Application of three different sets of explicit criteria for assessing inappropriate prescribing in older patients: a nationwide prevalence study of ambulatory care visits in Taiwan

Chirn-Bin Chang,1,2 Shu-Yu Yang,3,4 Hsiu-Yun Lai,5 Ru-Shu Wu,4 Hsing-Cheng Liu,4,6 Hsiu-Ying Hsu,4 Shinn-Jang Hwang,7,8 Ding-Cheng Chan2,9,10

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ABSTRACT

Objective: To investigate the national prevalence of potentially inappropriate medications (PIMs) prescribed in ambulatory care clinics in Taiwan according to three different sets of regional criteria and the correlates of PIM use.

Design: Cross-sectional study.

Setting: This analysis included older patients who visited ambulatory care clinics in 2009 and represented half of the older population included on the Taiwanese National Health Insurance Research Database.

Participants: We identified 1164,701 subjects who visited ambulatory care clinics and were over 65 years old in 2009.

Primary and secondary outcome measures: PIM prevalence according to the 2012 Beers criteria, the PIM-Taiwan criteria and the PRISCUS criteria was estimated separately, and characteristics of PIM users were explored. Multivariate logistic regression analysis was used to determine patient factors associated with the use of at least one PIM. Leading PIMs for each set of criteria were also listed.

Results: The prevalence of having at least one PIM at the patient level was highest with the Beers criteria (86.2%), followed by the PIM-Taiwan criteria (73.3%) and the PRISCUS criteria (66.9%). Polypharmacy and younger age were associated with PIM use for all three sets of criteria. The leading PIMs detected by the PIM-Taiwan and PRISCUS criteria were all included in the 2012 Beers criteria. Non-COX-selective non-steroidal anti-inflammatory drugs in the Beers criteria and benzodiazepines in the PIM-Taiwan and PRISCUS criteria accounted for most leading PIMs.

Conclusions: The prevalence of PIMs was high among older Taiwanese patients receiving ambulatory care visits. The prevalence of PIM and its associated factors varied according to three sets of criteria at the population level.

Strengths and limitations of this study

- This is a population-based study providing evidence on a high prevalence of potentially inappropriate medication (PIM) use in older Taiwanese patients. Efforts should be made to reduce polypharmacy and PIMs.
- By comparing three sets of PIM criteria, the 2012 Beers criteria are the most comprehensive for PIM identification for older Taiwanese patients. PIM prevalence and associated factors varied among the three sets of criteria at the population level.
- Drug availabilities in Taiwan are different for three sets of PIM criteria. For analysis of associated factors of PIMs, several potential confounders such as detailed information of comorbidities, functional status or living situation were not included because of limitations of the study database.

INTRODUCTION

The National Health Insurance (NHI) system enrols more than 99% of residents in Taiwan.1 Healthcare services are paid by the National Health Insurance Administration (NHIA) on a fee-for-service basis with a global budget cap. Patients are free to see any healthcare specialty without restrictions and no referral from a primary care physician is needed.2 Patients are also free to visit physicians in other residential regions. Older adults use more healthcare resources than younger adults because of more chronic diseases. Moreover, older adults in Taiwan use more healthcare resources than those in other countries.3

Drugs are often prescribed to treat chronic diseases in older adults. The pharmacokinetics and pharmacodynamics of these drugs change...
during the aging process and due to chronic disease associated alterations of renal or liver function. Therefore, the incidence of adverse drug events is higher in older adults than in younger adults. These adverse events are important causes of hospitalisation, morbidity and mortality in older adults.\textsuperscript{8-10} Nearly half of these events are considered preventable,\textsuperscript{7} and avoiding prescribing potentially inappropriate medications (PIMs) is an important strategy. Recent studies showed that PIMs were associated with adverse drug reactions (ADRs), ADR-associated medical errors (MEs) and negative clinical outcomes,\textsuperscript{8, 9} even though the PIMs were not directly responsible for these events.\textsuperscript{10} Explicit criteria of PIMs are defined as a list of drugs which are considered inappropriate in general or for older adults with certain chronic conditions. In contrast, implicit criteria of PIMs are statements that are used to evaluate the appropriateness of individual drugs prescribed for older patients.

