Absence of an Association between Macular Degeneration and Young-Onset Dementia

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Abstract: A few population-based studies have reported an association between prior age-related macular degeneration and senile dementia. No study has explored a possible link between prior macular degeneration and young-onset dementia (YOD). This case–control study aimed to evaluate the association of YOD with prior macular degeneration diagnosed in the 5-year period before their index date. Data for this retrospective observational study were retrieved from Taiwan’s National Health Insurance (NHI) dataset. A total of 36,577 patients with newly diagnosed YOD from January 2010 to December 2017 were identified as the study cohort, assigning their diagnosis date as their index date. Comparison patients were identified by propensity score-matching (three per case, n = 109,731 controls) from the remaining NHI beneficiaries of the period, their index date being the date of their first ambulatory care claim in the year of diagnosis of their matched YOD case. Chi-square test revealed no significant difference in the prevalence of prior macular degeneration between cases and controls (1.1% vs. 1.0%, p = 0.111). Conditional logistic regression analysis also showed an unadjusted odds ratio (OR) for prior macular degeneration of 1.098 among cases relative to controls (95% CI: 0.979–1.232). Adjusted analysis confirmed that YOD was not associated with prior macular degeneration, adjusted odds ratio 1.098 (95% CI = 0.979–1.232). We conclude that patients with macular degeneration are not at increased risk for YOD.

Keywords: young onset dementia; macular degeneration; epidemiology

1. Introduction

Dementia is an increasingly prevalent public health problem. Over 55 million people are estimated to have dementia worldwide, which is expected to increase to 78 million by 2030 [1]. The prevalence of dementia in the United States is 14.4% among the elderly aged over 68 years [2]. The dementia burden continues to grow due to an ageing society worldwide [3]. Dementia causes serious functional impairment at any age, and can be disastrous for patients and families when it sets in at a young age. Young-onset dementia (YOD) is usually defined as dementia with onset before the age of 65 years. The estimated
prevalence rates of YOD in various studies are much lower, about 42–77 per 100,000 population in the 30–65 age-group, and 98–163.1 per 100,000 in the 45–64 group [4–7]. Senile dementia and YOD share many risk factors, such as age [8,9], sex [10,11], smoking [12–14], alcohol use [15–17], stroke [18,19], traumatic brain injury [14,20,21], cardiovascular diseases [22,23], diabetes mellitus [24,25], obesity [26,27], dyslipidemia [28,29], and hereditary factors [30,31].

Age-related Macular Degeneration (AMD), the third leading cause of severe vision loss, is a disease that involves the macular region of the retina, resulting in progressive loss of central vision. In 2016, the number of people with AMD globally was estimated at about 176 million, and this is expected to increase to nearly 288 million by 2040 [32]. Therefore, AMD is a major public health problem causing substantial socioeconomic burden.

A few empirical studies have attempted to investigate the associations between AMD and dementia. One study from the Korean National Health Insurance Service—Health Screening Cohort found that AMD patients had a higher risk for Alzheimer disease (AD) compared to non-AMD participants. They also observed that the association between AMD and AD was sustained even among those with healthy lifestyle behaviors [33]. Another large-scale, population-based study reported that AMD, especially nonexudative AMD, is independently associated with an increased risk of subsequent AD or senile dementia [34]. One study also observed that dementia was associated with a higher likelihood of prior neovascular AMD than comparison patients without dementia [35]. Finally, one systematic review of studies documented during the period of 1959–2018 captured from a wide range of indexing databases (MEDLINE, EMBASE, Web of Knowledge, PsycInfo and the Cochrane database) concluded that AMD was significantly associated with increased risk of AD/cognitive impairment [36].

In contrast, a population-based, cross-sectional study of 1179 participants aged 60 to 80 years from the Singapore Malay Eye study failed to detect a significant independent association between AMD and cognitive dysfunction [37]. Another study from the English National Health Service also found that the likelihood of receiving a diagnosis of AD or other dementia after having had a prior diagnosis of AMD was no different from that expected by chance [38]. In addition, one study using a random sample of the US Medicare population found no association of low scores on the Modified Mini-Mental State Examination, dementia, or AD with early AMD [39]. Therefore, the association between AMD and dementia remains unclear.

