Evaluation of the treatment patterns and economic burden of dysmenorrhea in Japanese women, using a claims database

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Purpose: This study aimed to describe treatment patterns and estimate health care resource utilization and associated costs among Japanese women with dysmenorrhea, using a claims database.

Methods: This was a retrospective analysis using health insurance data from the Japan Medical Data Center, assessing female patients aged 18–49 years with newly diagnosed primary or secondary dysmenorrhea. Treatment patterns analyses focused on hormonal medications, analgesics, hemostatic agents, traditional Chinese medicine (TCM), and gynecological surgeries. Data were collected on health care resource utilization and costs associated with medications, imaging procedures, and inpatient and outpatient care in both patients and matched controls.

Results: The analysis included 6,315 women with dysmenorrhea (3,441 primary; 2,874 secondary). The most commonly prescribed initial therapies were low-dose estrogen progestins (LEPs, 37.7%) and TCM (30.0%), with substantial differences between primary (LEPs: 27.4%, TCM: 38.8%) and secondary (LEPs: 50.2%, TCM: 19.5%) dysmenorrhea cohorts. Surgery was conducted in <5% of all patients. Both primary and secondary cohorts of dysmenorrhea had significantly higher mean total health care costs compared to controls within the 1-year period following diagnosis (Case-primary: 191,680 JPY [1,916 USD]; secondary: 246,488 JPY [2,465 USD], Control-primary: 83,615 JPY [836 USD]; secondary: 90,711 JPY [907 USD]) (p<0.0001). After adjusting for baseline characteristics, these costs were 2.2 and 2.9 times higher for primary and secondary dysmenorrhea cohorts, respectively, compared with matched controls, (both p<0.0001). The main driver of these excess costs was outpatient care, with eight additional physician visits per year among dysmenorrhea patients compared to controls (p<0.0001).

Conclusion: Considerable heterogeneity in treatment patterns was observed, with relatively low utilization of LEPs in patients with primary dysmenorrhea and those treated by internal medicine physicians. Total annual health care costs were approximately 2–3 times higher in patients with dysmenorrhea compared to women without the condition.

Keywords: dysmenorrhea, women’s health, treatment patterns, resource use and costs, economic burden, database analysis

Introduction

Many Japanese women experience health issues associated with menstruation; these can include menstrual pain also referred to as dysmenorrhea, heavy menstrual bleeding (HMB) also referred to as menorrhagia, and premenstrual syndrome (PMS).¹–³ Dysmenorrhea is the most common gynecological complaint associated with menstruation with a prevalence of 25% among all women, and reaching up to 90% among adolescents.⁴ It has also been reported that one-third of Japanese women require analgesic therapy for dysmenorrhea.⁵
In some women, dysmenorrhea cannot be attributed to any specific cause and is referred to as primary or idiopathic. It was reported that 47% of patients who consulted a physician for menstrual cramping had primary dysmenorrhea, based on a survey in 2000. In other cases, it is associated with a preexisting gynecological disorder, and the disease is referred to as secondary or organic dysmenorrhea. Pre-existing disorders include endometriosis, adenomyosis, and fibroids. Collectively, these are common gynecological complications in women. Regardless of the cause, dysmenorrhea can have a substantial impact on patient quality of life, yet many patients do not seek treatment. In a patient survey, some untreated women have expressed feelings of resistance or aversion toward seeking therapy, and many suggested that gynecologist consultations were unnecessary for their disorder. However, a substantial proportion of women who did seek medical treatment agreed that their daily lives were significantly improved after therapy, and it was also estimated that gynecologist visits saved over 7,000 JPY (70 USD) monthly costs per-patient, occurring due to time off work.

According to the guidelines for gynecological practice in Japan, by the Japan Society of Obstetrics and Gynecology (JSOG) and Japan Association of Obstetricians and Gynecologists (JAOG) (2011 edition), low-dose estrogen progestins (LEPs) and nonsteroidal anti-inflammatory drugs (NSAIDs) are primarily recommended for primary dysmenorrhea, and traditional Chinese medicines (TCMs) could be used for primary dysmenorrhea. Other current clinical practice guidelines for the treatment of dysmenorrhea include the use of over-the-counter analgesics, NSAIDs, and oral contraceptives such as LEPs, progestin-only therapies, and the levonorgestrel-releasing intrauterine system. Two kinds of combined oral contraceptives (COCs) are available on the Japanese market: LEPs, (norethisterone/estrogen and drospirenone/estrogen), which are reimbursed for dysmenorrhea, and other COCs for contraceptive purposes, which are not reimbursed. Some of these options have demonstrated efficacy in alleviating symptoms among patients with dysmenorrhea. For example, LEPs have been reported to be effective in alleviating symptoms in up to 80% of women. Additional studies have shown efficacy of both gonadotropin-releasing hormone (GnRH) analogs and testosterone derivatives for treating underlying causes of secondary dysmenorrhea, such as endometriosis.

