Risk of sexually transmitted infections following depressive disorder
A nationwide population-based cohort study

Sheng-Yun Huang, Jeng-Hsiu Hung, Li-Yu Hu, Min-Wei Huang, Shyh-Chyang Lee, Cheng-Che Shen

Abstract
Depressive disorder is a severe mental disorder associated with functional and cognitive impairment. Numerous papers in the literature investigated associations between sexually transmitted infections (STIs) and psychiatric illnesses. However, the results of these studies are controversial.

We explored the relationship between depressive disorder and the subsequent development of STIs including human immunodeficiency virus (HIV) infection, primary, secondary, and latent syphilis, genital warts, gonorrhea, chlamydial infection, and trichomoniasis.

We identified patients who were diagnosed with the depressive disorder in the Taiwan National Health Insurance Research Database. A comparison cohort was constructed of patients without the depressive disorder who were matched according to age and sex. The occurrence of subsequent new-onset STIs was evaluated in both cohorts.

The depression cohort consisted of 5,959 patients, and the comparison cohort consisted of 23,836 matched control patients without depressive disorder. The incidence of subsequent STIs (hazard ratio [HR] 1.54, 95% confidence interval [CI] 1.34–1.76) was higher among the depressed patients than among the patients in the comparison cohort. Furthermore, female gender compared to male (HR 1.58, 95% CI 1.24–2.01) and young age <40-year-old (HR 1.79, 95% CI 1.38–2.32) are both risk factors for acquisition of STIs in depression patients. For individual STI, the results indicated that the patients with depressive disorder exhibited a markedly higher risk for subsequent STIs including HIV infection, syphilis, genital warts, gonorrhea, chlamydial infection, and trichomoniasis.

Depressive disorder might increase the risk of subsequent newly diagnosed STIs including HIV infection, syphilis, genital warts, gonorrhea, chlamydial infection, and trichomoniasis in Taiwan population. Clinicians should pay particular attention to STIs in depression patients. Depression patients, especially those with the history of high-risk sexual behaviors, should be routinely screened for STIs.

Abbreviations: CI = confidence interval, HIV = human immunodeficiency virus, HR = hazard ratio, ICD-9-CM = International Classification of Diseases, Ninth Revision, Clinical Modification, LHD 2000 = Longitudinal Health Insurance Database 2000, NHI = National Health Insurance, NHIRD = National Health Insurance Research Database, NHRI = National Health Research Institutes, STIs = sexually transmitted infections.

Keywords: chlamydia, depression, gonorrhoeae, human immunodeficiency virus, sexually transmitted infections, syphilis, warts

1. Introduction
The depressive disorder is a commonly occurring, severe and recurrent disorder linked to diminished role functioning and quality of life, medical morbidity, and mortality.[1] Convergent evidence indicates that depressive disorder is one of the leading cause of disability among patients in both developed and emerging economies.[2] The principal source of cost and illness-associated morbidity is due to a significant decrease in role-function among affected individuals.[3,4] As for the relationships between depression and sexually transmitted infections (STIs), depression may impair cognitive function and memory,[5] decrease impulse control,[6] which contribute to risky sexual practices. These depression-related effects may inhibit the clear perception of STIs risk and the ability to prevent risk behavior.[7,8]

The STIs are infections that are commonly spread by sexual behaviors. In 2015, about 1.1 billion people had STIs other than human immunodeficiency virus (HIV) infection.[9] STIs other than HIV infection resulted in 108,000 deaths in 2015.[10] In the United States, millions of cases of STIs occur each year, resulting in substantial medical costs to the nation. A study indicated that the total lifetime direct medical cost for STIs in 2008 in the United States was $15.6 billion and the burden of STIs would be even greater in the absence of STIs prevention and control efforts.[11]
Therefore, it is essential to identify risk factors of STIs and arrange appropriate screening protocols for people at risk of these infections.

Numerous papers in the literature investigated associations between STIs and depressive disorders. While STIs are risk factors for depression, depression also may increase susceptibility to risk behaviors and infection. However, most of these studies focused on HIV infection only, and few studies were designed to investigate the association between depressive disorder and other STIs, such as syphilis, genital warts, gonorrhea, chlamydial infection, and trichomoniasis. In addition, the results of these studies are controversial. For example, a study investigated of 10,783 young adults in the United States revealed that depression was associated with increased risk of STIs only among black men and there is no significantly higher risk among women. However, another study investigated a similar group with 18,142 cases in the United States found that a major depressive episode was associated with increased risk of STIs only in females. Furthermore, most of these study results are based on cross-sectional study design and lack a longitudinal perspective.

In response to the few longitudinal studies concerning the association between depressive disorder and the subsequent risk of STIs, and based on the hypothesis that depressive disorder, a psychiatric illness related to impaired cognitive function and poor impulse control, might have a higher risk for developing subsequent STIs, we designed a nationwide population-based retrospective cohort study to investigate the possible link between these 2 illnesses.