A few studies have explored some possible underlying pathogenic mechanisms shared between AMD and senile dementia. One study examining the sequence of a septet of snRNAs (miRNA-7, miRNA-9-1, miRNA-23a/miRNA-27a, miRNA-34a, miRNA-125b-1, miRNA-146a and miRNA-155) and their abundance reported that these were significantly increased and abundant in both the AD-affected superior temporal lobe neocortex and the AMD-affected macular region of the retina [40]. Another study found extensive similarity in the prevailing immune and inflammatory degenerative mechanisms between AMD and AD [41]. One study suggested that AD and AMD share common features such as vitronectin and amyloid-β accumulation, increased oxidative stress, and the apolipoprotein and complement activation pathways [42]. In addition, one study summarized recent findings on the shared characteristics and perspectives between AMD and AD. They reported that an important characteristic common to both diseases is the presence of amyloid β (Aβ) in the senile plaques of the brain among patients with AD and in the drusen of AMD patients [43].

Some uncertainty continues about potential associations between AMD and dementia. Further, although several studies report significant associations between AMD and dementia, to our knowledge, no study has documented the relationship, if any, between macular degeneration and YOD to date. In particular, the noticeably increased incidence of YOD has drawn much attention during the past decade. Uncovering such an association, if any, may contribute to clinical guidelines for follow up care of younger patients with macular degeneration. The present study aimed to document YOD occurrence among a population-
wide cohort of patients diagnosed with macular degeneration using a population-based, retrospective case–control study design.

2. Methods
2.1. Database

Data were retrieved from the Taiwan National Health Insurance (NHI) Research Database (NHIRD). Taiwan implemented its NHI program in 1995. Each year, the Bureau of NHI collects and curates claims data from the NHI program into the NHIRD, followed by de-identification of both patients and medical facilities, rendering it impervious to identification of subjects or institutions by researchers who are provided the data for research. The NHIRD consists of registration files and medical claims data of approximately 99% of Taiwan’s population (about 24.02 million as of December 2021) covered by the NHI program. The registration files include the registries of beneficiaries, contracted medical facilities, board-certified specialists, and medical personnel. The medical claims data include diagnoses, inpatient expenditures by admissions, details of inpatient orders, ambulatory care expenditures by visits, and details of ambulatory care orders. Many researchers in Taiwan have used data from the NHIRD for clinical-epidemiologic studies to identify associations between diseases and treatment outcomes. The NHIRD offers a unique opportunity to explore the association of young-onset dementia with macular degeneration.

The study was approved by the institutional review board of Taipei Medical University (TMU-JIRB N202009055), which complies with the Declaration of Helsinki. Informed consent was waived because the study used retrospective administrative claims data.

2.2. Identification of Cases and Controls

We retrieved 43,582 patients aged <65 years old with a first-time diagnosis of young-onset dementia during an ambulatory care visit between 1 January 2010 and 31 December 2017. We identified them using ICD-9-CM codes 290.0 (senile dementia, uncomplicated), 290.10 (presenile dementia, uncomplicated), 290.11 (presenile dementia with delirium), 290.12 (presenile dementia with delusional features), 290.13 (presenile dementia with depressive features), 290.20 (senile dementia with delusional features), 290.21 (senile dementia with depressive features), 290.3 (senile dementia with delirium), 290.4 (arteriosclerotic dementia), 294.1 (dementia in conditions classified elsewhere), 331.0 (Alzheimer’s disease), 331.1 (Pick’s disease), or ICD-10-CM codes F02.80 (dementia in other diseases classified elsewhere without behavioral disturbance), F02.81 (dementia in other diseases classified elsewhere with behavioral disturbance), F01.50 (vascular dementia without behavioral disturbance), F01.51 (vascular dementia with behavioral disturbance), F03.90 (unspecified dementia without behavioral disturbance), F03.91 (unspecified dementia with behavioral disturbance), and G30 (Alzheimer’s disease). We included patients with a dementia diagnosis in at least two claims during the sample selection period (n = 42,120) in order to increase diagnostic validity, often a concern with administrative datasets. We assigned the first date of dementia diagnosis as the index date. Of them, we further selected 36,577 patients who had at least one ophthalmologist visit within 5 years prior to the index date in order to reduce potential misclassification bias—persons with AMD identified as not having AMD due to not visiting an ophthalmologist.

