HIV STATUS AND LABOR MARKET PARTICIPATION IN SOUTH AFRICA

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Abstract—We use econometric methods based on the propensity score to estimate the causal effect of HIV status on employment outcomes in South Africa. Relying on rich data from a national survey, which included HIV testing, we control for systematic differences between HIV-positive and HIV-negative individuals. We provide the first nationally representative estimates of the impact of HIV status on employment outcomes for southern Africa. Being HIV positive is associated with an increase of 6 to 7 percentage points in the likelihood of unemployment overall and 10 to 11 percentage points for those who are less educated. This disadvantage reinforces existing inequalities in South Africa.

I. Introduction

This paper employs an especially rich South African data set to estimate the causal impact of HIV status on an individual’s labor market outcomes. The paper contributes to a better understanding of the economic impact of HIV/AIDS and the role HIV/AIDS plays in reinforcing inequality in South Africa.

The existence of an impact of HIV on labor market participation is hotly debated in South Africa. Some argue that HIV is a severe constraint on current economic growth because those who are too ill to work represent lost economic output. Others argue that because unemployment is so high—in the 30% to 40% range according to the broad definition, which includes individuals who desired employment but had no job search activity within the past month (i.e., discouraged workers)—HIV has minimal economic impact today. Proponents of this view note that the sex-age cohorts with the highest unemployment are essentially the same as those with the highest HIV prevalence rates. Our estimates speak directly to this issue. This paper provides the first nationally representative estimates of the impact of HIV on labor market status, certainly for southern Africa but probably also more broadly for nonrich countries.

There are at least two reasons for the paucity or outright lack of evidence: stringent data requirements and econometric challenges. Obtaining estimates of the impact of HIV on labor market participation clearly requires data on both labor market participation (which is generally available) and on an individual’s HIV status (which is collected fairly rarely). Our econometric methods also require extensive data on correlates of an individual’s HIV status.

Even with our rich data set, econometric endogeneity poses challenges. In our context, endogeneity arises because an individual’s labor market status might affect his or her HIV status. For example, migrating for employment opportunities may put individuals at greater risk for HIV (reverse causality bias). Alternatively, there may be factors that affect both HIV status and labor market status, such as an individual’s age or willingness to invest in the future (selection bias, omitted variable bias).

Two of the standard approaches to addressing these issues are infeasible in this instance. The traditional solution is to apply an instrumental variables approach. For the application at hand, this would require an instrument that is correlated with the individual-level HIV status and is orthogonal to shocks to labor market status (conditional on observed covariates.) This is a tall order to fill. One class of correlates with HIV status—socioeconomic variables such as income, education, sex, age, and even household size—is likely correlated with labor market status, while another class of correlates—health-related variables such as sexual practices and knowledge about HIV—is likely to be correlated with the socioeconomic covariates and, by association, labor market status. A closely related approach uses a control function with an exclusion restriction, and the same concerns that preclude good instruments similarly preclude a convincing exclusion restriction (see Heckman & Navarro-Lozano, 2004).

The other approach to addressing sample selection, which has gained currency in development economics, is to conduct interventions such that the bias vanishes by design. In a study of HIV status and labor market participation, this would involve randomly assigning HIV status to individuals and comparing labor market status across the two groups. This is one context for which the experimental approach is impossible.

We turn to methods based on the propensity score to address the issues of selection and reverse causality. We investigate the impact of HIV status on labor market participation using two methods: propensity score reweighting and control functions. While both have been used to address the

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1 There are other data sets available for which the techniques employed in this paper are potentially applicable. These include some of the Demographic Health Surveys (DHS) and the Botswana AIDS Impact Survey (see Levinsohn & McCrary, 2008).

2 These challenges are discussed in detail in Strauss & Thomas (1998).

3 With country-level data, the fraction of the male population that has been circumcised has been used as an instrument (see Ahuja et al., 2008). At the individual level, this instrument is problematic. The protective effect may not transfer to women (Wawer et al., 2008), and there is often virtually no variation within large categories of the male population like religion, race, and ethnic or tribal affiliation (see McKelvey, 2010).

4 For example, Thomas et al. (2006) use experimental methods to obtain a causal estimate of the effect of improved health (in this case, from increased iron intake) on labor market outcomes.

5 These methods are drawn mostly from the program evaluation literature, but they have also been used in other fields of economics. For example, a similar control function approach is used in Olley & Pakes (1996) for solving the endogeneity problem in the context of production functions. Levinsohn & McCrary (2008) use the propensity score reweighting technique to address a missing data problem in the context of HIV prevalence.

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endogeneity issue in the literature, there have been few direct comparisons of multiple estimators for the same problem. We use both methods as a marginal methodological contribution and, more important, a robustness check.

Propensity score reweighting and the control function approach both rely on the assumption that sample selection into being HIV positive is random, conditional on attributes that are observable to the econometrician (selection on observables). Given the richness of the data set we use, this assumption is weaker than it might be under other circumstances. The survey questionnaire included over 175 questions, many with multiple subsections, yielding over 400 individual covariates. The data set contains detailed information on sexual practices and knowledge of HIV transmission in addition to household characteristics and other factors that might influence HIV status. We examine the validity of the assumption of selection on observables in section V by testing whether a parsimonious specification succeeds in balancing variables not used in the estimation (balancing tests) and experimenting with multiple specifications to confirm that the inclusion of additional controls does not change the results (robustness checks).

In the next section, we briefly discuss the related literature. Section III discusses the data that are employed. In section IV, we present the two approaches to estimation. Section V presents results, and section VI provides both caveats and conclusions.

II. Related Literature

While understanding the relationship between HIV status and labor market participation is important from multiple perspectives, few studies present evidence on the impact of HIV and fewer still address issues of endogeneity.

A number of studies develop macroeconomic simulation models to examine the effect of HIV/AIDS on economic growth; however, these models may be extremely sensitive to assumptions about life expectancy for HIV-positive individuals (Cuddington & Hancock, 1994; Kambou, Devarajan, & Over, 1992; Arndt & Lewis, 2000). Instead of assuming an HIV/AIDS mortality rate, Bloom and Mahal (1997) estimate the impact of AIDS on growth by exploiting cross-country variation in HIV prevalence rates. They find that HIV has an insignificant effect on per capita GDP growth for their sample of 51 countries.

