Sexual and reproductive health and human rights of women living with HIV: a global community survey

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Abstract

Objective To determine the sexual and reproductive health priorities of women living with human immunodeficiency virus (HIV) and so allow the values and preferences of such women to be considered in the development of new guidelines.

Methods A core team created a global reference group of 14 women living with HIV and together they developed a global community online survey. The survey, which contained mandatory and optional questions, was based on an appreciative enquiry approach in which the life-cycle experiences of women living with HIV were investigated. The same set of questions was also used in focus group discussions led by the global reference group.

Findings The study covered 945 women (832 in the survey and 113 in the focus groups) aged 15–72 years in 94 countries. Among the respondents to the optional survey questions, 89.0% (427/480) feared or had experienced gender-based violence, 56.7% (177/312) had an unplanned pregnancy, 72.3% (227/314) had received advice on safe conception and 58.8% (489/832) had suffered poor mental health after they had discovered their HIV-positive status.

Conclusion The sexual and reproductive health needs and rights of women living with HIV are complex and require a stronger response from the health sector. The online survey placed the voices of women living with HIV at the start of the development of new global guidelines. Although not possible in some contexts and populations, a similar approach would merit replication in the development of guidelines for many other health considerations.
Introduction
In the development of any of its global guidelines, the World Health Organization (WHO) places importance on the values and preferences of the population or individuals that could be affected by the recommendations made within the guidelines.\(^1\) WHO has guidelines on the care, treatment and support for women living with HIV and their children in resource-constrained settings,\(^2\) but these guidelines were published in 2006 and require updating. As an initial step in the updating process, WHO commissioned a global survey to listen to the voices of women living with HIV and determine these women’s sexual and reproductive health priorities. A main aim of the survey was to ensure that the values and preferences of the women covered by the new guidelines would inform the development of the new guidelines from the very onset of the development process. The methods and key outcomes of the global survey are described and discussed below.

Methods
As no single network is likely to represent the wide diversity of women living with HIV, the investigation was based on a survey tool that (i) had been developed by women living with HIV in collaboration with a range of organizations and (ii) reflected a broad range of expertise, geographical perspectives and key affected populations.\(^3\) A core team, which had been established to coordinate and lead the survey development, created a global reference group of women living with HIV. The 14 women in this reference group represented a wide range of national, regional and global organizations of women living with HIV – including adolescents and elderly women and transgender, lesbian, bisexual and other women who have sex with women – and a wide range of experiences – including perinatal transmission, sexual violence or rape, comorbidities of tuberculosis and/or hepatitis C, current or previous use of drugs and prison or detention.\(^4\)

Pre-survey consultation
In a pre-survey consultation based on a quality of life and dimensions of well-being exercise,\(^5\) each member of the global reference group explored how the material, physical, psychological, sexual and spiritual dimensions of her quality of life – including some deeply personal issues – affected her sexual and reproductive health and vice versa. The aim was to reveal the key sexual and reproductive health priorities in the lives of women living with HIV.
Survey

Subsequently, the members of the global reference group and the core team worked together to develop a global online survey. The overarching human-rights-related priority issues that emerged from the comprehensive pre-survey consultation, formed the basis of the first – mandatory – section of the survey whereas other key priorities formed the basis of the other sections, all of which were optional. The optional sections related to (i) a healthy sex-life, (ii) pregnancy and fertility, (iii) violence against women living with HIV, (iv) mental-health issues, (v) women living with HIV in all of their diversity, (vi) puberty, menstrual issues and menopause, (vii) HIV treatment and side-effects and (viii) financial issues affecting access to services. The whole survey was based on an appreciative enquiry life-course approach that included both quantitative and qualitative aspects of the lives and experiences of women living with HIV. The survey platform was designed not only to enable the surveyed women to tell their stories and reflect on all stages of their lives, but also to establish the values–preferences input for the development of new WHO guidelines.