Because countries vary in their specific approved drugs and national therapeutic guidelines, many countries have developed their own PIM lists.\textsuperscript{11-13} The Beers criteria\textsuperscript{1} established in 1991 were the first to be used to reduce prescribing PIMs in nursing homes. The instrument was revised several times for general use among older adults, with the latest version published in 2012.\textsuperscript{13} In the same year, the PIM-Taiwan criteria, which were derived from seven sets of criteria established in different countries, were also published.\textsuperscript{14} In Europe, the German PRISCUS (Latin for ‘old and venerable’) criteria were shown to discourage PIM use.\textsuperscript{15-17} Many studies based on PIM criteria focus on the prevalence and correlates of PIMs in limited samples, such as inpatient populations or nursing home residents; fewer studies apply these criteria in a general, nationwide population. To the best of our knowledge, few criteria have been used for secondary health insurance claims data analysis studies.\textsuperscript{17-20} 35

The aim of this study was to estimate the prevalence of PIMs in a nationwide population prescribed during ambulatory care visits and to investigate the ability of PIM-Taiwan to identify PIMs versus the 2012 Beers criteria and the PRISCUS criteria. We also investigated the factors associated with PIM use and list the leading medications detected by three sets of criteria.

**METHODS**

**Source of data**

We conducted a secondary data analysis of medical claims from the National Health Insurance Research Database (NHIRD) in Taiwan for 2009. This study was approved by the Research Ethics Committee of National Taiwan University Hospital in 2013. NHIRD collected all claims data of hospitalisation, ambulatory care visits, emergency department visits and home healthcare. In all settings, physicians could prescribe oral, intravenous injected medications or those administered through other routes. Also, physicians could order other therapeutic procedures or programmes, such as surgical interventions or rehabilitation programmes. In this study, only drug prescriptions for the oral route in 2009 covered by NHI were collected for analysis. The patient records and information in this database were anonymised and de-identified before our analysis. We enrolled all patients who were aged 65 years or older who had at least one ambulatory care visit in 2009 if their birthday was an odd number. For confidentiality control, NHI would not release the records for the entire population of older patients for analysis. Thus, this study represents half the population of older Taiwanese patients enrolled in NHI who attended ambulatory care for their acute or chronic illness without considering their place of residence.

**Study design and identification of PIM users**

Each brand name oral medication covered by NHI has a unique code. For clarity of medication assignment and to avoid ambiguity, the NHI code was converted to the Anatomical Therapeutic Chemical (ATC) code to efficiently identify PIMs on the database. (This mapping strategy was created by Yea-Huei Kao Yang, Ching-Lan Cheng and Ya-Chuan Chang; their study was called ‘To establish therapeutic classification of the pharmaceutical preparations’, which was the final report of project no. 102-TFDA-P-044). Information on the pharmacological or chemical subgroups for each individual drug was also available on the WHO Collaborating Center for Drug Statistics Methodology website.\textsuperscript{21} We identified the PIMs independent of chronic conditions because the NHIRD did not contain an exhaustive list of diagnoses for their patients. Therefore, we identified PIMs based on statements that listed PIMs without considering chronic conditions, including table 1 of the original PIM-Taiwan criteria,\textsuperscript{14} the entire PRISCUS criteria\textsuperscript{22} and table 2 of the original 2012 Beers criteria.\textsuperscript{18} Individuals who were prescribed at least one PIM in 2009 based on the three sets of PIM criteria were considered as PIM users. The prevalence and associated factors of PIMs were analysed separately for the three sets of criteria. Age, gender, diagnoses, number of ambulatory care visits, physician visits and hospital visits, and generic names of oral medications were collected for all patients. Also, only three major International Classifications of Diseases, 9th edition Clinical Modification (ICD-9-CM) codes were collected for each ambulatory care visit on the database because of NHI administration specifications. We collected all the diagnoses made during the study period for each participant. The Charlson Comorbidity Index (CCI)\textsuperscript{23} which has been validated for higher scores associated with higher risk of death, was calculated based on all diagnoses for each participant with ICD-9-CM codes. Other patient data, such as weight, height, blood pressure, functional status and over-the-counter drug use, were not available due to the limitations of NHIRD.

