or targets for access to HIV testing information within families. They reported a three-factor solution accounting for 85% of the variance. The “extended family” factor included the aunt, uncle, cousin, and mother-, father-, sister- and brother-in-law targets. The “nuclear family” factor included mother, father, son, and daughter targets. The “marital family” factor included spouse, former spouse and lover targets. Four targets loaded on two factors (nuclear and extended): sisters, brothers, grandmothers, and grandfathers. Overall, participants reported most support for others’ access to HIV testing information to the marital system, moderate to the nuclear family, and least support for access to the extended family.

Taken together, the findings of these studies support the notion that people create informational subsystems in making decisions about who should have access to results of HIV tests. What has not been thoroughly examined is how HIV-positive people perceive access to HIV testing information. It is not known if those most directly affected by disclosure of this information (PLWAs) create boundaries or subsystems in similar ways. Given that previous researchers found these informational boundaries with other samples, it would be expected that PLWAs would create subsystems in similar ways.

**Hypothesis 1:** HIV-positive individuals create boundaries based on others’ access to results of HIV tests; specifically, those closer to the discloser (marital and nuclear family) are viewed as more appropriate recipients of HIV test results than those not as intimate (extended family, general public, and community).

**Privacy and HIV Positive Individuals**

Although there has been little research on disclosure and privacy in the context of HIV testing, there has been even less work specifically with HIV-positive individuals. Some research on attitudes of HIV-positive individuals has been published, but some of these studies are more qualitative in nature, using, for example, methodologies of interviews (e.g., Lang, 1991) or case studies (e.g., Carlson, Greeman, & McClellan, 1989; Fleck, 1991). These studies have provided valuable preliminary information that should be further explored.

There are tremendous risks for HIV-positive individuals in disclosing their serostatus. Yep (1993) hypothesized that the stigma associated with HIV was similar to what Goffman (1963) described as “spoiled identity.” People with HIV/AIDS are more stigmatized than people with other diseases, such as toxic shock or legionnaires’ disease (Hughes, 1985), cancer and coronary disease (Walkey, Taylor, & Green, 1990), or leukemia (St. Lawrence, Husfeldt, Kelly, Hood, & Smith, 1990). The disclosure process for HIV-positive persons may be similar to what has been identified as the risk of disclosing information perceived as negative, such as homosexuality, being raped, or being sexually abused. People with direct experience with any of these sensitive issues may have experienced negative reactions from disclosure of this information. However, disclosure is necessary to receive support and to acquire needed information and assistance. Pennebaker (1989) argues that disclosure of traumatic events is associated with lower overall
stress and increased psychological health. Thus, HIV-positive persons must make choices to try to disclose for the benefits yet avoid the negative consequences.

HIV-positive individuals may have more desire to restrict others' access to results of HIV tests because they fear potential negative consequences. For example, they may fear discrimination, such as loss of employment, eviction, losing child custody, and cancellation of insurance (Anderson, 1989). Documentation of this type of discrimination has been difficult, but there is widespread anecdotal evidence. PLWAs also may fear physical abuse, "bashing," particularly PLWAs identified as homosexuals (Anderson, 1989).

HIV-positive individuals also may have a high desire to restrict others' access to results of HIV tests because they are aware of perceptions of HIV/AIDS as a gay disease. According to public perception, many HIV infections are acquired through association with stigmatized activities (e.g., homosexual or bisexual sexual contact and IV drug use). HIV/AIDS, however, has been associated primarily with the homosexual/bisexual population (Robinson, Walters, & Skeen, 1988). Hence, negative attitudes toward HIV/AIDS have been clearly linked to negative attitudes toward homosexuality (Cohen & Grace, 1988; Kelly, St. Lawrence, Smith, Hood, & Cook, 1987; Larsen, Serra, & Long, 1990; Ross, 1988; Stipp & Kerr, 1989). Marks et al. (1992) reported that virtually no one revealed HIV infection to parents who did not already know their gay/bisexual orientation. People with negative attitudes toward homosexuality also are more likely to support increased access to information about HIV tests (Greene et al., 1993), and increased access might be a potential threat to HIV-positive persons.