2. Patients and methods

2.1. Data sources

Instituted in 1995, the National Health Insurance (NHI) program is a mandatory health insurance program that offers comprehensive medical care coverage, including outpatient, inpatient, emergency, and traditional Chinese medicine, to all residents of Taiwan, with a coverage rate of up to 98%. The NHI Research Database (NHIRD) contains comprehensive information regarding clinical visits, including prescription details and diagnostic codes based on the A code and International Classification of Diseases, Ninth Revision, Clinical Modification (ICD-9-CM). The NHIRD is managed and publicly released by the National Health Research Institute (NIHR) for research purposes, and confidentiality is maintained according to the directives of the Bureau of NHI. The data source for our study was the Longitudinal Health Insurance Database 2000 (LHID 2000), which is a data set of NHIRD. Data for the LHID were collected systematically and randomly sampling from the NHIRD; the database included the data of 1 million individuals. The NHRI of Taiwan reports that there were no significant differences in gender distribution, age distribution, or average insured payroll-related amount between the patients in the LHID and those in the original NHIRD.

2.2. Ethics statement

The Institutional Review Board of Taichung Veterans General Hospital approved this study. Written consent from the study patients was not obtained, because the NHI data set consists of de-identified secondary data for research purposes and the institutional review board approved that consent was not needed. The Institutional Review Board of Taipei Veterans General Hospital issued a formal written waiver for the need for consent.

2.3. Study population

Using data extracted from the LHID 2000, we conducted a retrospective cohort study of patients who were newly diagnosed with depressive disorder between January 1, 2000 and December 31, 2004. The patients with the depressive disorder were defined as ICD-9-CM code: 296.2, 296.3, 300.1, and 311. We excluded patients who were diagnosed with depressive disorder between January 1, 1996 and December 31, 1999. We also excluded patients who were diagnosed with STIs (HIV infection, primary, secondary, and latent syphilis, genital warts, gonorrhea, chlamydial infection, and trichomoniasis) before they were diagnosed with depressive disorder. For each depressive patient included in the final cohort, 4 age- and sex-matched control patients without STIs were randomly selected from the LHID 2000. The random assignment procedures were performed by SAS statistical software and were based on the random numbers which were generated from the uniform distribution. All depression and control patients were observed until diagnosed with HIV infection (ICD-9-CM codes: 042 or V08, A049), primary, secondary, and latent syphilis (ICD-9-CM codes: 091-097, A060), genital warts (ICD-9-CM codes: 078.11), gonorrhea (ICD-9-CM codes: 098, A061), chlamydial infection (ICD-9-CM codes: 078.8, 078.88), or trichomoniasis (ICD-9-CM codes: 131, A079), or until death, withdrawal from the NHI system, or December 31, 2013. The primary clinical outcomes assessed were newly diagnosed HIV infection, primary, secondary, and latent syphilis, genital warts, gonorrhea, chlamydial infection, and trichomoniasis. Common comorbidities including hypertension, diabetes mellitus, dyslipidemia, coronary artery diseases, congestive heart failure, chronic pulmonary diseases, and cerebrovascular diseases were also compared between depression and control patients.

2.4. Statistical analysis

The occurrence of newly diagnosed HIV infection, primary, secondary, and latent syphilis, genital warts, gonorrhea, chlamydial infection, or trichomoniasis in the depression and control patients were considered as the primary outcome in this study. We first compared the distribution of demographic characteristics between the depression and control patients by using the independent t tests and Chi-squared test. To investigate potential surveillance bias, subgroups were stratified according to the duration since enrollment. In addition, a Cox proportional-hazards regression model was performed to identify the risk factors associated with STIs in the whole sample and patients with depressive disorder. Furthermore, the Cox proportional-hazards regression model was also constructed to calculate the hazard ratio (HR) of 6 different kinds of STIs, including HIV infection, primary, secondary, and latent syphilis, genital warts, gonorrhea, chlamydial infection, and trichomoniasis of the depression cohort and control cohort.

The SAS statistical software for Windows, Version 9.3 (SAS Institute, Cary, NC), was used for data extraction, computation, linkage, processing, and sampling. All other statistical analyses were performed using the SPSS statistical software for Windows, Version 20 (IBM, Armonk, NY). The results of comparisons with a P-value <.05 were considered to indicate a statistically significant relationship.
3. Results

Our study sample comprised 5959 depression patients and 23,836 control patients without depressive disorder. The comparisons of the demographic and clinical variables between the depression and control patients are presented in Table 1. The median age of the patients was 42.6 years (interquartile range, 29.5–52.4 years) and the median follow-up duration of depression and control cohorts was 11.08 and 11.36 years, respectively. A higher percentage of patients with the depressive disorder were observed in the group of people aged 20 to 39 years. Baseline difference in comorbidities demonstrated the higher prevalence of hypertension, diabetes mellitus, dyslipidemia, coronary artery diseases, congestive heart failure, chronic pulmonary diseases, and cerebrovascular diseases among the depressed patients. During the follow-up period, 298 (5.0%) depression patients and 774 (3.2%) control patients were diagnosed with STIs (P < .001). The most common STIs in the patients with the depressive disorder were trichomoniasis in 121 patients (2.0%) and chlamydial infection in 68 patients (1.1%). Overall, significantly higher incidences of all STIs included HIV infection (P = .001), syphilis (P < .001), genital warts (P = .001), gonorrhea (P < .001), chlamydial infection (P = .006), and trichomoniasis (P = .004) were observed in the depression patients than in the control patients.

