Longitudinal Assessment of Pain Management Among the Employed Japanese Population with Knee Osteoarthritis

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Purpose: To assess comorbidity burden and pain-management patterns among working-aged patients with knee osteoarthritis only (KOA/O) and patients with knee osteoarthritis plus osteoarthritis at another site (KOA/+) in Japan.

Patients and Methods: Retrospective claims data analysis was conducted using the Japan Medical Data Center database. Working-aged adults (aged 40 to 71 years) with 5 years of follow-up and diagnosed with knee osteoarthritis (KOA) between January 1, 2011, and December 31, 2012, were evaluated. The first claim with a KOA diagnosis defined the index date. Patients were divided into two mutually exclusive cohorts: KOA/O and KOA/+.

Results: A total of 2542 patients met study criteria: 1575 KOA/O and 967 KOA/+. Mean age and number of comorbidities were higher among the KOA/+ versus KOA/O cohort. Pharmaceutical treatment was received by 91.5% of patients in the KOA/+ compared with 85.1% of patients in the KOA/O cohort during the first year of follow-up. The most common pharmacological treatment received during the first year of follow-up was either topical or oral nonsteroidal anti-inflammatory drugs for both cohorts. During each year of follow-up, the KOA/+ cohort had greater proportion of patients with at least one health-care encounter (ie, hospital admissions, outpatient and pharmacy visits) and higher direct medical costs compared with the KOA/O cohort.

Conclusion: This study demonstrates that a greater proportion of the working population with KOA/+ received pain-related treatment compared with patients with KOA/O. Further studies are necessary to evaluate appropriate pain management for both KOA only and KOA with other sites.

Keywords: knee osteoarthritis, pain management, Japan, working-aged, HCRU

Introduction

Osteoarthritis (OA) is a widely prevalent chronic and progressive joint disease characterized by cartilage degradation, bone remodeling, osteophyte formation, joint inflammation, and loss of normal joint function, which commonly affects weight-bearing joints such as the knees, hips, and spine as well as hands and feet.1-3 OA can affect single or multiple joints, causing acute and chronic pain, swelling, stiffness, restriction of movement, and, in extreme cases, even permanent disability, with corresponding impacts on overall quality of life and functionality.1,2,4 Prevalence of radiographically confirmed knee OA (KOA) in Japan has been reported to be 42.0% in males and 61.5% in females aged over...
40 years. Previous epidemiological studies have shown that patients aged 50 years and older often experience pain in multiple joints (median of six joints), with pain due to KOA being the most common complaint. Previous studies have reported that the risk of pain increases dramatically for those patients with KOA with concomitant feet, back, or hip problems compared with patients with KOA only. A UK-based postal survey of 18,474 adults aged ≥50 years who registered with primary care general practices also reported that as the number of joints affected by OA increases, the likelihood that the disease is debilitating also increases.

At the present time, no definitive treatment option exists for OA and the severity of the condition differs across individual patients. Current options for OA-related pain management may include pharmacological and nonpharmacological modalities, alone or in combination. Historically, guidelines for the nonsurgical treatment of KOA have stratified patients into groups with single- or multiple-joint OA or with or without the presence of comorbidities; these patient groups highlight the need to assess pain management and comorbidity patterns separately among patients with single-joint and multiple-joint KOA. Core nonpharmacological pain-management modalities may include weight reduction, rehabilitation, and physical therapy, hydrotherapy, aerobic exercises, and yoga. Correspondingly, pain relievers, anti-inflammatory drugs, corticosteroids, and intra-articular hyaluronic acid (IAHA) injections are some of the most commonly used pharmacological modalities in Japan.

Despite multiple treatment options available for the management of pain associated with OA, treatment satisfaction remains low. A web-based survey evaluated treatment satisfaction in Japanese patients with KOA and reported a medication treatment satisfaction rate of 50 as measured by Treatment Satisfaction Questionnaire for Medication global score (on a scale of 0 to 100; higher scores indicate higher satisfaction). Early diagnosis and treatment of early-stage KOA at a younger age become critical to delay the progression of highly degenerative KOA. While surgical treatment (eg, total knee arthroplasty) may significantly decrease pain symptoms and improve range of motion for patients with KOA, it is recommended only for advanced KOA or when conservative treatment is ineffective considering benefit-risk balances. Early stage KOA, as is commonly seen in the working population, currently has both a lack of evidence and a lack of options for effective pain-management.