Matched controls were retrieved out of the remaining NHI beneficiaries with a claim during the study period without a dementia diagnosis. Propensity score matching was used to select three controls per case. We first calculated a propensity score for each enrollee with and without a YOD diagnosis using selected demographic variables potentially associated with the diagnosis (age, sex, monthly income (NTD 0–15,840, NTD 15,841–25,000, ≥ NTD 25,001; the average exchange rate in 2011 was USD 1 ≈ NTD 29), geographic location (Northern, Central, Southern and Eastern) and urbanization level of the patient’s residence (5 levels, 1 most urbanized and 5 least urbanized), and relevant medical comorbidities (hyperlipidemia, diabetes, coronary heart disease, traumatic brain injury, tobacco use disorder, alcohol dependency, obesity, and stroke). We used these selected variables in
a multivariable logistic regression model to predict each selected beneficiary’s expected probability of receiving a dementia diagnosis. Because the calculated propensity score of a control patient may not exactly match that of a YOD subject, we used the method of nearest neighbor within calipers to match controls (a priori value for the calipers is ±0.01). Matching was carried out by defining the first index year of YOD patients as the year of their first YOD diagnosis. We matched three controls to a given patient with YOD based on utilization of any medical service in the index year of the YOD case. We defined each control patient’s index date as the date of their first ambulatory care claim during the matched subject’s year of diagnosis as their index date. The final study sample consisted of 36,577 YOD cases and 109,731 control patients.

2.3. Exposure Assessment

We identified patients with a prior diagnosis of macular degeneration using ICD-9-CM code 362.50 (macular degeneration (senile), unspecified), 362.51 (nonexudative senile macular degeneration), 362.52 (exudative senile macular degeneration), or 362.57 (Drusen (degenerative)) or ICD-10-CM code H35.30 (unspecified macular degeneration), H35.31 (nonexudative age-related macular degeneration) or H35.32 (exudative age-related macular degeneration). We defined patients as having prior macular degeneration if they had at least one claim with a diagnosis of macular degeneration during the 5 years before the index date.

2.4. Statistical Analysis

We carried out statistical analyses using the SAS statistical software (SAS System for Windows, vers. 9.4, SAS Institute, Cary, NC, USA). Descriptive statistics on demographics and comorbidities were summarized by counts and percentages for the YOD cases and control patients. We used chi-square tests to assess differences between cases and controls in demographic characteristics (age, sex, monthly income, geographic location and urbanization level of the patient’s residence) and medical comorbidities (hyperlipidemia, diabetes, coronary heart disease, traumatic brain injury, tobacco use disorder, alcohol dependency, obesity, and stroke). We used multivariable logistic regressions to estimate the odds ratios (ORs) and 95% confidence intervals (CI) for prior macular degeneration among patients with YOD vs. controls after accounting for age, sex, monthly income, geographic location and urbanization level of the patient’s residence, hyperlipidemia, diabetes, coronary heart disease, traumatic brain injury, tobacco use disorder, alcohol dependency, obesity, and stroke. We used two-sided p < 0.05 to determine statistical significance.

3. Results

Table 1 shows the sociodemographic characteristics and medical comorbidities among 36,577 cases and 109,731 propensity score-matched controls. Among the total 146,308 sampled patients, 64.5% were male, the majority (38.3%) resided in northern Taiwan, and only 4.4% were in southern Taiwan. Rural–urban distribution showed that the majority resided in urbanization level 2 communities (28.0%). In addition, most sample patients (45.5%) had a monthly income less than NTD 115,841.

Because we used the propensity score method to match cases to controls, we found as anticipated no statistically significant differences among most of the matching variables: age (p = 0.653), sex (p = 0.925), monthly income (p = 0.913), geographic location (p = 0.981), urbanization level (p = 0.970), hyperlipidemia (21.2% vs. 21.3%, p = 0.509), diabetes (18.2% both groups, p = 0.079), coronary heart disease (7.8% both groups, p = 0.906), traumatic brain injury (22.9% p = 0.980), tobacco use disorder (16.3% vs. 16.4%, p = 0.683), alcohol dependency (3.8% vs. 3.6%, p = 0.111), obesity (18.2% both groups, p = 0.922), and stroke (23.6% both groups, p = 0.725). These results support the appropriateness of the propensity score matching process used.
Table 1. Demographic characteristics of patients with young-onset dementia and control patients in Taiwan \((n = 146,308)\).

