Treatment Persistence Between Long-Acting Injectable Versus Orally Administered Aripiprazole Among Patients with Schizophrenia in a Real-World Clinical Setting in Japan

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ABSTRACT

Introduction: Persistence with antipsychotic treatment is critical in managing patients with schizophrenia. To evaluate whether aripiprazole long-acting injection (aripiprazole once-monthly, AOM) can contribute to longer treatment persistence compared with daily orally administered aripiprazole (OA) in real-world clinical settings in Japan, treatment persistence in patients with schizophrenia was compared between patients treated with AOM and those with OA, using a claims database compiled by JMDC Inc., Tokyo, Japan.

Methods: Data of patients with schizophrenia who newly initiated AOM or OA treatment between May 2015 and November 2017 were analyzed. The Cox proportional hazard model was used to estimate the hazard ratio (HR) for treatment discontinuation of AOM vs. OA treatment, adjusted for age, sex, chlorpromazine-equivalent dose of antipsychotics, and the number of psychiatric hospitalizations.

Results: The analysis included 198 patients in the AOM group and 1240 patients in the OA group (mean age 38.4 ± 11.9 years and 39.3 ± 12.4 years, respectively). The AOM group was significantly less likely to discontinue treatment than the OA group (adjusted HR 0.54, 95% confidence interval [CI] 0.43–0.68). When using the tolerable patients extracted from the OA group (i.e., patients with at least two OA prescriptions; n = 983) vs. the whole AOM group, AOM users were again significantly less likely to discontinue treatment (adjusted HR 0.67, 95% CI 0.53–0.86).

Conclusion: AOM was associated with longer treatment persistence than OA in the antipsychotic treatment of patients with schizophrenia in real-world clinical settings in Japan, suggesting that the use of AOM may contribute to longer antipsychotic treatment.

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Keywords: Antipsychotic; Aripiprazole once-monthly; Continuation rate; Japan; Long-acting injectable; Schizophrenia

Key Summary Points

Why carry out this study?

Poor adherence leads to worse prognosis, longer time to remission, and increased risks of relapse and hospitalization in patients with schizophrenia

Although the use of an antipsychotic long-acting injection (LAI) is recommended as a way of improving medication adherence, LAIs remain underused in Japan

This study evaluated whether patients with schizophrenia receiving the LAI formulation of aripiprazole (aripiprazole once-monthly, AOM) indeed have better treatment persistence than those receiving orally administered aripiprazole (OA) in real-world clinical settings in Japan

What was learned from the study?

Patients treated with AOM had significantly longer treatment persistence than those treated with OA

The use of AOM may contribute to prolonging treatment persistence among patients with schizophrenia, for whom such persistence is extremely important

INTRODUCTION

Antipsychotic treatment is the standard treatment in patients with schizophrenia, which aims not only to improve acute symptoms but also to maintain long-term stabilization of symptoms and prevent relapses. However, nonadherence is quite common among patients with schizophrenia [1], with the prevalence of nonadherence being 19.2–73% in 21 studies [2]. As poor adherence leads to worse prognosis, longer time to remission, and increased risks of relapse, hospitalization, and suicide [1, 3, 4], improvement of adherence to antipsychotic treatment or achievement of treatment persistence is important for the treatment of patients with schizophrenia.

Practice guidelines recommend the use of long-acting injectable (LAI) antipsychotics as a way of improving medication adherence [5, 6]. LAIs, which are commonly administered on a biweekly or monthly basis, reduce the burden of daily oral antipsychotic use and make early detection of nonadherence easier [7]. LAIs have at least similar treatment outcomes (e.g., relapse prevention) to oral antipsychotics [8, 9], with some studies reporting superiority of LAIs over oral treatments [10–12]. In addition, patients who initiated LAIs were reportedly more adherent and had longer time to treatment discontinuation than those receiving oral antipsychotics in several non-randomized studies [7, 13–17]. Although LAIs used to be reserved for patients with frequent relapses or poor adherence [5], the early use of LAIs is increasingly being advocated [10, 18, 19] and recommended for consideration as a first-line treatment for any patient for whom long-term antipsychotic treatment is needed [20]. Nevertheless, LAIs have been underused as a result of lack of knowledge and negative attitudes about LAIs among clinicians and patients [10, 18].

