Knowledge, stigma, and HIV testing: An analysis of a widespread HIV/AIDS program

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Using randomized methodologies, we study a common community HIV/AIDS program that seeks to promote HIV testing by improving knowledge and reducing stigmatizing attitudes. Contrary to expectations, the program has a substantial negative effect on HIV testing rates. We provide evidence of likely mechanisms behind the program's negative effect: it inadvertently increased misinformation about HIV transmission methods, and worsened HIV-related stigmatizing attitudes. Subsequent household-level randomized treatments providing correct information and addressing stigma concerns counteract the program's negative effect on HIV testing. These findings highlight the importance of improving knowledge and alleviating stigma concerns when promoting HIV testing.

1. Introduction

HIV testing plays a central role in global programs combating the HIV/AIDS pandemic (Granich et al., 2009). Widespread HIV testing in areas of high prevalence is important because people with HIV infection are often asymptomatic for years before the disease progresses to AIDS. When individuals are found to be infected with HIV, it is recommended that they immediately start antiretroviral therapy (ART) (WHO, 2017). Rapid initiation of ART lowers HIV plasma viral loads, providing private benefits for infected individuals in the form of better health outcomes (Ford et al., 2018). In addition, early initiation of ART has public health benefits due to a positive externality (Greenwood et al., 2019). During the asymptomatic phase, those with HIV infection can transmit HIV to others. Initiation of ART leads to reduced HIV viral loads and a much lower risk of transmitting HIV to sexual partners (Rodger et al., 2019).

While there has been substantial progress in expanding HIV testing around the world, testing rates remain far from achieving targets set by public health officials. The UN’s 90-90-90 goals are widely-adopted...
objectives: 90% of people with HIV infection should be diagnosed, 90% of those diagnosed should be in treatment, and 90% of individuals in treatment should have an undetectable HIV viral load (United Nations, 2016). However, UNAIDS estimates that in 2019, of the 38 million people infected with HIV globally, about 7.1 million (19%) are undiagnosed (UNAIDS, 2020).

We seek to shed light on the impact of a major type of HIV/AIDS program on HIV testing rates, and to understand the mechanisms underlying its effectiveness (or lack thereof). We focus on two mechanisms: alleviating imperfect information related to HIV and reducing HIV-related stigmatizing attitudes. We start with a simple theoretical model. Individuals decide whether to have an HIV test, trading off health benefits with social stigma costs of revealing one’s HIV risk type to others. Individuals revealed to have high HIV infection risk are stigmatized (excluded from social interactions). Beliefs about HIV transmission affect the extent to which people are stigmatized if they are observed getting an HIV test and reveal themselves as having high risk of HIV infection. The less people believe that HIV is transmissible, the lower the stigma, and the higher the HIV testing. Conversely, if people believe that HIV is more transmissible, stigmatizing attitudes rise, and HIV testing falls.

With this model as a framework, we study a program in Mozambique, *Força à Comunidade e Crianças* (FCC, “Strengthening Communities and Children”), that aims to raise HIV testing rates by improving knowledge about HIV/AIDS and reducing HIV-related stigmatizing attitudes. FCC is a community-level program that implements home visits to households, as well as complementary interventions in communities and schools. The program is representative of a broad category of HIV/AIDS interventions, known as programs for “orphans and vulnerable children” (OVCS), that are funded by the U.S. Presidential Emergency Plan for AIDS Relief (PEPFAR).2

We designed a randomized controlled trial to estimate the causal impact of the FCC program on HIV testing rates and the mechanisms through which it operates. We specified our analyses in advance in a pre-analysis plan (PAP). The research design involves three stages of randomization, as presented in Fig. 1. The sample is composed of 3700 households that we have been following from a 2017–18 baseline through a 2019 endline survey. First, we randomized half of 76 communities to treatment (receiving the program) and half to the control group. Second, motivated by concerns about statistical power, we randomized a subset of households within treatment communities to a strong encouragement to participate in the FCC program (“FCC-enrolled” households). FCC-enrolled households received home visits by FCC community workers and were assessed for inclusion in various FCC components. FCC enrollment led them to have higher participation rates in the program than other households in treatment communities.3

An endline survey collected data on a range of household outcomes, including self-reported HIV testing in the household. As pre-specified, all treatment effects reported in this paper are the effect of being an FCC-enrolled household in a treatment community, with the comparison group being households in control communities.

Immediately after the endline survey, our research staff then randomly assigned households to a set of “minitreatments” aimed at encouraging further HIV testing, or a minitreatment control group. The different minitreatments provide HIV-related information, seek to alleviate concerns about HIV-related stigma, and provide additional financial incentives for HIV testing.4 Our research staff implemented the minitreatment to which a household was assigned (if any), and then offered coupons to encourage household members to get HIV tests at the nearest health clinic.5

Our primary outcome of interest in this study is whether anyone in the household received an HIV test in the 14 days after the endline survey, measured by redemption of these encouragement coupons. As an administrative outcome, this measure of HIV testing is not subject to survey reporting biases.6

This research design yields several treatment effects. The treatment effect of primary interest is the impact of FCC enrollment on the coupon-based HIV testing measure. This is Comparison A in Fig. 1, the testing rate of FCC-enrolled households in treatment communities who did not receive any minitreatments, minus the testing rate of households in control communities who also did not receive any minitreatments. This is the “pure” effect of the FCC program that is not clouded by any effects of minitreatments.

We find that the FCC program has a negative effect on HIV testing: −10.9 percentage points, relative to a base of 26.3 percent in the control group. This result is contrary to our pre-specified expectation of a positive impact on HIV testing. It is also contrary to the positive impact expectations of 73 experts surveyed in advance by DellaVigna et al. (2020) before our results were publicly known.

In pre-specified secondary analyses, we shed light on mechanisms behind the FCC program’s negative effect on HIV testing. These analyses estimate the Comparison B treatment effect in Fig. 1, comparing FCC-enrolled households in treatment communities with all households in control communities. Outcomes are potential mechanisms measured in the endline survey.7 We find that the program did not improve HIV-related overall knowledge, and in fact increased misinformation. Treated respondents became more likely to believe “myths” about HIV transmission (e.g., that HIV can be spread by shaking hands or by witchcraft). In addition, the program actually worsened HIV-related stigmatizing attitudes, measured by answers to survey questions on HIV-related stigma (such as whether one would buy vegetables from an HIV-positive vendor, or thinks that an HIV-positive person should be a teacher).

These findings are suggestive that the FCC program’s negative impacts are due to worsened information and increased stigma. However, simply showing that the treatment leads to worsened information and increased stigma does not establish with certainty that these are mechanisms behind the program’s impacts, since these outcomes could co-move with HIV testing without being mechanisms in the causal chain linking FCC enrollment with testing. The minitreatments we implemented after the endline survey provide more direct evidence that changes in information and stigma are mechanisms behind the FCC program’s negative impact on testing.

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2 PEPFAR is the world’s largest source of funding for HIV/AIDS programs in developing countries (PEPFAR, 2020), with an annual budget ranging from $6.6 to $6.9 billion in 2015–2020 (US State Department, 2019).

3 Other households not randomly selected for direct enrollment were exposed to the FCC program as well, but at lower rates. We pre-specified treatment effects on these “FCC-ambient” households as of secondary interest; results are presented in the Populated Pre-Analysis Plan (PAP). The Populated PAP can be found in our AEA RCT Registry record (AEARCTR-0003990).

4 The initial motivation for these minitreatments was to examine complementarity between the FCC program and more targeted interventions to raise HIV testing. As it turns out, these minitreatments end up revealing mechanisms through which the FCC program’s (negative) effects on testing operate.

5 Conditional on the individual getting an HIV test, the coupons were redeemable at the clinic up to 14 days later for a financial reward of 50 meticais (PPP US$2.42).

6 As pre-specified, we prioritize the coupon-redemption-based measure of HIV testing because treatment effects using this measure differ from treatment effects on self-reported HIV testing from the endline survey. Treatment effects on self-reported HIV testing are significantly more positive, likely due to experimenter demand effects.

7 Comparison B is the appropriate comparison for the outcomes in the endline survey, as it maximizes sample size; the minitreatments were implemented after the endline survey and therefore cannot affect endline survey outcomes.
The minitreatments provided information about HIV; information about HIV treatment (ART); both HIV and ART information; information to reduce concerns about HIV-related stigma; and a higher financial incentive to receive an HIV test. There was also a control group that got no minitreatment. In analyses that were also pre-specified, we examine how these treatments affect the coupon-based measure of HIV testing. In Fig. 1, these analyses involve comparing households in the gray-shaded “Pure Control” box among FCC-enrolled households (FCC-enrolled households receiving no minitreatment) with households in the five boxes just below it (FCC-enrolled households assigned to some minitreatment).

The minitreatments counteract the negative effect of the FCC program. Among FCC-enrolled households, those getting an information minitreatment or the anti-stigma minitreatment have substantially higher financial incentives for this minitreatment was 100 meticais per coupon, double the incentive offered to all other households.
higher HIV testing rates than the minitreatment control group. These findings further support the interpretation that the FCC program caused lower HIV testing by worsening knowledge and increasing stigmatizing attitudes; minitreatments targeting these mechanisms helped reverse the FCC program’s negative effects on HIV testing.

Overall, our theoretical model encapsulates the mechanisms behind our empirical findings. The FCC program led to misinformation about the transmission of HIV, which worsened stigmatizing attitudes, and led to lower HIV testing rates. Minitreatments aimed at improving HIV-related information and reducing stigmatizing attitudes raised HIV testing rates.

Before our empirical results were known, we submitted the methods and data analysis plan for this study as a pre-results review paper to the Journal of Development Economics (JDE). The JDE accepted our pre-results review paper, committing to publish the complete paper and data analysis plan for this study as a pre-results review paper, committing to publish the complete paper and this paper is the complete paper with empirical results.