The HIV pandemic not only affects current economic growth but can have lasting effects on growth rates into the future. Kalemli-Ozcan (2012) and Bell, Devarajan, and Gersbach (2006) model the impact of current HIV prevalence rates on future economic growth. Because HIV-positive parents are likely to die before reaching old age, they may invest less in their children’s human capital acquisition, lowering the stock of human capital and contributing to lower growth rates in the future. But, if the AIDS epidemic causes a reduction in fertility that dominates this human capital effect, higher future living standards may result (Young, 2005).

Most papers that examine the impact of HIV on economic growth look exclusively at the effect of AIDS mortality, overlooking the effects of the illness on employment and productivity. There are some exceptions. Murray et al. (2005) find that the rate of minor work-related injuries was 30% higher for HIV-positive miners in South Africa than for HIV-negative miners. The correlation was observable within one year of seroconversion, which suggests that the acute initial infection or the psychological shock has an effect long before AIDS symptoms appear. This is likely an underestimate of the impact of HIV because miners who are most affected are more likely to take on easier tasks at work or leave employment altogether.

Habyarimana, Mbakile, and Pop-Eleches (2010) use the antiretroviral (ARV) therapy inception date as an instrument for health status to examine the effect of health on productivity for mine workers in Botswana. They document an inverse-V-shaped pattern of absenteeism for HIV-positive workers in the two years around the inception of ARV therapy. Workers who subsequently enrolled in ARV therapy missed about five times as many days of work as nonenrolled workers in the year prior to ARV therapy inception, but absenteeism rates returned to prepeak levels after a year of therapy. Nonenrolled workers appear to be a valid control group because the two groups had similar levels of absenteeism from five years to one year prior to therapy inception. These findings suggest that health can have large effects on employment outcomes and that ARV therapy is effective in reducing the disparity in productivity between HIV+ and other workers.6 Fox et al. (2004) examine differences in on-the-job productivity between workers who subsequently died of AIDS and other workers on tea plantations in Kenya. They find that in the year before death, AIDS victims are less productive (in this case, measured by the quantity of tea leaves picked), are more likely to be reassigned to less strenuous but less lucrative tasks, and are more often absent from work.

One way to estimate the causal effect of HIV status on employment is to use a plausibly exogenous instrument for HIV status. Variation in circumcision has been used to instrument for HIV status because circumcision has been found to be associated with a reduced risk of HIV in both regression analyses (Weiss, Quigley, & Hayes, 2000) and randomized controlled trials (Auvert et al., 2005). Ahuja, Wendell, and Werker (2006) find that HIV/AIDS did not have a measurable effect on economic growth, savings, or fertility behavior in African countries but that there was weak evidence that HIV/AIDS reduced youth literacy and increased malnutrition. McKelvey (2010) finds that across nine African countries and Haiti, HIV-positive individuals are significantly less likely to have been employed or to have earned enough to contribute more than half of household expenditures. However, specification checks using two populations that would not be expected to benefit from circumcision (men under 20

6 Thirumurthy, Zivin, and Goldstein (2008) also found that in western Kenya, labor supply increased within six months of initiating ARV therapy.
years old and those who have never had sex) suggest that unobservable differences that vary with circumcision may be driving the results.

Transitory economic shocks can have short-term effects on an individual’s propensity to engage in risky behavior but potentially long-term effects on health. This is one avenue through which employment outcomes may have an impact on HIV status and one reason that reverse causality is a salient issue. Women who are economically vulnerable may become active with multiple sexual partners (Dinkelman, Lam, & Leibbrandt, 2007) or turn to sex-for-gift exchanges to smooth consumption (Dunkle et al., 2004; LeClerc-Madlala, 2002). Migration is another potential response to transitory shocks that can put both men and women at greater risk for HIV (Zuma et al., 2005).

III. Data

Our data come from the nationally representative South African National HIV Prevalence, HIV Incidence, Behaviour and Communication Survey (SABSSM II) conducted in 2005 by the Human Sciences Research Council (HSRC), the Centre for AIDS Development, Research and Evaluation (CADRE), and the Medical Research Council.7 The survey asked adult respondents questions about demographics, knowledge of HIV, sexual history, knowledge of voluntary counseling and testing (VCT) services, health, mental health, and drug and alcohol use. It also included a household module that asked for basic demographic data for all household members, in addition to questions about household infrastructure and participation in government programs. Although no questions directly addressed income or expenditures, the individual survey did query labor market participation. Respondents were asked to classify their “present employment situation” as one of the following thirteen categories: homemaker not looking for work, homemaker looking for work, unemployed not looking for work, unemployed looking for work, informal sector not looking for permanent work, old age pensioner, sick/disabled and unable to work, student/pupil/learner, selfemployed full time (40 or more hours per week), self-employed part time (less than 40 hours per week), and employed part time (if none of the above), and employed full time or other.

The sample consisted of 23,275 individuals in 10,584 households.8 Not everyone in a household was sampled. Using the roster of all household members (including those who usually live in the household, whether or not they were present at the first interviewer visit, and any guests who stayed in the household the previous night), field workers randomly selected at most one person from each of three age groups (2–14 years, 15–24 years, and 25 and up) to be interviewed. While this sampling strategy is an efficient way to obtain a measure of HIV prevalence, it is problematic for any analysis of household dynamics. For example, we are unable to directly measure the impact of having an HIV-positive spouse on adult labor supply since only one household member above the age of 25 was surveyed.

The survey also included an opt-out HIV test for respondents age 2 and older. The response rate for testing was 65.4% overall and 73.3% in the adult sample (age 15 and older) used for analysis (see table 1). HIV incidence data were collected from all HIV-positive specimens using an enzyme immunoassay that measures the ratio of HIV antibodies to other antibodies to determine the elapsed time since HIV infection.