**Statistical analysis**

Patient demographic characteristics (age and gender), comorbidities (CCI) and healthcare resource utilisation
(frequency of clinic visits, number of hospital visits and number of oral medications prescribed) for 2009 were investigated for their association with PIM use. CCI was stratified as a score of 1, 2 or more than 2. We used step-wise multivariate logistic regression models to identify the correlates of having at least one PIM at the patient level. Each set of criteria was analysed separately to determine its associated factors. The statistical significance \( \alpha \) was set at \( p<0.05 \). We also ranked the leading PIMs from each set of PIM criteria based on PIMs accounting for more than 1% of total prescriptions. Data were analysed using SAS for Windows V.9.2 (SAS Institute Inc, Cary, North Carolina, USA).

**RESULTS**

**Basic characteristics of study subjects**

As shown in table 1, a total of 1 164 701 older patients were enrolled in this study. More than half (55%) were aged 65–74 years and 48% were men. Nearly half of the patients had more than 10 diagnoses and nearly 40% had a score of at least 2 on the CCI, which indicates a high disease burden for our study population. The group of patients aged over 85 years had the highest proportion (49%) scoring at least 2 on the CCI among three age stratifications. Healthcare resource use was also high: 43% of patients exceeded 24 ambulatory care visits and 32% of patients visited more than four hospitals in 1 year. Among three age groups, patients aged 75–84 years had the highest rate of clinic visits, but those aged 65–74 years visited the most hospitals. All patients in our study population had been prescribed at least one drug during their ambulatory clinic visits. Forty-three percent of all patients were prescribed more than 20 different medications in 1 year, especially those aged 74–85 years. Those who were aged over 85 years had the lowest rates of hospital visits and the lowest number of medications.

**Description of PIM users**

The characteristics of three sets of PIM criteria are listed in table 2. The 2012 Beers criteria included the highest

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### Table 1 Basic characteristics of Taiwan study population in 2009

| N (%) | Overall 1 164 701 (100%) | 65–74 years 639 634 (54.92%) | 75–84 years 414 265 (35.57%) | ≥85 years 110 802 (9.51%) | p Value |
|-------|--------------------------|-------------------------------|-------------------------------|--------------------------|--------|
| Gender |                          |                               |                               |                          | <0.01  |
| Male   | 558 904 (47.99)          | 296 436 (46.34)              | 210 250 (50.75)               | 52 218 (47.13)          |        |
| Female | 605 797 (52.01)          | 343 198 (53.66)              | 204 015 (49.25)               | 58 584 (52.87)          |        |
| Number of diagnoses in 1 year |                          |                               |                               |                          | <0.01  |
| 1–10   | 633 846 (54.42)          | 362 956 (56.74)              | 209 045 (50.46)               | 61 845 (55.82)          |        |
| >10    | 530 816 (45.58)          | 276 657 (43.26)              | 205 027 (49.54)               | 48 952 (44.18)          |        |
| Charlson Comorbidity Index |                          |                               |                               |                          | <0.01  |
| Score 1| 268 123 (23.02)          | 148 125 (23.16)              | 94 506 (22.81)                | 25 492 (23.01)          |        |
| Score 2| 183 247 (15.73)          | 94 274 (14.74)               | 69 395 (16.75)                | 19 578 (17.67)          |        |
| Score >2| 299 585 (25.72)         | 140 397 (21.95)              | 124 567 (30.07)               | 34 621 (31.25)          |        |
| Number of outpatient clinic visits in 1 year |                          |                               |                               |                          | <0.01  |
| 1–24  | 656 878 (56.40)          | 383 403 (59.94)              | 210 536 (50.82)               | 62 999 (56.80)          |        |
| >24   | 507 823 (43.60)          | 256 231 (40.06)              | 203 729 (49.18)               | 47 863 (43.20)          |        |
| Number of hospital visits in 1 year |                          |                               |                               |                          | <0.01  |
| 1–4   | 783 351 (67.26)          | 419 052 (65.51)              | 279 393 (67.44)               | 84 906 (76.63)          |        |
| >4    | 381 350 (32.74)          | 220 582 (34.49)              | 134 872 (32.56)               | 25 896 (23.37)          |        |
| Number of medications in 1 year |                          |                               |                               |                          | <0.01  |
| 1–20  | 659 452 (56.62)          | 375 455 (58.70)              | 218 320 (52.70)               | 65 677 (59.27)          |        |
| >20   | 505 249 (43.38)          | 264 179 (41.30)              | 195 945 (47.30)               | 45 125 (40.73)          |        |