In this paper, we do depart from the pre-specified set of analyses by presenting a subset rather than all the analyses detailed in the JDE Stage 1 Proposal. We report all primary analyses detailed in the JDE Stage 1 Proposal, but only a selection of secondary analyses. We concisely overview the other secondary analyses in Section 5.3, and more fully in a Populated PAP (following Duflo et al. (2020)).

We present only a subset of pre-specified results because of the unexpected negative treatment effect on our primary outcome of interest (HIV testing), which led us to focus on explaining potential mechanisms behind this unexpected result. We therefore focus in the paper on analyses of those potential mechanisms — HIV/AIDS-related information and stigmatizing attitudes. We report results for other secondary outcomes that are unlikely to represent potential mechanisms in the Populated PAP.9

The main contribution of this paper is to provide evidence on the importance of two intermediating mechanisms behind the efficacy of HIV/AIDS public health initiatives. We highlight the importance of (1) public health information, and (2) concerns about disease-related stigma. Our work is related to a large existing literature studying the role of information in health decision-making, and a much smaller literature on how stigma affects health decisions.

Studies of the role of information imperfections in health decision-making are reviewed by Duflo and Miguel (2016). In the HIV/AIDS context, provision of general HIV/AIDS information has been shown to affect knowledge, health behaviors, and demand for health goods (Duflo et al., 2015; Duflo, 2011; Godlonton et al., 2016; Ciancio et al., 2020; Kim et al., 2017; Chong et al., 2013; Banerjee et al., 2020). Smith et al. (2021) find that providing information that HIV treatment helps prevent transmission leads to higher HIV testing rates (Bor et al. (2021) provide a review of “treatment-as-prevention” interventions). Other studies have examined the impact of learning one’s own HIV infection status (Delavande and Kohler, 2012; Gong, 2015). Our study is novel in finding that a major type of HIV/AIDS program can create misinformation. As such, our findings concord with the smaller number of studies that find that informational HIV/AIDS interventions can lead to misinformation or have harmful effects (Jamison et al., 2013; Godlonton et al., 2016; Friedman, 2018).

While there has been a great deal of interest in stigma in the context of the HIV/AIDS pandemic,10 economics research on the topic is scarce.11 Prior work using randomized or experimental methodologies has shown that interventions can reduce HIV-related stigmatizing attitudes (Hoffmann et al., 2016; Lubega et al., 2019), thereby leading to higher rates of HIV testing (Derkson et al., 2020; Yu, 2021). Relative to this literature, our work is novel in finding that a major type of widely implemented HIV/AIDS program can actually worsen HIV-related stigmatizing attitudes, and that a simple anti-stigma intervention implemented later can offset the program’s negative impacts on HIV testing.

This research also contributes to economic research on large-scale HIV/AIDS programs, of which PEPFAR is the largest funder worldwide. FCC is a representative example of PEPFAR programs for orphans and vulnerable children (OVCs). U.S. law requires PEPFAR to spend 10% of its budget on OVC programs (US State Department, 2015). Among OVC programs, FCC is emblematic in key ways. The program delivers a multifaceted array of services through schools, as corner-stone institutions in communities. These school-based programs operate alongside household-level case management visits to households by program workers who deliver informational and anti-stigma messaging, encourage HIV testing, assess household needs, and connect households and individuals with appropriate program components (USAID, 2012; US State Department, 2019).

Prior studies of PEPFAR programs have not exploited prospectively randomized research designs. In addition, past studies have not tracked defined households or individuals over time (from before to after program implementation), raising concerns about sample selection biases (Bryant et al., 2012). Bendavid et al. (2012) examine the impact of PEPFAR funding at the country level using a difference-in-difference approach, finding substantial reductions in adult mortality in Africa. A number of past studies have used randomized controlled trials to examine the impact of more targeted interventions related to HIV/AIDS, such as Thornton (2008), McCoy et al. (2017), Sewamala et al. (2009), Ivers et al. (2014), Baird et al. (2011), Kiene et al. (2017), and Yotebieng et al. (2017). None of these have studied PEPFAR or community-level programs, or examined the interplay between knowledge, stigma, and HIV testing as we do.

2. Theoretical model

We model the HIV testing decision as a one-sided signaling game with two players over two periods. Our model is a simplified version of the model in Derksen et al. (2020). Rational individuals decide whether to test for HIV. The benefit of testing includes revealing to others that one has a high probability of having HIV, and experiencing stigma as a result. We model stigma as being excluded from social interactions with others. Beliefs about HIV transmission affect the extent to which people are stigmatized (excluded from social interactions) if they are believed to be at high risk of having HIV. If people start to believe that HIV is less transmissible, this leads to less stigma and higher HIV testing.

Consider two players, A and B, from two continua of individuals $A = [0,1]$ and $B = [0,1]$, respectively. Each player A has private information about his own type $(\theta_A, y_A)$. $\theta_A$ represents A’s probability of being HIV-positive and it takes two values: $0 < \theta_L < \theta_H < 1$ (low and high risk, respectively). The fraction of A with $\theta_A = \theta_H$ is $r$. $y_A$ is player A’s valuation of social engagement. Each player B is known to be HIV-negative and has a private valuation of social engagement, $y_B$. $y_B$ follow the same distribution $\mathcal{G}$ with a positive support $(\infty, \infty)$. $r$ and $\mathcal{G}$ are common knowledge.

In the first period, player A decides whether to take an HIV test. A’s action can be observed by individuals in B. Player A is then randomly

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9 We find zero treatment effects on these other secondary outcomes; discussing those findings only in the Populated PAP therefore also helps streamline the paper.

10 Mahajan et al. (2008) and Stangl et al. (2013) review public health research on HIV/AIDS stigma.

11 Economists have studied stigma in other contexts (Moffitt, 1983; Vishwanath, 1989; Akerlof et al., 1996; Furuya, 2002; Ivers, 2002; O’Flaherty and Sethi, 2008; Bhargaw et al., 2017); see Durlauf and Blume (2008) for a review.
matched with a player $B$ in a social activity. In the second period, player $B$ decides whether to interact with player $A$ (social avoidance) or not (social avoidance). We denote $A$’s action as $t_A \in \{0 : \text{do not test}, 1 : \text{test}\}$ and $B$’s action as $m_B \in \{0 : \text{do not interact}, 1 : \text{interact}\}$.

In period 1, player $A$’s direct cost for taking a test is $c$. This summarizes time and monetary costs as well as physical and psychological costs associated with the trip to taking a test. If testing positive, player $A$ receives ART treatment which brings a health benefit of $v$.12 We denote $A$’s real HIV status as $h_A$, which takes value 1 if positive and 0 if negative. The payoff for $A$ in period 1 is given by

$$u_A^*(t_A) = t_A(h_A v - c).$$

The expected gain in period 1 for taking a test, $\mathbb{E}_h[u_A^*(1) - u_A^*(0)]$, is equal to $\theta_A v - c$.

We make the following assumptions to restrict the parameter values:

$$\theta_A v < c < \theta_H v < \Sigma \quad (1)$$

Assumption (1) states that a low-risk ($\theta_h = \theta_A$) person does not have a high enough health-related incentive to take a test, but a high-risk person does. Assumption (2) means that the health benefit of testing is higher than the value of social engagement for at least some high-risk $A$.

In period 2, player $B$ forms a belief about $\theta_S$ based on the observed $t_A$, $\hat{\theta}_S(t_A)$. If $B$ chooses to interact with $A$, both players will receive utility from social engagement but $B$ also faces a cost due to potential HIV transmission from $A$. We denote $B$’s perceived HIV transmission probability as $\hat{\tau} \in [0, 1]$, and her health cost from HIV infection as $z$. If $B$ chooses not to interact, both players obtain no utility in this period.

To sum up, in period 2, player $A$’s payoff is

$$u_A^*(m_B) = m_B h_A,$$

and player $B$’s payoff is

$$u_B^*(m_B) = m_B \left[y_B - \hat{\theta_B}(t_A)z\right].$$

We make further assumptions to bound parameter values:

$$\Pr(y_B > \hat{\theta} z) = 1 \text{ and } \Pr(y_B < \hat{\theta_H} z) > 0, \quad (4)$$

which means that no one in $B$ would reject an unknown person from the general population, $A$, for fear of HIV transmission; but at least some would avoid social interactions with a person of high risk.

Overall, the payoff function for player $A$ is13

$$U_A = u_A^*(t_A) + u_A^*(m_B).$$

**Proposition 1.** There are two classes of pure strategy Perfect Bayesian Equilibria in this game. The first is a partially separating equilibrium. A fraction $S$ of individuals in $B$ discriminate: they avoid $A$ who has sought an HIV test. The remaining fraction $(1 - S)$ of individuals interact with any $A$. The second is a pooling equilibrium. A fraction $P$ of individuals in $B$ interact if and only if $A$ has sought an HIV test.

In what follows, we characterize the partially separating equilibrium of the game where stigmatizing behavior (social avoidance) occurs, and defer the formal proof of Proposition 1 to Appendix A.

First, consider player $A$’s optimization problem given $S$. $A$ chooses to take an HIV test or not to maximize his expected utility $\mathbb{E}_{h_A,m_B}[U_A]$:

$$\max_{t_A \in \{0, 1\}} \left(\theta_A v - c - SY_A h_A + y_A\right).$$

His optimal strategy is to test if and only if

$$y_A < \frac{\theta_A v - c}{S}.$$

Combining Assumption (1), (2), and that $y_A > 0$, we know that a player $A$ with $\theta_A = \theta_L$ never takes a test; and that some $\theta_A = \theta_H$ will take a test. Player $B$, with Bayesian updating, forms the belief of $\hat{\theta}_S(t_A)$ accordingly:

$$\hat{\theta}_S(t_A) = \theta_B; \quad \hat{\theta}(0) < \hat{\tau}.$$  \quad (5)

Given $A$’s strategy, now consider $B$’s optimization problem. $B$ chooses $m_B \in \{0, 1\}$ to maximize the payoff as in (3):

$$\max_{m_B \in \{0, 1\}} m_B \left[y_B - \hat{\theta}_B(t_A)z\right].$$

By Assumption (4) and belief (5), we know that when $t_A = 0$, $B$’s best response is $m_B = 1$; while when $t_A = 1$, $B$’s best response is to interact if and only if

$$y_B > \hat{\theta}_B z.$$  \quad (6)

Thus,

$$S = \int_{\Sigma} \hat{\theta}_B z \, d\mathcal{G}.$$  \quad (7)

Expression (7) leads to the following proposition.