A subanalysis based on the duration of follow-up revealed that most of the STIs developed beyond the first year following a depression diagnosis and that the risk of all STIs included HIV infection, syphilis, genital warts, gonorrhea, chlamydial infection, and trichomoniasis remained significantly elevated when the patients diagnosed with STIs within 1 year of depression diagnosis were excluded. The results of the subanalysis are summarized in Table 2.

Besides, a Cox proportional-hazards regression analysis was conducted to calculate the HR of the newly diagnosed STIs for the patients with the depressive disorder compared with the matched controls and revealed patients with depressive disorder exhibited a markedly higher risk for subsequent STIs (HR 1.54, 95% confidence interval [CI] 1.34–1.76). In addition, female gender compared to male (HR 2.20, 95% CI 1.91–2.54), young age <40 year old (HR 1.69, 95% CI 1.48–1.92), cerebrovascular disease (HR 1.42, 95% CI 1.14–1.78), and low income (HR 1.22, 95% CI 1.05–1.43) are risk factors for acquisition of STIs. Urbanization and other comorbidities including hypertension, diabetes mellitus, dyslipidemia, coronary artery diseases, congestive heart failure, and chronic pulmonary diseases are not risk factors for the subsequent STIs (Table 3).

In depression patients, female gender compared to male (HR 1.58, 95% CI 1.24–2.01) and young age less than the age of 40 (HR 1.79, 95% CI 1.38–2.32) are both risk factors for acquisition of STIs (Table 4). For individual STI, the results indicated that the patients with depressive disorder exhibited a markedly higher risk for subsequent STIs included HIV infection (HR 2.93, 95% CI 1.62–5.30), syphilis (HR 1.95, 95% CI 1.32–2.88), genital warts (HR 1.88, 95% CI 1.27–2.77), gonorrhea (HR 1.97, 95% CI 1.33–2.92), chlamydial infection (HR 1.44, 95% CI 1.08–1.90), and trichomoniasis (HR 1.33, 95% CI 1.08–1.64) (Table 5).

4. Discussion

The key findings of our study are listed as below: the prevalence of STIs, including HIV infection, primary, secondary, and latent syphilis, genital warts, gonorrhea, chlamydial infection, and trichomoniasis, in patients with depressive disorder were demonstrated in our work; depressive disorder was significantly associated with an increased sequential risk of all STIs included HIV infection, syphilis, genital warts, gonorrhea, chlamydial infection, and trichomoniasis in Taiwan population; female gender compared to male (HR 1.58) and young age <40 year old (HR 1.79) are both risk factor for acquisition of STIs in depression patients.

In our study, it revealed that depressive disorder was significantly associated with an increased sequential risk of STIs in Taiwan population. There are several possible explanations for the increased risk of STIs in patients with depressive disorder. First, depression may influence their ability to negotiate and engage in safer sexual behavior. Depression patients also lack adequate self-esteem and the assertiveness skills needed to implement safer sex practices. Second, the impaired cognitive function was noted in depression, and it may result in neglect of risky sexual behavior. Depressive symptoms may also be linked to poor sexual decision-making and make them engage in risky sexual practices. Third, depressed patients were more likely to have sex for money or drugs, to have had sex with an intravenous

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Table 1

| Characteristics of depressive disorder and sexually transmitted diseases (STIs) and control subjects. | Depressive disorder | Control cohort | P-values |
|---|---|---|---|
| No. | 5959 | 23,836 |  |
| Age, y | 42.6 (29.5–52.4) | 42.6 (29.5–52.4) | > .001 |
| Distribution of age |  |
| 20–29 | 3053 (51.2) | 12,212 (51.2) |  |
| 30–39 | 2127 (35.7) | 8508 (35.7) |  |
| ≥40 | 779 (13.1) | 3116 (13.1) |  |
| Sex |  |
| Female | 3534 (59.3) | 14136 (59.3) |  |
| Male | 2425 (40.7) | 9700 (40.7) |  |
| Comorbidities |  |
| Hypertension | 1416 (23.8) | 4069 (17.1) | < .001 |
| Diabetes mellitus | 859 (14.4) | 2343 (9.8) | < .001 |
| Dyslipidemia | 1058 (17.8) | 2852 (12.0) | < .001 |
| Coronary artery disease | 937 (15.7) | 2207 (9.3) | < .001 |
| Congestive heart failure | 173 (2.9) | 361 (1.5) | < .001 |
| Cerebrovascular disease | 744 (12.5) | 1651 (6.9) | < .001 |
| Chronic pulmonary disease | 711 (11.9) | 1798 (7.5) | < .001 |
| Income |  |
| High income | 661 (11.1) | 2665 (11.2) | < .001 |
| Medium income | 1044 (17.5) | 4702 (19.7) |  |
| Low income | 2982 (50.0) | 11,505 (48.3) |  |
| No income | 1272 (21.3) | 4964 (20.8) |  |
| Degree of urbanization |  |
| Urban | 3767 (63.2) | 14,606 (61.3) | < .001 |
| Suburban | 1707 (28.6) | 7686 (32.2) |  |
| Rural | 485 (8.1) | 1544 (6.5) |  |
| Newly diagnosed STIs, n (%) | 206 (5.0) | 774 (3.2) | < .001 |
| HIV infection | 19 (0.3) | 27 (0.1) | < .001 |
| Syphilis | 40 (0.7) | 78 (0.3) | < .001 |
| Genital warts | 38 (0.6) | 81 (0.3) | < .001 |
| Gonorrhea | 39 (0.7) | 74 (0.3) | < .001 |
| Chlamydial infection | 68 (1.1) | 183 (0.8) | < .001 |
| Trichomoniasis | 121 (2.0) | 357 (1.5) | .004 |
| Follow-up, y | 11.08 (11.23–12.89) | 11.36 (11.32–12.92) | < .001 |