Furthermore, with the majority of publications focusing on only single-joint OA, data regarding patients with KOA with multiple-site joint problems are limited. The hypothesis of this study is that patients with multiple-joint KOA will have a greater treatment burden compared with patients with single-joint OA. To address the noted literature gap and to provide data for patients with both single- and multiple-site joint OA, we conducted a real-world retrospective database study in which the primary objectives were to assess the longitudinal use of pain-management options among working-aged patients with KOA only (KOAO) compared with patients with KOA plus OA at another site (KOAO/) in Japan. Patient demographic and clinical characteristics as well as longitudinal health-care resource utilization and costs were also reported.

### Patient Selection and Study Cohorts

The cohort was selected from the population of patients aged over 40 years with an “International Classification of Diseases, Tenth Revision, Clinical Modification” (ICD-10-CM) diagnosis of KOA (ICD-10-CM code M17) between January 1, 2011, and December 31, 2012. The date of the first observed KOA diagnosis between January 1, 2011, and December 31, 2012, defined the index date. All patients were required to have a minimum of 1 year of enrollment before the index date (ie, baseline period) and 5 years of enrollment after the index date (ie, follow-up period). To ensure sample specificity, patients were further required to have at least one additional KOA diagnosis at any point during the follow-up period and no diagnoses for KOA during the baseline period. To avoid the likelihood of skewed health-care resource use (HCRU) and cost estimates in the two cohorts of interest, patients with...
a diagnosis of cancer during the 5-year follow-up period were excluded from the study.

This study compared the cohort of patients with KOA/O to the cohort of patients with KOA/+. Definitions for these cohorts are as follows:

KOA/+ cohort: patients were included in this cohort if they had at least one diagnosis of OA at another site in the 12 months before or after the index date. ICD-10-CM codes for other OA sites were M16, M18, M19, M40, M41.8, M43.0, M43.1, M47.8, and M47.9.

KOA/O cohort: patients were included in this cohort if they did not have any diagnosis of OA at another site in the baseline and follow-up periods.

As the patient data used in this study were anonymized, the Ethical Guidelines for Medical and Health Research Involving Human Subjects (Ministry of Education, Culture, Sports, Science and Technology and Ministry of Health, Labour and Welfare of Japan) were not applicable to this study.

Study Measures and Definitions

The study evaluated patient demographics, clinical characteristics, pain-management patterns, HCRU, and direct medical costs. Baseline characteristics measured at the index date included age, sex, and Charlson Comorbidity Index (CCI). Other comorbidities that may have affected OA-related treatment defined by Osteoarthritis Research Society International (OARSI) guidelines are described in Supplementary Table A-1 and were examined during each year of the follow-up period. The distribution of physician department (eg, orthopedic surgeons, rheumatologists, anesthesiologists, internal medicine/general practitioners) and the type of setting (eg, public hospital, university hospital, clinic) for the KOA diagnosis on the index date was reported. Among patients with OA at other sites within 1 year before and after the index date, the site of other OA diagnosis was reported. Longitudinal pain-management patterns were assessed for the two cohorts of interest. The number and percentage of patients with broad categories of treatment (ie, nonpharmacological treatment modalities, pharmaceutical treatment, injectable treatment, surgery) at least once during each year of the follow-up period were evaluated. Nonpharmacological treatments included physical therapy and manual/instrumental therapy. Pharmaceutical and injectable (ie, corticosteroids, intra-articular hyaluronic acid, trigger point injection) treatments were identified based on product generic and brand names, as well as therapeutic class descriptions as recorded in the study database. Medical procedures were identified based on unique Japanese procedure codes.

For each patient, all-cause and OA-related HCRU and costs were documented. All-cause HCRU included all claims, regardless of the diagnosis. OA-related hospital admissions were identified by searching for inpatient hospital confinements in which OA was recorded as the primary discharge diagnosis or in which an OA procedure occurred (ie, arthroscopic surgery, osteotomy, arthroplasty, arthrodesis). Osteoarthritis-related outpatient visits were identified by searching for medical claims with any diagnosis (ie, primary or secondary) of OA, an OA procedure, a nonpharmacological treatment modality, or administration of an injectable treatments.