Table 2 presents HIV prevalence rates by race. While HIV affects all races, Africans have by far the highest prevalence rates. In our sample, 17.34% of Africans are HIV-positive, while the population group with the next highest prevalence rate, Coloured, has a rate of 2.73%.9 For the remainder of this paper, we restrict our analysis to Africans, who comprise about 80% of the population. The HIV prevalence rate, the employment rate, the age profile of

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8 When matching individuals to households based on household identification numbers, approximately 14% of the individual observations (3,413) could not be matched to household data. Household variables were imputed to 0 for these unmatched observations. A specification check suggests that the data were consistent with a pattern of “missing at random”; a dummy variable for being unmatched was not significant in any specification. As with all variables that are imputed, an indicator variable was created that took a value of 1 if the value was imputed and 0 otherwise. These indicator variables are included in the empirical specifications. Individual survey records were matched to HIV incidence and viral load data using the bar code number that identified each specimen.
9 The prevalence rates in table 2 do not correct for sample selection due to nonrandom opt-out of the HIV test. We have replicated our analysis on the impact of HIV on labor market participation with weights that do correct for nonrandom opt-out (as computed by HSRC), and the results are virtually identical. This suggests that selection into testing is independent of the effect of HIV on employment. These rates apply to our sample, which includes only individuals aged 15 and over. Hence they differ from the national prevalence rates reported in Shisana et al. (2005).
employment, and the education profile of employment are each markedly different for Africans.\footnote{A version of this paper that includes all races in all the results is available on Levinsohn’s website: http://levinsohn.commons.yale.edu.}

Figure 1 presents the age profile of HIV prevalence—a profile broadly comparable to those of other African countries. HIV prevalence peaks between ages 25 and 30 for women and between ages 35 and 40 for men. The age profile for women is about 5 percentage points lower than UNAIDS antenatal clinic statistics, which may be due to differences in the sampling frame (UNAIDS, 2005).

Table 3 reports employment status by HIV testing status.\footnotetext{11 Those who responded being in the “other” category are grouped with the employed to generate conservative estimates.}

IV. Methodologies

A. Overview

Generating a plausible counterfactual is the core challenge of identifying a causal effect of HIV. For the situation at hand, this approach entails creating a counterfactual in which one could compare individuals who were virtually identical except for their HIV status and then compare differences in labor market status. We present two methods, each based on the propensity score, to generate unbiased estimates of the effect of being HIV positive on labor market status.

Our “treatment” (as the public health and program evaluation literatures refer to it) is HIV status, denoted by \( D \), where \( D = 1 \) is HIV positive and \( D = 0 \) is HIV negative. If HIV status were independent of the untreated value of our outcome of interest, labor market status (denoted \( Y_0 \)), then there is no sample selection problem and we can use a simple naive estimator, a simple difference in means, to estimate the treatment effect. However, HIV status is not randomly assigned and is in all likelihood related to our outcomes of interest. We make the conditional independence assumption (CIA) that the untreated value of labor market status is independent of HIV status conditional on a vector of covariates (denoted \( X \)):

\[
Y_0 \perp D|X. \quad (1)
\]

B. The Propensity Score

In this context, the propensity score, denoted \( P(HIV+)|X \equiv p(X) \), is the probability that an individual is HIV positive conditional on a vector of observable exogenous covariates \( X \). These covariates must be exogenous to HIV status in that they cannot be affected by HIV status (i.e., it would be inappropriate to use sexual behavior that may be changed if HIV status is known).\footnote{It is important to note that these are not the same variables we would use as instruments in an instrumental variables (IV) regression. In that case, we would want instruments correlated with HIV status but not with labor market status; in our case, these instruments should not be included in the propensity score regression. Because actual HIV status is known by the econometrician and can therefore be controlled for in the regression, the covariates in \( X \) do not need to be good predictors of HIV status. In fact, selecting the attributes for \( X \) depending on predictive power can increase bias. See Heckman and Navarro-Loranzo (2004).} Rosenbaum & Rubin (1983) show that

\[
p(X) = \frac{e^x}{1 + e^x} \quad (2)
\]

The propensity score is estimated by logistic regression:

\[
\text{logit}(p) = \beta'X \quad (3)
\]

The logistic or logit function maps the linear predictor \( \beta'X \) to the probability \( p \) of being HIV positive. The estimated propensity score, \( \hat{p} \), is then the predicted probability of being HIV positive based on the estimated coefficients, \( \hat{\beta} \). The first stage of the two-stage least squares (2SLS) regression is then a regression of labor market status on the estimated propensity score:

\[
Y = \alpha + \beta \hat{p} + \gamma X + \epsilon \quad (4)
\]

where \( Y \) is the labor market status, \( \alpha \) is the intercept term, \( \beta \) is the estimated coefficient on the propensity score, \( \gamma \) is the coefficient on the covariates, and \( \epsilon \) is the error term.

The second stage of the 2SLS regression is then a regression of the estimated propensity score on the covariates:

\[
\hat{p} = \gamma'X + \mu \quad (5)
\]

where \( \gamma' \) is the estimated coefficient vector on the covariates and \( \mu \) is the error term.

The estimated coefficients on the covariates \( \hat{\gamma} \) from the second stage regression are then used to generate unbiased estimates of the effect of HIV positive status on labor market status.

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\]

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The estimated coefficients on the covariates \( \hat{\gamma} \) from the second stage regression are then used to generate unbiased estimates of the effect of HIV positive status on labor market status.
if the CIA is satisfied by conditioning on the vector \( X \), it is satisfied by conditioning on the propensity score (i.e., the propensity score is a sufficient statistic for the \( X \) vector in the CIA). Hence, the propensity score makes the problem of finding a comparable control group tractable by reducing the dimensionality of the comparison while still satisfying the CIA.

### C. Propensity Score Reweighting

Propensity score reweighting uses the propensity score to create a counterfactual distribution of \( X \) in the HIV– (control) population so as to match the distribution of \( X \) in the HIV-positive population. Essentially HIV-negative observations with \( X \) characteristics that are most like HIV-positive observations (i.e., that have a high \( \hat{p}(X) \)) receive the most weight, whereas HIV-negative observations that are very different from the HIV-positive population receive less weight.