HIV = human immunodeficiency virus, STIs = sexually transmitted diseases.

* Median (Interquartile range).

† Statistical significance.
### Table 2
Number of newly diagnosed sexually transmitted diseases between depressive disorder and control subjects which were stratified by follow-up duration.

| Follow-up duration, y | Depressive disorder | Control cohort | Risk ratio (95% CI) |
|-----------------------|---------------------|----------------|-------------------|
|                       | No of HIV infection | Per 1000 person-years | No of HIV infection | Per 1000 person-years | |
| Overall               | 19                  | 1.56            | 27                | 0.63                | 2.86 (1.50–5.33)* |
| 0–1                   | 3                   | 36.20           | 0                 | 0                   | N/A |
| ≥1                    | 16                  | 1.54            | 27                | 0.63                | 2.41 (1.21–4.63)* |

| No. of syphilis | Per 1000 person-years | No. of syphilis | Per 1000 person-years |
|-----------------|------------------------|-----------------|-----------------------|
| Overall         | 40                     | 2.96            | 78                    | 1.65                 | 2.08 (1.39–3.09)* |
| 0–1             | 10                     | 75.53           | 11                    | 32.14                | 3.01 (1.15–7.81)* |
| ≥1              | 30                     | 2.90            | 67                    | 1.63                 | 1.82 (1.14–2.84)* |

| No. of genital warts | Per 1000 person-years | No. of genital warts | Per 1000 person-years |
|----------------------|------------------------|----------------------|-----------------------|
| Overall              | 38                     | 3.60                 | 81                    | 2.19                 | 1.91 (1.26–2.83)* |
| 0–1                  | 0                      | 0                    | 0                     | N/A                  |               |
| ≥1                   | 38                     | 3.60                 | 81                    | 2.19                 | 1.91 (1.26–2.83)* |

| No. of gonorrhea | Per 1000 person-years | No. of gonorrhea | Per 1000 person-years |
|------------------|------------------------|-----------------|-----------------------|
| Overall          | 39                     | 2.64             | 74                    | 1.07                 | 2.14 (1.41–3.20)* |
| 0–1              | 7                      | 55.03            | 9                     | 25.86                | 2.62 (0.83–7.90)* |
| ≥1               | 32                     | 2.60             | 65                    | 1.05                 | 2.00 (1.27–3.10)* |

| No. of chlamydial infection | Per 1,000 person-years | No. of chlamydial infection | Per 1,000 person-years |
|----------------------------|-------------------------|-----------------------------|------------------------|
| Overall                    | 68                      | 8.06                        | 183                   | 4.47                 | 1.77 (1.32–2.35)* |
| 0–1                        | 5                       | 45.64                       | 9                     | 30.46                | 1.90 (0.50–6.30)* |
| ≥1                         | 63                      | 6.84                        | 174                   | 4.45                 | 1.47 (1.08–1.97)* |

| No. of trichomoniasis | Per 1000 person-years | No. of trichomoniasis | Per 1000 person-years |
|-----------------------|------------------------|-----------------------|-----------------------|
| Overall               | 121                    | 8.64                  | 357                   | 6.76                 | 1.38 (1.11–1.70)* |
| 0–1                   | 19                     | 140.02                | 34                    | 74.30                | 1.79 (0.97–3.23)* |
| ≥1                    | 102                    | 8.52                  | 323                   | 6.71                 | 1.27 (1.01–1.60)* |

CI = confidence interval, HIV = human immunodeficiency virus.
* Statistical significance.

