Lower risk of prostate cancer in schizophrenia – Fact or artifact?

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ABSTRACT – Background and Objectives: Reduced risk for prostate cancer in persons diagnosed with schizophrenia has been reported repeatedly. While genetic factors and antipsychotic medications were imputed, a methodological hypothesis could be raised: given the reported disparities of health services with regard to service users with schizophrenia these persons may be screened less often compared to schizophrenia-free counterparts resulting in undetected prostate tumors. The study objective was to compare the rates of performance of the prostate specific antigen (PSA) screening test among persons diagnosed with schizophrenia to matched-comparison subjects.

Methods: A prospective historical epidemiological study which included service users with and without a diagnosis of schizophrenia (N = 52,131). Linkage was conducted between databases of the nation-wide psychiatric register and Israel’s largest health maintenance organization. Annual performance of PSA tests was recorded between the years 2002-2009.

Results: Service users with schizophrenia performed the PSA test 23% less than the comparison subjects. However, among service users who were diagnosed with prostate cancer only 44% performed the PSA test during the year prior to diagnosis.

Conclusions: In line with our hypothesis, lower rates of PSA screening tests were conducted among service users with schizophrenia. However, the relatively low rate of this test prior to prostate cancer diagnosis ought to be noted.

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Introduction

Several studies found a reduced risk for cancer in persons diagnosed with schizophrenia\textsuperscript{1-7}. Among cancer sites the most robust finding was associated with prostate cancer. Torrey\textsuperscript{8}, who reviewed five large epidemiological studies with a total of 98,951 individuals with schizophrenia, noted a combined estimated standardized incidence ratio (SIR) of 0.59 (0.49 - 0.76) for prostate cancer. Importantly, most studies were based on nation-wide registries in countries with national health insurances (Denmark, England, Finland, and Israel). A subsequent review by Catts et al confirmed these findings, SIR 0.59 (0.48-0.73)\textsuperscript{2}.

Torrey\textsuperscript{8} raised several substantive hypotheses to explain these lower rates: 1. Genetic mechanisms; 2. Anti-tumor action of psychotropic medications; and 3. Limited heterosexual activity with a subsequent decreased risk of sexual-transmitted infections. However, a methodological hypothesis cannot be excluded: given the disparities in the health care of persons with schizophrenia\textsuperscript{9-11}, if service-users with schizophrenia are less often screened with prostate-specific antigen (PSA) tests compared to schizophrenia-free men the identification of prostate cancer may fall.

We had the rare opportunity to explore the latter hypothesis in a study of medical services provided to persons with schizophrenia in a country with a national health insurance. In Israel, primary medical services are either entirely gratis (e.g., visits to general practitioners, laboratory tests, including PSA) for all legal residents and citizens, or a minimal co-payment is required to access a specialist.

In this study, we merged data from the National Psychiatric Case Registry (PCR), that records all psychiatric admissions and discharges since 1950, with the database of Israel’s largest health maintenance organization (HMO) that provided information on the PSA tests performed, and compared their rates of performance among persons with the diagnosis of schizophrenia to matched comparison subjects.

Objective: To study the rate of PSA screening tests performed by service users with schizophrenia compared to matched comparison subjects in a country with a national health insurance, using a prospective historical design.

Methods

The study was based on a prospective historical design where data on the participants were collected between the years 2002-2009. The study was approved by the IRB of Clalit Health Services (CHS).

Identification of persons diagnosed with schizophrenia

We used the PCR to identify persons diagnosed with schizophrenia. The PCR is legally mandated to maintain a cumulative record of all psychiatric hospitalizations\textsuperscript{12-13}. The PCR provided the subjects’ diagnoses upon admission and discharge as well as socio-demographic information. Diagnoses are based on the ICD-10; those made prior to its introduction have been updated. A test of the agreement between research diagnoses and those recorded in the PCR found a satisfactory match\textsuperscript{14-15}. The following inclusion criteria were applied: 1. Persons with last discharge diagnosis of schizophrenia (F20 - F29); and 2. Persons born before 1960 (age 40 and above at year 2000) and were alive by the year 2000.