Existing evidence on treatment patterns, health care resource use, and associated costs in Japanese patients with dysmenorrhea is limited. Information on treatment patterns and the economic burden of disease would be useful to guide the allocation of health care resources for the treatment of dysmenorrhea. Furthermore, the analysis of treatment patterns and resource utilization may shed light on any potential challenges related to the current diagnosis and management of dysmenorrhea in Japan.

The objectives of this study were to describe treatment patterns and estimate health care resource use and costs among Japanese women with newly diagnosed dysmenorrhea in a real-world setting. This included a detailed description of the baseline characteristics and comorbidities of these patients, their initial and subsequent therapies for dysmenorrhea, and the probability of surgical procedure related to dysmenorrhea.

To assess health care resource utilization and costs among patients, a comparison was made between patients with dysmenorrhea (cases) and those without dysmenorrhea (matched control group). Furthermore, all results were differentiated between women with primary and secondary dysmenorrhea and treatment patterns were evaluated by prescriber’s specialty. The reason why we took prescriber’s specialty into consideration is that the Japanese health care system allows patients free access to any clinic irrespective of specialty.

**Methods**

**Study design and population**

This was a retrospective claims analysis of the Japan Medical Data Center (JMDC) database, which includes deidentified medical (inpatient and outpatient) and pharmacy claims from up to 3.0 million beneficiaries (both employees and their dependents) from 2005 onward. The JMDC database covers nearly 10% of the 30 million individuals in the Japanese Social Insurance System, equating to approximately 2.5% of the total Japanese population. Data were extracted for patients with records between July 1, 2009, and June 30, 2014, and included information on patient demographics, diagnoses, drug prescriptions, medical procedures, medical facility characteristics, and reimbursement costs.

Included patients were women between ages 18 and 49 years who had two separate diagnoses of dysmenorrhea within 3 months. In Japan, patients with dysmenorrhea are generally asked to come back to the clinic within the 3 months following a first medical consultation, as it was considered that only one visit would not be enough for selecting patients requiring medical care for dysmenorrhea. Diagnoses were identified by ICD-10 (International Classification of Diseases 10th revision) codes N944 (primary dysmenorrhea) or N946 (dysmenorrhea, unspecified) or one of the following Medical Information System Development Center (MEDIS-DC)
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standard disease names: functional dysmenorrhea, menstrual pain, or dysmenorrhea. MEDIS-DC consists of standardized names of diseases and injuries for reimbursement by the National Health Insurance. The index date was defined as the date of the first diagnosis of dysmenorrhea. Primary dysmenorrhea was defined as patients without underlying causes (endometriosis, adenomyosis, and fibroids) during the preindex period and from 60 days after the index date, whereas secondary dysmenorrhea was defined as patients presenting at least one diagnosis of one of these conditions during this period. This criterion was defined based on the publications by Copher et al and an assumption that imaging procedures for detecting underlying causes would be performed within 2 months following the first visit. Patients were also required to have insurance records for at least 6 months with a diagnosis of dysmenorrhea prior to their initial diagnosis (preindex period) and for a continuous postindex period of at least 1 year, to be included in the study. Patients were excluded if they had been diagnosed with gynecological cancers or bleeding disorders during the study period (including both pre- and postindex), if they underwent gynecological surgery (hysterectomy, endometrial ablation, or myomectomy) during the preindex period, or if they were treated with hormonal agents during the preindex period (Figure 1).

For the resource utilization analysis, matched controls were identified among females in the JMDC database who did not suffer from dysmenorrhea (ie, no diagnosis between July 1, 2009, and June 30, 2014). Controls and cases were matched 2:1 by year of birth (within 5 years), Charlson Comorbidity Index (CCI) at baseline (±15 days from index date), and at least the same 18 months of follow-up period.

Treatment patterns
The study assessed therapies that fit into one of the following categories: hormonal treatments (LEPs, progestin, testosterone derivatives, and GnRH analogs), NSAIDs, hemostatic agents, and TCM therapies. Relevant gynecological surgeries, including hysterectomies, endometrial ablations, and myomectomies, were also considered.

The treatment patterns were analyzed based on prescription records, with a treatment line defined as a group of consecutive prescriptions of the same agent or combination of agents (combination therapy), with no interruption exceeding 90 days. For each treatment line, the start date, end date, duration, and treatment category were determined. When a period exceeding 90 days was observed between the end of one treatment line and the beginning of a new one, a treatment discontinuation was deemed to have occurred. Therapy switches were defined when an overlap shorter than 21 days between the end of one therapy and the beginning of a new one was observed and the interval did not exceed 90 days. If an overlap occurred for 21 days or more, however, this was considered a combination therapy. Because surgery was considered as a permanent treatment, a separate combination therapy type (surgery combination) was assigned. The drug categories assessed in this study are described in Table 1.