### Table 3
Analyses of risk factors for sexually transmitted diseases in patients with and without depressive disorder.

| Predictive variables | Univariate analysis | Multivariate analysis |
|----------------------|---------------------|-----------------------|
|                      | HR (95% CI)         | P-value               | HR (95% CI)          | P-value               |
| Depressive disorder  | 1.58 (1.38–1.80)    | <.001                 | 1.54 (1.34–1.76)     | <.001*                |
| Age (<40 = 1, ≥40 = 0) | 1.64 (1.44–1.85)   | <.001                 | 1.69 (1.48–1.92)     | <.001*                |
| Sex (female = 1, male = 0) | 2.23 (1.93–2.57) | <.001                 | 2.20 (1.91–2.54)     | <.001*                |
| Comorbidities        |                     |                       |                       |                       |
| Hypertension         | 0.87 (0.74–1.03)    | .112                  |                       |                       |
| Diabetes mellitus    | 0.98 (0.80–1.20)    | .814                  |                       |                       |
| Dyslipidemia         | 0.91 (0.75–1.09)    | .307                  |                       |                       |
| Coronary artery disease | 0.86 (0.69–1.07) | .182                  |                       |                       |
| Congestive heart failure | 1.03 (0.61–1.75)   | .909                  |                       |                       |
| Cerebrovascular disease | 1.23 (0.99–1.53)  | .059                  | 1.42 (1.14–1.78)     | .002*                 |
| Chronic pulmonary disease | 0.88 (0.70–1.12) | .310                  |                       |                       |
| Degree of urbanization |                     |                       |                       |                       |
| Urban                | Reference            |                       |                       |                       |
| Suburban             | 0.97 (0.85–1.11)    | .665                  |                       |                       |
| Rural                | 1.17 (0.93–1.48)    | .171                  |                       |                       |
| Income group         |                     |                       |                       |                       |
| No income            | Reference            |                       |                       |                       |
| Low income           | 1.10 (0.94–1.28)    | .224                  | 1.22 (1.05–1.43)     | .012*                 |
| Medium income        | 0.86 (0.71–1.04)    | .117                  | 0.90 (0.74–1.06)     | .287                  |
| High income          | 0.67 (0.53–0.86)    | .002                  | 0.84 (0.65–1.07)     | .162                  |

CI = confidence interval, HR = hazard ratio.
drug user, to have early sex age, and to have a greater number of lifetime sex partners. Also, the population of depression patient is in a more complicated environment of sexual initiation, continued sexual activity, choice and number of partners, and contextual prevalence of STDs. Finally, substance use disorders are common comorbidities in patients with depressive disorder and previous works have demonstrated that substance abuse and dependence increase the risk of STIs.

Reviewing the previous study, most of the research focuses on the association between depression and risky sexual behavior, which lack evidence about the relationships of biologically verified STIs. As for our study, we demonstrated the risk of biologically diagnosed STIs following depressive disorder. Besides, the results of past research are controversial, some revealed increased risk of STIs only among depression male, some revealed the increased risk of STIs only among depression female and even some showed no growing risk among depression patient. Reviewing their research method, most of the study focus on the specific groups such as STIs clinical patient, military, black race or Homeless adolescent. The controversial result may be due to the selection of different groups and our study survey the general population of Taiwan which is more extensive than those studies.

In the present study, we conducted a subgroup analysis stratified according to the duration between the diagnosis of depressive disorder and new-onset STIs (Table 2). The results indicated that most STIs included HIV infection, genital warts, gonorrhea, chlamydial infection, and trichomoni-asis remained high for the depression cohort, and the ratios were increased beyond the first year following a depression diagnosis. Patients with depressive disorder are likely to exhibit a higher frequency of outpatient and inpatient visits than the general population, leading to an earlier diagnosis of STIs which cause surveillance bias. However, in our work, the risk ratio for the newly diagnosed HIV infection, genital warts, gonorrhea, chlamydial infection, and trichomoni-asis reached statistically significant within 1 year of depression diagnosis for the depression cohort. Moreover, when patients diagnosed with STIs within 1 year of depression diagnosis were excluded, the risk ratio for the all newly diagnosed STIs included HIV infection, syphilis, genital warts, gonorrhea, chlamydial infection, and trichomoni-asis remained high for the depression cohort, and the ratios were all statistically significant. Thus, this result suggests that the increased risks of HIV infection, genital warts, gonorrhea, chlamydial infection, and trichomoni-asis in depression patients were not caused by surveillance bias.

Our study revealed that female gender compared to male (HR 1.58) and young age <40 years old (HR 1.79) are both risk factors for depression patients to acquire STIs. There are some possible reasons that female gender is a risk factor for STIs in depression patient. According to the previous study, higher rates of sexual risk behaviors in female depression patients than male depression patients was noted. Furthermore, a lack of female-controlled protective devices and the initial phase of STIs is often asymptomatic in women also add to the vulnerability of these patients. As for young age, there are several studies discussed risk factor of STIs in general population and age