**Proposition 2.** Stigmatization $S$ becomes more (less) severe as individuals’ perceived HIV transmission risk $\tau$ increases (decreases).

In this equilibrium, the testing rate among the population $A$ is given by

$$R = r \int_{\Sigma} \hat{\theta}_B z \, d\mathcal{G}.$$  \quad (7)

**Proposition 3.** A higher (lower) rate of stigmatization $S$ suppresses (encourages) HIV testing.

In sum, with knowledge (beliefs about the transmission of HIV) having such a central role in raising HIV testing, much can go wrong if a program inadvertently fails in its knowledge-raising objective, and instead creates misinformation. For example, a program could lead people to mistakenly believe that HIV is transmissible via mechanisms such as shaking hands or sharing food (which is not the case). If a program thereby leads people to believe that HIV is more transmissible, more people come to stigmatize those infected with HIV, and HIV testing falls. We now turn to the empirical analysis of the FCC program, which we interpret in the context of this model.

3. Research design

3.1. Country and program context

Out of an estimated 36.9 million people living with HIV worldwide in 2017, 25.7 million are in Sub-Saharan Africa. The region also accounts for a dominant share of new HIV infections: 1.17 million out of a global 1.8 million in that year. In Mozambique in 2017, 2.1 million people out of a population of 29.7 million were living with HIV (7.1%), out of which 170,000 were children (aged 14 or below). The country has an estimated 130,000 new HIV infections annually, of which 13.8% are children. Mozambique recorded 70,000 AIDS-related deaths in 2017, likely because only slightly more than half of HIV-infected patients have access to antiretroviral therapy (ART). Poor access and adherence to ART contributes to AIDS-related morbidity and mortality, as well as HIV transmission (to other adults as well as from mothers to children) (UNAIDS, 2019). Only an estimated 77% of
Mozambicans with HIV infection know their status [UNAIDS, 2020]. In our own Mozambican sample, nearly half of adults and 90% of children are reported to have never been tested for HIV.

The U.S. Government’s most important program responding to the HIV/AIDS crisis is the President’s Emergency Plan for AIDS Relief (PEPFAR). PEPFAR mandates that 10% of its funding be devoted to programs benefiting children orphaned or made vulnerable by HIV/AIDS (“orphans and vulnerable children”, or OVCs). In 2016, PEPFAR OVC programs supported 6.2 million OVCs and their caregivers worldwide (PEPFAR, 2017).

3.2. The program

The program we study, Força à Comunidade e Crianças (FCC, “Strengthening Communities and Children”), is a representative example of PEPFAR OVC programs. Its high-level aim is to improve families’ and communities’ ability to support, protect, and care for orphans and vulnerable children, their caregivers, and their households more generally.

The FCC program is composed of a number of interrelated components and is implemented in study districts by local implementing partner (LIP) organizations under contract to the international NGO World Education Inc. (WEI/Bantwana). Several FCC program components are school-based, and so programs are implemented in local communities surrounding a focal school. In each community, activities take place with the collaboration and advice of a Community Child Protection Committee (CCPC) whose membership includes community leaders, volunteers, and local government officials.

The most widespread FCC program component is home visits by LIP staff known as “Case Care Workers” (CCWs) to households. Roughly 700 CCWs work across the study communities. LIPs hire CCWs locally, in part based on recommendations by the CCPC and community leaders. In common with the local populations they serve, they typically have no more than primary school education. Roughly 80% of CCWs are female. They range in age from 18 to 48, with most falling between 25 and 40 years of age. CCWs receive a stipend of 3,100 MZN per month (roughly US$150), as well as in-kind compensation in the form of a bicycle, a work uniform, and cellphone airtime.

CCWs conduct home visits of OVC households, based on personal knowledge and recommendations of the CCPC. The home visit itself is a conduit for the dissemination of information and advice by CCWs. Household members may then participate in other FCC components, based on the results of the home visit. In home visits, CCWs conduct systematic vulnerability assessments, and households (and individuals therein) are then linked to appropriate programs and services in communities, schools, and health facilities. One of the most important results of these home visits is the referrals of individuals for HIV testing at the local PEPFAR-funded health clinic. The expectation is that CCWs refer all FCC program beneficiaries (both adults and children of all ages) who do not know their HIV status for HIV testing, and that testing should be repeated every twelve months even upon a negative test result. The number of individuals referred to HIV testing is a key outcome indicator for the FCC program, monitored by PEPFAR in the context of achieving the UNAIDS 90-90-90 global goals (90% of those with HIV diagnosed, 90% of those on ART, and 90% of those virally suppressed by 2020) (PEPFAR, 2017). Those testing positive for HIV are then referred to initiate antiretroviral therapy (ART) through the local clinic. CCWs in the community then follow up with individuals initiating ART to promote ART adherence on an ongoing basis. Because of the centrality of encouraging HIV testing in the FCC program, it is the primary outcome of interest in this study.

During home visits, CCWs seek to increase HIV testing rates via two mechanisms we examine explicitly: improving information and reducing stigma concerns. CCWs seek to improve FCC beneficiaries’ information related to HIV/AIDS, such as methods of disease transmission, progression of the disease, treatment, HIV testing, and locations of health clinics providing testing and treatment. Information is conveyed verbally and, at the LIP’s discretion, on printed material given to the household. In addition, CCWs are expected to engage program beneficiaries in “sensitization” to address stigma related to HIV (both one’s own stigmatizing attitudes, and fear of stigma from others). CCWs engage in discussions to reduce stigmatizing attitudes among program beneficiaries. They provide psychosocial support (PSS) and gradually gain program beneficiaries’ trust over time in repeated interactions, with the expectation that reductions in fear of stigma will encourage people to be open to HIV testing, voluntarily disclose HIV-positive status to CCWs, and be open to future CCW follow-up promoting ART initiation and adherence.

Households are connected to other FCC program components after the home visits, based on needs assessments conducted by CCWs. These program components include Village Savings and Loan (VSL) groups and nutritional screenings of children. Many components are school-based, so children can also be included in these components through their schools. Relatively few households report participating in these other program components. For further details on these other program components, see Appendix B.

3.3. Sample and data

The study timeline is shown at the bottom of Fig. 1, illustrating key points in data collection and the randomized interventions. We provide an overview of data collection here, and provide details in Appendix C.

In the 76 study communities, we selected households for inclusion in the sample using random-route sampling, with route starting points at the focal school in each community. In keeping with the focus on the FCC program, we administered a “vulnerability assessment” (designed jointly with WEI/Bantwana) to identify OVC households. 71.68% of households were determined to be OVC households and communities’ ability to support, protect, and care for orphans and vulnerable children, their caregivers, and their households more generally.

14 The UN defines an “orphan” as a child who has lost one or both parents. An estimated 13.4 million children and adolescents (0–17 years of age) worldwide had lost one or both parents to AIDS as of 2015. More than 80% of these children (10.9 million) live in sub-Saharan Africa (UNICEF, 2016). PEPFAR’s 2008 reauthorization mandated it to spend 10% of its funds on OVC programs. PEPFAR OVC funding amounted to $3.58 billion in total from 2015–2020 (PEPFAR, 2017, 2020).

15 Reviews of research on OVCs include Bryant and Beard (2016), Goldberg and Short (2016), Nyberg et al. (2012), and Shann et al. (2013). Also see Evans and Miguel (2007), Case et al. (2004), Larson et al. (2013), and Whetten et al. (2015).

16 LIPs are local non-government organizations (NGOs) operating in study areas.

17 Program provinces and districts are: Manica province (Manica, Chimoio, and Gondola districts), Sofala province (Dondo and Nhamatanda districts), and Zambezia province (Namacura and Nicoaodala districts).

18 CCWs were provided with a detailed “Home Visit Guide” detailing information they are expected to convey to households, covering health (HIV/AIDS in particular), education, nutrition, infant care, legal rights, savings, and psychosocial help. The home visit guide is included as part of our pre-analysis plan (PAP) at our AEA RCT Registry entry (https://doi.org/10.1257/rct.3990-5.0).
is the head of household. For some questions (in particular, self-reported HIV testing, HIV knowledge, and HIV-related stigmatizing attitudes), we also surveyed other adults who were present at the time of the survey. The sample is composed of 3658 households included in the endline survey. Household-level outcomes are reported by the primary household respondent or by aggregating reports of surveyed individuals.

The primary outcome of interest in this study is HIV testing at the household level. We focus on an objective, administrative measure of HIV testing by our study participants. At the end of the endline survey, our survey team recommended that individuals in the household be tested for HIV (if they had not had a test performed within the past three months) at their local health clinic within the next 14 days. Local health clinics in each study community collaborated with us to conduct the HIV testing and facilitated collection of testing data. To allow tracking of those who followed through with testing, households were given coupons redeemable for a financial incentive (50 MZN) at the health clinic after having the HIV test. Coupons had a unique code for each household, allowing us to track their redemption. Households were given as many coupons as needed, for however many individuals did not know their status or who reported being HIV negative but were tested more than three months in the past. To receive the financial incentive after having their HIV test, individuals had to present the coupon to our research staff (also stationed at the health clinic), along with a form signed by clinic staff that the individual had just gotten an HIV test.

An indicator for at least one of a household’s coupons being used (indicating at least one household member had an HIV test in the 14-day window after the endline survey) is our primary outcome variable. Because this outcome is a directly-observed, administratively recorded health behavior, it avoids potential reporting biases associated with survey-reported HIV testing. Conceptually, it captures a household’s receptiveness to a recommendation to be HIV tested. Programs such as FCC aim to not only facilitate an individual’s first HIV test, but also to encourage individuals to be receptive to regular, repeated testing (consistent with Mozambican public health recommendations).