Aripiprazole is a second-generation antipsychotic with a high affinity partial agonism of dopamine D2 receptors and serotonin 5-HT1A receptors and an antagonism of 5-HT2A receptors [21], with efficacy in the acute and long-term treatment, and low incidence of side effects such as weight gain, hyperprolactinemia, and metabolic adverse events [22]. A once-monthly LAI formulation of aripiprazole (aripiprazole once-monthly, AOM) was developed to improve patients’ adherence, which was first approved in the USA in 2013 [23] and later marketed in Japan in May 2015 [24]. AOM was shown to have comparable efficacy and tolerability to orally administered aripiprazole (OA) in an Asian randomized study [8]; however, there was no significant difference in treatment...
persistence between the two formulations. Outside Asia, a few studies previously suggested the advantage of AOM for the longer treatment persistence over oral antipsychotics in real-world clinical settings [15, 16], but these were not direct comparisons with OA. If real-world evidence suggests such an advantage of AOM over OA in long-term treatment persistence, it would add to clinicians’ knowledge and may change their attitudes toward the use of the formulation, which could potentially encourage its use in clinical practice. Therefore, in this study, we compared the treatment persistence between patients with schizophrenia who initiated AOM and those who initiated OA in real-world clinical settings in Japan, using a large-scale claims database.

METHODS

Study Design and Data Source

This was a retrospective, observational cohort study in patients with schizophrenia who newly initiated treatment with AOM or OA, using a claims database compiled by JMDC Inc., Tokyo, Japan. The JMDC database is based on insured persons’ claims (in-patient, outpatient, and pharmacy) received from medical institutions, and it includes data of about four million persons as of May 2019. The individuals included in the database are company employees and their dependents insured by health insurance societies. Contained data, which were securely anonymized, included patient demographics (age, sex), diagnoses, prescriptions, medical procedures, costs, and medical institutions (e.g., bed size). A unique identifier is assigned to each person to link their claims records across institutions, which enables one to track their movement and treatment across institutions as long as they are covered by health insurance societies. The JMDC database has been used in various areas of medical research in Japan, including studies in patients with schizophrenia [25, 26]. The present study used data from May 1, 2014 to November 30, 2018 (entire data period). This study was approved by the ethics committee of the Research Institute of Health Data Science (RI2019001) and was conducted according to the principles of the Declaration of Helsinki.

Patient Selection

Patients aged 18 years or more with a confirmed diagnosis of schizophrenia (International Statistical Classification of Diseases and Related Health Problems 10th Revision [ICD-10] code F20) who had a prescription record of AOM or OA (index drug) between May 1, 2015 and November 30, 2017, with no prescription records of the index drug in the 365 days prior to index date were identified within the database. This time frame, beginning in May of 2015, was chosen to compare the two treatments after AOM became available in Japan. Index date was the first date of index drug prescription with a diagnosis record of schizophrenia (Fig. 1). To target patients who had been receiving antipsychotic treatment continuously, patients were required to have both a diagnosis record for schizophrenia and a prescription record for antipsychotics on admission to hospital or at two or more outpatient visits in the 365-day pre-index period. In this study, both the diagnosis and the prescription records were considered conditions for patient identification. This approach was adopted to reduce possible misclassification in a previous claims database study of schizophrenia [27].

The following patients were excluded: (1) patients with less than a 150 mg/day chlorpromazine (CP)-equivalent dose of antipsychotics and also with a prescription record of antidepressants in the 365-day pre-index period; and (2) patients with a diagnosis record of dementia (ICD-10 codes: F00–03, F05.1, G30, G31.1), autism (F84), or intellectual disability (F70–79) in the month of index date or during the 12-month pre-index period. Patients with a prescription record of AOM any time in the entire data period were also excluded from the OA group. The concomitant use of multiple antipsychotics, irrespective of the formulation, was not an exclusion criterion in this study, excepting the use of AOM in the OA group.
Outcomes

The primary outcome was the duration of index drug treatment. For OA, patients were assumed to have taken aripiprazole for the number of days prescribed recorded on claims. For AOM, patients were assumed to have been on medication for 28 days per prescription. Discontinuation was defined as at least 60 days absence of subsequent prescriptions following the last day of the days prescribed. Treatment persistence was defined as the number of days from index date to the last day of the days prescribed by the latest prescription before discontinuation (Fig. 1). When the 60-day gap could not be confirmed because of the lack of data (e.g., end of the data period or withdrawn from the database), treatment was censored at the last day of data observed, and treatment persistence was defined to end on that day. In sensitivity analyses, the at least 30-day and at least 90-day gaps were used as definitions of discontinuation. For AOM, 28 days per prescription. For OA, the number of days prescribed recorded on claims. AOM aripiprazole once-monthly, OA orally administered aripiprazole.