| Variable                      | Patients with Young-Onset Dementia \((n = 36,577)\) | Controls \((n = 109,731)\) | \(p\) Value |
|-------------------------------|------------------------------------------------------|----------------------------|-------------|
|                               | Total No. | Percent | Total No. | Percent |                               |
| Males                         | 23,583 | 64.5    | 70,719 | 64.5    | 0.925                          |
| Age                           |          |         |          |         | 0.653                          |
| 18–44                         | 15,218 | 41.6    | 45,507 | 41.5    |                                |
| 45–64                         | 21,359 | 58.4    | 64,224 | 58.5    |                                |
| Monthly income                |          |         |          |         | 0.913                          |
| <NTD 1–15,841                 | 16,637 | 45.5    | 49,909 | 45.5    |                                |
| NTD 15,841–25,000             | 12,226 | 33.4    | 36,578 | 33.3    |                                |
| ≥NTD 25,001                   | 7714    | 21.1    | 23,244 | 21.2    |                                |
| Geographic region             |          |         |          |         | 0.981                          |
| Northern                      | 13,991 | 38.3    | 42,016 | 38.3    |                                |
| Central                       | 9696    | 26.5    | 29,080 | 26.5    |                                |
| Eastern                       | 11,265 | 30.8    | 33,712 | 30.7    |                                |
| Southern                      | 1625    | 4.4     | 4923   | 4.5     |                                |
| Urbanization level            |          |         |          |         | 0.970                          |
| 1 (most urbanized)            | 7865    | 21.5    | 23,735 | 21.6    |                                |
| 2                             | 10,249 | 28.0    | 30,795 | 28.1    |                                |
| 3                             | 6633    | 18.1    | 19,697 | 18.0    |                                |
| 4                             | 5922    | 16.2    | 17,835 | 16.3    |                                |
| 5 (least urbanized)           | 5908    | 16.2    | 17,669 | 16.1    |                                |
| Hyperlipidemia                | 738     | 21.2    | 23,393 | 21.3    | 0.509                          |
| Diabetes                      | 6661    | 18.2    | 20,008 | 18.2    | 0.079                          |
| Coronary heart disease        | 2850    | 7.8     | 8529   | 7.8     | 0.906                          |
| Traumatic brain injury        | 8380    | 22.9    | 25,147 | 22.9    | 0.980                          |
| Stroke                        | 8624    | 23.6    | 25,971 | 23.6    | 0.725                          |
| Alcohol abuse                 | 1382    | 3.8     | 3990   | 3.6     | 0.211                          |
| Tobacco use disorder          | 5953    | 16.3    | 17,959 | 16.4    | 0.683                          |
| Obesity                       | 6661    | 18.2    | 20,008 | 18.2    | 0.922                          |

Note: In 2017, the average exchange rate was USD 1 \(\approx\) New Taiwan Dollar (NTD) 30. SD, standard deviation.

The prevalence of prior macular degeneration among cases and controls is presented in Table 2, showing no significant difference (1.1% vs. 1.0%, \(p = 0.111\)). Univariable logistic regression analysis showed an unadjusted OR for prior macular degeneration of 1.098 among cases relative to controls (95% CI: 0.979–1.232).

Table 2. Prevalence, crude odds ratios (ORs), and 95% confidence intervals (CIs) for prior age-related macular degeneration among patients with young-onset dementia vs. controls.

| Prior Diagnosis of Macular Degeneration | Patients with Young-Onset Dementia \((n = 36,577)\) | Controls \((n = 109,731)\) | \(p\) Value |
|----------------------------------------|------------------------------------------------------|----------------------------|-------------|
| Yes                                    | 399 | 1.1 | 1091 | 1.0 | 0.111 |
| No                                     | 36,178 | 98.9 | 108,640 | 99.0 | 1.000 |
| OR (95% CI)                            | 1.098 (0.979–1.232) | | | |

Notes: The OR was calculated by a logistic regression.