All-cause and OA-related HCRU and costs were estimated during each year of follow-up. HCRU was reported for hospitalizations, outpatient visits, and pharmacy visits and cost components included hospitalizations, medication use, and outpatient visits. Specifically, proportions of patients with an inpatient or outpatient visit were reported. Similarly, the mean number of outpatient visits during the relevant period was reported. Furthermore, medical costs by hospital department (inpatient and outpatient) and medication costs were reported. The costs represent payments for medical services and prescription drugs and included the insurance payment and patient copayment amount. All cost data were reported in Japanese yen (¥) and were inflated to 2017 using the medical care component of the Japanese Consumer Price Index.

Statistical Analysis

Patient demographics, clinical characteristics, pain-management patterns, and all-cause and OA-related HCRU and costs were reported for KOA/O and KOA/+ cohorts. Descriptive analyses entailed the tabular display of mean values and standard deviations of continuous variables of interest (eg, HCRU) and frequency distributions for categorical variables (eg, sex). The statistical significance of descriptive differences in patient characteristics, pain-management patterns, and HCRU between KOA/O and KOA/+ cohorts were tested using the Student t test, chi-square test, and Fisher exact test, as appropriate, with results of significance reported. A critical value of 0.05 was used to determine statistical significance. All analyses were conducted using SAS version 9.3 or later (SAS Institute, Inc., Cary, North Carolina).

Results

A total of 2542 patients met the study criteria: 1575 in the KOA/O cohort and 967 in the KOA/+ cohort (Figure 1). Patient demographies are presented in Table 1. Mean age
Patients aged ≥40 years with a KOA diagnosis between January 1, 2011, and December 31, 2012

Patients without malignancy post-index date

Patients with at least 5 years continuous health plan enrollment post-index date

Patients with at least 1 year continuous health plan enrollment pre-index date

Patients with no diagnoses of KOA pre-index date (1 year)

Patients with at least 1 additional KOA diagnosis post-index date

KOA patients excluding late other OA diagnosed cohort*

KOA cohort:
No other OA diagnosis during baseline and follow up period

KOA/+ cohort:
At least 1 other OA diagnosis ±1 year from the index date

Table 1 Patient Demographic and Clinical Characteristics

| Characteristics                                         | KOA/O (n=1575) | KOA/+ (n=967) |
|---------------------------------------------------------|----------------|---------------|
| Age at index date, mean (SD) [min:max]                  | 52.0 (6.9) [40:71] | 54.7 (7.3) [40:71] |
| Female, n (%)                                           | 791 (50.2) | 612 (63.3) |
| CCI score, mean (SD) [min:max]                          | 0.5 (1.0) [0.0:7.0] | 0.8 (1.1) [0.0:7.0] |
| At least one comorbidity, n (%)                         | 392 (24.9) | 362 (37.4) |

Physician department for index knee OA diagnosis, n (%)

| Department                          | KOA/O (n=1575) | KOA/+ (n=967) |
|-------------------------------------|----------------|---------------|
| Orthopedic surgery                  | 1206 (76.6) | 728 (75.3) |
| Anesthesiology                      | 7 (0.4) | 4 (0.4) |
| Internal medicine (general physician)| 315 (20.0) | 205 (21.2) |
| Other specialties                    | 47 (3.0) | 30 (3.1) |

Type of setting for the index KOA diagnosis, n (%)

| Setting                          | KOA/O (n=1575) | KOA/+ (n=967) |
|----------------------------------|----------------|---------------|
| Public hospital                   | 22 (1.4) | 28 (2.9) |
| University hospital               | 16 (1.0) | 7 (0.7) |
| Other hospital                    | 222 (14.1) | 149 (15.4) |
| Clinic                            | 1315 (83.5) | 783 (81.0) |