Fig. 1 Study time frame. Index date is the earliest date of prescription of the index drug (AOM or OA) for the treatment of schizophrenia within the identification period. *In sensitivity analyses, at least 30 days and at least 90 days were used as definitions of discontinuation. bFor AOM, 28 days per prescription. For OA, the number of days prescribed recorded on claims. AOM aripiprazole once-monthly, OA orally administered aripiprazole.

Analysis Sets

The analysis sets in this study were defined as follows:

1. Overall analysis set (AOM, \( n = 198 \); OA, \( n = 1240 \)), consisting of all patients who met all inclusion criteria without meeting any exclusion criteria.

2. Tolerable OA subset (\( n = 983 \)), consisting of patients with at least two prescription records of OA among the overall analysis set of the OA group.
3. No-antidepressant subset (AOM, n = 152; OA, n = 837), consisting of patients without prescriptions for antidepressants in the 365-day pre-index period.
4. Maintenance dose subset (AOM, n = 193; OA, n = 917), consisting of patients with at least a 150 mg/day CP-equivalent dose of antipsychotics in the 365-day pre-index period. A dose of 150 mg/day is the minimum maintenance dose of OA for schizophrenia in Japan.

Statistical Analysis

Patient characteristics were descriptively summarized for each group. The Cox proportional hazard model was used to compare the treatment persistence between AOM and OA, and hazard ratios (HRs) for treatment discontinuation and 95% confidence intervals (CIs) were computed. The multivariable model included the following background characteristics of patients as covariates: age (less than 40, at least 40 years), sex, CP-equivalent dose of antipsychotics (less than 600, at least 600 mg/day), and the number of psychiatric hospitalizations (0, at least 1). In addition, adjusted Cox regression survival curves were generated to depict the treatment continuation rate in each treatment group.

The AOM group was assumed to include tolerable patients because tolerability of aripiprazole must be established with OA prior to the initiation of AOM [24], whereas the tolerability was unknown among patients initiating OA. To reduce potential bias, we also conducted the same analysis as above using the tolerable OA subset and compared their treatment persistence against the overall AOM group. In this subset, the treatment persistence was defined to start at the second prescription date of OA, not at index date.

In addition, we performed two subgroup analyses to further examine the treatment persistence between AOM and OA by eliminating potential confounding effects from the unintended use of the index drug. First, the no-antidepressant subset was analyzed to eliminate the potential use of aripiprazole as an augmentation therapy for depression. Second, the maintenance dose subset was analyzed to eliminate patients on a dose below the maintenance dose of antipsychotics for schizophrenia treatment. This suggests a potential use of aripiprazole for depression or other treatment purposes.

The level of significance was set at \( p < 0.05 \) (two-sided). All statistical analyses were conducted using SAS 9.4 (SAS Institute Inc., Cary, NC). Statistical analyses were independently conducted by JMDC Inc. (Tokyo, Japan).

RESULTS

Patient Characteristics

The present study included 198 patients for the AOM group and 1240 patients for the OA group (= overall analysis set; Fig. 2). The background characteristics of patients are summarized in Table 1. The age and sex distribution was similar between the AOM and OA groups (mean age 38.4 ± 11.9 years vs. 39.3 ± 12.4 years, respectively; proportion of women, 63.1% vs. 59.4%, respectively). In the index month, OA users had comorbid depressive disorders more commonly than AOM users (41.1% vs. 27.8%, respectively) and used more antidepressants (25.0% vs. 12.6%, respectively). The mean CP-equivalent dose of antipsychotics during the 365-day pre-index period was 626.5 ± 481.6 mg/day in the AOM group and 346.3 ± 369.0 mg/day in the OA group. During the pre-index period, patients in the AOM group had experienced psychiatric hospitalizations more than those in the OA group (41.9% vs. 25.0%, respectively). The proportions of patients who were hospitalized at index date were similar between the AOM and OA groups (20.2% vs. 17.2%, respectively).