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### Table 2 Characteristics of the three sets of explicit criteria and their performance in detecting potentially inappropriate medications (PIMs) in Taiwanese study patients in 2009

|                  | Beers criteria | PIM-Taiwan criteria | PRISCUS criteria |
|------------------|----------------|---------------------|-----------------|
| Year of criteria | 2012           | 2012                | 2010            |
| Country          | USA            | Taiwan              | German          |
| Statements*      | 34             | 24                  | 15              |
| Number of medications† | 137 84 83 | 84 (100%)          | 83 (84.15%)     |
| Availability of PIMs listed in each set of criteria in Taiwan | 105 (76.64%) | 84 (100%)          | 69 (84.15%)     |
| Number of patients with any PIMs | 1 004 234 (86.22%) | 853 915 (73.32%) | 778 825 (66.87%) |

*Statements are those for potentially inappropriate medications (PIMs) without considering drug–disease or drug–syndrome interactions.
†Medications are considered as potentially inappropriate without considering drug–disease or drug–syndrome interactions.
number of statements and individual drugs, and the numbers of individual drugs in the PIM-Taiwan and PRISCUS criteria were only about 60% of the Beers’ lists. The availability of PIMs is highest in Taiwan but lowest in the 2012 Beers criteria (100% vs 76%). After considering medication availability, the 2012 Beers criteria still had the highest number of individual drugs in three sets of criteria. The prevalence of patients receiving at least one PIM during the year was 86%, 73% and 67% using the Beers criteria, the PIM-Taiwan criteria and the PRISCUS criteria, respectively.

By applying the 2012 Beers criteria, 87% of women had been prescribed at least one PIM (table 3). Among the patients receiving PIMs from the three sets of criteria, those aged 74–85 years had the highest risk of being prescribed at least one PIM using the 2012 Beers criteria and the PRISCUS criteria. However, those aged 65–74 years had the highest risk of being prescribed medications listed in the PIM-Taiwan criteria. Patients having a higher number of diagnoses were at high risk of being prescribed PIMs compared with those with a lower number of diagnoses. In accordance with their high disease burden, when patient visits exceeded 24 ambulatory clinics and more than four hospitals, they had a higher risk of receiving PIMs. The percentage of patients having PIMs listed in the 2012 Beers criteria increased to over 95% if patients exceeded 24 clinic visits, four different hospital visits and 20 different medications.

### Association factors of PIMs

In multivariate analysis, male patients were more likely than female patients to be prescribed PIMs listed in the Beers and PRISCUS criteria (table 4). Patients aged over 75 (75–85, and 85 and older) were more likely to receive PIMs listed in the PRISCUS criteria only. Patients who were prescribed more medications in 2009 were more likely to have PIMs across all three sets of criteria. When a patient was prescribed one more drug, the odds of having PIMs increased the most (OR=1.6) when the Beers criteria were used for assessment. Inverse relationships were found between CCI score and the use of at least one PIM, except when using the Beers criteria.