The content of the survey tool was developed following broad stakeholder outreach, several teleconferences and meetings and a consultation with the Joint United Nations Programme on HIV/AIDS’s dialogue platform of women living with HIV and other United Nations partners. By following existing recommendations for mixed methods research in the health sciences and using Survey Monkey software (Survey Monkey, Palo Alto, United States of America), the tool was adjusted and finalized over a period of three months. The survey tool began with background information on the purpose, aims, content and background of the survey, a list of key definitions and a description of how the survey findings would be used. A woman invited to complete the online survey had to answer “Are you a woman living with HIV?” before she could see and answer any further questions. As well as quantitative, closed-answer questions, all of the survey’s nine sections included spaces in which respondents could elaborate on their answers – e.g. they could give details of their own experience, context and/or perceived barriers to accessing services and/or achieving rights. Most of the closed-answer questions were worded positively and either solution-oriented or human-rights-oriented but the sections exploring violence and mental health included questions about negative experiences. The survey included follow-up questions – either free-text or closed – about positive experiences and/or examples of best practice, sources of support, resilience and solution-oriented, rights-affirming recommendations.
Face-to-face focus group discussions using the same set of questions as the online survey were designed to run concurrently, for participants with no or limited computer access or literacy. It was hoped that these discussions and the online survey would create a safe virtual or physical space for women living with HIV to share their experiences. The survey tool, which was created in English, was translated, by volunteers, first into French, Russian and Spanish and subsequently into Bahasa Indonesian, Chinese and Portuguese. The translators were all women living with HIV or affected by the virus, who perceived the importance of this work. All considered themselves to be activists who were familiar with the language and content of the survey and were sensitive to using rights-based, gender-equitable and culturally sensitive language. Although a professional translator synthesized the French, Portuguese and Spanish responses, Google Translate software (Google, Mountain View, USA) was used to convert the Bahasa Indonesian, Chinese and Russian responses into English.

Focus group discussions in English, French and five further languages (Table 1) were facilitated by 10 women – including six women living with HIV – who were either members of the global reference group or women who already had established relationships with – and were trusted by – the participants. All of these facilitators worked professionally within the HIV, gender and human-rights sectors and had experience in conducting focus group discussions, workshops, training sessions and advocacy meetings. Most were already known to the core team and had been involved in previous related studies. The core team and facilitators adapted and amended an initial discussion guide to match the particular population of women involved in each discussion. The first focus group discussion, which was undertaken in Thailand, served as a pilot. Facilitators identified the key themes of each discussion in a short report. These reports were returned to a coordinator and shared with the core team. Any requests for clarification or more detail were collated and sent back to the facilitators. The findings were then used to supplement and enrich the qualitative data collected through the online survey, with particular reference to women in specific contexts. The online survey data were, however, the primary source used to articulate, shape and structures the main findings listed in the overall report.

Ethics
Ethical considerations were made in line with international guidelines on research on women living with HIV or domestic violence against women. Potential respondents who still
wished to participate in the online survey after reading the introductory text were required to give their informed consent, by clicking on the relevant box, before they could proceed.

A potential participant in a focus group discussion gave her explicit written or verbal consent before the discussion began. Participants were told that they could drop out of the discussion at any time or choose not to answer any individual questions. Participants in the online survey were informed about how the data would be handled and that any identifying information would be removed if quotes were used. The name of each participant, the organization she belonged to – if any – and her within-country location were removed before the results were collated. Any paperwork that held such details was shredded.

**Timing and promotion**

The online survey and focus group discussions took place between the February and June of 2014 and were entirely managed by and for the community of women living with HIV. Survey announcements were disseminated widely by the global reference group and core team, women’s groups and support groups, on Facebook and Twitter and via the relevant United Nations partners, donor agencies and nongovernmental organizations. In addition, the global reference group and core team sent emails and made phone calls to individual women and activists with broad networks and to individual clinicians and doctors who treat women living with HIV. Outreach clinics were visited so that their clients could be invited to participate. Weekly reminder emails, showing the numbers of respondents recruited, were sent to the electronic mailing lists of women living with HIV to show progress, build a sense of owned momentum and encourage further outreach.

**Data analysis**

The analysis of the survey responses started with the structured thematic headings. Quantitative data under each heading were examined to identify major trends. The qualitative free-text responses were read in relation to the key findings derived from the quantitative data. Two members of the core team coded responses to explore key themes in relation to the barriers to – and enablers for – achieving various desired outcomes in terms of sexual and reproductive health and human rights.