We identified 28,579 persons diagnosed with schizophrenia. Each one was identified
based on his ID number. In addition to diagnoses, we gathered information on the year of first and last hospitalization, the number of hospitalizations, and the length of each hospital stay (Table 1). The latter was done annually between the years 2000-2009 to control for the possible artifact caused by periods when the person was not in the community.

General health services

Data from CHS, with 3.8 million insured persons, were collected from its electronic database. The target population included subjects meeting the second inclusion criterion (above). A total of 1,040,000 subjects were identified. Information on the annual performance of PSA tests were gathered for the years 2002-2009, since CHS started to apply a central database in 2002 only. In addition, medical diagnoses, including prostate cancer were recorded. Two measures were calculated based on the performance of PSA screening tests: a. sum of tests performed during the follow up period (2002-2009); b. since the performance of PSA tests is expected to rise in persons diagnosed with prostate cancer we have calculated an additional measure which accounts for the year of such diagnosis. The adjusted measure was based on the sum of tests performed during the follow up period before prostate cancer diagnosis was recorded. For example, if a person was diagnosed in 2006 the sum of PSA tests during the years 2002-2005 was calculated.

Linkage procedure and selection of a matched comparison group

The data extracted from the PCR and from the CHS databases were merged according to the personal ID numbers. The number of identified persons in both databases was 17,377. To generate a matched comparison group (2:1 ratio) we defined an algorithm based on age (groups of three consecutive years), sex, birth place (Israel, Asia, Africa, Europe-America), socioeconomic status (high, mid, low), and ethnic-national sector (Jews and Arabs). We omitted cases: 1. That had missing information on one of the matching variables; 2. That died before 2002; 3. That had an average yearly hospitalization period of 270 days or more. The matching procedure was tested and found satisfactory (Table 1). The data set included 17,377 service users with schizophrenia and 34,754 matched comparison subjects (n = 52,131).

Data analysis

The analysis was based on 8,285 men diagnosed with schizophrenia, and 16,570 matched comparison subjects. To account for the effects of matching, we applied the complex samples design of the SPSS 21.0 software (IBM Inc.) with case-control triads as cluster. Sum and adjusted sum of PSA tests were analyzed using general linear models (GLM), with the number of follow up years as a covariate. Rates of PSA tests (yes/no) and prostate cancer were analyzed using logistic regression models, reported in terms of odds ratios (OR) and 95% confidence intervals (95% CI).

Results

The PSA test was performed at least once during the follow up period (2002 - 9) among 26.2% of the service users with schizophrenia and 30.4% of their counterparts (OR = 0.83, 95% CI 0.78 - 0.88; p < 0.001). The performance of PSA tests during the follow up period was significantly lower among
service-users with schizophrenia (mean 0.76) compared to their counterparts (mean 1.03) \((F = 48.7, df = 1; 23,958, p < 0.001)\), reflecting a reduced performance of 26% (Table 2).

Similar effects were observed when the length of stay of the psychiatric hospitalizations was controlled.

We identified a total of 523 service users diagnosed with prostate cancer. The rate of prostate cancer diagnosis was lower among service-users with schizophrenia (1.5%) compared to their matched counterparts (2.4%); odds ratio 0.61 (95% CI 0.50 - 0.75; \(p < 0.001\)). The adjusted PSA performance measure accounting for prostate cancer diagnosis was significantly lower among service-users with schizophrenia (mean 0.73) compared to their matched counterparts (mean 0.95) \((F = 40.3, df = 1; 23,824, p < 0.001)\), reflecting a reduced performance of 23% (Table 2).

Among service users diagnosed with prostate cancer, 329 were diagnosed during the years 2003 - 2010. The PSA test during the year prior to cancer diagnosis (2002-9) was performed in 44% of these cases, with slight variation between those with schizophrenia (39.7%) and comparison subjects (46.4%) \(p = 0.3\).

### Table 1
Sociodemographic characteristics of the sample and clinical data of service users diagnosed with schizophrenia.