### Leading medications in three sets of PIM criteria

Overall, 27 485 169 prescriptions were written for our study population in 2009. The leading PIMs varied in the three sets of criteria. Twenty-three medications had a prevalence higher than 1% of total prescriptions. Nineteen medications were identified by the Beers criteria (table 5), while eight were identified by the PIM-Taiwan criteria and nine by the PRISCUS criteria. In the Beers and PRISCUS criteria, psychotropic drugs and non-steroidal anti-inflammatory drugs (NSAIDs) accounted for the most frequent PIMs. In contrast, the most frequent PIMs according to the PIM-Taiwan criteria were first-generation histamine H1-receptor antagonists. Diclofenac accounted for 5% of total study population prescriptions and was ranked first by the Beers criteria.

### Table 3

| Description of PIM users based on three sets of criteria in study population | N (%) | 2012 Beers | p Value | PIM-Taiwan | p Value | PRISCUS | p Value |
|---|---|---|---|---|---|---|---|
| Patient number | 1 004 243 (86.22) | 853 915 (73.32) | <0.01 | 778 825 (66.87) | <0.01 | 778 825 (66.87) | <0.01 |
| Gender | | | | | | | |
| Male | 476 549 (85.26) | 397 196 (71.07) | <0.01 | 368 560 (65.94) | <0.01 | 368 560 (65.94) | <0.01 |
| Female | 527 694 (87.11) | 456 719 (75.39) | <0.01 | 410 265 (67.72) | <0.01 | 410 265 (67.72) | <0.01 |
| Age (years) | | | | | | | |
| 65–74 | 548 368 (85.73) | 473 753 (74.07) | <0.01 | 407 054 (63.64) | <0.01 | 407 054 (63.64) | <0.01 |
| 75–84 | 361 763 (87.33) | 305 697 (73.79) | <0.01 | 294 762 (71.15) | <0.01 | 294 762 (71.15) | <0.01 |
| ≥85 | 94 112 (84.94) | 74 465 (67.21) | <0.01 | 77 009 (69.50) | <0.01 | 77 009 (69.50) | <0.01 |
| Number of diagnoses | | | | | | | |
| 1–10 | 486 807 (76.80) | 372 386 (58.75) | <0.01 | 326 566 (51.52) | <0.01 | 326 566 (51.52) | <0.01 |
| >10 | 517 433 (97.48) | 481 526 (90.71) | <0.01 | 452 258 (85.20) | <0.01 | 452 258 (85.20) | <0.01 |
| Charlson Comorbidity Index | | | | | | | |
| Score 1 | 232 362 (86.66) | 195 198 (72.80) | <0.01 | 181 090 (67.54) | <0.01 | 181 090 (67.54) | <0.01 |
| Score 2 | 162 592 (88.73) | 137 535 (75.05) | <0.01 | 132 979 (72.57) | <0.01 | 132 979 (72.57) | <0.01 |
| Score >2 | 279 879 (93.42) | 241 659 (80.66) | <0.01 | 242 536 (80.96) | <0.01 | 242 536 (80.96) | <0.01 |
| Number of outpatient clinic visits in 1 year | | | | | | | |
| 1–24 | 514 117 (78.27) | 406 031 (61.81) | <0.01 | 349 443 (53.20) | <0.01 | 349 443 (53.20) | <0.01 |
| >24 | 490 126 (96.52) | 447 884 (88.20) | <0.01 | 429 382 (84.55) | <0.01 | 429 382 (84.55) | <0.01 |
| Number of hospital visits in 1 year | | | | | | | |
| 1–4 | 532 047 (76.57) | 501 371 (76.57) | <0.01 | 457 348 (58.38) | <0.01 | 457 348 (58.38) | <0.01 |
| >4 | 372 196 (98.82) | 352 544 (98.82) | <0.01 | 321 477 (84.3) | <0.01 | 321 477 (84.3) | <0.01 |
| Number of medications in 1 year | | | | | | | |
| 1–20 | 504 939 (76.57) | 379 265 (57.51) | <0.01 | 336 536 (51.03) | <0.01 | 336 536 (51.03) | <0.01 |
| >20 | 499 304 (98.82) | 474 650 (93.94) | <0.01 | 442 289 (87.54) | <0.01 | 442 289 (87.54) | <0.01 |

PIM, potentially inappropriate medication.
DISCUSSION

This is the first nationally representative evaluation of PIM use with an instrument developed in Taiwan. We were able to compare the results of the PIM-Taiwan criteria with the results of two other sets of well established PIM criteria. All three sets of PIM criteria identified high percentages of older adults who had been prescribed at least one PIM in the ambulatory care setting. PIM prevalence estimated from the PIM-Taiwan criteria ranked in the middle among the three sets of criteria. Generally, polypharmacy and being aged 65–74 years were associated with more PIM use.