**Results**

A full report of the study, including an executive summary, was made available online in January 2015 – i.e. following a WHO Stakeholder Consultation in which the results of the online survey were presented. Some of the results of the study have also already been
published in two peer-reviewed articles – on the impacts of gender-based violence\textsuperscript{13} and poor mental health\textsuperscript{14} on the sexual and reproductive health of women living with HIV – and other analyses of the survey data are in preparation. Here we focus on the methods, approach and main findings. The study covered 945 women living with HIV: 113 from the focus group discussions (Table 1) and 832 from the online survey (Table 2). These women came from 94 countries and were aged 15–72 years.\textsuperscript{4} The key issues raised by the community are summarized in Table 3.

Of the 480 participants in the online survey who answered the optional question on gender-based violence, only 53 (11.0\%) reported never having experienced such violence. According to the respondents, gender-based violence from intimate partners increased after the diagnosis of the women’s HIV positivity. Gender-based violence from family members – other than intimate partners – and community members were often reported to start only after the women had received the diagnosis. Of the 489 women who answered some or all of the optional questions on mental health in the online survey, most reported that they had suffered the symptoms of depression and rejection after they received their diagnosis (Table 3) but only 98 (20.0\%) claimed to have suffered mental health issues before the diagnosis. The respondents recommended that policy-makers and clinicians should address the issues of gender-based violence and poor mental health as a part of a comprehensive global package of care for women living with HIV. The findings from the focus group discussions echoed and expanded upon these findings from the online survey.

In general, the women living with HIV who responded to the online survey appeared to trust the data recorders – as reflected in the large numbers of respondents and the positive comments received (Box 1) – and had a strongly-felt desire to inform the relevant policy-makers of their visions and challenges. Some participants revealed that they had shared experiences – either in the online survey or the focus group discussions – that they had never revealed to anyone before.

Participants responded positively to the online survey both as a reflective, therapeutic process and as an advocacy and learning tool. In both the online survey and focus group discussions, participants called for women living with HIV – in all of their diversity and at all of their different life stages – to be at the centre of all decision-making that affects their lives, so that all of their human and sexual and reproductive health rights can be realized. They called for clinical approaches to be underpinned by a strong human-rights framework, with
laws that actively promote and protect human rights and gender equality instead of simply addressing violations of such rights and equality.

**Discussion**

Here we describe the process of hearing the voices of the community of women living with HIV before the start of the process of developing a guideline. The approach used – of interweaving issues relating to sexual and reproductive health and human rights across the lifespans and diversities of women with HIV around the world – appears to have been welcomed among participants. Although such community participation has been supported elsewhere, here the values and preferences of the key affected population have been elicited for the first time at the start of guideline development. Although such an approach may not always be possible in other contexts or populations, it would merit replication, where possible, in the development of guidelines for many other health considerations. The online survey also provided an opportunity to strengthen institutional capacity and to build understanding – among women living with HIV – of policy and guideline development.

Women living with HIV often suffer from gender-based violence and poor mental health, and these problems may be exacerbated following disclosure of the woman’s HIV status. The outcomes highlight how the lives of women living with HIV are complex – partly as the result of interconnections between issues that often extend beyond the women’s sexual and reproductive health needs. For example, financial insecurity was not only seen as a substantial barrier to obtaining the care needed to protect the women’s sexual and reproductive health, but was also interwoven with the women’s decision-making about having children, more children or no children.

The study had several limitations. The prevalence we recorded was self-reported and, for most sections of the online survey, participants could choose whether to answer or ignore the questions. We have no way of knowing if the women who answered a question were representative of all the participants in the survey. For example, we cannot tell if, compared with the other participants, those with poor mental health or a history of poor mental health were more likely to answer the questions on mental health – perhaps because they were more interested in the topic – or less likely – perhaps because they were too embarrassed to admit they had mental health issues. More rigorous research, with direct observation or other forms of validation, is required. There is also some uncertainty about the quality and accuracy of the electronically translated responses – although most non-English responses were translated, unaided, by a professional translator. There may also have been selection bias, as most of the
women engaging with the survey had access to support networks or groups and/or were advocates. In general, as the participants noted, the survey results are liable to underestimate the impacts of living with HIV, especially in terms of poverty, human rights abuses and violence. Policy-makers need to be more understanding of the complexity of the lives of women living with HIV and the interconnectedness of their problems. All women – including those living with HIV – need humane, holistic services that take into account the different stages of a woman’s life.