Treatment patterns were analyzed among the entire cohort and according to the type of dysmenorrhea (primary or secondary). A subgroup analysis was also conducted according to the prescriber’s specialty for the first treatment line.

Resource utilization and cost calculations
The resources used and their associated costs were calculated based on the 1-year follow-up period after the index date for each patient. Data were collected on inpatient costs, outpatient costs, and costs associated with both prescriptions and imaging procedures. From this information, total health care costs were calculated.

Resource utilization data collected in this analysis included the number of inpatient admissions, cumulative length of stay, number of outpatient visits, and number of imaging procedures. Imaging procedures, which included echography, computed tomography scan, and magnetic resonance imaging, were determined to be related to a diagnosis of dysmenorrhea when they occurred within a 1-month period around the date of the diagnosis (±15 days). The use of imaging procedures in controls was evaluated during the same 1-month period as it was for their corresponding matched cases.

Statistical analysis
Nonadjusted comparisons were conducted for outcomes with respect to primary vs secondary dysmenorrhea and cases vs controls. For continuous variables, a Fisher’s $F$-test was conducted to test the assumption of homoscedasticity. Mean values were then compared using a Student’s $t$-test (for variables showing homoscedasticity) or Satterthwaite test (heteroscedasticity). For categorical variables, a $\chi^2$ or Fisher’s exact test (in cases of insufficient theoretical size of subgroups) was performed to compare the distributions.

Time-to-event variables were assessed using Kaplan–Meier curves and log-rank tests for between-group comparisons. Time to “competing events” (ie, end-of-treatment events such as treatment switch, discontinuation of treatment, surgery, add-on therapy, and step-down therapy) were analyzed using the competing risks survival method, estimating the cumulative
Population of health insurance societies contributing to JMDC claims database between July 1st, 2009 and June 30th, 2014

N=2.5m.

Women with a diagnosis of dysmenorrhea* between July 1st, 2009 and June 30th, 2014

N=37,115

Women patients aged 18 through 49 years

N=34,118

Patients with continuous enrollment during 6 months preindex*

N=22,423

Patients with continuous enrollment during 12 months postindex

N=13,849

Patients with a second diagnosis of dysmenorrhea within 3 months

N=7,412

No diagnosis of dysmenorrhea during the preindex period*

N=7,412

Diagnosis of cancer during the pre- and postindex periods

N=6,843

Hysterectomy or endometrial ablation during the preindex period

N=6,815

Prescription of hormonal treatment during the preindex period

N=6,501

Bleeding disorder during the pre- and post-index periods

N=6,315

Dysmenorrhea cohort

N=6,315

Primary dysmenorrhea

N=3,441

Secondary dysmenorrhea

N=2,874

Endometriosis

n=2,071

Uterine Fibroids

n=1,116

Adenomyosis

n=296

* Dysmenorrhea defined by ICD10 code “N944”, “N946” and MEDIS-DC standard disease name “functional dysmenorrhea”, “menstrual pain”, “dysmenorrhea”

* Index date: first diagnosis of dysmenorrhea

* This criteria was added for clarity but is already included in the 2nd inclusion criteria with the definition of the index date as the first diagnosis.

Figure 1 Flow chart of selection process for patients with dysmenorrhea.

Abbreviations: JMDC, Japan Medical Data Center; ICD10, International Classification of Disease 10th revision; MEDIS-DC, Medical Information System-Development Center.

incidence. Differences between groups were assessed using a Gray’s test.

The analyses of costs and resources were conducted through Generalized Linear Models (GLMs), testing normal, Poisson, negative binomial, zero-inflated Poisson, and zero-inflated negative binomial distributions. The identity and log link functions were investigated. The goodness of fit of the models was assessed with the deviance, and the dispersion with the Pearson $\chi^2$ value divided by the number of degrees of freedom. The models minimizing the deviance or with the best compromise deviance/dispersion (in case of high dispersion) were retained. Models were adjusted for age, the category (based on quartiles) of total health care costs over the preindex period, the quantity corresponding to the
Table 1 Type of treatment evaluated in the analysis