More formally, Dehejia & Wahba (1997) and DiNardo, Fortin, and Lemieux (1996) show that

\[
\Delta^{ATET} = \frac{1}{N_F} \sum_{i=1}^{N_F+N_C} \left( D_i Y_i - (1 - D_i) Y_i \right) \left( \frac{p(X_i)}{1 - p(X_i)} \right) \tag{2}
\]

is a consistent estimator for the average treatment effect on the treated (ATET), provided the CIA and the common support condition hold. In the calculation, HIV-positive observations receive a weight of 1 (the first term in parentheses) and HIV-negative observations are weighted with \( \omega_i = \hat{p}(X_i)/(1 - \hat{p}(X_i)) \) (the second term in parentheses). This estimator does not impose a functional form on the relationship between HIV status and the outcome of interest (Dehejia & Wahba, 1999).

This approach requires overlap in the support of the propensity scores for the HIV-positive and HIV-negative groups (the common support condition). In section V, we present density plots to verify the validity of this assumption.\(^{13}\) Busso, DiNardo, and McCrary (2010) show that reweighting outperforms propensity score matching in settings likely to be encountered in empirical work.

### D. Control Functions

The control function approach is an alternative econometric strategy for addressing selection bias. The reason one obtains biased estimates from a simple regression of labor market status on HIV status is that the disturbance term in such a regression is correlated with HIV status (the independent variable). The essence of the control function approach is to control for the portion of the disturbance term that is correlated with HIV status. Once the portion of the disturbance term that is responsible for the correlation is expunged, the new error term is uncorrelated with HIV status, and the regression yields unbiased estimates of the impact of HIV on employment status. The control function approach was developed in Heckman & Robb (1985) and has been used to estimate the impact of training on earnings (Heckman & Hotz, 1989), returns to education (Card, 1999) and the capitalization of pollution into housing values (Chay & Greenstone, 2005), among other applications. As with propensity score reweighting, we maintain the assumption of selection on observables.\(^{14}\)

The data-generating process is given by

\[
Y = \Lambda (\beta_0 + \gamma D + f(X) + \epsilon), \tag{3}
\]

where \( f(X) \) is a function of observables \( X \).

Under the assumption of selection on observables, conditioning on \( f(X) \) results in a disturbance term, \( \epsilon \), that is independent of \( D \) (HIV status) and hence, the estimate of the parameter of interest, \( \gamma \) is unbiased. In practice, a polynomial in the estimated propensity score, as well as linear terms in \( X \), are used to flexibly model \( f(X) \). We estimate

\[
Y = \Lambda \left( \beta_0 + \gamma D + X' \phi + \sum_{i=1}^{k} \beta_i \hat{p}(X)' + \epsilon \right) \tag{4}
\]

separately for HIV-negative and HIV-positive groups and obtain predicted values, \( \hat{Y}_0 \) and \( \hat{Y}_1 \), respectively, from each regression. We calculate the ATET by averaging the difference in predicted values across HIV-positive observations.

### V. Results

Estimating the impact of HIV status on labor market participation is a three-step process. The first step is to estimate the propensity score. The second is to empirically examine the validity of the CIA and the common support condition to ensure that the appropriate observable variables are included in the propensity score regression. If the underlying assumptions hold, then one can proceed to the third step, estimating the impact of HIV on labor market participation using propensity score reweighting and a control function approach.

#### A. The Propensity Score

There are competing philosophies behind what constitutes a properly specified propensity score regression. One approach is to adopt a relatively parsimonious specification, albeit one still rich enough to plausibly satisfy the CIA, while another approach is to include most all plausible regressors. We adopt the former but experiment with the latter in sensitivity analyses.

\(^{13}\) Frölich (2004) demonstrates that an estimated propensity score close to 1 can cause problems for estimating the ATET. Our values of the estimated propensity scores are not large enough for this to be a concern.

\(^{14}\) A similar control function method can be used if there is selection on unobservables (see Heckman & Navarro-Lozano, 2004). However, allowing for selection on unobservables requires an exclusion restriction—a variable that is correlated with HIV status but uncorrelated with labor market status.
The plausibility of the selection on observables assumption clearly rests on having data that can, in our context, account for selection into HIV status. The SABSSM II data set has several hundred variables for most respondents. In addition to the usual demographic information, the survey collected extensive information on sexual practices and knowledge about HIV transmission, which is exactly the type of information needed to account for selection into HIV status. Information on either sexual practices or knowledge about HIV transmission can be misleading, so it is important to have information on both for predicting HIV status. For example, a respondent who has multiple partners but was well informed on how HIV is transmitted before his or her sexual debut and therefore practices safer sex may have a low probability of being HIV positive. Similarly, a respondent who knows very little about how HIV is transmitted but is abstinent will have a low likelihood of being HIV positive. Conditional on comprehensive information about an individual’s sexual practices and knowledge regarding HIV transmission, unobservables such as attitudes toward risk or moral beliefs may have little explanatory power. Hence, the selection on observables assumption seems especially appropriate given the specifics of our data. The selection on observables assumption is testable, and we investigate the reasonableness (or not) of this assumption by conducting balancing tests on covariates in the propensity score specification as well as plausible covariates that were not included in the specification (see section VB).

We estimate the propensity score as the predicted value of a logit regression. Table 4 reports the estimated coefficients (not marginal effects) and standard errors from our base case. The covariates include basic demographic characteristics, educational attainment, information about a respondent’s sexual debut, knowledge of HIV transmission, and whether there is a pensioner living in the household. We do not discuss the estimated coefficients because the main focus of this paper is on using the propensity score to correct for selection bias rather than on the correlates of seropositivity. Furthermore, it is quite difficult to have much intuition about the marginal impact of a single regressor conditional on the other.

### B. Examining the Validity of the Propensity Score Method

We next examine the validity of the CIA and common support assumption under our preferred (base case) propensity score specification. The former is done with balancing tests and the latter by examining empirical distributions.