Site of other OA diagnosis, ±1 year of index date, n (%)

| Site                              | KOA/O (n=1575) | KOA/+ (n=967) |
|-----------------------------------|----------------|---------------|
| Spine                             | 761 (78.7) |                |
| Hip                               | 120 (12.4) |                |
| Ankles                            | 63 (6.5) |                |
| Unspecified                       | 63 (6.5) |                |
| Hands                             | 40 (4.1) |                |
| Elbows                            | 32 (3.3) |                |
| Fingers                           | 19 (2.0) |                |
| Shoulders                         | 9 (0.9) |                |
| Feet                              | 4 (0.4) |                |
| Other                             | 3 (0.3) |                |

Number of other diagnosed sites, mean (SD) [min:max]

| Number of sites | KOA/O (n=1575) | KOA/+ (n=967) |
|-----------------|----------------|---------------|
| 1.2 (0.4) [1.0-5.0] |                |                |

Notes: Bold values indicate P < 0.05. Statistical comparisons were made using the Chi-square/Fisher exact test for proportions and Student t test for means.

Abbreviations: CCI, Charlson Comorbidity Index; KOA/O, knee OA only; KOA/+, knee OA plus OA at another site; OA, osteoarthritis; SD, standard deviation.

was higher among the KOA/+ (54.7 years) versus the KOA/O (52.0 years) cohort, with greater proportion of females in the KOA/+ versus the KOA/O (63.3% versus 50.2%) cohort. Across both cohorts, most patients

Figure 1 Sample attrition chart.

Note: *Late other OA diagnosed cohort is defined as no other OA diagnosis during selection window and at least one other OA diagnosis from the second to fifth year from the index date.

Abbreviations: KOA/O, knee OA only; KOA/+, knee OA plus OA at another site; OA, osteoarthritis.
visited orthopedic surgery clinics on the index date. Among patients in the KOA/+ cohort, spine (78.7%) and hip (12.4%) were the most common sites of other OA.

KOA/+ cohort showed higher mean CCI score than KOA/O cohort (0.8 versus 0.5). A higher proportion of patients had at least one comorbidity during the baseline period that may have been taken into account when deciding on OA treatment in the KOA/+ (37.4%) compared with patients in the KOA/O cohort (24.9%) (Table 1). The details of reported comorbidities during each year of follow-up period are presented in Figure 2. During the follow-up period, a higher percentage of patients in the KOA/+ than patients with KOA/O cohort had a diagnosis of diabetes, hypertension, cardiovascular disorders, and depression.

Pharmaceutical treatment was received by 91.5% of patients in the KOA/+ compared with 85.1% of patients in the KOA/O cohort during the first year of follow-up. The proportion of patients receiving pharmaceutical treatment decreased in the second year of follow-up and remained at approximately the same level for the remainder of the 5-year follow-up period (ie, years 3 through 5). Regardless of the cohort, topical nonsteroidal anti-inflammatory drugs (NSAIDs) (received by 82.5% of patients in the KOA/+ and 74.2% of patients in the KOA/O), followed by oral non-selective NSAIDs and cyclooxygenase-2 (COX-2) selective inhibitors were the most common pharmaceutical treatments received during the first year of follow-up as well as during the entire follow-up period. Weak opioids were received by 4.3% of patients in the KOA/O and 6.2% of patients in the KOA/+ cohorts during the first year of the follow-up period, and these percentages did not tend to drastically change over time. Strong opioids were rarely received, regardless of cohort and time period (Table 2).

Injectable treatment was received by more than 40.0% of patients during the first year of follow-up in both cohorts; IAHA and corticosteroids were the most common injectable treatments received during the first year of follow-up (Table 2). Overall, about 30% of patients with KOA/O compared with only 15% of patients with KOA/+ did not receive any treatment between years 2 and year 5 of the follow-up period.

Analysis of the overall treatment regimens and order of therapies received by patients revealed that the most common regimen first received was either topical or oral non-selective NSAIDs (data not shown). A slightly higher mean number of pharmaceutical treatments were received by patients with KOA/+ (3.7) compared with patients with KOA/O (3.0) during the entire 5-year follow-up period (data not shown).