<table>
<thead>
<tr>
<th>Hormonal treatments</th>
<th>Analgesic agents</th>
<th>Hemostatic agents</th>
<th>TCM therapies</th>
<th>Surgical procedures</th>
</tr>
</thead>
<tbody>
<tr>
<td>LEP:</td>
<td></td>
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</tr>
<tr>
<td>Low-dose norethisterone and estrogen</td>
<td>NSAIDs, prescribed with a diagnosis of pain*:</td>
<td>Tranexamic acid</td>
<td>Tokishakuyakusan extract</td>
<td>Complete ablation:</td>
</tr>
<tr>
<td>Low-dose drospirenone and estrogen</td>
<td></td>
<td></td>
<td></td>
<td>Hysterectomy</td>
</tr>
<tr>
<td>Progestin:</td>
<td></td>
<td></td>
<td></td>
<td>Partial ablation:</td>
</tr>
<tr>
<td></td>
<td>Dienogest</td>
<td></td>
<td></td>
<td>Endometrial ablation</td>
</tr>
<tr>
<td>Testosterone derivatives:</td>
<td></td>
<td></td>
<td></td>
<td>Myomectomy</td>
</tr>
<tr>
<td>Danazol</td>
<td></td>
<td></td>
<td></td>
<td>Uterine–artery embolization</td>
</tr>
<tr>
<td>GnRH analog:</td>
<td></td>
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<td></td>
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<tr>
<td>Buserelin</td>
<td></td>
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<tr>
<td>Leuprorelin</td>
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<tr>
<td>Goserelin</td>
<td></td>
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</tbody>
</table>

Notes: *As suggested by experts, physicians may prescribe NSAIDs to patients, associated with a diagnosis of headache, acute pain, pain in the abdomen (upper, lower, or unspecified), low back pain, and pelvic or perineal pain. Surgical procedure codes were identified based on the medical procedure index available on the MHLW website. Abbreviations: GnRH, gonadotropin-releasing hormone; LEP, low-dose estrogen progestin; NSAIDs, nonsteroidal anti-inflammatory drugs; TCM, traditional Chinese medicine.

This study was approved by the Ethics Committee of the nonprofit organization, Clinical Research Promotion Network Japan. The information available in the JMDC database is anonymized. The informed consent of patients was not applicable based on the Ethical Guidelines for Epidemiological Research issued by the Ministry of Health, Welfare and Labor.

Results

Patient characteristics

A total of 6,315 patients with dysmenorrhea, 3,441 (54.5%) primary and 2,874 (45.5%) secondary, were identified, with an average follow-up duration of 4.0±0.9 years. Patients with secondary dysmenorrhea were significantly older compared with patients with primary dysmenorrhea (35.7±8.0 vs 31.6±7.9 years; p<0.0001; Table 2), and over half of the secondary dysmenorrhea cohort were older than 35 years (59.3% vs 36.8% in the primary dysmenorrhea cohort). Primary ovarian dysfunction was the most common comorbidity (22.1%) at baseline and was more frequent in patients with primary dysmenorrhea than secondary dysmenorrhea (25.0% vs 18.6%; p<0.0001; Table 2). The other comorbidities were more frequently reported in patients with secondary dysmenorrhea than primary dysmenorrhea, including erosion and ectropion of cervix uteri (21.8% vs 15.6%; p<0.0001), anemia (19.7% vs 6.2%; p<0.0001), acute vaginitis (14.5% vs 10.2%; p<0.0001), and low back pain (11.2% vs 8.1%; p<0.0001), respectively. Over three-quarters of patients had a CCI score of 0, and less than 4% had a score greater than 1 (Table 2).

For both cohorts, first diagnoses of dysmenorrhea were reported in clinics (0–19-bed) (76.4%) and in hospitals (≥20-bed) (23.6%). In more detail, 81.3% of primary and 70.6% of secondary dysmenorrhea cases were reported in clinics and 18.7% of primary and 29.4% of secondary dysmenorrhea...
cases were reported in hospitals. Patients who were diagnosed in large hospitals (≥500-bed) represented 5.7% of the primary dysmenorrhea cohort and 11.7% of the secondary dysmenorrhea cohort. Thus, patients in the secondary dysmenorrhea cohort were significantly more likely to be diagnosed in hospitals \(p < 0.0001\) and very large \(\geq 500\) facilities \(p < 0.0001\) compared to the primary dysmenorrhea cohort. The majority of patients were diagnosed in obstetrics and gynecology facilities (53.0% and 64.6% for primary and secondary dysmenorrhea cohorts, respectively), whereas the remainder were typically diagnosed in general internal medicine facilities (32.1% and 27.6% for primary and secondary dysmenorrhea cohorts, respectively). At least one diagnostic imaging procedure within 2 weeks before or after diagnosis was reported for 38.7% of patients with primary dysmenorrhea and 69.2% of those with secondary dysmenorrhea; in most cases, this was echography. Among women with secondary dysmenorrhea, 4.9% also presented with a diagnosis of heavy menstrual bleeding at baseline.