Table 4

| Table 4 | Analyses of risk factors for sexually transmitted diseases in patients with depressive disorder. |
| Predictive variables | Univariate analysis | Multivariate analysis |
|----------------------|--------------------|----------------------|
|                       | HR (95% CI)        | P-value | HR (95% CI) | P-value |
| Age (<40=1, ≥40=0)   | 1.64 (1.29–2.98)   | <.001   | 1.58 (1.24–2.01) | <.001* |
| Sex (female=1, male=0) | 1.83 (1.41–2.36)  | <.001   | 1.79 (1.38–2.32) | <.001* |
| Comorbidities        |                    |         |            |         |
| Hypertension         | 0.79 (0.59–1.07)   | .126    |            |         |
| Diabetes mellitus    | 0.97 (0.70–1.39)   | .935    |            |         |
| Dyslipidemia         | 0.85 (0.62–1.17)   | .322    |            |         |
| Coronary artery disease | 0.77 (0.53–1.10) | .147    |            |         |
| Congestive heart failure | 1.12 (0.53–2.30) | .764    |            |         |
| Cerebrovascular disease | 0.77 (0.51–1.15) | .201    |            |         |
| Chronic pulmonary disease | 0.66 (0.43–1.02) | .60    | 0.82 (0.53–1.27) | .365 |
| Degree of urbanization |                  |         |            |         |
| Urban                | Reference          |         |            |         |
| Suburban             | 0.88 (0.68–1.15)   | .357    |            |         |
| Rural                | 1.07 (0.70–1.62)   | .762    |            |         |
| Income group         |                    |         |            |         |
| No income            | Reference          |         |            |         |
| Low income           | 1.17 (0.87–1.56)   | .306    |            |         |
| Medium income        | 0.91 (0.63–1.33)   | .641    |            |         |
| High income          | 0.74 (0.46–1.18)   | .200    |            |         |

**Table 5**

| Table 5 | Hazard ratios of time until sexually transmitted diseases between depressive disorder and control subjects during a 10-year follow-up period. |
|----------------------|----------------------|
|                      | Crude HR (95% CI)     | Adjusted HR (95% CI) |
| HIV infection        | 2.86 (1.59–5.14)†     | 2.93 (1.62–5.30)†    |
| Syphilis             | 2.08 (1.42–3.05)†     | 1.95 (1.32–2.88)†    |
| Genital warts        | 1.91 (1.30–2.81)†     | 1.88 (1.27–2.77)†    |
| Gonorrhea            | 2.13 (1.45–3.14)†     | 1.97 (1.33–2.92)†    |
| Chlamydial infection | 1.51 (1.14–2.00)†     | 1.44 (1.08–1.90)†    |
| Trichomoniasis       | 1.38 (1.12–1.69)†     | 1.33 (1.08–1.64)†    |

**Table 4**

- **Crude HR (95% CI)**: Crude hazard ratio. *Adjusted for age, sex, hypertension, diabetes mellitus, dyslipidemia, coronary artery disease, congestive heart failure, chronic pulmonary disease, cerebrovascular disease, income, and urbanization.

**Table 5**

- **Crude HR (95% CI)**: Crude hazard ratio. *Adjusted for age, sex, hypertension, diabetes mellitus, dyslipidemia, coronary artery disease, congestive heart failure, chronic pulmonary disease, cerebrovascular disease, income, and urbanization.

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- **Crude HR (95% CI)**: Crude hazard ratio. *Adjusted for age, sex, hypertension, diabetes mellitus, dyslipidemia, coronary artery disease, congestive heart failure, chronic pulmonary disease, cerebrovascular disease, income, and urbanization.

**Table 4**

- **Crude HR (95% CI)**: Crude hazard ratio. *Adjusted for age, sex, hypertension, diabetes mellitus, dyslipidemia, coronary artery disease, congestive heart failure, chronic pulmonary disease, cerebrovascular disease, income, and urbanization.

**Table 5**

- **Crude HR (95% CI)**: Crude hazard ratio. *Adjusted for age, sex, hypertension, diabetes mellitus, dyslipidemia, coronary artery disease, congestive heart failure, chronic pulmonary disease, cerebrovascular disease, income, and urbanization.

**Table 4**

- **Crude HR (95% CI)**: Crude hazard ratio. *Adjusted for age, sex, hypertension, diabetes mellitus, dyslipidemia, coronary artery disease, congestive heart failure, chronic pulmonary disease, cerebrovascular disease, income, and urbanization.

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**Table 5**

- **Crude HR (95% CI)**: Crude hazard ratio. *Adjusted for age, sex, hypertension, diabetes mellitus, dyslipidemia, coronary artery disease, congestive heart failure, chronic pulmonary disease, cerebrovascular disease, income, and urbanization.
is one of the significant risk factors for STIs. Young people are at high risk of infection because they may be more vulnerable biologically, lack the social skills to negotiate safe sex, and they lack appropriate health care delivery.

Consistent with previous studies in our work, it revealed that low income (HR 1.22) was significantly associated with an increased sequential risk of STIs. However, we also found that cerebrovascular disease (HR 1.42) was a risk factor of STIs. No epidemiologic studies have investigated the association between cerebrovascular disease and STIs. Further prospective clinical studies on the relationship between STIs and cerebrovascular disease are warranted.

There are several strengths of our study. First, our study design included an unbiased patient selection process. Because participation in the NHI is mandatory and all residents of Taiwan can access healthcare with low copayments, referral bias is low, and follow-up compliance is high. Second, our study is a population-based study including large sample sizes from all hospitals in the country. With small samples or single hospital case studies popular in existing literature, it is difficult to develop such an index with a greater acceptance across the healthcare industry. Third, the data used in the study were derived from the National Health Insurance system in Taiwan. As an observational database, these data reflect current real-world diagnostic patterns.

