Frequently reported adverse events of rebamipide compared to other drugs for peptic ulcer and gastroesophageal reflux disease

Eunkyeong Jang1,3, Minju Park1,3, Ji Eun Jeong1,3, Ji Young Lee1,3 & Myeong Gyu Kim1,2*

This study aimed to detect safety signals of rebamipide and search for adverse events (AEs) of rebamipide that are more common than those of other drugs for peptic ulcer disease (PUD) and gastroesophageal reflux disease (GERD) in the elderly population. A total of 101,735 AE reports for drugs used to treat PUD and GERD between 2009 and 2018 from the KIDS-KAERS database (KIDS-KD) were used. Disproportionality analysis was performed to calculate the proportional reporting ratio (PRR), reporting odds ratio (ROR), and information component (IC). Drug labels in Korea, Japan, and China were reviewed to identify signals that have been listed. AEs frequently reported in the elderly population were also analyzed. Seriousness and median time to AEs were evaluated for statistically significant AEs. A total of 14 signals were detected, and 4 signals (dry mouth, dermatitis, purpura/petechia, and fluid overload) were not listed on drug labels; however, they may be included as part of other listed AEs. In the elderly population, 11 AEs such as dyspepsia/indigestion/gastrointestinal distress, somnolence, dry mouth, and edema were common. These AEs were not serious and occurred within 2–9 days. This study identified possible AEs of rebamipide, a relatively safe drug.

Rebamipide is a mucoprotective drug for peptic ulcer disease (PUD) and gastroesophageal reflux disease (GERD). It induces prostaglandins, resulting in increased blood flow to the gastric mucosa, mucous secretion, and enhanced mucosal defense. It also scavenges free radicals and inhibits inflammatory reactions. In Korea, the size of the outpatient prescription market for rebamipide in 2020 was $90 million (approximately $0.08 a tablet). Rebamipide is known as a drug with few and mild adverse drug reactions (ADRs). The most common ADRs are gastrointestinal (GI) reactions such as nausea, vomiting, constipation, diarrhea, and bloating. Gastritis and GERD are the seventh and eighth most common diseases, respectively, in the Korean elderly population. Proton pump inhibitors (PPIs) and histamine receptor antagonists (H2RAs) are representative drugs used to treat PUD and GERD. However, PPIs can increase the risk of Clostridium difficile infection, bone loss, and fractures in elderly patients; thus, it is recommended not to use PPIs for more than eight weeks. In addition, H2RAs are generally avoided for patients with delirium even though the evidence for adverse cognitive effects is weak. For this reason, rebamipide has been frequently used, which is considered relatively safe for the elderly. As the elderly population increases, it is necessary to determine whether there are any unknown or incompletely documented adverse events (AEs) of rebamipide by examining large-scale AE reports and verify the safety of rebamipide.

Pharmacovigilance is a scientific study and activity involving the detection, evaluation, interpretation, and prevention of drug-related problems. Pharmacovigilance is crucial because it can identify a safety signal, defined as “reported information on a possible causal relationship between an AE and a drug, of which the relationship is unknown or incompletely documented previously”, and provide real-world evidence. The Food and Drug Administration (FDA) Adverse Event Reporting System (FAERS) has become an important resource for pharmacovigilance analysis. Similarly, the Korea Institute of Drug Safety and Risk Management (KIDS) developed the Korea Adverse Event Reporting System (KAERS) database (KIDS-KD). The KIDS-KD contains data for AEs

1College of Pharmacy, Ewha Womans University, Seoul 03760, Republic of Korea. 2Graduate School of Pharmaceutical Sciences, Ewha Womans University, Seoul 03760, Republic of Korea. 3These authors contributed equally: Eunkyeong Jang, Minju Park, Ji Eun Jeong, and Ji Young Lee. *email: kimmg@ewha.ac.kr
that were spontaneously reported and can be used for pharmacovigilance analysis. Renal neoplasm, dementia, acute kidney injury, chronic kidney disease, and hepatotoxicity have been evaluated for their causal relationship with PPIs in several pharmacovigilance studies. However, pharmacovigilance studies of rebamipide are limited. A study evaluated pulmonary AEs using the KIDS-KD. Another study evaluated the efficacy of rebamipide in preventing non-steroidal anti-inflammatory drug (NSAID)-induced lower GI tract injury using the KIDS-KD and the Japanese Adverse Event Reporting Database.