All-cause and OA-related resource utilization patterns are presented in Table 3. During each year of the 5-year follow-up period, a higher proportion of patients had at least one OA-related hospitalization among the KOA/+ cohort.
A higher number of OA-related outpatient visits were observed among the KOA/+ cohort (7.5, 5.3, 5.3, 5.4, 5.7) compared with the KOA/O cohort (4.7, 2.6, 2.4, 2.4, 2.5) during each year of the 5-year follow-up period. Similarly, total OA-related direct medical costs were higher among patients in the KOA/+ cohort (¥206,353, ¥174,444, ¥181,785, ¥188,829, ¥196,164) compared with the KOA/O cohort (¥78,350, ¥56,924, ¥55,816, ¥72,852, ¥81,821) during each year of the 5-year follow-up period.

Figure 3 presents data on all-cause and OA-related health-care resource costs.

Discussion
This study is one of the first to longitudinally assess pain-management options along with health-care resource use and direct medical costs for the working age population in Japan diagnosed with KOA. A considerable proportion of patients with KOA also had diagnosis of OA at other sites, and these patients (the KOA/+ cohort) had greater pain-related treatment use and comorbidity burden compared with patients with KOA/O. In our study, a majority of patients in both cohorts received pharmaceutical treatment during the first year of follow-up. The proportion of patients in our study receiving pharmaceutical treatment decreased between the first year of follow-up and the second year of follow-up and remained stable at this level for the remainder of the follow-up period (ie, years 3 through year 5). Topical and oral NSAIDs were the most frequently observed treatments received for pain management across both the cohorts. Additionally, this study observed that the direct medical cost burden was highest during the first year of follow-up.

Patients in the KOA/+ cohort were observed to have more comorbidities, which may have impacted treatment decisions, during the follow-up periods compared with patients in the KOA/O cohort. Diabetes, hypertension, cardiovascular disorders, and depression were observed in higher proportions of patients with KOA/+ than in those with KOA/O.

Table 2 Pain-Management Options After Index Date

| Treatment Modality, % | KOA/O (n=1575) | KOA/+ (n=967) |
|-----------------------|----------------|----------------|
|                       | 1 y 2 y 3 y 4 y 5 y | 1 y 2 y 3 y 4 y 5 y |
| Nonpharmacological treatment | 26.2 18.7 16.3 16.3 16.6 | 44.4 39.5 35.1 32.7 38.2 |
| Physical therapy | 12.3 7.3 7.2 6.9 7.4 | 24.0 20.0 16.6 16.4 18.9 |
| Manual/instrumental therapy | 16.6 12.8 10.8 11.3 11.1 | 27.9 25.4 24.2 21.5 25.0 |
| Pharmaceutical treatment | 85.1 64.4 63.0 64.1 66.0 | 91.5 81.3 79.7 81.7 82.8 |
| Non-selective NSAIDs – oral | 58.8 43.5 40.8 44.3 42.4 | 71.8 60.3 55.2 57.6 58.2 |
| NSAIDs – topical | 74.2 42.4 40.1 39.6 42.8 | 82.5 61.9 61.2 60.3 64.9 |
| COX-2 selective inhibitors | 21.8 9.2 7.7 8.0 8.8 | 22.9 13.9 12.4 14.6 15.0 |
| Acetaminophen | 12.5 13.0 15.5 19.4 22.3 | 17.7 22.3 22.0 26.3 28.2 |
| Weak opioid | 4.3 3.9 4.1 4.3 5.6 | 6.2 8.6 7.8 8.4 9.3 |
| Strong opioid | 0.0 0.0 0.0 0.1 0.1 | 0.4 0.2 0.2 0.2 0.4 |
| Other non-opioid drug | 3.1 3.6 2.9 2.4 2.7 | 9.7 8.2 7.6 7.8 7.8 |
| SNRI | 1.0 0.8 1.0 0.7 1.3 | 1.2 1.6 1.7 1.6 2.7 |
| Pregabalin | 1.0 1.5 1.1 1.5 1.8 | 6.5 6.5 6.9 6.6 8.1 |
| Injectable treatment | 42.0 31.7 28.6 29.0 30.8 | 43.5 41.6 38.5 39.5 39.0 |
| Corticosteroids | 24.3 18.9 18.7 19.6 21.2 | 27.1 28.2 27.7 28.5 26.6 |
| IAHA | 33.2 19.5 16.1 15.9 17.3 | 31.1 23.7 21.1 21.0 21.5 |
| Trigger point injection | 2.3 2.2 2.4 2.3 2.8 | 8.0 8.1 8.0 7.8 7.7 |
| No pain-related treatment | 12.0 30.7 32.6 31.6 30.1 | 7.1 15.3 16.9 15.3 14.4 |

Note: Bold values indicate P < 0.05.