Treatment Persistence

The analyses using the Cox proportional hazard models revealed that patients treated with AOM were significantly less likely to discontinue treatment (adjusted HR 0.54, 95% CI 0.43–0.68; Table 2 and Fig. 3a). When the tolerable OA
subset was used, AOM users were again significantly less likely to discontinue treatment (adjusted HR 0.67, 95% CI 0.53–0.86; Fig. 3b). In our sensitivity analyses using the overall analysis set, the alternative definitions of at least 30-day and at least 90-day gaps as treatment discontinuation both yielded similar results showing significantly lower risk for treatment discontinuation in the AOM group (Supplementary Table S1).

The subgroup analysis in the no-antidepressant subset demonstrated that patients treated with AOM were still significantly less likely to discontinue treatment than those treated with OA (adjusted HR 0.53, 95% CI 0.40–0.69). Another subgroup analysis using the maintenance dose subset also revealed that AOM users were significantly less likely to discontinue treatment than OA users (adjusted HR 0.54, 95% CI 0.43–0.69).

**DISCUSSION**

This is a unique study comparing the treatment persistence in patients with schizophrenia between AOM and OA in real-world clinical settings in Japan. A meta-analysis of randomized controlled trials (RCTs) on AOM previously reported the lower risk for all-cause discontinuation in AOM vs. OA (relative risk [RR] 0.78, 95% CI 0.64–0.95) [29]. However, because RCTs, which are conducted under strict conditions,
Table 1  Background characteristics of patients in the AOM and OA groups (overall analysis set)

|                        | AOM  (n = 198) | OA  (n = 1240) |
|------------------------|----------------|----------------|
| Age, years             | 38.4 ± 11.9    | 39.3 ± 12.4    |
| Female, n (%)          | 125 (63.1)     | 736 (59.4)     |
| Psychiatric comorbidities in index month, n (%) | | |
| Depressive disorder    | 55 (27.8)      | 510 (41.1)     |
| Bipolar disorder       | 35 (17.7)      | 331 (26.7)     |
| ADHD                   | 1 (0.5)        | 26 (2.1)       |
| Concomitant psychotropics in index month, n (%) | | |
| Antidepressants        | 25 (12.6)      | 310 (25.0)     |
| Mood stabilizers       | 43 (21.7)      | 316 (25.5)     |
| Anxiolytics            | 57 (28.8)      | 440 (35.5)     |
| Hypnotics              | 107 (54.0)     | 710 (57.3)     |
| DZP-equivalent of benzodiazepines at index date, mg/day | | |
| Mean ± SD              | 2.1 ± 5.4      | 9.6 ± 15.1     |
| > 0, n (%)             | 35 (17.7)      | 728 (58.7)     |
| Inpatients at index date, n (%) | 40 (20.2)     | 213 (17.2)     |
| During 365-day pre-index period | | |
| Comorbidity scorea    | 0.5 ± 1.0      | 0.5 ± 1.1      |
| CP-equivalent dose of antipsychotics, mg/day | | |
| Mean ± SD              | 626.5 ± 481.6  | 346.3 ± 369.0  |
| Dose category, n (%)   | | |
| < 150                  | 5 (2.5)        | 323 (26.0)     |
| ≥ 150, < 300           | 33 (16.7)      | 437 (35.2)     |
| ≥ 300, < 600           | 77 (38.9)      | 282 (22.7)     |
| ≥ 600                  | 83 (41.9)      | 198 (16.0)     |
| PDC for antipsychotics | | |
| Mean ± SD              | 0.79 ± 0.27    | 0.79 ± 0.26    |
| ≥ 0.8, n (%)           | 133 (67.2)     | 830 (66.9)     |
| Number of psychiatric hospitalizations | | |
| Mean ± SD              | 0.5 ± 0.7      | 0.3 ± 0.5      |
| ≥ 1, n (%)             | 83 (41.9)      | 310 (25.0)     |
would not reflect the reality of daily clinical practice, real-world evidence should confirm such a potential advantage of AOM over OA in treatment persistence. In this claims data study, we found that patients who initiated AOM were, indeed, less likely to discontinue treatment compared with those who initiated OA in routine clinical practice in Japan (adjusted HR 0.54, 95% CI 0.43–0.68). Our findings are consistent with trends observed in other claims data studies showing that AOM users in the USA are less likely to discontinue treatment (HR 0.69, 95% CI 0.60–0.79) [15] and have significantly longer time to discontinuation than oral antipsychotic users [16].