Finally, HIV-positive individuals may have more desire to restrict access to results of HIV tests because they are aware of generally negative attitudes toward PLWAs. Researchers have documented general negative attitudes toward people with HIV/AIDS and anxiety over potential contagion by medical personnel (Pleck, O'Donnell, O'Donnell, & Snarey, 1988; Sherr, 1987; Wallack, 1989). Researchers have reported that men generally have more negative attitudes toward HIV/AIDS and PLWAs than women (e.g., Eiser, Eiser, & Lang, 1989; Shrum, Turner, & Bruce, 1989; Walkey et al., 1990). Negative attitudes toward PLWAs, however, may be decreasing (Johnson, 1989), although people may not be developing positive or even neutral attitudes toward PLWAs, rather they may be becoming less negative.

HIV-positive individuals may fear three results from others' access to HIV testing information: behavioral consequences, perceptions of HIV/AIDS as a gay disease, and general negative attitudes toward PLWAs. Each of these may explain PLWAs' expected desire for less access to results of HIV tests.

Hypothesis 2: HIV-positive individuals report others should have less access to HIV testing information than other sample groups for marital, nuclear, and extended family, general public, and community targets.

Disclosure of HIV Infection

Previous research reviewed one part of the disclosure and HIV area—perceptions of who should know about HIV test results—yet little work has examined who actually is told about HIV infection. Attitudes toward disclosure provide significant information, and additional information about boundaries might be gained from examination of actual disclosure practices. Marks et al. (1992) examined disclosure
patterns in a sample of HIV-positive Hispanic men and proposed that people with HIV evaluate subjectively the potential consequences of informing a particular target person before a disclosure is made, a process that may be similar to the boundary formation in attitudes described earlier. They did find HIV-positive individuals' expectations about disclosing vary by relational type and level of intimacy. In this study, HIV-positive persons were more likely to disclose their HIV status to more significant others than to less significant others. The lowest rate of disclosure was to employers, landlords, and religious leaders, a higher rate of disclosure was to mothers, siblings, and friends, and the highest rate was to lovers and spouses. In interviews with five HIV-positive women, Kimberly, Serovich and Greene (1995) reported similar high disclosure to partners, parents, siblings, and children (depending on age). It seems clear from these findings that the creation of informational boundaries in disclosure practices may be quite similar to patterns observed in attitudes toward who should know about the results of HIV tests.

**Hypothesis 3:** HIV-positive individuals disclose their serostatus most to those closer to them (i.e., marital subsystem and nuclear family) and least to those further from them (i.e., the extended family).

**METHOD**

**Participants and Procedure**

Participants in this study consisted of 77 individuals who had been diagnosed HIV-positive and resided in the southwestern portion of the United States. Participants were recruited through two non-profit organizations aimed at servicing the informational, educational, counseling, and referral needs of PLWAs. Participants were recruited in two ways. Some received a questionnaire with a stamped return envelope (response rate was 34%), whereas others were asked to fill out the questionnaire at the center where they received services (no in-person refusals). The questionnaire took approximately 30 minutes to complete and was anonymous. Participants had known their HIV-positive status, on average, 48.13 months (range 0 to 132 months). The sample was 78% male and ranged in age from 23 to 50 (M = 33.96; SD = 7.05). In terms of religious preference, 27 reported “none,” 9 Catholic, 22 Protestant, and 13 “other” (6 not reported). The sample included 52 Caucasians, 5 African-Americans, 10 Hispanics, and 4 Asian (6 not reported). Fifty-four percent reported being currently unemployed, 24% reported full-time employment, and 22% reported losing their job due to their HIV-positive status.