Abbreviations: COX-2, cyclooxygenase-2; IAHA, intra-articular hyaluronic acid; KOA/O, knee osteoarthritis only; KOA/+, knee osteoarthritis plus osteoarthritis at another site; NSAID, nonsteroidal anti-inflammatory drug; SNRI, serotonin-norepinephrine reuptake inhibitor.
during the follow-up period. A meta-analysis of observational studies reported a higher proportion of comorbidities among patients with KOA compared to those with no KOA. A recent cross-sectional cohort study conducted in the Netherlands also reported significant comorbid burden among all patients with OA of knee or hip. Another retrospective database analysis study conducted in the US reported significant comorbidity burden among patients with OA, with approximately 54% of patients having hypertension and 10% having cerebrovascular disease.

**Table 3 All-Cause and OA-Related Health Care Resource Use**

|                      | KOA/O (n=1575) | KOA/+ (n=967) |
|----------------------|---------------|---------------|
|                      | 1 y  | 2 y  | 3 y  | 4 y  | 5 y  | 1 y  | 2 y  | 3 y  | 4 y  | 5 y  |
| **Patients with services, %** |
| Outpatient visit – OA related | 99.9 | 50.2 | 44.4 | 43.1 | 43.2 | 100.0 | 73.0 | 69.4 | 71.3 | 71.7 |
| Outpatient visit – All cause | 99.9 | 92.7 | 92.0 | 92.4 | 93.1 | 100.0 | 97.8 | 97.5 | 97.9 | 97.8 |
| Pharmacy visit – OA related | 71.8 | 48.6 | 46.3 | 47.1 | 49.1 | 78.5 | 64.8 | 63.8 | 65.5 | 65.1 |
| Pharmacy visit – All cause | 85.2 | 75.6 | 75.4 | 77.7 | 77.8 | 91.6 | 85.6 | 86.0 | 87.8 | 87.8 |
| Hospital admission – OA related | 2.4 | 2.3 | 2.2 | 2.3 | 3.2 | 4.9 | 2.7 | 4.1 | 4.8 | 4.2 |
| Hospital admission – All cause | 0.9 | 0.4 | 0.6 | 0.4 | 0.8 | 3.2 | 1.1 | 1.7 | 2.5 | 2.1 |

**Number of services, mean (SD)**

|                      | KOA/O | KOA/+ |
|----------------------|-------|-------|
| Outpatient visit – OA related | 4.7 (4.2) | 11.1 (7.8) |
| Outpatient visit – All cause | 2.6 (4.1) | 9.1 (7.9) |
| Pharmacy visit – OA related | 2.4 (4.2) | 9.2 (8.0) |
| Pharmacy visit – All cause | 2.3 (0.1) | 9.3 (7.9) |
| Hospital admission – OA related | 2.5 (4.4) | 9.7 (8.0) |
| Hospital admission – All cause | 7.5 (5.7) | 15.8 (9.6) |

**Notes:** KOA/O, n=1575; KOA/+, n=967.
**Abbreviations:** KOA/O, knee OA only; KOA/+, knee OA plus OA at another site; OA, osteoarthritis; SD, standard deviation.