Although our results showed the advantage of AOM in treatment continuation, there were some biases to be considered when interpreting the results of this study. For example, the tolerability of aripiprazole had not been confirmed prior to treatment initiation among the OA group, whereas it had most probably been confirmed before initiating AOM. Because of this, early dropouts due to intolerability must have contributed to the lower continuation rates among the OA group. However, our analysis using tolerable patients extracted from the OA group still demonstrated significantly lower risk for treatment discontinuation in the AOM treatment (adjusted HR 0.67, 95% CI 0.53–0.86). The presence of at least two prescriptions does not necessarily confirm the tolerability; however, this condition could at least exclude some OA-administered patients who discontinued treatment early, e.g., after initial administration. Thus, we consider that the results of this analysis corroborate our finding.

OA is also indicated for major depressive disorders with inadequate response to antidepressant therapy [30], and its efficacy as an augmentation therapy has been well documented [31, 32]. Considering the high

### Table 1 continued

|                                | AOM (n = 198) | OA (n = 1240) |
|--------------------------------|---------------|---------------|
| Total length of hospital-stay, days | 43.2 ± 78.3  | 20.7 ± 53.4  |
| Time intervals between antipsychotic prescriptionsb, days | 31.6 ± 34.6  | 28.9 ± 27.4  |
| Use of other LAI antipsychotics, n (%) | 33 (16.7)    | 38 (3.1)     |
| Time from the oldest OA prescription to index date, days | 204.3 ± 147.1| –             |
| Dose of OA at most recent prescription, mg/day | 13.2 ± 8.2   | –             |

Data are expressed as mean ± SD or n (%)

ADHD attention-deficit hyperactivity disorder, AOM aripiprazole once-monthly, CP chlorpromazine, DZP diazepam, LAI long-acting injectable, OA orally administered aripiprazole, PDC proportion of days covered, SD standard deviation

a Comorbidity score was calculated on the basis of the Charlson Comorbidity Index but excluded AIDS/HIV because of the unavailability of data

b Time intervals between outpatient prescriptions of antipsychotics during the 365-day pre-index period were calculated

### Table 2 Hazard ratios for treatment discontinuation (overall analysis set)

|                                | Unadjusted model | Adjusted modela |
|--------------------------------|------------------|-----------------|
|                                | HR (95% CI)      | p value         | HR (95% CI)      | p value         |
| AOM vs. OA                     | 0.50 (0.40–0.63) | < 0.0001        | 0.54 (0.43–0.68) | < 0.0001        |

AOM aripiprazole once-monthly, CI confidence interval, CP chlorpromazine, HR hazard ratio, OA orally administered aripiprazole

a Adjusted for age, sex, the CP-equivalent dose of antipsychotics (mg/day), and the number of psychiatric hospitalizations
prevalence of comorbid depressive disorders in the OA group (41.1%), the possible use of OA as augmentation therapy for depressive disorders may have been responsible, in part, for the shorter treatment persistence in the OA group. To address this point, we conducted a subgroup analysis using patients without prescriptions of antidepressants, for whom OA must have been administered exclusively for the treatment of schizophrenia. However, even among these patients, we found that AOM users still had a significantly lower risk for treatment discontinuation than OA users (adjusted HR 0.53, 95% CI 0.40–0.69). Similarly, patients with only less than the maintenance dose of antipsychotics (26.0% in the OA group vs. 2.5% in the AOM group), who had probably received aripiprazole for other treatment purposes, might also have

Fig. 3 Adjusted Cox regression survival curves for treatment discontinuation in each treatment group. a AOM and OA groups were compared. b The tolerable OA subset was compared against the AOM group. Adjusted for age, sex, the CP-equivalent dose of antipsychotics (mg/day), and the number of psychiatric hospitalizations. AOM aripiprazole once-monthly, CP chlorpromazine, OA orally administered aripiprazole

△ Adis
influenced the results of this study. However, again, the analysis after excluding these patients also showed that AOM users were still significantly less likely to discontinue treatment than OA users (adjusted HR 0.54, 95% CI 0.43–0.69). Altogether, we reasoned that these results supported the advantage of AOM over OA in treatment continuation even when considering these potential biases.