As a secondary outcome, we examine a self-reported measure of HIV testing: an indicator for anyone in the household having had an HIV test in the previous 12 months, reported in the endline survey (self-reported by adults, and reported by the primary caregiver for children under the age of 18). Prioritizing the coupon-based measure of HIV testing over the self-reported measure follows a decision rule we pre-specified in our PAP, stemming from the fact that treatment effects differ across the two measures (see Section 4.2 below).

We also examine secondary outcome variables related to information and stigma mechanisms, reported in the endline survey. These outcomes are at the individual level, for all adult respondents of the endline survey. The information questions assess objective knowledge about HIV: correct transmission methods, incorrect transmission methods (“transmission myths”), protection methods, and treatment. Stigma questions measure stigmatizing attitudes towards HIV-positive individuals (would buy food from an HIV-positive seller, would allow an HIV-positive teacher, would keep a relative’s HIV-positive status secret), and that there is no impact of treatment on in-migration into treatment communities.

After households were enrolled in the study, households in treatment communities were randomly assigned to FCC-enrolled vs. FCC-control communities. FCC beneficiary counts in WEI/Bantwana’s data indicate relatively smooth enrollment of FCC beneficiaries across quarters, from Q1 (Jan–Mar) 2017 to Q1 (Jan–Mar) 2018. The program then continued serving beneficiaries, remaining active through the period of our endline survey.

We administered informed consent and included households in the study sample from May to November 2017 for 98.7% of our eventual sample. Appendix Figure A.1 displays the number of households that were enrolled in the study sample by month. WEI/Bantwana’s data indicate that about two-thirds to three-quarters of eventual FCC beneficiaries (depending on program subcomponent) had received their first contact with the program by Q3 (Oct–Dec) 2017. The fact that household enrollment into the study occurred after initiation of FCC program activities raises potential concerns about selection into treatment communities. We show in Section 5.1.1 below that observable characteristics of households show no relationship with FCC treatment, and that there is no impact of treatment on in-migration into treatment communities.

Randomization stage 1: Assigning FCC and control communities

The FCC program is a community-level intervention, so the first stage was random selection of communities to receive or not receive the FCC program. FCC interventions are centered in primary and secondary schools, so geographic areas of interest are residential areas surrounding schools. (We refer to areas surrounding schools simply as “communities”, each of which has a “focal school” where school-based program components are implemented.) WEI/Bantwana consulted with local implementing partners (LIPs) and government officials in the three provinces and seven districts in which the FCC program was to be implemented to identify a set of 76 communities deemed eligible for participation.

3.4. Methodology

We aim to estimate causal effects of the FCC program using a randomized controlled trial (RCT) methodology. Random assignment allows estimated relationships to be interpreted as causal effects, rather than simply correlations. Our approach involves a three-stage randomized controlled trial methodology to estimate causal effects of the FCC program, and to shed light on some of the mechanisms through which its effects operate.

Fig. 1 displays the research design and timeline of the study. In November 2016, we randomly assigned 76 communities to be FCC treatment communities or control communities (Randomization Stage 1), after which WEI/Bantwana initiated the program in FCC treatment communities. FCC beneficiary counts in WEI/Bantwana’s data indicate relatively smooth enrollment of FCC beneficiaries across quarters, from Q1 (Jan–Mar) 2017 to Q1 (Jan–Mar) 2018. The program then continued serving beneficiaries, remaining active through the period of our endline survey.

Data indicate relatively smooth enrollment of FCC beneficiaries across quarters, from Q1 (Jan–Mar) 2017 to Q1 (Jan–Mar) 2018. The program then continued serving beneficiaries, remaining active through the period of our endline survey. 2019. At the end of the endline survey we randomly assigned households to the mini-treatments (Randomization Stage 3) and offered all households encouragement coupons for HIV testing. We collected the HIV testing encouragement coupons up to each community’s deadline (14 days after conclusion of the endline survey in a community). We now discuss each of the three stages of randomization in detail.
for the program. These communities were chosen on the basis of being geographically proximate to ART sites (health clinics offering HIV testing and treatment), having sufficient populations of orphans and vulnerable children (OVCs), and having no other active donor funded HIV/AIDS programs. These 76 communities were then sorted into stratification cells of matched community pairs, sets of two communities that were very similar in terms of distance to ART sites, school type (secondary or primary), and student enrollments.

Within each matched pair, treatment status was randomly assigned to one community, with the other school assigned to control status. Randomization of treatment status within matched pairs helps ensure balance in baseline characteristics between treatment and control units, so that treatment-control comparisons can then be credibly interpreted as causal effects of the program. This random assignment was carried out on the computer of one of the coauthors, one time, with no re-randomization. We communicated the result of the randomization to WEI/Bantwana in November 2016. The FCC program was then implemented in treatment communities, and not in control communities. Maps of the locations of FCC treatment and control communities can be found in Appendix E.

Randomization stage 2: FCC enrollment within FCC communities

The second stage of randomization, at the household level, was implemented only within treatment communities. This randomization stage was motivated by concerns that treatment effect estimates based on generally comparing households in treatment and control communities would have low statistical power (because of low penetration of the program in treatment communities). This stage of randomization creates a subgroup in treatment communities, FCC-enrolled households, with relatively high take-up or participation in the FCC program. Estimates of the impact of the FCC program comparing FCC-enrolled households to households in control communities, therefore, have higher statistical power.

Among households in FCC communities who consented and were included in the study sample, a subset was randomly assigned to “FCC enrolled” status. GPS coordinates and household head’s name and contact information of FCC-enrolled households were provided to WEI/Bantwana and their local implementing partners (LIPs). LIP staff (case care workers, CCWs) then conducted home visits to these households.

We carried out the randomization one FCC community at a time, with no re-randomization, on the computer of one of the co-authors, in November and December 2017. Seven-twelfths (58.33%) of study households in each FCC community were assigned to FCC-enrolled status. Other households not randomly selected for direct enrollment, which we refer to as “FCC-ambient”, end up being exposed to the FCC program at lower rates.

CCWs conducted home visits with FCC-enrolled households alongside their broader FCC responsibilities over the course of the following several months. Our research team supported LIPs with training in use of GPS devices to locate households using latitude and longitude coordinates. By November 2017, we had completed all enrollments of households in the study sample necessary for randomization of FCC-enrolled status (Randomization Stage 2). The implementation of Randomization Stage 2 was spread out over November and December 2017, as we completed data cleaning and finalized the study sample before implementing the randomization, community by community. The one community whose 46 household study enrollments were delayed until March 2018 happened to have been a control community, so there was no Randomization Stage 2 for that community.

Because the FCC-enrolled treatment was to be of primary interest, we chose to have the probability of FCC enrollment be slightly above half, to increase our statistical power to detect the FCC-enrolled treatment effect. The probability of seven-twelfths derived from the ratio of 35 out of 60 households in the targeted endline survey sample in each community. Due to attrition, we have fewer than 60 households per community in the ultimate sample for analysis in this paper.

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coordinates. WEI/Bantwana reported that CCWs had completed home visits to 44% of FCC-enrolled households by February 2018. This figure rose to 64% by May 2018, and 77% by their final update to our research team in November 2018 (six months before the endline survey).

We pre-specified in our PAP that the causal effect of FCC-enrollment would be of primary interest, while the causal effect of FCC-ambient status would be of secondary interest. In this paper, we focus on effects of FCC-enrollment, and report FCC-ambient effects in overview in Section 5.3 below (with full detail in the Populated PAP).

Randomization stage 3: The minitreatments

In the third part of our randomized methodology, after households completed the endline survey, we randomly assigned them to one of five treatment conditions or a control condition. We refer to these post-endline-survey treatments as “minitreatments”. The outcome variable of interest for the minitreatments is the redemption of the incentive coupon for HIV testing. This outcome is the only HIV testing outcome we observe after the minitreatments, because the self-reported measure is collected in the endline survey itself.

We originally conceived of these minitreatments as providing insight into whether the FCC program is complementary with or substitutable for more targeted interventions to promote HIV testing. The minitreatments turn out to help reveal likely mechanisms through which the FCC program’s (negative) effects operate.

The minitreatments were randomly assigned, and then implemented by the same research staff member who had just administered the endline survey to the respondent. For further information about these treatments (including links to videos), please see Appendix F.

1. Anti-Stigma: Individual-specific information aimed at reducing concerns about HIV-related stigma in the community. Past surveys in Mozambique have shown that many stigmatizing attitudes have fallen over time, as depicted in Fig. 2. However, individuals tend not fully realize this. In a companion paper to this study in an overlapping sample, Yu (2021) shows that correcting such overly-pessimistic beliefs about stigma can encourage HIV testing. We ask endline-survey respondents about the fraction of residents in their community they think hold specific stigmatizing attitudes towards people living with HIV. Respondents overestimating this fraction for any question are told the true (lower) value we collected from the baseline survey. Our theoretical model predicts that informing people that the rate of stigmatization against HIV-positive individuals is actually lower than they think can lead them to be more willing to get an HIV test.

2. HIV/AIDS Information: Factual information about HIV/AIDS, delivered through a short video presented on a computer tablet, in a language chosen by the respondent. The video covers the negative health consequences of leaving HIV infection untreated, how HIV infection transmits, and how infected people may look and feel normal before the infection develops into AIDS.

3. Antiretroviral Therapy (ART) Information: Factual information about ART, delivered through a short video presented on a computer tablet, in a language chosen by the respondent. The video stresses that HIV infection is no longer a death sentence because free ART treatment is available and effective in helping people stay healthy and preventing transmission. The point that ART helps reduce transmission is potentially important, because this fact is often not widely known (Kaler et al., 2016). It explains that taking a test opens the door to access to ART for infected people.