**Figure 3** Mean all-cause and OA-related costs estimated during each year of follow-up.

**Abbreviations:** KOA/O, knee OA only; KOA/+, knee OA plus OA at another site; OA, osteoarthritis.
To our knowledge, this is the first study reporting the pain-management burden comparing KOA/O and KOA/+ among the working-age population in Japan. Our research suggests that 55% of patients with KOA had at least one diagnosis of OA at another site at any time during the study period. A previous study conducted in a community-based cohort of patients with radiographically confirmed OA in the US also reported a similar proportion (42.8%) of patients having multiple-joint disease. Clinicians should consider focusing on treatments that would benefit multiple-joint OA along with KOA given the extent of the disease in and the greater health-care burden for the population with multijoint involvement observed in the present analysis. This recommendation is consistent with clinical guidelines that recommend a multidisciplinary approach should be considered for treatment of OA, specifically OA involving multiple sites. Multidisciplinary treatments as defined by the National Institute for Health and Care Excellence are interventions that target at least two different health disciplines (eg, exercise and manual therapy, weight loss, acupuncture, thermotherapy, analgesics, intra-articular injections, surgeries). Arthralgia in Japan is a very common complaint among the elderly population and constitutes a growing segment of Japanese demographics. Furthermore, comorbid chronic diseases and disability have become a major public health concern. Recent evidence has shown joint disease is the most common reason for entry into certified support level in the long-term care insurance system in Japan. Further studies are necessary to evaluate appropriate pain management of KOA including OA of other painful sites.

Our study reported that the proportion of patients receiving pharmaceutical treatment decreased between the first year of follow-up and the second year of follow-up. This observed decrease may be attributed to several factors, one of which may include low pain-related treatment satisfaction among KOA patients in the first year of follow-up. For example, a recent patient survey reported low treatment satisfaction among KOA patients in Japan. The results of the present analysis may potentially indicate that current treatments may not be providing desired outcomes among KOA patients in Japan, and there is a need for better pain-management options that may in turn improve patient satisfaction.

Overall, higher HCRU and costs were observed among patients with KOA/+ during the follow-up period compared with patients with KOA/O. Wang and colleagues also reported high medical costs among patients with OA, regardless of site. The present study reported the highest medical costs in the first year after index KOA diagnosis, which could be explained by the more frequent diagnostic and outpatient care visits during the initial year of treatment. Le et al conducted a retrospective analysis and reported that patients with newly diagnosed OA incurred higher medical costs compared with patients with existing OA. To our knowledge, few studies have reported medical costs among patients with OA. Le and colleagues reported mean total all-cause costs of $19,391 and OA-related costs of $6811 among patients with newly diagnosed OA. Another study conducted in Singapore reported annual direct costs ranging from S$1460–S$7477 among patients with KOA. Although the cost burden from other countries cannot be compared with Japan due to variability in health-care systems and differing time periods, in general, OA-related costs reported were high.

There are some limitations to be noted in our study. First, the study sample was selected from enrollees covered by the employees’ health insurance system. Because most enrollees are working adults, the proportion of elderly patients aged 65 years and older is low. The current study may be overestimating OA-related costs by incorporating costs attributed to OA-comorbidity-related pharmacy use (eg, serotonin-norepinephrine reuptake inhibitor treatment for depression). Additionally, we required 6 years of continuous health plan enrollment for all patients. Patients in our population may have been healthier than the overall OA population (and correspondingly have had less severe OA than those patients without continuous enrollment for 6 years) as they remained employed for a continuous period of time after their first observed OA diagnosis. Finally, the diagnoses listed in the claims were not validated, and it was not possible to confirm patient diagnoses.

Conclusion
This study is the first to demonstrate that the working-age population in Japan with KOA/+ received a greater proportion of pain-related treatment compared with patients with KOA/O. Also, this study reports detailed information on patient costs including outpatient visits, inpatient visits, and pharmaceutical treatments filled for patients with KOA/+. The results of the present analysis suggest that multidisciplinary and comprehensive treatment starting at an early stage for not only KOA but also OA at other sites may be important for effective resource utilization in an aging society coping with socioeconomic burdens.
Disclosure

Juliana Meyers and Saurabh P. Nagar are full-time employees of RTI Health Solutions, which received funding from Eli Lilly Japan to conduct this research. Kaname Ueda and Shinji Fujikoshi are the full-time employees of Eli Lilly Japan K.K. Hiroyuki Enomoto was a full-time employee of Eli Lilly Japan at the time the study was conducted and is a current employee of Gilead sciences K.K. Tomoyuki Takura received consultant fee from Eli Lilly Japan K.K. The authors report no other conflicts of interest in this work.

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