**Measurement Instruments**

One construct of interest in this study, perceived appropriateness of others’ access to HIV testing information, was measured by a series of Likert-type items. The 26 items consisted of statements such as, “Lovers should have access to information about results of AIDS tests.” The items were changed by manipulating the target/potential recipient, and the targets included the following (proposed subsystem in parentheses): spouses, former spouses, lovers, (marital subsystem); sons, daughters, mothers, fathers (nuclear subsystem); aunts, uncles, cousins, mothers-in-law, fathers-in-law, sisters-in-law, and brothers-in-law (extended family); employers, potential employers, co-workers, teachers (community subsystem);
community leaders, general public, and newspaper (general public subsystem); classmates, family members, sisters, brothers, grandmothers, grandfathers (previous split loadings). Responses to these items ranged from "Strongly Agree" to "Strongly Disagree," with a higher score indicating less perceived appropriateness of others' access to information about HIV testing or more support for privacy.

Actual disclosure of HIV infection also was measured. Participants were asked, "does x know you are HIV-positive?" A number of targets/recipient used in previous research were excluded due to the length of the questionnaire, and the ten targets retained in this study included the following: significant other, mother, father, first, second and third sibling, maternal grandmother and grandfather, and paternal grandmother and grandfather. Because of the length of the survey, the disclosure questions were limited to the marital and nuclear family targets. It was also necessary to ask if each target person was relevant to the participant (e.g., only children would not have siblings, or a grandparent might be deceased).

**Reliability.** Previous research indicated the attitude toward disclosure items formed reliable measures by subsystem. Serovich et. al (1993) reported, for a sample of college students and parents, low-to-acceptable reliabilities (alpha range .61–.85). Serovich and Greene (1993) reported, for a sample of married and dating couples, adequate reliabilities (alpha range .75–.98). The only questionable subsystem measure in previous research was the marital subsystem, but the three-item measure (Serovich & Greene, 1993) showed marked improvement over the two-item measure (Serovich et al., 1993). The reliabilities (Cronbach’s alpha) by subsystem for the present study were adequate, ranging from .87 to .98 (see Table 1).

There are several limitations to this study worth noting. First, the sample size is relatively small and over-represents males, potentially affecting the generalizability of findings. The moderate response rate might also be a concern, however, given that some of the target sample were in advanced stages of an AIDS-related disease process, it would be understandable if they did not return the questionnaire. Thus,

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<td>Sample of:</td>
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<tr>
<td>General Public (3 items)</td>
<td>alpha = .84</td>
<td>95% CI (4.01–4.29)</td>
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<tr>
<td></td>
<td>M = 4.14; SD = .95</td>
<td></td>
<td></td>
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<tr>
<td>Community (4 items)</td>
<td>alpha = .85</td>
<td>95% CI (3.09–3.45)</td>
<td></td>
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<tr>
<td></td>
<td>M = 3.27; SD = .95</td>
<td></td>
<td></td>
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<tr>
<td>Extended Family (7 items)</td>
<td></td>
<td>alpha = .98</td>
<td></td>
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<tr>
<td></td>
<td>M = 3.43; SD = 1.05</td>
<td>95% CI (3.28–3.58)</td>
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<tr>
<td>Nuclear Family (4 items)</td>
<td></td>
<td>alpha = .95</td>
<td></td>
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<tr>
<td></td>
<td>M = 2.32; SD = 1.05</td>
<td>95% CI (2.28–2.48)</td>
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<tr>
<td>Marital (2–3 items)</td>
<td>alpha = .61</td>
<td>alpha = .75</td>
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<td></td>
<td>M = 1.66; SD = .75</td>
<td>M = 1.63; SD = .73</td>
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<td></td>
<td>95% CI (1.54–1.78)</td>
<td>95% CI (1.48–1.73)</td>
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these data provide some useful insight, but results must be interpreted with these
limitations in mind.

RESULTS

Analyses

Data were analyzed using MANOVAs, factor analyses (varimax rotation), and
99% confidence intervals. The level of significance was set at $p < .05$ for most tests,
with an alpha adjustment for the number of tests in Hypothesis 2.