4. Both HIV/AIDS and ART Information: The combination of items 2 and 3 above. Respondents assigned to this minitreatment are shown both the HIV/AIDS and ART Information videos, in that order.

5. High Incentive for HIV Testing: Each HIV testing coupon offered to the household provides a financial incentive of 100 MZN or PPP $4.85, instead of the 50 MZN (PPP $2.42) coupons offered to all other households. This minitreatment helps scale the size of other minitreatment effects with respect to variation in financial incentives.

6. Control: None of the above minitreatments.

The minitreatments were randomly assigned on the computer of one of the co-authors one time, with no re-randomization. The randomization was stratified by unique combinations of community, FCC-enrolled status, and baseline asset level. Fig. 1 presents the full cross-cutting set of treatments, indicating the number of households per cell.

4. Hypotheses

4.1. Study registration, pre-analysis plan, and pre-results review

We registered this study on the AEA RCT Registry (ID AEARCTR-0003990) on March 8, 2019 (https://doi.org/10.1257/rct.3990-5.0). On that date, we uploaded our first pre-analysis plan (PAP) to the registry. This date was prior to the endline survey and HIV testing coupon redemption, which were carried out between May and November 2019.

We had previously submitted our study as a Pre-Results Review Paper to the Journal of Development Economics (JDE). The JDE refereeing process led to minor changes to our pre-specified analyses. The JDE accepted our Pre-Results Review Paper on July 22, 2019 (Yang et al., 2019). We then uploaded the JDE Pre-Results Review Paper to our AEA RCT Registry as our second (and final) PAP on July 24, 2019. Both time-stamped PAPs are available for public viewing at our AEA RCT Registry record.

20 The impacts of the anti-stigma treatment that we report in this paper differ from the results in Yu (2021) by focusing on the household level and examining interactions with the FCC-enrolled treatment. By contrast, Yu (2021) conducts analyses at the individual level and explores the anti-stigma minitreatment impacts (alone) in substantially greater depth.

31 For stratification by baseline asset levels, we grouped households into three groups: above median of a baseline asset index, below median baseline asset index, and missing data on baseline asset index.
The submission of the second PAP was two months into the seven-month period covering the endline survey and HIV testing coupon redemption. No changes to the PAP between our first and second PAP submissions were informed by any analyses of endline or HIV testing coupon redemption data. Prior to submitting the second PAP, we had only conducted data quality control checks for feedback to enumerators in the field.

The differences between the first and second PAPs are as follows. In the first PAP, we stated two final outcomes of primary interest: (1) the coupon-based HIV testing measure, and (2) school attendance of children. We classified self-reported HIV testing as an outcome of secondary interest. In response to JDE referee and editor feedback, in the second PAP, we stated only one final outcome of primary interest: the combined measure of HIV testing (equal to one if either the coupon-based or self-reported HIV testing measure was equal to one, and zero otherwise). The two components of this HIV testing measure (self-reported and coupon-based) were individually listed as of secondary interest. We also stated in the second PAP the decision rule that if treatment effects differed across the self-reported and coupon-based HIV testing outcomes, we would base conclusions on the coupon-based measure. We also relegated child school attendance to be of secondary interest.

This paper reports on all primary analyses pre-specified in our second and final PAP, as well as a selection of secondary analyses. Remaining secondary analyses are described briefly in Section 5.3 below, and more fully in our Populated PAP. All empirical analyses in this paper are conducted exactly as pre-specified.

In the remainder of this paper, “PAP” refers to the second and final PAP.

### 4.2. Primary hypotheses

HIV testing is the outcome variable of primary interest because it is a prerequisite for benefiting from the FCC program in the health domain. HIV testing opens the door to FCC interventions promoting ART treatment initiation and adherence. Our hypotheses related to HIV testing focus on household-level outcomes because the intervention components related to HIV testing are delivered at the household level (not at the individual level).

The primary question of interest in this study is: what is the impact of FCC enrollment on HIV testing? We address this question by estimating the causal effect of a household being randomly assigned to enrollment in the FCC program. In estimating this effect, all households in control communities serve as the control group.

We hypothesized in our PAP that FCC enrollment would have a positive effect on HIV testing.

Our primary outcome of interest is an indicator that at least one of a household’s HIV testing coupons has been redeemed. This is a household-level variable equal to 1 if at least one of a household’s encouragement coupons was presented at the local health clinic for the HIV testing incentive payment before the 14-day deadline, and 0 otherwise. Note that even if an individual had been tested before (early in the implementation of the FCC program in the community), it is desirable (and consistent with the country’s public health protocols) for them to be tested again later (and therefore follow our testing recommendation and redeem the coupon). Programs such as FCC aim not only to facilitate an individual’s first HIV test, but to encourage them to be open to regular, repeated testing.

In presenting this coupon-based HIV testing measure as our primary outcome of interest, we are following a decision rule pre-specified in our PAP (page 11): if treatment effects on the coupon-based HIV testing measure differ significantly from treatment effects on the self-reported HIV testing measure, we would base our conclusions on the coupon-based measure. We pre-specified this decision rule because of concerns that the self-reported testing measure may be subject to reporting biases due to experimenter demand effects (Orne, 1962; Rosenthal, 1966; Zizzo, 2010; De Quidt et al., 2018). Experimenter demand effects may lead testing to be differentially overstated in the endline survey by treated households. Because the coupon-based measure is an administrative outcome, it is immune from reporting biases. Coupons were only redeemable with written confirmation of having just had an HIV test.

While we focus on the coupon-based HIV testing measure, we also show treatment effects on the self-reported HIV testing measure. This outcome is an indicator equal to 1 if at least one household member is reported in the endline survey to have had an HIV test in the last 12 months, and 0 otherwise. In addition, we show treatment effects on the composite HIV testing measure that is equal to 1 if HIV testing (self-reported) is equal to 1 or HIV testing (coupon-based) is equal to 1, and 0 otherwise.

### 4.3. Secondary hypotheses

A number of pre-specified secondary hypotheses are of interest, in particular outcomes representing potential mechanisms. We hypothesized in the PAP that FCC-enrollment would lead to improvements in HIV-related knowledge and reductions in HIV-related stigmatizing attitudes.

In addition, we hypothesized that the minitreatments would have positive effects on HIV testing overall, and that these effects would be smaller among FCC-enrolled households (i.e., that the minitreatments and FCC enrollment would be substitutes).

### 4.4. Multiple hypothesis testing

To conduct correct statistical inference in the context of testing multiple hypotheses, we do the following. To reduce the number of hypotheses tested, following Finkelstein et al. (2010) and Almeida et al. (2014), we construct indices of HIV-related knowledge and HIV-related stigmatizing attitudes. Within sets of related coefficients, we report p-values adjusted for the familywise error rate on each coefficient, following the List et al. (2019) method, modified to allow inclusion of control variables by Barsbasi et al. (2020).

We pre-specified our multiple hypothesis test (MHT) adjustments incompletely (and with some errors) in the PAP. We describe here...
the MHT adjustments we carry out in each table, making clear which adjustments were pre-specified, which were not, and how we have rectified pre-specification errors. We follow the PAP whenever possible, and otherwise have sought to remain true to the spirit of the pre-specified MHT adjustments.

- Table A.3. As pre-specified, we report MHT-adjusted p-values within the set of the three coefficients on Treatment status in Columns 1–3.
- Table 1. As pre-specified: (1) we adjust p-values within the set of two coefficients on Treatment in column 1 (coupon-based HIV testing measure) and 3 (self-reported HIV testing measure), and (2) we do not adjust the p-value on Treatment in column 4 (combined HIV testing measure), since the outcome in that regression combines information from the outcomes in columns 1 and 3. We pre-specified this MHT adjustment assuming the primary outcome of interest would be the combined HIV testing measure. We did not pre-specify what MHT correction we would apply if we followed the pre-specified decision rule that leads us to prioritize the coupon-based HIV testing measure over the combined measure. Now that we are in this case, a more natural approach would be to apply the MHT adjustment among the three coefficients on Treatment in columns 1, 3, and 4.30 For coefficients in Column 2, we report p-values adjusted for MHT across all five coefficients in that column.
- Tables 2 and 3. First, we reduce the number of outcomes by creating indices of overall knowledge, knowledge subindices by topic, and stigmatizing attitudes. Second, when we examine multiple outcomes (knowledge subindices and the separate stigma questions), we apply MHT adjustments within outcome families (the knowledge and stigma families separately). In the PAP, we said that we would apply MHT adjustments within one family of the 33 knowledge questions, and separately within the family of four stigma questions. Due to an oversight, we did not pre-specify creation of indices. Analyses of the indices should therefore be taken as exploratory, but we note that analysis of such indices is a widely-used approach to addressing MHT concerns (Finkelstein et al. (2010) and Almeida et al. (2014)).

Table 2. We apply no MHT adjustment to the coefficient p-value in Column 1; the overall index incorporates information from all knowledge questions, so the single coefficient on Treatment in this regression reveals impacts on overall knowledge. We apply MHT adjustments within the set of five Treatment coefficients in Columns 2–6.

Table 3. MHT adjustments in this table are analogous to those we apply in Table 2: we do not adjust the coefficient p-value in Column 1, and we adjust coefficient p-values within the group of four coefficients in Columns 2–5.

Table 4. The MHT adjustment we pre-specified in our PAP was incomplete: when stating the set of coefficients we would consider as a group when adjusting p-values, we listed just three out of five of the minitreatment coefficients (and interaction terms),31 and we neglected to include the coefficient on the main effect of Treatment in the set of coefficients listed. We now conservatively adjust p-values within a larger set of coefficients than pre-specified: for Column 1, we apply MHT adjustment to p-values within the group of all six coefficients presented; for Column 2, we do the same within the group of all 11 coefficients presented.