### Treatment patterns

A total of 83.4% of patients in the primary dysmenorrhea cohort and 89.2% in the secondary dysmenorrhea cohort were treated with at least one pharmacological agent or surgical procedure. Median times to any treatment initiation after diagnosis of dysmenorrhea were 9 and 3 days, in primary and secondary dysmenorrhea cohorts, respectively. In the primary dysmenorrhea cohort, TCM therapies were the most frequently prescribed first-line treatment and were administered in 38.8% of all patients with primary dysmenorrhea (including treated and untreated patients), LEPs were prescribed in 27.4% of patients, and the use of hemostatic agents was reported in 12.8% (Figure 2). In the secondary dysmenorrhea cohort, the proportion of patients treated with LEPs was higher, with 50.2% receiving these agents as a first-line treatment, whereas 19.5% were treated with TCM therapies (Figure 3). Substantial differences in the treatment of dysmenorrhea were observed according to the specialty of the prescriber: LEPs were prescribed in first line by obstetricians and gynecologists in 57.2% and 67.7% of treated patients from the primary and secondary dysmenorrhea cohorts, respectively, while internal medicine physicians prescribed LEPs to 11.6% and 35.7% of treated patients, respectively.

**Table 2 Patient baseline characteristics at index date**

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Primary (N=3,441)</th>
<th>Secondary (N=2,874)</th>
<th>Total (N=6,315)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean (SD) age, years, at index date</td>
<td>31.64 (7.92)</td>
<td>35.65 (8.01)</td>
<td>33.47 (8.21)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Age, years, category, N (%)</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>18–34</td>
<td>2,174 (63.18)</td>
<td>1,170 (40.71)</td>
<td>3,344 (52.95)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>35–39</td>
<td>608 (17.67)</td>
<td>609 (21.19)</td>
<td>1,217 (19.27)</td>
<td></td>
</tr>
<tr>
<td>40–49</td>
<td>659 (19.15)</td>
<td>1,095 (38.10)</td>
<td>1,754 (27.78)</td>
<td></td>
</tr>
<tr>
<td>Type of member, N (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Individual</td>
<td>1,570 (45.63)</td>
<td>1,289 (44.85)</td>
<td>2,859 (45.27)</td>
<td>0.54</td>
</tr>
<tr>
<td>Family</td>
<td>1,871 (54.37)</td>
<td>1,585 (55.15)</td>
<td>3,456 (54.73)</td>
<td></td>
</tr>
<tr>
<td>Disease status, N (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dysmenorrhea only</td>
<td>3,385 (96.37)</td>
<td>2,734 (95.13)</td>
<td>6,119 (96.90)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Both dysmenorrhea and HMB</td>
<td>56 (1.63)</td>
<td>140 (4.87)</td>
<td>196 (3.10)</td>
<td></td>
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<tr>
<td>Underlying conditions, N (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Endometriosis</td>
<td>0</td>
<td>2,071 (72.06)</td>
<td>2,071 (72.06)</td>
<td>NA</td>
</tr>
<tr>
<td>Adenomyosis</td>
<td>0</td>
<td>296 (10.30)</td>
<td>296 (10.30)</td>
<td></td>
</tr>
<tr>
<td>Fibroids</td>
<td>0</td>
<td>1,166 (40.57)</td>
<td>1,166 (40.57)</td>
<td></td>
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<tr>
<td>CCI category, N (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>2,706 (78.64)</td>
<td>2,269 (78.95)</td>
<td>4,975 (78.78)</td>
<td>0.61</td>
</tr>
<tr>
<td>1</td>
<td>614 (17.84)</td>
<td>494 (17.19)</td>
<td>1,108 (17.55)</td>
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</tr>
<tr>
<td>2</td>
<td>104 (3.02)</td>
<td>95 (3.31)</td>
<td>199 (3.15)</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>15 (0.44)</td>
<td>11 (0.38)</td>
<td>26 (0.41)</td>
<td></td>
</tr>
<tr>
<td>4+</td>
<td>2 (0.06)</td>
<td>5 (0.17)</td>
<td>7 (0.11)</td>
<td></td>
</tr>
<tr>
<td>Top five comorbidities at baseline, N (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Primary ovarian dysfunction (ovarian insufficiency, luteal phase deficiency)</td>
<td>861 (25.02)</td>
<td>533 (18.55)</td>
<td>1,394 (22.07)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Erosion and ectropion of cervix uteri</td>
<td>538 (15.63)</td>
<td>625 (21.75)</td>
<td>1,163 (18.42)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Iron deficiency anemia, unspecified</td>
<td>214 (6.22)</td>
<td>566 (19.69)</td>
<td>780 (12.35)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Acute vaginitis</td>
<td>351 (10.20)</td>
<td>417 (14.51)</td>
<td>768 (12.16)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Low back pain</td>
<td>280 (8.14)</td>
<td>322 (11.20)</td>
<td>602 (9.53)</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

**Abbreviations:** CCI, Charlson Comorbidity Index; HMB, heavy menstrual bleeding; SD, standard deviation.
and 21.8% by internal medicine physicians, of treated patients in the primary and secondary dysmenorrhea cohorts, respectively. TCMs were prescribed in first line in 34.0% and 20.6% by obstetricians and gynecologists, and in 56.1% and 25.3% by internal medicine physicians, of treated patients in the primary and secondary dysmenorrhea cohorts, respectively.