The intuition behind balancing tests is appealingly clear. The idea of propensity score reweighting is to reweight the distribution of the observables ($X$’s) of the HIV-negative population so as to match the distribution of the $X$s for the HIV-positive population (see equation [2]). Balancing tests simply examine whether the difference in means of $X$s between HIV-positive and HIV-negative populations is reduced when the observations are reweighted. If the reweighted means are similar (i.e., not significantly different) for HIV-positive and HIV-negative populations, then the reweighting has achieved its goal, the data are balanced, and the CIA is appropriate.

Table 5 reports the results of balancing for variables that enter the propensity score regression (i.e., internal balancing). We also conducted balancing tests for each of the indicator variables for missing data, but these are not reported in the table. All were balanced. The results in table 5 apply to the entire sample. Balancing tests were also conducted for each of the four subsamples with comparable results. Row 1 of table 5 presents the results for age; HIV-positive individuals were on average 2.5 years younger than the HIV-negative population. This difference had a $t$-statistic of $-4.86$ so the difference was highly significant. After reweighting, the difference falls to 0.255 years and is not significantly different from 0. The last column of the table shows that one of the variables in

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coefficient</th>
<th>Coefficient (Base Case)</th>
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</thead>
<tbody>
<tr>
<td>Age</td>
<td>0.138**</td>
<td>(0.051)</td>
</tr>
<tr>
<td>Female</td>
<td>0.470***</td>
<td>(0.112)</td>
</tr>
<tr>
<td>Urban resident</td>
<td>0.305**</td>
<td>(0.127)</td>
</tr>
<tr>
<td>Never married</td>
<td>0.371***</td>
<td>(0.126)</td>
</tr>
<tr>
<td>Completed primary education</td>
<td>0.347</td>
<td>(0.216)</td>
</tr>
<tr>
<td>Completed secondary education</td>
<td>0.338*</td>
<td>(0.201)</td>
</tr>
<tr>
<td>Holds a matric qualification</td>
<td>0.096</td>
<td>(0.232)</td>
</tr>
<tr>
<td>Has some postmatric education</td>
<td>−0.514</td>
<td>(0.277)</td>
</tr>
<tr>
<td>Has had sex</td>
<td>2.454**</td>
<td>(1.066)</td>
</tr>
<tr>
<td>Age at first sex</td>
<td>−0.165*</td>
<td>(0.096)</td>
</tr>
<tr>
<td>Age at first sex squared</td>
<td>0.004*</td>
<td>(0.002)</td>
</tr>
<tr>
<td>Used a condom at first sex</td>
<td>−0.285*</td>
<td>(0.169)</td>
</tr>
<tr>
<td>Knows HIV transmitted through vaginal sex</td>
<td>−0.169</td>
<td>(0.209)</td>
</tr>
<tr>
<td>Believes HIV not transmitted through witchcraft</td>
<td>−0.139</td>
<td>(0.169)</td>
</tr>
<tr>
<td>Knows condoms prevent HIV transmission</td>
<td>−0.121</td>
<td>(0.175)</td>
</tr>
<tr>
<td>Knows reducing number of partners reduces risk</td>
<td>0.199*</td>
<td>(0.119)</td>
</tr>
<tr>
<td>Male of pension age in household</td>
<td>−0.002</td>
<td>(0.479)</td>
</tr>
<tr>
<td>Female of pension age in household</td>
<td>−0.233</td>
<td>(0.193)</td>
</tr>
<tr>
<td>Pseudo-$R^2$</td>
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<td></td>
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<tr>
<td>Observations</td>
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</tr>
</tbody>
</table>

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15 Elements of the $X$ vector of covariates were imputed to 0 for item nonresponse, and a dummy variable for imputation was included in the specifications. We include indicator variables for each province in South Africa. These coefficients are not reported in the table.

16 We also examine particular subsamples of the data (e.g., by sex, education, area of residence). For each of these subsamples, we reestimate the propensity score. These results are available on request.
the propensity score regression has a statistically significant difference between the HIV-positive and HIV-negative populations in the reweighted data.

The propensity score weight, by design, attempts to minimize differences between HIV-positive and HIV-negative groups for the variables included in the propensity score estimation (the \(X\) vector). A more stringent balancing test criterion is whether the propensity score weight also succeeds in balancing covariates that were not used in the estimation. External balancing tests were conducted on 56 variables. These external variables included measures of sexual activity and almost 50 variables that are plausibly related to economic well-being (e.g., source of water for the household, type of cooking fuel, type of toilet, and measures of privation.) In the unweighted data, 20 of these variables had means that were significantly different (at the 95% significance level) between the HIV-positive and HIV-negative populations. After reweighting, only three of those differences were still significant.

Based on internal and external balancing tests, we conclude that the propensity score specification is adequately reweighting the data to justify the CIA. While selection on observables is a strong assumption, we find that the richness of our data provides support for the assumption.

The balancing tests provide support for the CIA. The other assumption underlying the propensity score reweighting approach is the common support condition.\(^{17}\) We examine the appropriateness of the support condition by comparing the empirical distributions of the propensity scores for the HIV-positive and HIV-negative populations. These distributions are shown in figure 2. In the figure, it is clear that the densities have a common support.

\(^{17}\) The support condition is not required for the control function approach; however, confirming that it holds ensures that we are not relying solely on functional form assumptions for any values of the propensity score.

The second column presents the coefficient on HIV status when the variables in the propensity score (listed in table 4) are included as controls in a simple logit regression of employment status for the HIV-positive and HIV-negative populations, without the inclusion of any controls. Variables are defined such that the 0.079 figure in the first cell implies that, on average, HIV-positive individuals are 7.9 percentage points more likely to be unemployed.

The first column presents naive estimates of the impact of HIV on labor market participation. The table is organized such that each column presents estimates resulting from a different estimator and each row presents estimates using different samples of the data.

C. The Impact of HIV

Estimates. Table 6 presents estimates of the causal impact of HIV on labor market participation. The table is organized such that each column presents estimates resulting from a different estimator and each row presents estimates using different samples of the data.