5. Empirical analyses

We estimate impacts of Treatment (FCC-enrolled) status using ordinary-least-squares regression analyses, with the following regression equation:

\[ Y_{ij} = \alpha + \beta \text{Treatment}_{ij} + \lambda \text{FCCambient}_{ij} + \gamma_s + \epsilon_{ij} \]  (8)

\( Y_{ij} \) is the post-treatment outcome for individual or household i in community j in stratification cell (matched pair) s. \( \text{Treatment}_{ij} \) is the indicator that community j was randomly assigned as an FCC community, and that household i was randomly assigned to FCC-enrolled status in that community (1 if so, and 0 otherwise). \( \text{FCCambient}_{ij} \) is the indicator for a household being in a treatment community but not randomly assigned to FCC-enrolled status (1 if FCC-ambient, and 0 if not). (Both \( \text{Treatment}_{ij} \) and \( \text{FCCambient}_{ij} \) are equal to zero for anyone in a control community. In other words, \( \text{Treatment}_{ij} \) and \( \text{FCCambient}_{ij} \) partition households in treatment communities into two mutually exclusive subgroups.) \( \gamma_s \) is a fixed effect for stratification cell s.\(^{32} \epsilon_{ij} \) is a mean-zero error term. We cluster standard errors at the level of 76 communities (Moulton, 1986). For each coefficient we report p-values after making corrections for multiple hypothesis testing (MHT) (as described in Section 4.4).

The coefficient \( \beta \) is the primary treatment effect of interest. It is the intent to treat (ITT) effect of assignment to FCC-enrolled status. The coefficient \( \lambda \) is the corresponding effect of being in an FCC treatment community but not being assigned to FCC-enrolled status; as pre-specified, this is of secondary interest. Random assignment allows these coefficients to be interpreted as causal effects. In this paper we focus on the primary effect of FCC enrollment, the coefficient \( \beta \) on \( \text{Treatment}_{ij} \).

5.1. Treatment effect estimates

We first discuss balance with respect to treatment (including whether attrition is related to treatment), as well as intervention fidelity. Then we turn to treatment effect estimates.

5.1.1. Balance and attrition

In Appendix Section H, we test and discuss whether there is any relationship between our randomized treatments, on the one hand, and household characteristics at the time of study enrollment, in-migration, and endline survey attrition, on the other. Balance tests are important because, as mentioned in Section 3.4, enrollment of households into the study sample occurred slightly after the start of FCC program activities in communities (Randomization Stage 1).

In Appendix Table A.5, we find no imbalance of household characteristics at study enrollment, in-migration, or attrition with respect to our primary randomized treatment of interest, assignment to Treatment (FCC-enrolled) status. In Appendix Tables A.6 and A.7, we also show analogous tests for balance and differential migration with respect to the Randomization Stage 3 minitreatments. These regressions have large numbers of coefficients, so random variation would lead some coefficients to be statistically significant by chance. We also find no indication of imbalance in these analyses: the share of statistically significant coefficients is very similar to what would be expected to occur by chance.

\(^{32}\) The inclusion of the stratification cell fixed effects reduces standard errors by absorbing residual variation. Stratification is at the level of 38 matched pairs of communities within which treatment status was randomly assigned (so stratification cell fixed effects are equivalent to matched pair fixed effects).

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30 Such an adjustment would lead to even larger p-values, strengthening the conclusion of null Treatment (FCC-enrolled) effects in these regressions.

31 Column 2 estimates Eq. (9), which was not pre-specified in our PAP; it is a simplified version of pre-specified Eq. (11) below, and simply highlights Comparison A, the pure effect of the FCC program. The MHT adjustments we apply to the coefficient p-values in Column 2 (Eq. (9)), Table 1 are analogous to the MHT adjustments we apply to the coefficient p-values in Column 2 (Eq. (11)), Table 4.

32 Specifically, we failed to edit this part of the 2nd (final) PAP after adding two additional minitreatments between the 1st and 2nd PAPs.
### Table 1
HIV testing.

<table>
<thead>
<tr>
<th>Variables</th>
<th>(1) Coupon redemption for HIV testing</th>
<th>(2) Coupon redemption for HIV testing</th>
<th>(3) Self-reported HIV testing</th>
<th>(4) Combined HIV testing measure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treatment</td>
<td>−0.0212 (0.0182) [0.371]</td>
<td>−0.105*** (0.0386) [0.337]</td>
<td>0.0234 (0.0233) [0.420]</td>
<td>0.0222 (0.0193) [0.253]</td>
</tr>
<tr>
<td>Any minitreatment</td>
<td>−0.0256 (0.0259) [0.337]</td>
<td>0.103** (0.0403) [0.017]</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Observations: 3,658, 3,658, 3,489, 3,658
R-squared: 0.058, 0.060, 0.033, 0.031
Obs level: Household, Household, Household, Household
Control Mean Dep. Var.: 0.263, 0.263, 0.652, 0.721

Notes: Dependent variables are as follows. Columns 1–2: indicator equal to one if someone in the household got an HIV test at a local health clinic (based on redemption of encouragement coupon for HIV testing), and zero otherwise. Column 3: indicator equal to one if someone in the household self-reported in the endline survey having gotten an HIV test in last 12 months, and zero otherwise. Column 4: indicator that either the coupon-based or self-reported HIV testing measure is equal to one, and zero otherwise. “Treatment” defined in Table A.3. “Any Minitreatment” is indicator equal to one if the household was assigned to any minitreatment after the endline survey in Randomization Stage 3, and zero otherwise. See Section 5.2 for definition of minitreatments. All regressions control for indicator for “FCC ambient” status and matched pair fixed effects. Regression in Column 2 also includes an interaction term between “Any Minitreatment” and indicator for “FCC ambient” status. Standard errors clustered at the community level in parentheses. 𝑃-values adjusted for multiple hypothesis testing in square brackets.

### Table 2
HIV-related knowledge.

<table>
<thead>
<tr>
<th>Variables</th>
<th>(1) HIV knowledge index</th>
<th>(2) General HIV knowledge index</th>
<th>(3) Correct methods of transmission index</th>
<th>(4) Transmission myth index</th>
<th>(5) Protection methods index</th>
<th>(6) Knowledge about HIV treatment index</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treatment</td>
<td>−0.00598 (0.00828) [0.472]</td>
<td>−0.00602 (0.00961) [0.616]</td>
<td>−0.00788 (0.0134) [0.653]</td>
<td>−0.0302** (0.0148) [0.132]</td>
<td>0.000199 (0.00906) [0.859]</td>
<td>0.00234 (0.00989) [0.846]</td>
</tr>
</tbody>
</table>

Observations: 3,940, 3,940, 3,940, 3,940, 3,940
R-squared: 0.062, 0.052, 0.039, 0.071, 0.051
Obs level: Adult, Adult, Adult, Adult, Adult
Control Mean Dep. Var.: 0.263, 0.062, 0.831, 0.747, 0.823

Notes: Dependent variables are as follows. Column 1: fraction of all 33 HIV knowledge questions answered correctly. Columns 2–6: fraction of subsets of HIV knowledge questions answered correctly. For the full list of knowledge questions and groupings by subcategory, see Appendix Section D. “Treatment” defined in Table A.3. All regressions control for indicator for “FCC ambient” status and matched pair fixed effects. Standard errors clustered at the community level in parentheses. 𝑃-values adjusted for multiple hypothesis testing in square brackets.

### Table 3
HIV-related stigmatizing attitudes.

<table>
<thead>
<tr>
<th>Variables</th>
<th>(1) HIV stigma attitude index</th>
<th>(2) Buy groceries from infected person</th>
<th>(3) Not keep infected family member a secret</th>
<th>(4) Care for infected family member in own home</th>
<th>(5) Infected teacher should be allowed to teach</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treatment</td>
<td>−0.0135*** (0.00505) [0.009]</td>
<td>−0.0139 (0.00991) [0.272]</td>
<td>−0.0281 (0.0196) [0.288]</td>
<td>−0.00506 (0.00313) [0.176]</td>
<td>−0.00330 (0.00657) [0.708]</td>
</tr>
</tbody>
</table>

Observations: 3,820, 3,756, 3,777, 3,801, 3,748
R-squared: 0.025, 0.039, 0.048, 0.017, 0.028
Obs level: Adult, Adult, Adult, Adult, Adult
Control Mean Dep. Var.: 0.746, 0.858, 0.168, 0.993, 0.965

Notes: Dependent variables are as follows. Column 1: fraction of four questions on HIV-related stigma answered in a non-stigmatizing way. Columns 2–5: for each separate question on HIV-related stigmatizing attitudes, indicator equal to one if answered in a non-stigmatizing way, and zero otherwise. For full detail on each stigmatizing attitudes question, see Appendix Section D. “Treatment” defined in Table A.3. All regressions control for indicator for “FCC ambient” status and matched pair fixed effects. Standard errors clustered at the community level in parentheses. 𝑃-values adjusted for multiple hypothesis testing in square brackets.

### 5.1.2. Intervention fidelity

We now examine intervention fidelity — the extent to which the treatments were carried out respecting the randomized assignment. We examine intervention fidelity at three levels, reflecting the levels of randomization. Below is a summary, with detailed analyses in Appendix G.
Intervention fidelity for Randomization Stage 1, randomization of the FCC program at the community level, appears high. Reports by WEI/Bantwana to the USAID Mission in Mozambique indicate that the program was implemented in the 38 treatment communities, and not in the 38 control communities. We confirm this empirically by surveying principals of the study community schools, which were focal points for delivery of many FCC program components. There is a large positive treatment effect of being in an FCC treatment community on principal reports that they have been visited by the FCC local implementing partner (LIP) organization, and that they are receiving financial support from the LIP.