However, it has several limitations inherent to the use of claims databases that should be considered. First, the NHIRD does not provide detailed information on patients such as numbers of sex partners, sex with promiscuous partners, and condom use. They are all major risk factors for STIs. Thus, we were unable to control for these potentially confounding factors. Second, compared with previous studies regarding the prevalence of STIs in patients with mental disorders, the prevalence of STIs in our work is relatively lower. For example, in a study surveying the prevalence of STIs in psychiatric outpatients, the results showed the prevalence of gonorrhea was 1%, chlamydial infection 3.3%, and T trichomonia 15.7%. Underestimate of the prevalence of STIs in our work should be taken into consideration. Because STIs may be asymptomatic, some patients with STIs may not seek medical treatment and were not diagnosed. Moreover, STI patients may feel stigmatized because of their STIs and did not want to search for medical help. Third, the diagnostic accuracy and severity of the depressive disorder in our work could not be obtained. Whether the severity of the depressive disorder is related to STIs development should be further evaluated.

In conclusion, this nationwide cohort study indicated an increased risk of HIV infection, syphilis, genital warts, gonorrhea, chlamydial infection, and trichomonia among depression patients in Taiwan. Female gender and young age are both risk factors for acquisition of STIs in depression patients. Taking a sexual history in persons with the depressive disorder is, and those engaging in high-risk behavior should be routinely screened for STIs allowing for detection, treatment, and preventive education. Furthermore, prospective studies, particularly those with additional patient-level data, are warranted to confirm our findings.

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Author contributions

Conceptualization: Sheng-Yun Huang, Shyh-Chyang Lee, Cheng-Che Shen.

Data curation: Jeng-Hsiu Hung.

Formal analysis: Jeng-Hsiu Hung, Li-Yu Hu.

Investigation: Li-Yu Hu, Min-Wei Huang.

Methodology: Jeng-Hsiu Hung.

Resources: Min-Wei Huang.

Supervision: Min-Wei Huang, Shyh-Chyang Lee.

Writing – original draft: Sheng-Yun Huang.

Writing – review & editing: Min-Wei Huang, Cheng-Che Shen.

Cheng-Che Shen orcid: 0000-0002-8080-2841.

References

[1] Spijker J, Graaf R Fau - Buij RV, Bijl RV Fau - Beekman AT, Beekman At Fau - Armel J, Ormel J Fau - Nolen WA, Nolen WA Fau - Functional disability and depression in the general population. Results from the Netherlands Mental Health Survey and Incidence Study (NEMESIS). (0001-690X (Print)).

[2] Murray CJ, Lopez AD. Evidence-based health policy-lessons from the Global Burden of Disease Study. (0036-8075 (Print)).

[3] McIntyre RS, Xiao Hx Fau - Sysa K, Sysa K Fau - Vinberg M, et al. The prevalence, measurement, and treatment of the cognitive dimensions/ domain in major depressive disorder. (1179-1934 (Electronic)).

[4] Keefe RS, McClintock Sm Fau - Roth RM, Roth Rm Fau - Doraiswamy PM, Doraiswamy Pm Fau - Tiger S, Tiger S Fau - Madhoo M, Madhoo M. Cognitive effects of pharmacotherapy for major depressive disorder: a systematic review. (1555-2101 (Electronic)).

[5] Jackson JM, Seth P, DiClemente Rj, Lin A. Association of Depressive Symptoms and Substance Use With Risky Sexual Behavior and Sexually Transmitted Infections Among African American Female Adolescents Seeking Sexual Health Care. (1541-0048 (Electronic)).

[6] Lejoyeux M, Arbaretz M Fau - Mcloughlin M, Mcloughlin M Fau - Ades J, Ades J. Impulse control disorders and depression. (0022-3018 (Print)).

[7] Shrier LA, Harris Sk Fau - Sternberg M, Sternberg M Fau - Beardslee W, Beardslee W Fau - Associations of depression, self-esteem, and substance use with sexual risk among adolescents. (0091-7435 (Print)).

[8] Rohde P, Noell J Fau - Ochs L, Ochs L Fau - Seeley Jr, Seeley Jr Fau - Depression, suicidal ideation and STD-related risk in homeless older adolescents. (0140-1971 (Print)).

[9] King C, Feldman J Fau - Waihaka Y, Waihaka Y Fau - Ahan I, et al. Sexual risk behaviors and sexually transmitted infection prevalence in an outpatient psychiatry clinic. (1537-4521 (Electronic)).

[10] GBD 2015 Disease and Injury Incidence and Prevalence COGlobal, regional, and national life expectancy, all-cause mortality, and cause-specific mortality for 249 causes of death, 1980-2015: a systematic analysis for the Global Burden of Disease Study 2015. Lancet (London, England) 2016;388:1459–544.

[11] Owusu-Edusei Kj, Chesson HW, Gift Tl, et al. The estimated direct medical cost of selected sexually transmitted infections in the United States, 2008. Sex Transm Dis 2013;40:197–201.