The second column presents the coefficient on HIV status when the variables in the propensity score (listed in table 4) are included as controls in a simple logit regression of
investigate this heterogeneity in the remainder of table 6 by
heterogeneity in the impact of HIV on labor market status. We
raises the probability of unemployment by between 6 and 7
employment imply that, all else equal, being HIV positive

restricted to individuals aged 25 and older. Naive is OLS without controls. Controls used in logit are

causal impact of HIV on employment using propensity score
reweighting, while in column 4, the reported
estimate is the coefficient on HIV status (γ1,973) in the regression given by equation (4). Using the entire sample, propensity score reweighting indicates that being HIV-positive raises the probability of unemployment by 6.2 percentage points,
while the control function approach indicates an increase of 7.0 percentage points. Each of these impacts is precisely estimated.18 Our estimates of the causal impact of HIV on employment imply that, all else equal, being HIV positive raises the probability of unemployment by between 6 and 7 percentage points.

The estimates in the first row of table 6 apply to the entire sample and, as such, potentially hide substantial underlying heterogeneity in the impact of HIV on labor market status. We investigate this heterogeneity in the remainder of table 6 by

18 Both the reweighting and the control function approach use the estimated propensity score. Hirano, Imbens, and Ridder (2003) show that using the estimated propensity score rather than the true propensity score produces efficient estimates. They suggest bootstrapping to obtain standard errors. We do so.

employment status on HIV status. This is a simple regression
to run and is the same as the control function approach, but it
excludes the higher-order terms of the propensity score. The
logit result implies that, for the full sample, being HIV-positive lowers the probability of employment by 6.5 percentage
points.

The third and fourth columns present estimates of the causal impact of HIV on employment using propensity score reweighting and the control function approaches, respec-
tively. For the internal consistency of our results, both the propensity score weights and the control function are based on the propensity score, which is calculated using the set of control variables included in the logit in column 2. Each parameter estimate in columns 3 and 4 is the marginal effect of HIV status from a logit regression. For column 3, it is a simple logit regression of labor market status on HIV status using propensity score reweighting, while in column 4, the reported estimate is the coefficient on HIV status (γ) in the regression given by equation (4). Using the entire sample, propensity score reweighting indicates that being HIV-positive raises the probability of unemployment by 6.2 percentage points, while the control function approach indicates an increase of 7.0 percentage points. Each of these impacts is precisely estimated.18 Our estimates of the causal impact of HIV on employment imply that, all else equal, being HIV positive raises the probability of unemployment by between 6 and 7 percentage points.

The estimates in the first row of table 6 apply to the entire sample and, as such, potentially hide substantial underlying heterogeneity in the impact of HIV on labor market status. We investigate this heterogeneity in the remainder of table 6 by

restricting the sample to particular subpopulations. The second
to seventh rows in table 6 present results obtained from an analysis of subsamples of the data, using propensity score estimates calculated within the subsample alone. For men, propensity score reweighting and the control function approach give estimates of 0.061 and 0.080, respectively. The former is not precisely estimated, while the latter still is. For females, the point estimates from propensity score reweighting and the control function approach are 0.055 and 0.060. These are slightly lower than the estimates obtained with the entire sample.

The next two rows in table 6 restrict the analysis to respon-
dents aged 25 and older and divide the sample between individuals whose education level is a matric or higher and those whose education level is less than a matric. Respondents under age 25 may not have completed their schooling. A matric is about equivalent to a high school education. The message here is clear: the impact of HIV on labor market status is severe for those with lower levels of education and is negligible for those with higher levels. The causal impact of HIV status on labor market status for those with less than a matric is an increase in the likelihood of unemployment of 10 percentage points (with propensity score reweighting) and 10.9 percentage points (with the control function approach.) These are large and precisely estimated impacts. We repeated the analysis for subgroups defined by race as well as by groups defined by education and sex, age, and education and age. These results are not reported here but are available on request from the authors. The results in table 6 capture the gist of these divisions. Women, and especially women with lower levels of education, experience larger causal impacts of HIV on labor market status.

The last two rows highlight urban versus rural differences in the impact of HIV on unemployment. Although both HIV and unemployment are more prevalent in rural areas, the causal impact of HIV on unemployment is larger in urban areas where being HIV positive lowers the probability of employment by between 7 and 8 percentage points.

Discussion. A concern about the role of selection bias motivated our choice of methodologies. Comparing the naive estimates in column 1 of table 6 to the causal estimates in columns 3 and 4 speaks to this issue. The naive estimates are generally two to three percentage points higher than the causal estimates, and this highlights the importance of selection. Our results are consistent with the hypothesis that individuals who are HIV-positive are more likely to be unemployed than the average South African, regardless of their HIV status.

Because this study is probably the first to examine the causal impact of HIV on employment outcomes, it is difficult to place the magnitude of the estimated impact in context. There are no other estimates available for comparison. There are at least two economic arguments that one might have expected no causal impact. First, if unemployment were
so pervasive that HIV-positive individuals would be unemployed even in the absence of HIV, one would expect no impact. Second, if ARVs were sufficiently widely used, one might expect either no impact or a tiny impact. Our estimates indicate that these arguments, while perhaps ex ante plausible, are on average simply incorrect. Being HIV positive lowers the probability of employment.

A counterargument to the notion that HIV confers a negligible penalty in the face of extremely high unemployment is that in the presence of high unemployment, even a small disadvantage (e.g., the stigma sometimes associated with HIV), much less a large one (the adverse physical effects of HIV), means the difference between keeping a job and losing it. The results in table 6 are consistent with this counterargument.

One way to gauge the magnitude of our estimates (beyond noting that they are not 0) is to compare the estimated marginal effect of HIV status \(D\) with the estimated marginal effect of other respondent characteristics \(X\) in equation (4). For men, the magnitude of the labor market advantage of being HIV negative is approximately equal to the impact of three years of age, a matric qualification (compared to no education), or the absence of a female pensioner in the household. For women, it is equal to the impact of 1.5 years of age, some secondary education (compared to no education), or the absence of a male pensioner in the household. It is worth noting that our results are likely an underestimate of the true magnitude of the effect of HIV status on labor market outcomes because they do not account for intracategory changes in employment outcomes. As their disease progresses, HIV-positive workers may shift into less physically demanding occupations at reduced wages, which has implications for inequality.