For Randomization Stage 2, randomization of study households into FCC-enrolled status, both WEI/Bantwana administrative data and our household surveys provide positive indications of implementation fidelity. There is, however, a discrepancy regarding magnitudes. WEI/Bantwana reports that 77.0% of households assigned to FCC enrollment received a home visit by a LIP case care worker (CCW). In our survey data, we find that FCC enrollment leads to a doubling of the share reporting having been visited by the LIP from 5.6% to 12.1%. While we expected a positive effect of FCC-enrolled status on LIP household visits, there is a large difference – 12.1% vs. 77.0% – between the share of FCC-enrolled households reporting such visits in survey data compared to the WEI/Bantwana data. This discrepancy likely reflects substantial under-reporting by households of their contacts with LIPs. Households appear to imperfectly know or recall the specific NGOs with which they interact. Relatedly, household members responding to our survey may differ from the household members who interacted with the CCWs in household visits. Directly indicating such under-reporting, households report higher rates of services received (that would have been provided by LIPs), compared to rates of contact with LIPs. The nature of our survey data does not allow us to estimate more precisely the share of FCC-enrolled households actually visited by LIP CCWs.45

Finally, for Randomization Stage 3 (the minitreatments), implementation fidelity is high. These treatments were implemented by our research staff. Prompts to implement the correct treatment appeared on our staff members’ computer tablets, and the survey digital metadata on the timing of survey sections indicate high adherence to treatment assignment. Our survey data also show that the minitreatments are followed by improvements in knowledge and reductions in stigmatizing attitudes.

### 5.1.3. Impacts on HIV testing

We now test the primary hypotheses of impacts on HIV testing. Results are presented in Table 1. The coefficient on the pre-specified primary outcome of interest, the coupon-based HIV testing measure (Column 1), is negative but not statistically significantly different from zero at conventional levels. The point estimate indicates a 2.12 percentage point decline in testing rates, relative to the 26.3% rate in control communities. This treatment effect is based on Comparison B in Fig. 1, comparing testing rates of all FCC-enrolled households to all households in control communities.

That estimate is not the “pure” effect of FCC enrollment, because many FCC-enrolled households were assigned the Randomization Stage 3 minitreatments before being offered the HIV testing coupons. The treatment effect in Column 1 is an average treatment effect, pooling households who received minitreatments with those who did not. If there is any interaction between the minitreatments and FCC enrollment, the estimate in Column 1 will not be identical to the pure effect of FCC enrollment.

The pure effect of FCC enrollment is given by Comparison A in Fig. 1, the testing rate of FCC-enrolled households in treatment communities who did not receive any minitreatment, compared to the testing rate of control community households who also did not receive any minitreatment. While we pre-specified that we would estimate Comparison A in the context of analyzing the Randomization Stage 3 minitreatments (in Section 5.2 below, Table 4), we show a simplified version of it here as well in Table 1, to emphasize the contrast between Comparisons A and B.

We estimate the pure effect of FCC enrollment (Comparison A) using the following modification of Eq. (8):

\[ Y_{ij} = \alpha + \beta \text{Treatment}_{ij} + \gamma \text{AnyMinitreatment}_{ij} + \delta \text{FCCambient}_{ij} + \pi \text{AnyMinitreatment}_{ij} \times \text{FCCambient}_{ij} \]

AnyMinitreatment_{ij} is an indicator equal to 1 if a household received any of the minitreatments, and 0 otherwise. Eq. (9) includes this as a main effect as well as in interaction with Treatment_{ij} and FCCambient_{ij}.

We present coefficient estimates from this regression in Column 2.
These coefficient estimates in Column 2 indicate that FCC enrollment by itself (the negative coefficient $\beta$) has a large negative effect on HIV testing. Receiving some minitreatment offsets this negative effect of FCC enrollment (the positive coefficient $\theta$). $\beta$ and $\theta$ are about the same magnitude, indicating that receiving any minitreatment approximately counteracts the negative effect of FCC enrollment.

Which minitreatment(s) in particular is (are) having this positive counteracting effect? We defer this to Section 5.2 when we examine the separate minitreatments in detail.

To conclude the discussion of Table 1, we proceed to Columns 3 and 4. In Column 3 we estimate Eq. (8) where the outcome variable is self-reported HIV testing from the endline survey. The coefficient is positive, small in magnitude, and not statistically significantly different from zero. We can compare this treatment effect estimate to expert predictions elicited in advance. Prior to our results being known, DellaVigna et al. (2020) collected from subject-matter experts their forecasts of the treatment effect of being FCC-enrolled on self-reported HIV testing. The mean expert prediction was 11.36 percentage points. The actual treatment effect, 2.34 percentage points, is statistically significantly below the expert prediction ($p$-value $< 0.001$).

The coefficient on Treatment$_{ij}$ in Column 1 is more negative than the coefficient in Column 3, and the difference between them is marginally significant: the $p$-value of the F-test of equality of the coefficients is 0.1480. Even more striking, the coefficients on Treatment$_{ij}$ in columns 2 and 3 are statistically significantly different from one another at the 1% level ($p$-value 0.0043). These statistical tests lead us to follow our pre-specified PAP decision rule to base substantive conclusions on the coupon-based HIV testing measure, rather than the self-reported HIV testing measure.

For completeness, in Column 4 we show the coefficient on Treatment$_{ij}$ from estimation of Eq. (8) when the outcome variable is the composite HIV testing measure. The coefficient is positive, but modest in size and not statistically significantly different from zero.

In sum, FCC enrollment leads to lower rates of HIV testing. To explore the mechanisms through which this negative effect operates, we now turn to analyses of endline survey outcomes in Section 5.1.4, and detailed analyses of the minitreatments in Section 5.2.

### 5.1.4. Potential mechanisms behind negative impacts on HIV testing

We now shed light on mechanisms behind the negative impact of FCC enrollment on HIV testing. We focus on two key mechanisms we pre-specified: information on HIV/AIDS and HIV-related stigmatizing attitudes. As emphasized in the theoretical model, worsened misinformation could lead to heightened stigma and thereby lower HIV testing. The 33 information and four stigma questions are detailed in Appendix Section D.

In Table 2 we estimate Eq. (8) using individual-level data, examining impacts of treatment on knowledge regarding HIV/AIDS. These regressions implement Comparison B in Fig. 1, comparing outcomes of all FCC-enrolled households to all households in control communities. This is the appropriate comparison because outcomes of interest in the table are measured in the endline survey, and therefore cannot be affected by the minitreatments (which happened after the endline survey).

In Column 1, we examine impacts on an overall HIV knowledge index (the share of questions answered correctly). In Columns 2 to 6, we examine analogous knowledge sub-indices by topic: correct methods of HIV transmission, myths about HIV transmission, methods to protect against HIV, and treatments for HIV. Each index is defined so that higher values represent improvements in knowledge.

Treatment status has no large impact on the overall index or any of the subindices, except for the outcome in Column 4, the “transmission myth index”. These are questions about whether HIV can be transmitted in certain ways, all of which are not transmission channels (in other words, correct answers to these questions are all “no”): mosquito bites, shaking hands, kissing, sharing food, or witchcraft. Strikingly, the impact is negative, indicating that knowledge on this front worsens: respondents are more likely to believe myths about HIV transmission. This negative coefficient on treatment is marginally statistically significant ($p$-value $0.132$).

The other key mechanism through which HIV testing might be influenced by the FCC program is HIV-related stigmatizing attitudes. We examine impacts on responses to four yes/no questions from the AIDS Indicator Survey of the Demographic and Health Surveys (DHS), which have been fielded in Mozambique and other DHS countries since 2003 (INS, 2017). We examine treatment effects on an index of the four questions (Column 1), as well as on responses to each question separately (Columns 2–5). Each outcome variable is defined such that higher values indicate less stigma (Column 1 is the share of non-stigmatizing responses, and the remaining columns are indicators for giving a non-stigmatizing response.)

We report regression results in Table 3. Column 1 indicates that treatment worsens stigmatizing attitudes: the coefficient is negative and statistically significant ($p$-value $0.009$). Treatment reduces the share of non-stigmatizing responses by 1.35 percentage points. The standard deviation of this outcome variable is 14.63 percentage points, so this treatment effect is equal to 0.092 standard deviations — not an insubstantial magnitude.

Results in the other columns of the table indicate that treatment effects on each stigma question individually are also negative, but none are individually statistically significantly different from zero at conventional levels.

### 5.2. Minitreatments

The results presented so far are consistent with the theoretical model presented above. Increased misinformation about transmission may lead people to believe the probability of transmission is higher, leading to higher stigma against people with HIV, and thus to lower HIV testing rates. However, the above results do not prove conclusively that increased misinformation and higher stigma are mechanisms or intermediate links in the causal chain linking the FCC program with testing.

The minitreatments help reveal potential mechanisms behind the negative effects of FCC enrollment more directly. If FCC-enrollment inadvertently caused misinformation about HIV to rise, leading to increases in HIV-related stigma, and as a result depressing HIV testing, then interventions providing correct HIV information and alleviating stigma concerns should raise HIV testing differentially among FCC-enrolled households.

We first examine the effect of the minitreatments by estimating the following modification of Eq. (8):

$$
Y_{ij} = \alpha + \beta \text{Treatment}_{ij} + \lambda FC\text{Cambience}_{ij} + \delta M_{ij} + y_i + \epsilon_{ij}. \quad (10)
$$

$Y_{ij}$ is the coupon-based HIV testing measure (the only outcome available after the minitreatments). $M_{ij}$ is a vector of indicator variables for each of the five minitreatments. $\delta$ is the vector of coefficients
representing the intent to treat (ITT) effects of household assignment to the corresponding minitreatment.

Analyses of the minitreatments’ effects on the FCC-enrolled treatment effect are conducted using the following regression equation, which is a modification of Eq. (10):

\[
\gamma_{ij} = a + \beta_{\text{Treatment}_{ij}} + \alpha_{\text{FCC Ambient}_{ij}} + \tau_{M_{ijk}}
\]

\[
+ \mathbf{x}' \mathbf{\beta} \quad \text{Treatment}_{ij} \times \mathbf{M}_{ijk} + \psi \mathbf{\delta}_{\text{FCC Ambient}_{ij}} \times \mathbf{M}_{ijk} + \gamma_i + \epsilon_{ij}. \tag{11}
\]

This regression is similar to Eq. (10), but adds interaction terms between Treatment_{ij} and each of the minitreatments, as well as interaction terms between FCC Ambient_{ij} and each of the minitreatments. These interaction terms reveal whether the effects of the minitreatments differ for FCC-enrolled and FCC-ambient households, compared to the effect in control communities. Because of the inclusion of these interaction terms, the coefficients in the vector \( \mathbf{r} \) represent the ITT effects of assignment to the respective minitreatment in control communities.