The aim of this study was first to detect safety signals of rebamipide that are not listed on drug labels using the KIDS-KD in general population, second to identify AEs of rebamipide that are more common than those of other drugs used for PUD and GERD in subgroup analysis of the elderly population.

Results

Characteristics of AE reports. Figure 1 shows the study flow diagram. There were 101,735 AE reports for drugs used to treat PUD and GERD between January 2009 and December 2018. We used 173,637 A02B drug–AE pairs for signal detection regardless of causality and 59,505 A02B drug–AE pairs classified as certain, probable, or possible for signal detection considering causality. Moreover, 16,773 A02B drug–AE pairs were used to determine frequently reported AEs in the elderly population. The characteristics of AE reports can be found in our previous study.

Signal detection. We detected 11 signals from drug–AE pairs classified as certain, probable, or possible. Of the 11 signals, 8 signals were AEs listed on drug labels in three countries, which included somnolence, dyspepsia/indigestion/GI distress, face edema, generalized edema, peripheral edema, periorbital edema, and thirst. As edema was included on the drug labels without further details, various types of edemas were evaluated as previously mentioned. Dry mouth, dermatitis, and purpura/petechia were not listed on the drug labels under the corresponding names.

Regardless of causality, 15 rebamipide–AE pairs met the criteria of the PRR, ROR, and IC (Table 2). Of these AEs, 6 AEs were already listed on drug labels in Korea, Japan, and China. Among the other 9 AEs, dry mouth and fluid overload were clinically relevant.

A total of 14 signals were identified according to the results presented in both tables: somnolence, dyspepsia/indigestion/GI distress, dry mouth, face edema, generalized edema, peripheral edema, periorbital edema, thirst, dermatitis, purpura/petechia, vertigo, increased stool frequency, and fluid overload. Dry mouth, dermatitis, purpura/petechia, and fluid overload were not listed on the drug labels.

AEs frequently reported in the elderly population. Table 3 shows 11 AEs that were common in the elderly population. Dyspepsia/indigestion/GI distress (ROR = 2.24), somnolence (ROR = 2.23), dry mouth (ROR = 3.97), and face edema (ROR = 3.04) were representative AEs reported substantially more than other A02B drugs. All AEs, except for purpura/petechia, were significant in subgroup analysis. There was no serious AE, and the median time to AEs ranged from 2 to 9 days.
Discussion

We detected the safety signals of rebamipide in this study, which is the first to detect safety signals and compare the AEs of rebamipide with those of other drugs used to treat PUD and GERD in elderly patients.

A total of 14 signals were detected, and most were already included on drug labels. Signals not included on the drug labels (dry mouth, dermatitis, purpura/petechia, and fluid overload) may be included as part of other listed ADRs. Dry mouth is one of the peripheral signs and attributes of thirst, which is listed on drug labels\(^1\). Dermatitis is a general term that describes a common skin irritation and a group of drug hypersensitivity reactions involving the skin\(^17,18\). Hypersensitivity reactions such as urticaria, rash, itching, and eczema are known ADRs of rebamipide and can be regarded as dermatitis. Purpura and petechia can be associated with a decreased platelet count, which is indicated on drug labels\(^19\). Edema, which is listed on drug labels, can be regarded as fluid overload. Nevertheless, for clarity, these signals should be included on the drug labels.

Regardless of causality, some AEs met the criteria of signal detection; however, they may be explained by the patient’s underlying diseases or co-administered drugs. For example, apathy has been well described in patients with psychiatric diseases such as major neurocognitive disorders, schizophrenia, and major depressive disorder and taking drugs that act on the central nervous system (CNS), such as selective serotonin reuptake