The time to treatment discontinuation was longer for LEPs compared with other therapies. In patients with primary dysmenorrhea, 35.1% were still receiving LEP treatment at 1 year, compared to 10.4% for TCM therapies. Similarly, 33.9% of patients with secondary dysmenorrhea continued to receive LEPs at 1 year, while only 11.5% continued taking TCM therapies for this duration. Hemostatic agents and NSAIDs were administered only briefly, with a median treatment duration of 5 days for both agents, regardless of the cohort.

Of those patients who did not continue with their first-line treatment for the first 12 months postindex date, only a minority switched to a new therapy (17.1% and 21.3% for the primary and secondary dysmenorrhea cohorts, respectively) and most did not receive any second-line treatment (73.7% and 63.3%, respectively). The most widely used treatments in both second and third lines were also LEPs and TCM therapies. The probability of surgery at 1 year was 4% in the secondary dysmenorrhea cohort and 0.2% in the primary dysmenorrhea cohort.

### Resource utilization and costs associated with dysmenorrhea

Patients with dysmenorrhea and the matched control cohort of patients without dysmenorrhea had similar baseline characteristics and resource utilization levels prior to the index period, with all groups reporting a mean number of inpatient admissions of 1.1 for the pre-index period. The analysis of inpatient care demonstrated a significant increase in the number of hospital admissions and length of stay due to dysmenorrhea in the secondary dysmenorrhea cohort vs the controls (5.7 additional admissions per 100 persons over 12 months; 2.2 times longer cumulative length of stay; both \(p<0.0001\)), whereas there was no change in admissions compared to controls in the primary dysmenorrhea cohort. Outpatient care visits occurred substantially more often in patients with dysmenorrhea vs controls, with primary and secondary dysmenorrhea cohorts reporting 8.0 and 8.5 additional outpatient visits over 12 months, respectively, after adjusting for baseline characteristics (both \(p<0.0001\); Table 3).

Primary and secondary dysmenorrhea cohorts had significantly higher mean total health care costs than controls (mean ± SD: 191,680 JPY (1,917 USD) ± 261,226 JPY (2,612 USD) vs 83,615 JPY (836 USD) ± 246,093 JPY (2,461USD) for primary dysmenorrhea cohort and 246,488 JPY (2,465 USD) ± 295,936 JPY (2,959 USD) vs 90,711 JPY (907 USD) ± 297,513 JPY (2,975 USD) for secondary dysmenorrhea cohort; \(p<0.001\); Table 3). After adjusting for baseline characteristics, costs were 2.2 times and 2.9 times higher in patients in the primary and secondary dysmenorrhea cohorts, respectively, than controls (both \(p<0.0001\)). Furthermore, total costs in secondary cases were higher by 33.5% compared to primary cases (\(p<0.001\); Table 3).

### Discussion

This analysis demonstrated that the most commonly prescribed treatments for dysmenorrhea in Japan – excluding non-reimbursed agents – were LEPs and TCM therapies, and patients with secondary dysmenorrhea had the highest utilization rates of LEPs. Important differences in treatment patterns were observed according to the specialty of the prescriber of the first treatment line: obstetricians and gynecologists mainly prescribed LEPs, whereas internal medicine physicians prescribed TCM therapies most frequently. This might be due to differences in severity of dysmenorrhea among women who seek medical care from obstetricians and gynecologists vs internal medicine physicians and because women suffering from more severe dysmenorrhea may be more likely to consult an obstetrician or gynecologist and to receive LEPs. Our findings were generally compatible with the guidelines for gynecological practice by the JSOG and JAOG 2011 edition. Although such guidance suggests NSAID treatment for initial pain relief, our findings show relatively low levels of NSAID use. One of the reasons may be that many patients may have used over-the-counter therapies including NSAIDs, which are not captured by the database. Persistence on LEPs was higher in this analysis than for any of the other therapies assessed, including TCM therapies. It was found by adding up the proportions of patients without defined treatment (13.0% of both primary and secondary dysmenorrhea cases) and those with hemostatic agents (11.2%) and NSAIDs (1.6%) that around 25% of patients diagnosed with dysmenorrhea received no reimbursed treatment or short-term treatments only. Surgery was rarely used to treat dysmenorrhea.