Table 3 shows the results of multivariable logistic regression analysis. After adjusting for the demographic and comorbidity variables, YOD was not significantly associated with prior macular degeneration, adjusted odds ratio 1.103 (95% CI = 0.979–1.232).
Table 3. Covariate-adjusted odds of prior age-related macular degeneration (OR and 95% confidence interval, CIs) among patients with young-onset dementia vs. controls (n = 146,308).

| Variable                        | Presence of Young-Onset Dementia | Adjusted OR | 95% CI       | p Value |
|---------------------------------|---------------------------------|-------------|--------------|---------|
| Prior macular degeneration      |                                 | 1.103       | 0.982–1.238  | 0.535   |
| Males                           |                                 | 0.998       | 0.973–1.023  | 0.851   |
| Age 18–44                       |                                 | 1.000       |              |         |
| Age 45–64                       |                                 | 1.053       | 1.026–1.081  | <0.001  |
| Monthly income                  |                                 |             |              |         |
| <NTD 15,841 (reference group)   |                                 | 1.000       |              |         |
| NTD 15,841–25,000               |                                 | 1.011       | 0.984–1.039  | 0.439   |
| ≥NTD 25,001                     |                                 | 0.988       | 0.957–1.020  | 0.457   |
| Geographic region               |                                 |             |              |         |
| Northern (reference group)      |                                 | 1.000       |              |         |
| Central                         |                                 | 0.997       | 0.964–1.020  | 0.854   |
| Eastern                         |                                 | 0.957       | 0.928–0.987  | 0.005   |
| Southern                        |                                 | 1.007       | 0.946–1.073  | 0.820   |
| Urbanization level              |                                 |             |              |         |
| 1 (reference group)             |                                 | 1.000       |              |         |
| 2                               |                                 | 0.956       | 0.923–0.990  | 0.012   |
| 3                               |                                 | 0.970       | 0.932–1.009  | 0.130   |
| 4                               |                                 | 0.906       | 0.868–0.944  | <0.001  |
| 5                               |                                 | 0.963       | 0.912–1.017  | 0.172   |
| Hyperlipidemia                  |                                 | 0.988       | 0.957–1.020  | 0.459   |
| Diabetes                        |                                 | 0.995       | 0.962–1.029  | 0.763   |
| Coronary heart disease          |                                 | 0.894       | 0.854–0.934  | <0.001  |
| Traumatic brain injury          |                                 | 0.994       | 0.966–1.023  | 0.671   |
| Stroke                          |                                 | 0.985       | 0.956–1.014  | 0.312   |
| Alcohol abuse                   |                                 | 1.046       | 0.982–1.114  | 0.161   |
| Tobacco use disorder            |                                 | 1.006       | 0.973–1.040  | 0.730   |
| Obesity                         |                                 | 0.888       | 0.789–0.999  | 0.048   |

4. Discussion

To our knowledge, this is the first nationwide population-based study exploring a possible association between macular degeneration and subsequent YOD. We found that individuals with prior macular degeneration showed no different risk for YOD (odds ratio 1.103 (95% CI = 0.979–1.232) compared to propensity score-matched controls after adjusting for sex, age group, income, geographical location, urbanization level, hyperlipidemia, diabetes, coronary heart disease, traumatic brain injury, tobacco use disorder, alcohol dependency, obesity, and stroke.

Our findings are consistent with several empirical studies that assessed the association between AMD and dementia. In the Cardiovascular Health Study Cohort recruited from a random sample of the Medicare-eligible files from four US counties, Baker et al. reported no statistically significant associations of dementia or AD with early AMD after adjustment for age, sex, ethnicity, and study center [39]. In another study using data from English national hospital episode statistics, Keenan et al. found no significant difference in the risk of dementia among patients with AMD: the rate ratio was 0.91 (95% CI, 0.79–1.04) when comparing observed vs. expected cases in the AMD cohort based on the rate observed in the reference cohort [38]. Furthermore, in a population-based, cross-sectional study of 1179 participants aged 60 to 80 years from the Singapore Malay Eye study, Ong et al. did not observe a significant association between AMD and cognitive dysfunction after adjusting for age, sex, education level, income category, and type of housing [37]. The above studies all consistently reported no association between AMD and dementia. The
present study confirms these findings of a lack of association between macular degeneration and YOD.

As noted earlier, some contradicting findings are also documented. The study from Korea referenced earlier (Choi et al.) reported that AMD was associated with higher risk of subsequent AD (aHR 1.48, 95% CI 1.25–1.74) when compared to participants without AMD [33]. They suggested that patients with AMD be closely monitored for subsequent development of AD. In another case–control study using the Taiwan National Health Insurance Research Database, Tsai et al. reported that the incidence of AD or senile dementia was higher in patients with AMD than among control patients ($p = 0.044$), with an adjusted hazard ratio (HR) of 1.44 (95% CI, 1.26–1.64) after adjusting for potential confounding factors [34]. Similarly, another study from Taiwan also found that patients with AMD had a 1.23-fold increased risk of developing AD (aHR = 1.23, 95% CI = 1.04–1.46).