The coefficients in the vector \( \mathbf{x} \) represent the difference in the ITT effect of the respective minitreatments for FCC-enrolled households, compared to the effect of the minitreatments for households in control communities. Alternately, they represent how the respective minitreatment changes the effect of FCC-enrollment, compared to the effect of FCC-enrollment for households receiving no minitreatment. (There are analogous coefficients related to the effects for FCC-ambient households.)

Both Eqs. (10) and (11) are as described in our pre-analysis plan. Results from estimating Eqs. (10) and (11) are displayed in Table 4.

Estimation of the average effects of minitreatments in the full sample (Eq. (10), Column 1) reveals that the high-value coupon has a positive effect on HIV testing rates, 7.24 percentage points, that is statistically significantly different from zero (p-value 0.044).

Estimation of differential effects of the minitreatments across treatment groups (Eq. (11), Column 2) provides explanations for the effects found in prior results tables. The coefficient on the treatment main effect (top row of Column 2) represents the impact of treatment status for individuals who did not get any of the minitreatments. The coefficient is negative, large in magnitude (10.5 percentage points), and statistically significantly different from zero (p-value 0.032).

Coefficients on the interaction terms between treatment status and each minitreatment (coefficient rows 7–11 of Column 2) indicate how the minitreatments modify the main effect of FCC enrollment. Nearly all the interaction term coefficients are positive, and most are large in magnitude and statistically significantly different from zero at conventional levels. Providing HIV-related information, providing ART-related information, countering concerns about HIV-related stigma, and providing higher financial incentives all improve the impact of FCC-enrolled status on HIV testing. The exception to this pattern is the coefficient on the interaction term with the combined HIV and ART information treatment, which is negative, much smaller in magnitude, and not statistically significantly different from zero at conventional levels. It is possible that providing too much information to respondents reduces the effectiveness of all information provided, perhaps by causing lapses in respondents’ attention or retention.

5.3. Other pre-specified analyses

In the PAP, we pre-specified some secondary analyses not reported in this paper. The Populated Pre-Analysis Plan (Populated PAP) contains results of all pre-specified analyses, including those presented in this paper. We briefly discuss them here.

We pre-specified that the impacts of FCC-ambient status were of secondary interest. FCC-ambient households are those in treatment communities who were not randomly assigned to FCC enrollment in Randomization Stage 2. FCC-ambient households are included in all samples analyzed in this paper. All regressions include an indicator for FCC-ambient status (and corresponding interaction terms with minitreatments, as appropriate) on the right-hand-side. FCC-ambient effects are generally smaller in magnitude and less statistically significant than FCC-enrolled effects. No effects of minitreatments among FCC-ambient households are statistically significantly different from effects of minitreatments in control communities. These patterns reflect that

\footnote{This coefficient is nearly the same as the coefficient in Table 1, Column 2, discussed previously, but is not exactly identical because Eq. (11) replaces the “Any Minitreatment” indicator (and associated interaction terms) with the full set of five separate minitreatment indicators.}

\footnote{In pre-specified secondary analyses, we show that the finding that the information minitreatments offset the negative “pure” effect of the FCC program on HIV testing is robust to pooling the information minitreatments rather than examining their effects individually. We estimate a modified version of Eq. (11) in which we pool the HIV information, ART information, and combined HIV/ART information minitreatment indicators into one “Pooled HIV and ART information” minitreatment indicator. In this analysis, which we show in the Populated PAP, the coefficient on the Treatment * “Pooled HIV and ART information” interaction term is negative and statistically significantly different from zero.}

The main effects of the minitreatments in Column (2) (coefficient rows 2–6) represent impacts in control communities. Most effects are negative, small in magnitude, and not statistically significantly different from zero. The exception is the coefficient on the anti-stigma treatment, which is negative and statistically significantly different from zero (p-value 0.025) in control communities (2nd row, Column 2). This may reveal that in control communities, where people may not have HIV-related stigma at top of mind, simply raising the topic of stigma may increase its salience. In control communities, people may not pay as much attention to the actual quantitative information we provide on HIV-related stigmatizing attitudes, and only increase their worry that going for an HIV test may reveal to others that they are at high risk of having HIV in their household. This then leads the anti-stigma treatment to have a negative effect on testing in control communities.

The positive interaction term on “Treatment * Anti-Stigma” in the same column could then indicate that once concerns about HIV-related stigma had previously been raised (by FCC enrollment itself), the anti-stigma treatment can have a positive effect by revealing to people that stigmatizing attitudes are not as bad as they previously thought.

The regressions in Table 4 examine impacts on an indicator for anyone in the household getting an HIV test, and so the unit of observation is the household. Because HIV testing is an individual decision, it is also possible to conduct the analysis at the individual level. We do this in Appendix Table A.8, where the unit of observation is the individual, and the outcome variable is an indicator for the individual-level HIV testing coupon redemption. All key patterns in the household-level analyses hold in the regression for all individuals (column 1), as well as subsamples of individuals (adults, primary household respondents, children, adults aged 18–49, adults aged 49+, adult men, and adult women). Perhaps the most apparent pattern in these results is that coefficients on the interaction terms between treatment and the minitreatments are larger in magnitude for the subsamples of children and adults aged 49+, compared to adults aged 18–49.

All told, this analysis of the minitreatments provides additional support for the interpretation of our prior results. The negative effect of FCC enrollment on HIV testing likely stems from the unintended consequence that the program worsened HIV-related misinformation, leading to increases in stigmatizing attitudes and a large decline in HIV testing. From this starting point of a negative effect of FCC enrollment, the minitreatments providing correct HIV information and countering concerns about HIV-related stigma raise HIV testing among FCC-enrolled households.

ART information” minitreatment indicator. In this analysis, which we show in the Populated PAP, the coefficient on the Treatment * “Pooled HIV and ART information” interaction term is negative and statistically significantly different from zero.
FCC-ambient households had less contact with the FCC program than FCC-enrolled households. In other pre-specified analyses, we present treatment effect estimates from estimation of Eq. (8) where the dependent variables are indicators for correct answers to each of 33 HIV knowledge questions, positive responses to three questions on positive attitudes related to HIV, and “safe” responses to eight questions on sexual behavior. Among these results, one that stands out is that FCC enrollment leads to a reduction in reported number of sexual partners in the last 12 months (p-value 0.060). We view this finding as related to increased fears of transmission of HIV (due to increases in beliefs in transmission myths), as well as fears of stigma should one become infected with HIV.49

We also find no large or statistically significant effect of FCC enrollment on child school attendance, life satisfaction, or household asset ownership. We examine impacts on anti-retroviral therapy (ART) initiation and adherence among individuals who self-report being HIV positive, and also find no large or statistically significant treatment effects. Finally, we find no evidence of spillovers (via geographic or social proximity) from FCC-enrolled to FCC-ambient households within FCC communities.

6. Conclusion

We study the impacts of a widespread community health program on HIV testing in Mozambique. We exploit a multilevel randomized research design to identify causal effects. We find that the program Força à Comunidade e Crianças (FCC) had a negative effect on HIV testing rates. Analysis of rich survey data on secondary outcomes suggests that the program’s negative impacts are due to increased misinformation about HIV, and worsened HIV-related stigmatizing attitudes. This interpretation is bolstered by additional treatments we administered at the household level. These additional treatments providing correct HIV-related information and countering HIV-related stigma concerns make the treatment effects of the FCC program on HIV testing rates more positive, suggesting that the FCC program was deficient in these areas.

We provide a rare glimpse into the impacts and mechanisms of a widespread community-level program seeking to raise HIV testing. Our results point to a thus-far neglected possibility: programs seeking to raise HIV testing may fail due to deficiencies in information provision and in countering HIV-related stigma. Indeed, such programs may inadvertently worsen HIV knowledge and increase stigmatizing attitudes, leading to less voluntary HIV testing.

These results suggest priority directions for future research. A key question is how exactly the program led to increased misinformation on HIV transmission, and to worsened HIV-related stigmatizing attitudes. It is possible that providing basic information that HIV is transmitted via bodily fluids could lead people to believe in transmission myths, since some of these also involve possible sharing of bodily fluids (e.g., shaking hands, kissing, sharing food). FCC home visits could have also increased the salience of HIV and of HIV-related stigmatizing attitudes in the community, if it was known in the community that home visits by program staff (CCWs) were targeted towards households with HIV infection.49

Because we did not anticipate these negative findings, we did not collect information needed to understand how the program could have increased misinformation and worsened stigmatizing attitudes. In future studies, it will be important to collect detailed data on exactly what program staff do, what kind of information they convey, and how people react to and interpret the information. It is also important to understand how people respond to and interpret home visits, either as recipients of visits or observers of others’ visits. This could require high-frequency household surveys as the program is taking place, in combination with spot checks and direct observation of program staff. Controlled lab-in-the-field studies could also reveal mechanisms through which people come to believe incorrect information (such as transmission myths), and how they come to acquire HIV-related stigmatizing attitudes and concerns about such stigma.

A related area for future exploration is ways to improve people’s knowledge about HIV and to reduce HIV-related stigmatizing attitudes. Our own findings suggest that there are simple and cheap ways to improve knowledge and to reduce stigma, as evidenced by the effects of our minitreatments (at least among households who had suffered worsened information and heightened concerns about stigma beforehand). Future work should seek to pursue these and other related research directions.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Data will be made available on request.

Appendix A. Supplementary data

Supplementary material related to this article can be found online at https://doi.org/10.1016/j.jdeveco.2022.102958.

References

[References are not listed in the provided text.]

Some beneficial effects, but too little data, and programs spread thin. Health Aff. 31 (7), 1508–1518.