Although we found the survey results and the respondent’s attitudes to the survey approach encouraging, the future impact of the survey process and results and the cost–effectiveness of this approach remain unknown. However, in 2015, in response to demand from policy-makers and stakeholders in Latin America, the survey report was translated into Spanish and the survey results were presented in a web conference in Spanish. Furthermore, in December 2014, the Positive Women charitable organization in Ukraine launched a nationwide survey – of the sexual and reproductive rights of women living with HIV in Ukraine – using the same survey tool as used in this study (S Moroz, personal communication, 2015).

The results from the global study reported here should be very useful in the development of new guidelines on the sexual and reproductive health and human rights of women living with HIV and should play an important role in facilitating evidence-based recommendations. Many of the respondents included in our online survey called for further research into (i) the sustained and meaningful engagement of the populations that are likely to be affected by new guidelines in the whole development process for such guidelines and (ii) the benefits and unintended consequences of such engagement.

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Competing interests:
AW and MV are women living with HIV who work professionally as advocates for the rights of women living with HIV globally.
References


Box 1. Selected quotes recorded during the online survey and focus group discussions of women living with HIV

From a member of the global reference group

“Thank you to those of you who have sent others and me words of support in solidarity – your sisterhood is so appreciated! Sometimes it feels like life is unrelenting but I take comfort in knowing that all things, no matter how hard, pass. This is what keeps me going. Thanks for sharing what you are going through and have gone through – it takes such huge courage to share our own hurt and it is so hard. I have no words of wisdom but just sending you so much love and huge amounts of respect. I feel so privileged to know all of you. Thank you! Thank you!”

From a survey respondent in the United States of America

“…I found the experience to be cathartic, much to my surprise. Surveys don’t generally have that effect. It was meaningful to answer questions that truly reflected my experiences both as a girl and young woman before HIV and since my diagnosis. Even though there were questions about violence and trauma that could have felt difficult, the fact that the survey was written by and for women living with HIV and in a tone that is empowering rather than victimizing, made my participation feel good and made me feel that I could be really honest in my answers.”

From a facilitator of the focus group discussion in Myanmar

“[The discussion participants] became active and spoke out, though they dared not speak at first. I shared my experiences and feelings so that they would open up their feelings. When we discussed about discrimination, they said there was no discrimination before their friends knew [that they had HIV infection]. But after discovering that they had the infection, their friends discriminated [against them] by not playing with them. One participant said that her friend’s aunty did not let her friend play with her. Another told how her friend stopped playing with her after discovering that her mother had died of AIDS. Then I asked, who comforts you when you feel bad, and two of them said that only their grandmother comforts because both of their parents had passed away.”

From a survey respondent in Australia

“We are not a one-size-fits-all [community]. Guidelines need to be responsive to individual needs and circumstances, they somehow need to take into account psychosocial factors that impact on my whole-of-life experience and not just my HIV.”
Table 1. Details of the focus group discussions held, with 113 women living with HIV, in seven countries, 2014

<table>
<thead>
<tr>
<th>Country</th>
<th>Language</th>
<th>Participantsa</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ethiopia</td>
<td>Amharic</td>
<td>One discussion with 20 rural women living with HIV and one with 10 sex workers</td>
</tr>
<tr>
<td>Jamaica</td>
<td>English</td>
<td>Eight women aged 41–63 years living with HIV</td>
</tr>
<tr>
<td>Myanmar</td>
<td>Myanma bhasa</td>
<td>10 young female teenagers who were born with HIV</td>
</tr>
<tr>
<td>Nepal</td>
<td>Nepali</td>
<td>One discussion with 10 widows, aged 28–55 years, whose husbands were migrant workers, one with four sex workers aged 17–25 years, and one with five transgender women</td>
</tr>
<tr>
<td>Senegal</td>
<td>French and Wolof</td>
<td>One discussion with 20 sex workers aged 17–65 years and one with 10 women – some of whom were physically disabled</td>
</tr>
<tr>
<td>Thailand</td>
<td>Thai</td>
<td>Five women – two of whom were drug users</td>
</tr>
<tr>
<td>United Kingdom</td>
<td>English</td>
<td>11 migrant women – three of whom had been in prison or detention</td>
</tr>
</tbody>
</table>