[12] Dembo R, Krupa J, Warren J, Schneider J, DiClemente Rj, A Multigrouplongitudinal Study of Truant Youths, Marijuana Use, Depression, and STD-Associated Sexual Risk Behavior. (1067-828X (Print)).

[13] Harbertson J, Grillo M Fau - Zimulinda E, Zimulinda E Fau - Murego C, et al. Prevalence of PTSD and depression, and associated sexual risk factors, among male Rwanda Defense Forces military personnel. (1365-3156 (Electronic)).

[14] Holden AE, Shain Rn Fau - Miller Wb, Miller Wb Fau - Piper Jm, et al. The influence of depression on sexual risk reduction and STD infection in a controlled, randomized intervention trial. (1537-4521 (Electronic)).
[15] Hutton HE, Lyketsos Cg Fau - Zenilman JM, Zenilman Jm Fau - Thompson RE, Thompson Re Fau - Erbelding EJ, Erbelding Ej. Depression and HIV risk behaviors among patients in a sexually transmitted disease clinic. (0002-953X (Print)).

[16] Jenkins WD, Botchway A. Young adults with depression are at increased risk of sexually transmitted disease. (1096-0260 (Electronic)).

[17] Khan MR, Kaufman Js Fau - Pence BW, Pence Bw Fau - Gaynes BN, et al. Depression, sexually transmitted infection, and sexual risk behavior among young adults in the United States. (1538-3628 (Electronic)).

[18] Kiene SM, Lule H, Sileo KM, Silmi KP, Wanyenze RK. Depression, alcohol use, and intimate partner violence among outpatients in rural Uganda: vulnerabilities for HIV, STIs and high risk sexual behavior. (1471-2334 (Electronic)).

[19] Shrier LA, Harris Sk Fau - Beardslee WR, Beardslee WR. Temporal associations between depressive symptoms and self-reported sexually transmitted disease among adolescents. (1072-4710 (Print)).

[20] Taniguchi T, Shacham E Fau - Onen NF, Onen Nf Fau - Grubb JR, Grubb Jr Fau - Overton ET, Overton ET. Depression severity is associated with increased risk behaviors and decreased CD4 cell counts. (1360-0451 (Electronic)).

[21] Pratt LA, Xu F Fau - McQuillan GM, McQuillan Gm Fau - Robitz R, Robitz R. The association of depression, risky sexual behaviours and herpes simplex virus type 2 in adults in NHANES, 2005-2008. (1879-0046 (Electronic)).

[22] Lai HM, Cleary M, Sitharthan T, Hunt GE. Prevalence of comorbid substance use, anxiety and mood disorders in epidemiological surveys, 1990-2014: a systematic review and meta-analysis. (1070-4351 (Electronic)).

[23] Zenilman JM, Hook EW3rd, Shepherd M, et al. Alcohol and other substance use in STD clinic patients: relationships with STDs and prevalent HIV infection. Sex Transm Dis 1994;21:220–5.

[24] Tetrault JM, Fiellin DA, Nicolai LM, et al. Substance use in patients with sexually transmitted infections: results from a national U.S. survey. Am J Addict 2010;19:504–9.

[25] Cook RL, Clark DB. Is there an association between alcohol consumption and sexually transmitted diseases? A systematic review. Sex Transm Dis 2005;32:156–64.

[26] DiClemente RJ, Wingood Gm Fau - Crosby RA, Crosby Ra Fau - Sionean C, et al. A prospective study of psychological distress and sexual risk behavior among black adolescent females. (1098-4275 (Electronic)).

[27] Coverdale JH, Bayer TL, McCullough LB, et al. Sexually transmitted disease prevention services for female chronically mentally ill patients. Community Ment Health J 1995;31:303–15.

[28] HIV and STDs: improving services and reducing risk. An interview with AIDS CAP Associate Director Gina Dallabetta, M.D. about the connection between HIV and other STDs, the risk to adolescents, and improving STD services.

[29] Cingolani A, Zona S, Girtardi E, et al. Incidence and factors associated with the risk of sexually transmitted diseases in HIV-infected people seen for care in Italy: data from the Icona Foundation cohort. (1468-1293 (Electronic)).

[30] Falasinnu T, Gilbert M, Hottez TS, Guatathon P, Ogilvie G, Shoveller J. Predictors identifying those at increased risk for STDs: a theory-guided review of empirical literature and clinical guidelines. (1758-1032 (Electronic)).

[31] Ameli N, Bacchetti P, Morrow RA, et al. Herpes simplex virus infection in women in the WIHS: epidemiology and effect of antiretroviral therapy on clinical manifestations. AIDS (London, England) 2006;20:1051–8.

[32] Kelly JD, Cohen J, Grimes B, et al. High rates of herpes simplex virus type 2 infection in homeless women: informing public health strategies. J Women Health (Larchmt) 2016;25:840–5.

[33] King C, Feldman J, Wathaka Y, et al. Sexual risk behaviors and sexually transmitted infection prevalence in an outpatient psychiatry clinic. Sex Transm Dis 2008;35:877–82.