We have estimated the causal impact of HIV status on the probability of unemployment, and in this sense HIV contributes to unemployment in South Africa. It would be wrong, though, to think that HIV was substantially responsible for South Africa’s dire unemployment. The numbers simply do not add up.

Our results are conditional on the availability of ARVs as of 2005. We do not have data on which HIV-positive respondents were on ARV therapy. Access to ARVs in South Africa is far from universal. One estimate is that in 2005, the year of our sample, only about 18% of those who needed ARVs were actually using them (Dorrington et al., 2006). As the availability of ARVs changes, the impact of HIV on labor market status, as we estimate it, will change. Because ARVs are more widely available today than they were in 2005, ceteris paribus, the impact of HIV on unemployment would be lower today than it was in 2005. In addition, ARV use is nonrandom, and this may in part contribute to the pattern of results in table 6. For example, ARVs are much more likely to be employer provided in the formal sector than they are in the informal sector. Our findings are consistent with the fact that women and the less educated tend to be more heavily represented in the informal sector and in domestic (housekeeper) work and hence less likely to receive employer-provided ARVs.

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**Table 7.—Sensitivity Check Using Narrow Definition of Unemployment: Marginal Effect of Being HIV-positive on Likelihood of Being Unemployed**

<table>
<thead>
<tr>
<th>Sample</th>
<th>Naive</th>
<th>Logit</th>
<th>RW</th>
<th>CF</th>
<th>N</th>
</tr>
</thead>
<tbody>
<tr>
<td>Full sample</td>
<td>0.068***</td>
<td>0.025</td>
<td>0.034</td>
<td>0.038***</td>
<td>4,620</td>
</tr>
<tr>
<td>Male</td>
<td>−0.012</td>
<td>0.021</td>
<td>0.031</td>
<td>0.049</td>
<td>1,761</td>
</tr>
<tr>
<td>Female</td>
<td>0.086***</td>
<td>0.031</td>
<td>0.027</td>
<td>0.040</td>
<td>2,859</td>
</tr>
<tr>
<td>Matric and up</td>
<td>0.017</td>
<td>−0.049</td>
<td>−0.007</td>
<td>0.015</td>
<td>839</td>
</tr>
<tr>
<td>Less than matric</td>
<td>0.091***</td>
<td>0.044</td>
<td>0.051</td>
<td>0.063***</td>
<td>2,581</td>
</tr>
<tr>
<td>Rural</td>
<td>0.063</td>
<td>0.002</td>
<td>0.009</td>
<td>0.039**</td>
<td>1,973</td>
</tr>
<tr>
<td>Urban</td>
<td>0.082**</td>
<td>0.038</td>
<td>0.049</td>
<td>0.040**</td>
<td>2,647</td>
</tr>
</tbody>
</table>

Full sample and results by sex include individuals aged 15 and older. Results by education level are restricted to individuals aged 25 and older. Naive is OLS without controls. Controls used in logit are same as those used in propensity score estimation. Bootstrapped standard errors reported for propensity score reweighting (RW) and control function (CF). Significant at ***99% and **95% level.

We also found that being HIV positive had virtually no employment impact for better-educated workers (and recall that this result already accounts for the fact that highly educated individuals are less likely to be HIV positive). ARVs may also be contributing to this finding. Employers have greater incentive to invest in ARVs for workers who are more difficult to replace, such as highly educated workers.

In sum, HIV appears to reinforce the already existing inequalities in South Africa.

**D. Sensitivity Analyses**

We investigate the sensitivity of our results to both alternative definitions of unemployment and specifications of the propensity score regression.

As described in section 3, individuals were divided into three groups in our base case specification: unemployed, employed, and not economically active (NEA). The first two groups are included in our analysis; the third is not. In the base case results using the broad definition of unemployment, so-called discouraged workers—those who were not actively seeking a job but desired employment—were classified as unemployed. The narrow definition of unemployment classifies discouraged workers as NEA. We repeat the analysis using this narrow definition of unemployment. Results are reported in table 7. The results are very similar to those in table 6. The estimated coefficients are about one-half to two-thirds the size of those reported in table 6. We also repeated the analysis by combining the NEA with the employed to create a group that might (awkwardly) be called “not unemployed”. Results are reported in table 8. Compared to table 6, coefficients tend to increase, usually by about one and one-quarter to one and a half times. Results tend to be more precisely estimated, and this is driven in part by the increase in the sample size when NEA individuals are included in the analysis. From tables 7 and 8, we conclude that our findings are robust to alternative definitions of labor market status.
The average impact hides important heterogeneity. HIV's penalty exists despite very high unemployment rates. We find that being HIV positive causes, on average, an increase in the likelihood of unemployment of 6 to 7 percentage points. This penalty exists despite very high unemployment rates. The average impact hides important heterogeneity. HIV's causal impact on unemployment is larger (10 to 11 percentage points) for less educated South Africans. The results are robust to multiple alternative econometric specifications.