This study also demonstrated that affected patients have more frequent physician visits, equating to roughly one additional visit every 45 days that can be attributed to dysmenorrhea. This high frequency suggests that the disease has a strong impact on the quality of life of these patients. Furthermore, dysmenorrhea does require inpatient care in some cases, with approximately six additional inpatient
Primary dysmenorrhea (n=3,441)

No treatment (570; 16.56%)

First treatment line
N=2,871
173.47–82

Second treatment line
N=1,525
91.23–30

Third treatment line
N=784
97.49–31

No treatment
1,346 (46.88%)

LEP
941 (27.35%)

TCM
50 (5.31%)

Hemo. agents
11 (8.46%)

Progestin
2 (0.06%)

GnRH analog
6 (0.17%)

NSAIDs
51 (1.49%)

Hemo. agents
144 (12.82%)

TCM
136 (38.83%)

Combination
90 (2.62%)

No treatment
741 (48.59%)

No treatment
19 (50.00%)

No treatment
16 (55.17%)

No treatment
5 (41.67%)

No treatment
8 (61.54%)

No treatment
12 (31.58%)

No treatment
9 (91.82%)

No treatment
179 (48.91%)

No treatment
136 (49.64%)

No treatment
93 (72.66%)

No treatment
86 (57.33%)

No treatment
128 (13.60%)

No treatment
19 (14.84%)

No treatment
81 (29.56%)

No treatment
52 (67.53%)

No treatment
262.79–248

No treatment
130 (13.33%)

No treatment
116 (18.99%)

No treatment
10 (12.99%)

No treatment
74 (13.60%)

No treatment
12 (13.33%)

No treatment
12 (13.33%)

No treatment
16 (55.17%)

No treatment
10 (9.43%)

Note: Boxes in the treatment patterns are colored according to the drug used in the first treatment line.

Abbreviations: GnRH, gonadotropin-releasing hormone; Hemo. agents, hemostatic agents; LEP, low-dose estrogen progestin; NSAID, nonsteroidal anti-inflammatory drug; TCM, traditional Chinese medicine; Test. deriv., testosterone derivatives.

Legend:
N (%)
Avg: Average treatment duration (days)
Med: Median of treatment duration (days)
--- ≤10% >10%

Figure 2 Treatment patterns in primary dysmenorrhea cohort (n=3,441).
### Treatment Patterns in Secondary Dysmenorrhea Cohort (n=2,874)

**Legend:**
- N (%): Number and percentage of patients in each treatment group.
- Avg-Med: Average and median treatment duration (days).
- <10%: Patients with treatment duration less than 10 days.
- >10%: Patients with treatment duration greater than 10 days.

**Note:** Boxes in the treatment patterns are colored according to the drug used in the first treatment line.

**Abbreviations:** GnRH, gonadotropin-releasing hormone; Hemo. agents, hemostatic agents; LEP, low-dose estrogen progestin; NSAID, nonsteroidal anti-inflammatory drug; TCM, traditional Chinese medicine; Test. deriv., testosterone derivatives.

---

**Table 1**

<table>
<thead>
<tr>
<th>Treatment Line</th>
<th>No treatment</th>
<th>Progestin</th>
<th>Test. deriv.</th>
<th>LEP</th>
<th>GnRH analog</th>
<th>NSAIDs</th>
<th>Hemo. agents</th>
<th>TCM</th>
<th>Surgery</th>
<th>Combination</th>
<th>LEP</th>
</tr>
</thead>
<tbody>
<tr>
<td>First treatment line</td>
<td>2,871</td>
<td>173.47–82</td>
<td>150.26–51</td>
<td>1424</td>
<td>73.25–91</td>
<td>52</td>
<td>185</td>
<td>143.84–70</td>
<td>30</td>
<td>70</td>
<td>150.34–82</td>
</tr>
</tbody>
</table>

**Figure 3**

Treatment patterns in secondary dysmenorrhea cohort (n=2,874).

Note: Boxes in the treatment patterns are colored according to the drug used in the first treatment line.

Abbreviations: GnRH, gonadotropin-releasing hormone; Hemo. agents, hemostatic agents; LEP, low-dose estrogen progestin; NSAID, nonsteroidal anti-inflammatory drug; TCM, traditional Chinese medicine; Test. deriv., testosterone derivatives.
Table 3 Health care resource utilization and costs in women with vs without dysmenorrhea