<table>
<thead>
<tr>
<th>Variables</th>
<th>Comparison sample ((n = 16,570))</th>
<th>Persons with schizophrenia ((n = 8,285))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age at year 2009</td>
<td>61.8 (SE 0.09)</td>
<td>61.8 (SE 0.09)</td>
</tr>
<tr>
<td>Birthplace</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Israel</td>
<td>45.9%</td>
<td>45.9%</td>
</tr>
<tr>
<td>Asia</td>
<td>14.3%</td>
<td>14.3%</td>
</tr>
<tr>
<td>Africa</td>
<td>16.8%</td>
<td>16.8%</td>
</tr>
<tr>
<td>Europe-America</td>
<td>23.0%</td>
<td>23.0%</td>
</tr>
<tr>
<td>Ethnic-national sector</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jewish</td>
<td>91.0%</td>
<td>91.0%</td>
</tr>
<tr>
<td>Arab</td>
<td>9.0%</td>
<td>9.0%</td>
</tr>
<tr>
<td>Age at first diagnosis</td>
<td>–</td>
<td>31.9 (SE 0.14)</td>
</tr>
<tr>
<td>Mean number of psychiatric hospitalizations</td>
<td>–</td>
<td>10.6 (SE 0.11)</td>
</tr>
<tr>
<td>Mean hospitalization days (2002-2009)</td>
<td>–</td>
<td>41.9 (SE 1.2)</td>
</tr>
</tbody>
</table>

### Table 2
Number of PSA tests performed between the years 2002-2009.

<table>
<thead>
<tr>
<th></th>
<th>Comparison sample</th>
<th>Persons with schizophrenia</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>SD</td>
</tr>
<tr>
<td>Number of tests</td>
<td>1.03</td>
<td>2.5</td>
</tr>
<tr>
<td>Adjusted number of tests*</td>
<td>0.95</td>
<td>2.3</td>
</tr>
</tbody>
</table>

*Accounting for the year of prostate cancer diagnosis.
Discussion

The current study compared the performance of PSA test in a large sample of adult service users with schizophrenia and matched comparison subjects. We found 23% (adjusted) decreased performance of PSA tests among service users diagnosed with schizophrenia compared to their matched counterparts. In line with previous reports\(^1\)\(^-\)\(^8\) we found that the risk of prostate cancer among service users with schizophrenia was lower than matched comparison subjects (in about 61%). According to our hypothesis, the lower rate of PSA tests could suggest that this negative association may reflect more an issue of method than an issue of substance. Since service-based disparities were noted repeatedly in the literature\(^9\)\(^-\)\(^11\), as well as in our data (Gal et al., submitted), it is possible that the lower rate of PSA tests among service users diagnosed with schizophrenia result in less biopsies, and the consequent lower rate of cancer diagnosis.

This interpretation should be carefully weighted given the relatively lower rate of performance of PSA test preceding the cancer diagnosis. According to our data, 44% of the persons with confirmed cancer diagnosis had a PSA test during the year previous to it, suggesting that for slightly more than half of the service users an alternative screening method (e.g., digital rectal examination (DRE)) was applied. This notion could be supported by the reservations raised regarding the PSA as a screening test\(^16\)\(^-\)\(^18\), which is recommended currently by some but not all professional organizations\(^16,19\).

The major limitation of this study is the reliance on the PSA test as the sole screening procedure for the identification of prostate cancer. Our findings would have been fully confirmed if we had data on the performance of DREs, and, especially biopsies, but such information was not available to us. Indeed, results of prostatic biopsies are considered a gold standard, but biopsies are ordered generally when the results of a PSA test or DRE are found positive. Note, however, that the lower performance of PSA screening tests among the service users with schizophrenia in our study would have resulted in an equally lower rate of biopsies. Among the US general population, 65% of the persons with high PSA levels are referred for biopsy\(^20\).

In conclusion, given the disparities prevailing in the health services provided to service users with schizophrenia, the lower rate of PSA tests might lead to lesser performance of biopsies which may be manifested in the underestimation of prostate cancer in this population group.

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Conflict of interest

Co-author Hanan Munitz is employed by Clalit Health Services. All other authors declare that they have no conflicts of interest.

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References


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