The annual prevalence of PIMs in our population is higher than that reported in previous community-based or health insurance studies (Beers range 28–65% and PRISCUS 22–25%).17 24–26 Only one study applying the Beers criteria in an Indian hospitalised population showed a high rate of PIM exposure similar to Taiwan’s.3 Several factors are responsible for the relatively high prevalence in our study. First, the 2012 version of the

Table 4  Multivariate logistic regression analysis for having at least one drug as potentially inappropriate (N=1,164,701)

|                          | 2012 Beers | PIM-Taiwan | PRISCUS |
|--------------------------|------------|------------|---------|
| Gender                   |            |            |         |
| Male                     | 1          | 1          | 1       |
| Female                   | 0.93 (0.93 to 0.93)* | 1.02 (1.02 to 1.02)* | 0.87 (0.87 to 0.88)* |
| Age (years)              |            |            |         |
| 65–74                    | 1          | 1          | 1       |
| 75–84                    | 0.94 (0.94 to 0.94)* | 0.93 (0.93 to 0.93)* | 1.05 (1.04 to 1.05)* |
| ≥85                      | 0.89 (0.89 to 0.90)* | 0.82 (0.82 to 0.83)* | 1.03 (1.02 to 1.03)* |
| Charlson Comorbidity Index |          |            |         |
| Score 1                  | 1          | 1          | 1       |
| Score 2                  | 0.88 (0.88 to 0.88)* | 0.85 (0.85 to 0.85)* | 0.89 (0.88 to 0.89)* |
| Score ≥2                 | 1.03 (1.03 to 1.04)* | 0.89 (0.89 to 0.91)* | 0.90 (0.90 to 0.91)* |
| Number of medications†   | 1.60 (1.60 to 1.60)* | 1.56 (1.56 to 1.56)* | 1.46 (1.46 to 1.46)* |

*All variables in table 1 were included in logistic regression and only those with significance were listed.  
*p<0.01.  
†Numbers of medications were analysed as continuous variables in multivariate logistic regression analysis.

Table 5  The leading PIMs* identified in 27,485,169 prescriptions in a Taiwanese study population in 2009

| Drug name and anatomical (ATC) code | 2012 Beers | PIM-Taiwan | PRISCUS |
|-------------------------------------|------------|------------|---------|
| 1 Diclofenac (M01AB05)/5.07%        | ✓          | ✓          |         |
| 2 Cimetidine (A02BA01)/3.69%        |            | ✓          |         |
| 3 Clorzoxazone (M03BB03)/2.88%      | ✓          | ✓          |         |
| 4 Zolpidem (N05CF02)/2.80%          | ✓          | ✓          |         |
| 5 Alprazolam (N05BA12)/2.51%        | ✓          | ✓          |         |
| 6 Dipyridamole (B01AC07)/2.45%      | ✓          | ✓          |         |
| 7 Mefenamic acid (M01AG01)/2.30%    | ✓          | ✓          |         |
| 8 Lorazepam (N05BA06)/2.08%         | ✓          | ✓          |         |
| 9 Nifedipine (B01AC05)/2.01%        | ✓          | ✓          |         |
| 10 Ibuprofen (M01AE01)/1.90%        | ✓          | ✓          |         |
| 11 Metoclopramide (A03FA01)/1.76%   | ✓          | ✓          |         |
| 12 Doxazosin (C02CA04)/1.74%        | ✓          | ✓          |         |
| 13 Meloxicam (M01AC06)/1.56%        | ✓          | ✓          |         |
| 14 Dextroamphetamine (R06AB02)/1.55%| ✓          | ✓          |         |
| 15 Piretrem (N06BX03)/1.46%         | ✓          | ✓          |         |
| 16 Mequitazine (R06AD07)/1.43%      | ✓          | ✓          |         |
| 17 Diazepam (N05BA01)/1.31%         | ✓          | ✓          |         |
| 18 Clonazepam (N03AE01)/1.28%       | ✓          | ✓          |         |
| 19 Chlorpheniramine (R06AB54)/1.24% | ✓          | ✓          |         |
| 20 Terazosin (G04CA03)/1.18%        | ✓          | ✓          |         |
| 21 Estazolam (N05CD04)/1.17%        | ✓          | ✓          |         |
| 22 Fluoxetine (N05BA17)/1.10%       | ✓          | ✓          |         |
| 23 Glibenclamide (A10BB01)/1.07%    | ✓          | ✓          |         |