Unlike the prior studies, our study focused on young-onset dementia and found no association between macular degeneration and YOD. There are several possible explanations. Some studies have suggested that chronic oxidative stress and neuroinflammation, derangement of the processing and degradation of dysfunctional cellular components, and alterations of neuronal homeostasis are biological pathological mechanisms common to both macular degeneration and dementia. However, the temporal sequence remains unclear. Neuronal cell degeneration in the brain may be followed by retinal degeneration via unrelated pathways, and it remains unknown whether genetic factors, amyloid-β deposition, oxidative stress and the associated mitochondrial and lysosomal dysfunction, or complement-related systems drive the neurodegenerative changes seen in the retina of AMD patients. Additionally, heterogeneous pathogenic mechanisms may contribute to YOD, distinct from those driving the development of senile dementia.

Our study has several strengths. Taiwan’s NHI is a highly accessible and affordable health care system for every citizen (due to negligible copayments and a widely dispersed network of physicians, almost all of them affiliated with the NHI). This minimizes the potential for diagnosis bias arising out of socioeconomic status or residential location. Further, because the NHIRD covers every medical care episode of citizens, about 23 million Taiwanese people, covering outpatient visits, emergency department visits, and inpatient admissions, NHI claims data capture the diagnoses of prior macular degeneration and YOD from all sources. Macular degeneration and YOD are highly concerning conditions that usually cause affected patients to seek medical help, which, in turn, is facilitated by affordable and accessible health care enabled by NHI. Therefore, misclassification bias is unlikely, via differential identification of macular degeneration and YOD based on socioeconomic status. Use of claims data preempts potential recall bias and social desirability bias associated with self-reported data. The case–control study design, selecting controls by propensity score matching, strengthens the study validity by minimizing selection bias and misclassification bias.

There are some study limitations. First, the diagnoses of macular degeneration or YOD were identified from the NHIRD database through the ICD-9-CM or ICD-10-CM codes captured in claims. Therefore, the study sample would have missed those who were not coded accurately. Second, the NHIRD claims lack certain critical items of data, such as physical activity, diet, cognitively challenging life activities, stress, and genetic parameters which are thought to be associated with the development of dementia. Third, the NHIRD database only includes patients who had sought care for symptoms of macular degeneration or YOD. Symptoms of early macular degeneration or YOD are often not recognized until the disease is fairly advanced. Bias from underdiagnosis and undertreatment of macular degeneration or YOD may challenge the validity of findings. Potential non-differential misclassification of YOD would have biased the results toward the null hypothesis. Fourth, patients with macular degeneration may have a great likelihood of being diagnosed with YOD because of surveillance bias, i.e., increased exposure to medical checkups and, therefore, more scrutiny of patient status and referrals (referral bias in clinical populations). As a result, the present study does not rule out the possibility that YOD detection may be enhanced among those
with prior diagnosis of macular degeneration. However, our study shows no association despite possible referral bias, which adds validity to our finding. Finally, the NHIRD lacks data on the severity of macular degeneration, and it is possible that the risk of YOD may be associated with the severity of macular degeneration.

5. Conclusions

Based on medical claims data, our study found no evidence of an association between prior macular degeneration and YOD in Taiwan after adjusting for sex, age group, income, geographical location, urbanization level, hyperlipidemia, diabetes, coronary heart disease, traumatic brain injury, tobacco use disorder, alcohol dependency, obesity, and stroke. Despite being a population-based study, the present findings may not generalize to other countries because of differences in ethnicity and living environment. Our study suggests the need for clinical-epidemiological studies among other ethnicities and regions to extend the findings of our study.

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Informed Consent Statement: Patient consent was waived because this study used administrative data.

Data Availability Statement: Data from the National Health Insurance Research Database, now managed by the Health and Welfare Data Science Center (HWDC), can be obtained by interested researchers through a formal application process addressed to the HWDC, Department of Statistics, Ministry of Health and Welfare, Taiwan (https://dep.mohw.gov.tw/DOS/lp-2506-113.html. accessed on 2 January 2022).

Conflicts of Interest: The authors declare no conflict of interest.

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