Hypothesis 1

Hypothesis 1 predicted HIV-positive individuals create boundaries among sub-
systems (general public, community, extended, nuclear and marital) based on
desire to allow others’ access to information about results of HIV testing. This
hypothesis was partially supported. The means by subsystem were all in the
expected order, but not all subsystems were significantly different (see Table 1).
MANOVAs also indicated the presence of three (not five) significantly different
subsystems for this sample of HIV-positive individuals. A MANOVA indicated a
significant within-subject effect for target or recipient ($F (73) = 52.3$; Hotell-
ings = 2.92; $p < .001$); that is, there were perceived differences in appropriateness
of disclosure by target.²

An initial factor analysis of all 21 targets (those cleanly loading in previous
research) also indicated a three-factor solution, consistent with results of the
MANOVA, accounting for 79% variance. The first factor (eigenvalue = 11.53; %
Var. = 54.9) loaded seven extended family items above .8 (e.g., in law, cousin). The
second factor (eigenvalue = 3.67; % Var. = 17.5) loaded seven nonfamily items
above .6 (e.g., coworker, community leader). The third factor (eigenvalue = 1.47; %
Var. = 7.0) loaded three nuclear family items above .5 (e.g., spouse, lovers). Four
items (son, daughter, mother and father) loaded on both factors 1 and 3, but
additional evidence indicated they were better explained as part of the immediate
family subsystem (factor 3).

The previous marital and nuclear family subsystems were not significantly
different (by factor analyses, confidence interval, or MANOVA), and these items
were combined to form a subsystem labeled “immediate family.” The “extended
family” was significantly different from all other systems, consistent with previous
research. The final subsystem, labeled “non-family,” was a combination of previous
general public and community subsystems. As expected and found in previous
research, participants reported most desire for others’ access to results of HIV
testing information to the immediate family subsystem, next most to the extended
family, and least access to non-family members.

Hypothesis 2

Hypothesis 2 predicted HIV-positive individuals, as compared to the general
population, report less desire for access to HIV testing information (or more privacy)
across all subsystems. This hypothesis was generally supported, as most confidence
intervals did not overlap. Table 1 presents the means for the present study along
with results from two comparable previous studies using different samples. Across
the subsystems examined, HIV-positive individuals perceived others should have significantly less access to results of HIV testing information than did members of the general population. There was only one subsystem (extended family) where HIV-positive individuals did not desire more privacy than other groups (confidence intervals were not significantly different). For each of the other four subsystems, HIV-positive individuals reported significantly less desire for access to results of HIV testing information.

Hypothesis 3

Hypothesis 3 predicted more HIV-positive persons report disclosure of their serostatus to the marital and nuclear family and fewer report disclosure to extended family members. This hypothesis was supported. Table 2 presents the frequencies and percentages for disclosure of HIV infection by target. It was not possible to make direct comparisons with all subsystems because of what data were collected (focus on actual disclosure to family members). According to these data, there is most disclosure of HIV infection to the immediate family (marital and nuclear subsystems previously) and less to the extended family. Percentages of who had disclosed to the immediate family were extremely high (73–85%), with disclosure to mothers (85%) slightly higher than to fathers (73%). Disclosure to siblings was moderate (range 40–55%), and disclosure of infection to grandparents was even lower still (range 29–44%).

The general pattern of these disclosure percentages reveals three groupings similar to the kind of pattern found in attitudes toward disclosure. The highest disclosure was to partners and parents (range 73–85%), next highest disclosure to siblings (range 40–55%), with lowest disclosure to grandparents (range 29–44%). In order to test if these differences in groupings were significant, z-tests for proportions were performed (Hayslett, 1968). If the maternal grandmother was ignored as an outlier, the z-tests between the closest targets in adjacent subsystems approached the critical value of 1.645. Specifically, fathers were significantly different from first siblings (z = 1.67), but second siblings were not significantly different from paternal grandmothers (z = .64).