In Japan, LAIs remain underused [27] probably because of negative attitudes toward the use of LAIs among Japanese psychiatrists, such as little experience with LAIs, concerns for switching, fear of pain, and high drug costs [33]. However, a recent Japanese study reported that patients treated with LAIs had favorable attitudes toward LAIs regarding side effects, relapse prevention, efficacy, pain, and costs [34]. Considering that LAI antipsychotics offer a simpler mode of administration than oral antipsychotics, with the latter having more barriers to treatment adherence (e.g., hospital visit for a prescription form, pharmacy visit for drug purchase, daily drug intake), some patients may favor LAI formulations for simplicity. Thus, the use of LAIs should be encouraged among both psychiatrists and patients, beyond the cases of poor adherence and high risks for recurrence [18, 35, 36]. We expect that the results of this study add to the knowledge on AOM among Japanese psychiatrists, thereby contributing to its broader use in clinical settings.

The focus of this study was on the difference in treatment continuation between two formulations of aripiprazole; however, the observed trend favoring an LAI formulation over its oral counterpart was consistent with a previous report that included haloperidol, perphenazine, and risperidone [37]. This also paralleled the trend of better adherence with risperidone LAI compared with orally administered risperidone [12]. Our results would also add to the benefits of LAIs compared with oral antipsychotics in terms of a longer treatment duration, previously reported for paliperidone, risperidone, and haloperidol [7, 14, 15]. Now that the advantages of LAIs over oral antipsychotics in treatment persistence seem to be evident, the next research topic may be a comparison among LAIs. Since there are several LAI antipsychotics available in Japan, it would be worth investigating the benefits of an AOM relative to other LAI antipsychotics in a future study.

There were several limitations in this study. First, because the JMDC database only includes insured individuals covered by health insurance societies, patients with different socioeconomic background, who may be covered by other health insurance policies, and those aged 65 years and older were quite limited in our study population. Therefore, the generalizability of our results to the overall patients with schizophrenia in Japan may be limited. Second, as data were originally recorded for the purpose of claims rather than research, it is possible that recorded diagnosis and treatment may be inaccurate, influencing the results of this study. Third, it has been suggested that an AOM is a more direct indicator of medication adherence, as the prescription record most probably indicates actual injection of the AOM. In contrast, the process of drug administration for the OA group is complicated [38]. Given the potential failures of drug intake in the OA group, the between-group gap in continuation rates may have been greater. Fourth, there were some differences in patients’ background characteristics between the AOM and OA groups, probably reflecting the selection bias in real-world clinical settings, which suggest the potential of confounding. Our subgroup analyses showed that the results were consistent even after excluding patients with antidepressants and the extremely low CP-equivalent dose of antipsychotics, demonstrating that the impact of differences in these two variables was not substantial. The same trend was also observed when other additional, potentially relevant, patient characteristics were considered in our exploratory analysis (see Supplementary Table S2). However, there might have been other factors influencing the results of this study, such as patients’ severity or psychiatrists’ attitudes toward the use of LAIs. We could not assess the effects of these factors because of the absence of such data within the database; these are worth investigating in future research to confirm the result of this study. Finally, and most importantly, we were unable to evaluate the reasons for treatment discontinuation or
switching because of the absence of data. A future study should clarify such reasons or causes, which would help develop strategies to improve patients’ adherence to treatment.

CONCLUSIONS

This large-scale claims data study showed that patients with schizophrenia receiving AOM had significantly longer treatment persistence with an adjusted HR for treatment discontinuation of 0.54, compared with those receiving OA in real-world clinical settings in Japan. Although there were some important biases and study limitations to be considered, additional analyses, including sensitivity and subgroup analyses, yielded consistent results, supporting the advantage of AOM over OA in treatment continuation in routine practice. The use of AOM may contribute to prolonging treatment persistence among patients with schizophrenia, for whom such persistence is extremely important.

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Compliance with Ethics Guidelines. This study was approved by the ethics committee of the Research Institute of Health Data Science (reference no. RI2019001) and was conducted according to the principles of the Declaration of Helsinki. Informed consent was not required for this retrospective database study because this study exclusively used de-identified claims data without involving any human participants.

Data Availability. The dataset generated during and/or analyzed during the current study are not publicly available because it was purchased from a commercial provider, but is available from the corresponding author on reasonable request.

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