### Table 8. Sensitivity Check Including Not Economically Active (NEA) Individuals in the Sample: Marginal Effect of Being HIV-positive on Likelihood of Being Unemployed

<table>
<thead>
<tr>
<th>Sample</th>
<th>Naive</th>
<th>Logit</th>
<th>RW</th>
<th>CF</th>
<th>N</th>
</tr>
</thead>
<tbody>
<tr>
<td>Full sample</td>
<td>0.180***</td>
<td>0.086***</td>
<td>0.093***</td>
<td>0.104***</td>
<td>7,445</td>
</tr>
<tr>
<td></td>
<td>(0.022)</td>
<td>(0.023)</td>
<td>(0.022)</td>
<td>(0.021)</td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>0.103***</td>
<td>0.048</td>
<td>0.063</td>
<td>0.101***</td>
<td>2,672</td>
</tr>
<tr>
<td></td>
<td>(0.039)</td>
<td>(0.040)</td>
<td>(0.044)</td>
<td>(0.039)</td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>0.209***</td>
<td>0.096***</td>
<td>0.100***</td>
<td>0.087***</td>
<td>4,771</td>
</tr>
<tr>
<td></td>
<td>(0.026)</td>
<td>(0.028)</td>
<td>(0.027)</td>
<td>(0.025)</td>
<td></td>
</tr>
<tr>
<td>Matric and up</td>
<td>0.056</td>
<td>−0.007</td>
<td>0.013</td>
<td>0.023</td>
<td>932</td>
</tr>
<tr>
<td></td>
<td>(0.056)</td>
<td>(0.064)</td>
<td>(0.067)</td>
<td>(0.056)</td>
<td></td>
</tr>
<tr>
<td>Less than matric</td>
<td>0.178***</td>
<td>0.090***</td>
<td>0.096***</td>
<td>0.096***</td>
<td>3,804</td>
</tr>
<tr>
<td></td>
<td>(0.029)</td>
<td>(0.030)</td>
<td>(0.029)</td>
<td>(0.029)</td>
<td></td>
</tr>
<tr>
<td>Rural</td>
<td>0.177***</td>
<td>0.073**</td>
<td>0.084***</td>
<td>0.099***</td>
<td>3,538</td>
</tr>
<tr>
<td></td>
<td>(0.034)</td>
<td>(0.035)</td>
<td>(0.035)</td>
<td>(0.031)</td>
<td></td>
</tr>
<tr>
<td>Urban</td>
<td>0.185***</td>
<td>0.100***</td>
<td>0.106***</td>
<td>0.110***</td>
<td>3,905</td>
</tr>
<tr>
<td></td>
<td>(0.029)</td>
<td>(0.030)</td>
<td>(0.030)</td>
<td>(0.028)</td>
<td></td>
</tr>
</tbody>
</table>

*Full sample and results by sex include individuals aged 15 and older. Results by education level are restricted to individuals aged 25 and older. Naive is OLS without controls. Controls used in logit are same as those used in propensity score estimation. Bootstrapped standard errors reported for RW and CF. Significant at ***99% and **95%.

We also experimented with alternative specifications of the propensity score regression. The results obtained from regressions using additional variables in the calculation of the propensity score are virtually identical to those obtained from the preferred specification. These results are not reported here. When including ten variables with information on the household, no point estimate changed by more than 2 percentage points, and only three of the fifteen-point estimates changed by more than 1 percentage point. The additional household variables chosen to indicate socioeconomic level were type of toilet facility, source of energy for cooking, access to electricity, presence of a land line, and a dummy for whether the household information was missing for the observation. Adding more behavioral variables did not change any of the point estimates by as much as 1 percentage point. The additional behavioral variables were condom use at last sex, number of current sexual partners, number of partners in the past year, whether the respondent had been tested for HIV before, whether he or she had received the test result, and whether he or she had heard of ARVs.

### VI. Conclusion and Caveats

#### A. Conclusion

Identifying the causal impact of HIV on labor market status requires addressing the issue of selection into being HIV positive. In the absence of plausible instruments, we exploit the richness of the data and assume that selection is on observables. External balancing tests support the validity of this assumption. Employing two estimation strategies, we find that being HIV positive causes, on average, an increase in the likelihood of unemployment of 6 to 7 percentage points. This penalty exists despite very high unemployment rates. The average impact hides important heterogeneity. HIV’s causal impact on unemployment is larger (10 to 11 percentage points) for less educated South Africans. The results are robust to multiple alternative econometric specifications.

#### B. Caveats

These results are the first nationally representative estimates of the causal impact of HIV on employment in South Africa. Although they are informative, they are not dispositive. Rather, the results should be interpreted with caution for at least five reasons.

First, our analysis does not account for any general equilibrium effects. In particular, it would be misleading to think that if ARVs were made universally available or a cure for HIV/AIDS were found, that HIV-positive individuals would see their likelihood of employment rise on average by about 7 percentage points. Rather, the labor market would adjust, and these adjustments would depend on supply and demand elasticities. Complicating this analysis, the ability of the labor market to immediately absorb additional healthy workers is questionable.

Second, data limitations preclude an analysis of the indirect labor market impact of having multiple HIV-positive adult household members. Recall that the structure of the SABSSM II survey is such that only one adult age 25 or older is sampled from each household. It is unclear in which direction our results may be biased. An HIV-negative worker could be unemployed because he or she is caring for an HIV-positive spouse, resulting in downward bias (i.e., the true impact is larger than our estimates suggest), or an HIV-positive worker might be more motivated to obtain employment to financially support another HIV-positive household member, resulting in upward bias.

Third, our results are conditional on the time profile of HIV and prevalence rates as of 2005. Given an approximately nine-year period (on average) of latent HIV infection before AIDS conversion, and HIV prevalence rates that increased from 5% in 1996 to 12% in 2001,19 we would expect the “stock” of individuals with AIDS to increase quite sharply between 2003 and 2008. This implies that, ceteris paribus, the impact of HIV on unemployment would rise in coming years as the number of HIV-positive individuals who are too ill to work increases. However, it is unclear how this effect would interact with any increase in the availability of ARVs.

Fourth, our results are not structural, so we are unable to convincingly address the particular avenues through which HIV affects labor market status. Relatedly, we are unable to conduct detailed policy analysis. For example, increased access to ARVs and successful programs to destigmatize HIV might each increase the likelihood of employment, but our approach cannot conduct the counterfactual experiments to estimate the likely impacts of these potential policies.

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19 ASSA demographic model predictions cited in Natassa (2004). The model on which these estimates are based predicted a 14% prevalence rate for 2004—about the same as that of our 2005 data.
Finally, our results are generally not applicable to other countries. South Africa has a stunningly high rate of unemployment, high HIV prevalence rates, and a troubled history with the distribution of ARVs—three factors that suggest that it may be misguided to generalize the results of this study to other countries.

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