Some items in the present study were designed to measure perceptions of others’ access to results of HIV tests, consistent with the theoretical approach. It must be

<table>
<thead>
<tr>
<th>Target</th>
<th>Knows*</th>
<th>Does Not Know</th>
<th>Unsure</th>
<th>Not Relevant</th>
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<tr>
<td>Sig. Other</td>
<td>36/84%</td>
<td>5</td>
<td>2</td>
<td>34</td>
</tr>
<tr>
<td>Mother</td>
<td>52/85%</td>
<td>8</td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td>Father</td>
<td>38/73%</td>
<td>13</td>
<td>2</td>
<td>24</td>
</tr>
<tr>
<td>Sibling 1</td>
<td>28/54%</td>
<td>24</td>
<td>0</td>
<td>25</td>
</tr>
<tr>
<td>Sibling 2</td>
<td>19/40%</td>
<td>29</td>
<td>0</td>
<td>29</td>
</tr>
<tr>
<td>Sibling 3</td>
<td>16/55%</td>
<td>13</td>
<td>0</td>
<td>48</td>
</tr>
<tr>
<td>Paternal grandfather</td>
<td>4/31%</td>
<td>8</td>
<td>1</td>
<td>64</td>
</tr>
<tr>
<td>Paternal grandmother</td>
<td>7/33%</td>
<td>12</td>
<td>2</td>
<td>56</td>
</tr>
<tr>
<td>Maternal grandfather</td>
<td>2/29%</td>
<td>3</td>
<td>2</td>
<td>70</td>
</tr>
<tr>
<td>Maternal grandmother</td>
<td>8/44%</td>
<td>6</td>
<td>4</td>
<td>59</td>
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*Percentages are calculated after subtracting those who did not have target person/not relevant.
noted, however, that this does not assess actual disclosure of HIV infection. In order to examine this relation, correlations were run between actual disclosure (measured as number of target persons disclosed to) and the five subsystems (see Table 3). Correlations between the five subsystems and actual disclosure indicated the relations varied in both direction and strength. Correlations between actual disclosure and marital and nuclear family subsystems were inverse; that is, people who had disclosed their HIV infection to more people were less likely to support access to HIV tests for the marital or nuclear family. It may be that disclosure to intimates and immediate family are still viewed as private choices, not sharing that should be automatic. The nonrelation between actual disclosure and extended family is not surprising given that the extended family was the most ambiguous target (neutral) in previous research; thus, we still know little about the extended family. Correlations between actual disclosure and general public and community subsystems were direct; that is, the more a person had disclosed his/her HIV infection the more likely she or he was to support others’ access to HIV test results. This relation is understandable when considering the extremes in disclosure: a person who had disclosed to few if any others would be unlikely to support access for non-intimates, but a person who had disclosed to many might have confronted fears of negative consequences and might not see others’ access as so threatening. In sum, actual disclosure and perceptions of others’ access to HIV tests can inform each other.

**DISCUSSION**

The purpose of this study was to investigate PLWA’s perceptions of appropriateness of others’ access to information about HIV and disclosure of their serostatus. Also of interest was how these perceptions might be different from perceptions of people who were not HIV-positive. Results of this study suggest that differences do exist, with PLWAs being more conservative or restrictive concerning others’ access to HIV testing information. This is consistent with findings that HIV-positive persons are highly selective in choosing targets of disclosure (Marks et al., 1992).

In this study, PLWAs constructed informational boundaries concerning who should have access to results of HIV testing information. PLWAs grouped targets into three distinct subsystems, with immediate family members (spouses, former spouses, lovers, sons, daughters, mothers, fathers) being viewed as more appropriate recipients of HIV test results than extended family members (aunts, uncles, cousins, mothers-, fathers-, sisters-, and brothers-in-law), who, in turn, were
viewed as more appropriate recipients than non-family members (employers, potential employers, co-workers, teachers, community leaders, general public, and newspaper). These results are consistent with the types of boundaries reported by previous researchers (Marks et al., 1992; Serovich & Greene, 1993; Serovich et al., 1992). There are several possible reasons for these particular groupings or creation of informational boundaries.