HIV: human immunodeficiency virus.

a Ages are shown only when they were recorded.

Table 2. Numbers of respondents in the online survey of women living with HIV in 94 countries, 2014

<table>
<thead>
<tr>
<th>Language</th>
<th>No. of respondents</th>
<th>No. of respondents who were women living with HIV</th>
</tr>
</thead>
<tbody>
<tr>
<td>English</td>
<td>568</td>
<td>480</td>
</tr>
<tr>
<td>Russian</td>
<td>135</td>
<td>99</td>
</tr>
<tr>
<td>Spanish</td>
<td>128</td>
<td>104</td>
</tr>
<tr>
<td>Chinese</td>
<td>80</td>
<td>57</td>
</tr>
<tr>
<td>French</td>
<td>46</td>
<td>42</td>
</tr>
<tr>
<td>Portuguese</td>
<td>44</td>
<td>28</td>
</tr>
<tr>
<td>Bahasa Indonesian</td>
<td>27</td>
<td>22</td>
</tr>
<tr>
<td>Total</td>
<td>1038</td>
<td>832</td>
</tr>
</tbody>
</table>

HIV: human immunodeficiency virus.
Table 3. **Key responses of the women living with HIV who participated in the online survey covering 94 countries, 2014**

<table>
<thead>
<tr>
<th>Response</th>
<th>No. of women answering relevant question</th>
<th>No. of women answering yes (%)&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Had experienced violence or fear of violence&lt;sup&gt;b&lt;/sup&gt;</td>
<td>480</td>
<td>427 (89.0)</td>
</tr>
<tr>
<td>Always or usually had a healthy libido and/or feeling of sexual desire</td>
<td>479</td>
<td>154 (32.2)</td>
</tr>
<tr>
<td>Found service providers to be well trained, knowledgeable, friendly and supportive</td>
<td>589</td>
<td>297 (50.4)</td>
</tr>
<tr>
<td>Had been supported by service providers to make fertility-related choices</td>
<td>318</td>
<td>169 (53.1)</td>
</tr>
<tr>
<td>Had had an unplanned pregnancy</td>
<td>312</td>
<td>177 (56.7)</td>
</tr>
<tr>
<td>Had accessed family-planning counselling</td>
<td>273</td>
<td>122 (44.7)</td>
</tr>
<tr>
<td>Had received advice on safe conception</td>
<td>314</td>
<td>227 (72.3)</td>
</tr>
<tr>
<td>Had received practical support on safe conception</td>
<td>304</td>
<td>168 (55.3)</td>
</tr>
<tr>
<td>Had had a mental health issue after the diagnosis:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Depression</td>
<td>486</td>
<td>360 (74.0)</td>
</tr>
<tr>
<td>Shame</td>
<td>459</td>
<td>325 (70.8)</td>
</tr>
<tr>
<td>Self-blame</td>
<td>478</td>
<td>334 (70.0)</td>
</tr>
<tr>
<td>Feelings of rejection</td>
<td>468</td>
<td>327 (69.9)</td>
</tr>
<tr>
<td>Insomnia</td>
<td>459</td>
<td>314 (68.4)</td>
</tr>
<tr>
<td>Other&lt;sup&gt;c&lt;/sup&gt;</td>
<td>473</td>
<td>285 (60.3)</td>
</tr>
</tbody>
</table>

<sup>a</sup> Of the women answering the relevant question.

<sup>b</sup> Before, since and/or because of positivity for human immunodeficiency virus.

<sup>c</sup> Anxiety, body-image problems, loneliness/isolation and/or very low self-esteem.