* A potentially inappropriate medication (PIM) was defined as ‘leading’ if it accounted for more than 1% of total prescriptions.  
ATC, Anatomical Therapeutic Chemical.
Beers criteria added more medications as PIMs, including all non-COX-selective NSAIDs and benzodiazepines regardless of their half-life. Second, under the NHIA policy to control the growth of drug expenditure, physicians may prefer to prescribe cheaper drugs. Therefore, cheaper drugs including generic first antihistamine and non COX-II selective NSAIDs accounted for the high prevalence of PIMs in Taiwan. Third, our NHI offered unrestricted access to physicians and almost all prescription medications. Therefore, Taiwanese patients made more frequent visits to ambulatory care clinics than older American patients and polypharmacy is highly prevalent in Taiwan compared with western countries, which certainly increases the risk of PIM use. There were only limited studies of polypharmacy prevalence in Asia but this problem was also apparent in Korea. Although the PIM-Taiwan criteria were established for local use in Taiwan, most individual PIMs in our criteria were included in the Beers criteria and other country-specific PIM criteria. It is convenient for other regions without their own PIM criteria to apply this tool to measure prescription quality at the population level. Nevertheless, PIM prevalence in Taiwan only represented local prescription preference and health resource utilisation under the coverage of the NHI of Taiwan. Also, the availabilities of drugs listed in the 2012 Beers criteria and the PRISCUS criteria were different, which suggests that the drug markets and prescription patterns are different in the USA, Germany and Taiwan. Our findings would be applicable to healthcare systems with universal health insurance and relatively unrestricted medication prescriptions.

Education of physicians, pharmacists and patients about PIM lists will increase awareness and possibly decrease PIM use. Also, computer-assisted prescription alert programmes incorporated in PIM lists would provide possible therapeutic alternatives, such as the PIM-Taiwan and PRISCUS criteria, and may help to reduce use of these high-risk medications. In the near future, a computer-assisted prescription warning system will be incorporated into our NHI identification card based on table 1 of the PIM-Taiwan criteria. Physicians will be discouraged from prescribing these medications for older adults at each clinical encounter. Lastly, including fewer PIMs in a hospital pharmacy, for example, purchase famotidine rather than cimetidine, is also an effective strategy to reduce PIM use.

For factors associated with PIMs, we found different patterns of risk between three sets of PIM criteria. Several studies applying the 2012 Beers and PRISCUS criteria found that female gender was associated with use of more PIMs. Various clinical settings and study populations can account for these differences. In our study population, acute upper respiratory infections, benign prostate hypertrophy and contact dermatitis were leading diagnoses and also more prevalent in our male patients (data not shown). The 2012 Beers criteria and PRISCUS criteria included more individual drugs such as α-1 blockers and first-generation antihistamines; therefore, the risk of being prescribed PIMs was increased in male patients. For the PIM-Taiwan criteria, the literature was limited. Contrary to our previous study on home care service recipients, female patients were more likely to be prescribed PIMs using the PIM-Taiwan criteria in the current study. A possible explanation is that cimetidine, muscle relaxants and short-acting benzodiazepines were major PIMs in the PIM-Taiwan criteria. The prevalence of functional digestive disorders, disease of the musculoskeletal system and mental disorders was higher among women; therefore, their risk of receiving PIMs increased. As for the association between the CCI and PIM use, it is difficult to reach a conclusion. In previous studies, positive and negative associations between the CCI and PIMs were observed. We found that older patients with a CCI score of 1 had the highest risk when we applied the PIM-Taiwan and PRISCUS criteria. However, on applying the Beers criteria, patients with a CCI of 3 had the highest risk. The only explanation is a cautious prescribing attitude of physicians to reduce prescribing medications that have a higher risk of side effects for older patients with multiple comorbidities. Our database only included patients’ diseases which can be classified using ICD-9-CM codes; therefore, the results should be interpreted with caution because other important risk factors for PIM use, such as functional status, were not included in the analysis. Further prospective studies investigating the attitude of physicians towards PIMs are necessary.