First, it is plausible that PLWAs need the support of immediate family members beyond the marital subsystem. In previous studies, participants reported a distinction between the marital and nuclear family subsystems. A re-examination of these data revealed only 39% of respondents in this study reported having a significant other or partner. If a PLWA is not in a significant non-family relationship, he or she might be more likely to see other targets, such as parents and/or siblings, as equally appropriate or necessary targets for disclosure. PLWAs may recognize a need for social support and seek more sources for assistance. Kimberly et al. (1995) provide anecdotal evidence for this need for social support, for example, in child care or reading information about HIV. In order to receive this assistance, PLWAs may need to disclose their HIV-positive status to marital and nuclear family members (immediate family).

It is equally plausible that PLWAs feel uncomfortable portioning out immediate family members to tell, preferring that if one knows, they all know. This might serve several functions. First, it eliminates re-telling family members over time and might save emotional energy needed to deal with this type of disclosure. Second, telling multiple family members provides a mechanism for support for the family members themselves. Thus, PLWAs might see it as a necessity for others to know at the same time. This might be especially important for persons who fear they might have to be the ones providing support for the PLWA. In this instance, not disclosing might be a form of self-preservation. PLWAs also might want to avoid putting family members in awkward positions by having information other family members do not, for example telling a mother but not a father. HIV-positive persons (especially those nonsymptomatic) have described not wanting to worry others or cause stress as additional reasons for not disclosing (Kimberly et al., 1995; Marks et al., 1992).

One possible explanation for PLWAs being more conservative in attitudes toward disclosure (Hypothesis 2) is that they may base their decision on who should have access to HIV testing information on their own need to tell (which includes the goal of disclosure to gain support). Non-PLWAs, however, might base their decision on who should have access to HIV testing information on the other person’s need to know. Thus, non-PLWAs may perceive sexual partners as significantly different from other immediate family members because of their potential risk for infection. PLWAs may not make this distinction, or may make it in a different way. The framing of “should have access” items focuses on who has a right to know, not if a person intends to disclose. This also may fail to take into account indirect disclosure, through a third person, inappropriate access to results, or by observation of health conditions (all of which have become foundations for discrimination suits). It is also possible that more conservative attitudes toward disclosure could merely be a reflection of perceived risk, and PLWAs may clearly be in a better position than non-PLWAs to assess risks of disclosing HIV infection.

What is potentially disturbing about these results is that PLWAs are more conservative about disclosure to the marital subsystem, as seen in the results of Hypothesis 2. PLWAs do not perceive spouses, former spouses, or lovers as
appropriate recipients of disclosure in the same way as other sample groups. This is
disturbing because it would be hoped that disclosure of HIV infection to intimate
others would happen automatically. In fact, a major goal of partner notification
programs is to reduce the spread of HIV. PLWAs may have difficulty developing
and/or maintaining primary relationships; specifically, they may fear their partner
would leave if their HIV status were disclosed. Thus, PLWAs may put off disclosure
until the relationship has developed further. Studies have generally reported
disclosure by HIV-positive persons to sexual partners to be around 50% or less (e.g.,
Marks, Richardson, & Maldonado, 1991), and Marks et al. (1991) reported disclo-
sure of HIV infection to sexual partners decreased in direct proportion to number of
partners. In the present study, 84%, a much higher percentage, of those PLWAs who
reported having significant others had disclosed their serostatus. It is also possible
that figures from previous studies were not adjusted for whether the respondent had
a significant other (the unadjusted figure for this study is 47%, much like previous
reports).