<table>
<thead>
<tr>
<th>Resource</th>
<th>Primary</th>
<th>Secondary</th>
<th>Primary vs Secondary</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Controls (N=6,882)</td>
<td>Cases (N=3,441)</td>
<td>Adjusted* difference (Δ) or ratio (r)</td>
</tr>
<tr>
<td></td>
<td>N=5,746</td>
<td>N=2,873</td>
<td>Adjusted* difference (Δ) or ratio (r)</td>
</tr>
<tr>
<td>Inpatient care</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of admissions</td>
<td>1.13 (0.46)</td>
<td>1.23 (0.65)</td>
<td>Δ=0.0002 (NS)</td>
</tr>
<tr>
<td>Number of inpatient days</td>
<td>12.64 (35.11)</td>
<td>13.48 (18.61)</td>
<td>Δ=1.254 (NS)</td>
</tr>
<tr>
<td>Total inpatient costs, JPY</td>
<td>21,998 (180,505)</td>
<td>29,307 (174,684)</td>
<td>r=1.419***</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Outpatient care</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of visits</td>
<td>8.41 (10.71)</td>
<td>16.81 (15.46)</td>
<td>Δ=7.995*</td>
</tr>
<tr>
<td>Number of imaging procedures</td>
<td>1.78 (1.59)</td>
<td>2.28 (2.82)</td>
<td>Δ=0.872*</td>
</tr>
<tr>
<td>Total outpatient costs, JPY</td>
<td>42,911 (104,127)</td>
<td>111,386 (111,232)</td>
<td>r=51.544*</td>
</tr>
<tr>
<td>Pharmacy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total pharmacy costs, JPY</td>
<td>18,706 (64,620)</td>
<td>50,987 (77,456)</td>
<td>Δ=3.627*</td>
</tr>
<tr>
<td>Total health care costs, JPY</td>
<td>83,615 (246,093)</td>
<td>191,680 (261,226)</td>
<td>Δ=21.185*</td>
</tr>
</tbody>
</table>

Notes: Data reported as mean (SD) unless otherwise stated. *Analyses adjusted by age category, total preindex medical costs, presence of preindex hospitalization, type of insurance, quantity corresponding to the modeled variable computed over the preindex period; Generalized Linear Models with negative binomial distribution with identity link function (Δ) or log link function (r). Remark: the annual pharmacy costs attributable to dysmenorrhea mentioned in the discussion (JPY 31,840) were determined as a weighted average of differences in costs between cases and controls across primary and secondary cases. <p><0.01; **p<0.001; ***p<0.0005.

Abbreviations: NS, not statistically significant; SD, standard deviation.
Study limitations include those related to any retrospective claims data analysis, including the lack of data supporting the specific reason for each treatment choice. Furthermore, dysmenorrhea is certainly underreported in claims databases and women included in this study may represent severe cases consulting a practitioner. It was previously estimated that 64.6% of women with menstrual symptoms do not seek medical care. In addition, treatment patterns observed in this analysis may not fully reflect all medications taken to treat dysmenorrhea if they are not reimbursed by insurance, such as COCs or over-the-counter pain medications such as NSAIDs. As a result, women who were identified as having ‘no treatment’ may have received nonreimbursed or over-the-counter therapies during the study period; this could have artificially increased the observed ‘no treatment’ rate. Because the JMDC is an administrative database, its diagnostic reporting may be incomplete as physicians may report codes for which reimbursement is provided instead of the full or actual diagnoses. For example, NSAIDs prescribed for diagnosis with other pain symptoms not associated with menstruation were not included in this study. Patients who require surgery may not receive a diagnosis of dysmenorrhea as they will instead receive a diagnostic code for the underlying condition requiring surgery (e.g., fibroids). The burden of the disease was estimated over the first year of follow-up and not over the whole available period. Building on this research, future analyses may look at costs beyond 12 months, which are the whole available period. Further research is recommended for a maximum of 3 months.

**Conclusion**

Considerable heterogeneity in treatment patterns was observed among patients with dysmenorrhea, with relatively low utilization of LEPs in patients with primary dysmenorrhea compared to secondary dysmenorrhea, and those treated by internal medicine physicians compared to obstetricians and gynecologists. Total health care costs were significantly higher among women with dysmenorrhea compared with similar women who do not suffer from this condition, and excess costs are primarily driven by outpatient care. Further research is recommended to evaluate whether a different allocation of resources, for example with higher utilization of LEPs, may yield better health outcomes and reduce the economic burden of dysmenorrhea.

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**Author contributions**

Yutaka Osuga, Erika Tanaka, and Yoshie Onishi designed the study and wrote the protocol. Olivier Cristeau and Yoshie Onishi wrote the statistical analysis plan. Olivier Cristeau performed the statistical analyses. Yutaka Osuga, Erika Tanaka, and Sayako Akiyama interpreted the results. All authors contributed to critically revising the manuscript and have approved the final manuscript.

**Disclosure**

This study was funded by Bayer Yakuhin Ltd. Sayako Akiyama and Erika Tanaka are employees of Bayer Yakuhin Ltd. Olivier Cristeau and Yoshie Onishi are employees from Creativ-Ceutical, which received funding from Bayer Yakuhin Ltd. Yutaka Osuga has received funding from Bayer Yakuhin, Ltd. during the conduct of the study. He has received funding outside the study from Bayer Yakuhin, Ltd., Nippon Shinyaku Co., Ltd., Fuji Pharma Co., Ltd., and Mochida Pharmaceutical Co., Ltd. Marcia Reinhart provided writing assistance in the production of this manuscript. The authors report no other conflicts of interest in this work.

**References**