NSAIDs, benzodiazepines and first-generation antihistamines were major offending PIMs in this study, and these results are consistent with previous studies using the 2012 Beers and PRISCUS criteria. Almost all PIMs identified by the PIM-Taiwan and PRISCUS criteria are included in the 2012 Beers criteria. Only certain non-COX-selective NSAIDs and long-acting benzodiazepines are listed in the PIM-Taiwan criteria, and similarly in the PRISCUS criteria. In contrast, the 2012 Beers criteria include all non-COX-selective NSAIDs and long, intermediate and short-acting benzodiazepines. This comprehensive inclusion of benzodiazepines was supported by several studies. Almost all PIM criteria were established based on expert consensus; therefore, they should be used to support the prescribing decision not to substitute the physician’s decision on prescription. Although some studies show that drugs in the PIM criteria contributed to only a small percentage of ADRs, a recent study with a different design demonstrated an association between PRISCUS drugs and ADRs. More importantly, Koyama and colleagues found that patients taking PIMs and drugs with high anticholinergic effects are at higher risk of functional and cognitive impairment in a 5-year prospective cohort study. Their study implied that the adverse events related to PIMs may be difficult to identify in cross-sectional studies. Further study is needed to confirm the
impact of these PIM lists on the quality of life, healthcare resource use and mortality of older adults.45

This study has several strengths. First, it was a population-based study including half of the total population of older patients included on the Taiwanese NHIRD. Therefore, the data are highly representative. Limited studies have included national population data to evaluate PIM criteria. Second, we selected the evidence-based Beers criteria and well studied PRISCUS criteria for comparison with the PIM-Taiwan criteria. Although the 2012 Beers criteria seem to be more comprehensive, the PIM-Taiwan criteria which only contains half the number of individual drugs as the 2012 Beers criteria can identify 80% of PIMs prescribed for older Taiwanese patients. PIM-associated factors also varied among the three sets of criteria. This suggests that region-specific criteria are better tools for evaluating prescription quality in specific drug markets and health insurance systems. Third, all prescribed medications are covered by the NHI, which means our database contains detailed information of medications, including combination drugs.

Our study is a secondary data analysis with fundamental limits. First, only three diagnoses were made for one ambulatory care visit. Therefore, we only adapted PIM criteria independent of chronic diseases. Second, some drugs listed in the Beers criteria (meprobamate, trimethobenzamide, oxaprozin etc.) and the PRISCUS criteria (e.g. metildigoxin, tetrazepam, briotizolam etc.) are not available in Taiwan, but all drugs in the PIM-Taiwan criteria are available. Drug availability is one of the major determinants of PIM prevalence.32 Even so, we still found that the Beers criteria detected more PIMs than the other criteria, mainly due to the inclusion of more medications on the list. Third, to analyse associated factors of PIMs, not all potential confounders such as detailed information of comorbidities, functional status or living situation were included because of limitations of the study database.

In conclusion, the prevalence of PIM users in our population is over 65%. The prevalence was highest for the 2012 Beers criteria, which also included most leading PIMs in the PIM-Taiwan and PRISCUS criteria. The prevalence of PIMs and associated factors varied among the three sets of criteria at the population level.
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