For the purposes of this study, it would be erroneous to conclude that intimate
others are not told or to speculate about the time frame in which PLWAs might
decide their HIV-positive status to the marital subsystem or others. What these
findings might better represent is a recognition of the unpleasantness associated
with this kind of disclosure. This is especially true considering PLWAs consistently
saw other subsystems as less appropriate targets than other sample groups (cf.
Serovich et al., 1992; Serovich & Greene, 1993). Thus, PLWA’s hesitancy to disclose
HIV-positive test results is not isolated with the marital subsystem but permeates all
subsystems.

The patterns for actual disclosure of serostatus (Hypothesis 3) were similar to
patterns in attitudes toward who should have access to this information in this
study. This provides some evidence that attitudes, behavioral intentions, and
behavior are closely related in this context. Most PLWAs in this study had disclosed
their infection to their significant other, mother, and father (if they had these
persons in their lives). Interestingly, disclosure to mothers (85%) was higher than to
fathers (73%), consistent with differences reported by Marks et al. (1992), though
percentages in this study were much higher. This finding is consistent with reports
that women receive more disclosure overall than men (Jourard, 1961; Jourard &
Lasakow, 1958; Jourard & Richman, 1963), and that people disclose more to
mothers than fathers (Daluiso, 1972; Komarovsky, 1974; Riverbark, 1971). Gender
differences in disclosure, however, are dependent on the dimension of disclosure
studied, such as amount versus intimacy (cf. Dindia & Allen, 1992).

Roughly half of the participants in this study had disclosed their infection to
siblings (if they had siblings). This is slightly higher than the 30–34% disclosure to
siblings reported by Marks et al. (1992), but this also may be a function of the
passage of time between the studies. Roughly one third of these participants had
disclosed their infection to grandparents, but extended family members have not
been included in previous studies addressing actual disclosure. These targets form
only rough approximations of targets or recipients in families (e.g., they do not fit
well for people with non traditional family forms or “families of choice” dominated
by extensive friendship networks); however, the same kinds of informational
boundaries described by other researchers in disclosure and attitudes toward
disclosure of HIV infection are clear (Marks et al., 1992; Serovich & Greene, 1993;
Serovich et al., 1992).
Implications

One widely publicized health recommendation has been for HIV-positive persons to disclose their serostatus, especially to those who might be or have been at risk for infection; this, however, is a more complex process than previously recognized. Results of this study provide useful information for people designing messages to persuade HIV-positive persons to disclose their serostatus and reduce their risk behavior. First, results confirm that PLWAs have more restrictive attitudes than other sample groups; that is, they are more protective of privacy in this context. Message designers may choose to focus not only on the marital subsystem but the nuclear family to encourage disclosure. In this study, there were no differences in attitudes toward disclosure to the immediate family (between the marital and nuclear systems). It may be useful to target parent(s) or children as sources of social support because these decisions are not generally tied to questions of the recipient’s risk for infection.

Currently, health messages to promote disclosure of HIV infection are infrequent (with the exception of those at HIV testing sites); these messages simply recommend that HIV-positive persons disclose to those who might be at risk for infection. These current messages are simplistic and fail to recognize the complexity of this process. Messages, perhaps utilizing a PLWA or the spouse/partner of a PLWA as a source, could use role plays or depict people in the process of disclosing their serostatus. Such messages might be most effective if they emphasize both the positive and negative consequences of disclosure, and this could be done by focusing on the intrapersonal process of deciding to disclose before an actual disclosure scene. Reframing messages from a duty to disclose (public health) frame to what can be gained by disclosing (social support) would be another avenue for message designers.

Results of this study also have implications for professionals in contact with PLWAs, for example, therapists and social workers. Clearly, PLWAs have difficult decisions about disclosing their serostatus, and they are reluctant to do so in some cases. Therapists may want to encourage discussions about disclosure decisions and the process of obtaining social support. Therapists and social workers must be aware that disclosure to spouse/intimate other should not be assumed. It may be that the extended family holds the most difficult decisions, since PLWAs overall reported that immediate family members generally should know and non-family members generally should not know results of HIV tests; attitudes toward disclosure to extended family members were ambiguous (siblings and children might be especially problematic).

Implications for theory. This study provides information for examination of the utility of boundary management theory. One significant component of boundary management theory concerns the expectations of the discloser. Disclosure implies the anticipation of a response from a partner (the disclosee). In this study, the relationship with the discloser (e.g., mother, sister, grandparent) was a significant determinant of whether PLWAs thought others should have access to information about HIV tests. From the perspective of the discloser, Petronio’s (1991) description of the role of intimacy and relational level in expectations of the discloser is a valid one. Greene and Serovich (1995) reported the best predictors of willingness to disclose HIV infection to various family members were anticipated response and relational quality, both components of boundary management theory. PLWAs do
Indeed create informational boundaries in their perceptions of access to sensitive information and in actual disclosure practices. Additional work to test components of Petronio's theory and Yep's (1993) application would be fruitful.

There are other theories that could be useful for understanding the process of disclosure of HIV. Uncertainty reduction theory (Berger & Calabrese, 1975) is a way to explain how people are motivated to communicate in order to understand. This theory may be a unifying perspective for explaining how communication affects individuals' perceptions of support (Albrecht & Adleman, 1984): disclosure of HIV could simply be a way to reduce anxiety. One fundamental impact of support has been to reduce uncertainty over difficult experiences. Other theories recognize social support as critical to handling life stress, crisis and illness (Albrecht & Adleman, 1984), and people may disclose HIV infection to gain social support. From this perspective, the most effective support may come from those who share the context (cf. Albrecht, 1982), and this would lend credence to the use of support groups for PLWA (or partners, children, etc.) as a place to vent and validate feelings.

Future Research

There are a number of ways in which further research could add to understanding of PLWA's perceptions of others' access to HIV testing information and their disclosure practices. PLWAs (and their families) must be included in future samples as they are most directly affected by others' access to this information. It would be useful to continue to study just who PLWAs do disclose to, and this may be informed by structured interview methods. Other information about PLWA's discussions about the disclosure process itself also would be beneficial. For example, it would be worthwhile to study reasons why certain persons were/were not told, their responses, and who told them. Ideally, research could be undertaken from a developmental perspective to see if disclosure patterns change through the disease process. Looking at the timing of the disclosure (e.g., who was told first and how she or he reacted) might reveal relationships between timing and adjustment to the diagnosis. Kimberly et al. (1995) indicated, for HIV-positive women, there may be an initial adjustment period after the diagnosis when no disclosure occurs. Marks et al. (1992) reported disclosure increased with severity of disease (independent of length of time since testing seropositive), perhaps indicating a need for more instrumental kinds of social support.

Further research should be grounded in coherent theoretical perspectives. Boundary management theory, for instance, suggests message strategy and message content influence disclosure. Other theoretical approaches that focus on relationships, such as social network theory, social penetration theory, or uncertainly reduction theory, assert that relationship with target (rather than simply target role) and frequency of interaction with target influence disclosure. It might also be helpful to compare disclosure of HIV testing information to disclosure of other information (e.g., homosexuality, being raped) to see if the disclosure processes are similar.

END NOTES

1. It is difficult to estimate a precise return rate with these data. The mailing list used was drawn from the client lists of these two organizations. Thirty-four percent of the surveys returned were usable. An additional 16% were returned but not used because they were either incomplete or not filled out
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properly. In addition, an unknown percentage of surveys were sent to persons who were deceased, who were hospitalized, or who were too ill to complete the survey.

2. An additional analysis was conducted to see if age of participant and/or length of time known HIV-positive were significant covariates in attitudes toward disclosure of HIV test results. The MANCOVA indicated significant effects for both age and length of time known positive. Both coefficients were positive, indicating older participants and those who knew their HIV status longer were more likely to view disclosure as appropriate. These variables will continue to be factors that should be considered in research about attitudes toward disclosure.

REFERENCES