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### Table 1

| AEs                        | Cases (rebamipide) | PRR  | ROR  | IC 95% CI | Label | Clinical relevance |
|----------------------------|--------------------|------|------|-----------|-------|--------------------|
| Somnolence                 | 1194               | 2.66 | 2.86 | 0.94      | Yes   |                    |
| Dyspepsia/indigestion/GI distress | 1115         | 2.13 | 2.26 | 0.73      | Yes   |                    |
| Dry mouth                  | 482                | 4.34 | 4.50 | 1.29      | No    |                    |
| Face edema                 | 247                | 2.78 | 2.82 | 0.86      | Yes\(^1\) |                |
| Generalized edema          | 166                | 3.44 | 3.48 | 1.00      | Yes\(^1\) |                |
| Malaise/feeling queasy     | 116                | 2.18 | 2.19 | 0.53      | Yes   |                    |
| Peripheral edema           | 62                 | 2.28 | 2.29 | 0.46      | Yes\(^2\) |                |
| Periorbital edema          | 54                 | 2.07 | 2.08 | 0.33      | Yes\(^1\) |                |
| Thirst                     | 43                 | 5.16 | 5.18 | 1.02      | Yes   |                    |
| Dermatitis                 | 37                 | 2.01 | 2.01 | 0.20      | No    |                    |
| Purpura/petechia           | 31                 | 2.14 | 2.15 | 0.21      | No    |                    |

### Table 2

| AEs                        | Cases (rebamipide) | PRR  | ROR  | IC 95% CI | Label | Clinical relevance |
|----------------------------|--------------------|------|------|-----------|-------|--------------------|
| Somnolence                 | 1285               | 2.94 | 3.04 | 1.11      | Yes   | -                 |
| Dry mouth                  | 551                | 2.69 | 2.73 | 0.97      | No    | Yes               |
| Generalized edema          | 198                | 2.03 | 2.04 | 0.58      | Yes   | -                 |
| Malaise/feeling queasy     | 144                | 2.20 | 2.20 | 0.63      | Yes   | -                 |
| Apathy                     | 56                 | 2.02 | 2.02 | 0.36      | No    | No\(^1\)          |
| Thirst                     | 51                 | 2.89 | 2.89 | 0.70      | Yes   | -                 |
| Vertigo                    | 47                 | 2.39 | 2.39 | 0.49      | Yes   | -                 |
| Micturition disorder       | 43                 | 2.08 | 2.08 | 0.33      | No    | No\(^1\)          |
| Herpes simplex             | 25                 | 2.12 | 2.13 | 0.18      | No    | No\(^1\)          |
| Pyelonephritis             | 22                 | 2.30 | 2.30 | 0.21      | No    | No\(^1\)          |
| Increased stool frequency  | 15                 | 2.23 | 2.23 | 0.01      | Yes   | -                 |
| Fluid overload             | 10                 | 7.44 | 7.44 | 0.69      | No    | Yes               |
| Aortic stenosis            | 10                 | 3.30 | 3.31 | 0.10      | No    | No\(^1\)          |
| Vaginal pain               | 8                  | 15.86| 15.87| 0.82      | No    | No\(^1\)          |
| Rheumatoid arthritis       | 6                  | 5.10 | 5.10 | 0.04      | No    | No\(^1\)          |

\(^1\) Labels including edema.

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\(^1\)43 out of 56 (76.8%) used drugs that act on the central nervous system. \(^2\)35 out of 43 (81.4%) used anticholinergic agents. \(^3\)14 out of 25 (56.0%) used anticancer agents, and 6 out of 25 (24.0%) used immunosuppressive agents. \(^4\)22 out of 22 (18.2%) used antibiotics, 6 out of 22 (27.3%) used non-steroidal anti-inflammatory drugs (NSAIDs), 4 out of 22 (18.2%) used both antibiotics and NSAIDs, and 3 out of 22 (13.6%) used diuretics. \(^5\)All had more than two risk factors for aortic stenosis (diabetes, dyslipidemia, and hypertension). \(^6\)5 out of 8 (62.5%) used topical povidone-iodine, 1 out of 8 (12.5%) used estriol, and 1 out of 8 (12.5%) used a topical antifungal agent. \(^7\)4 out of 6 (66.7%) had a history of rheumatoid arthritis.
**Methods**

**Data processing.** We obtained spontaneous AE reports including drugs used to treat PUD and GERD between January 2009 and December 2018 from the KIDS. The study protocol was exempted from review by the institutional review board of Ewha Womans University (institutional review board number: ewha-202102-0009-0).
The data consisted of ASCII format tables: basic information (ADR_REPORT_BASIC), drug information (DRUG_INFO_ADR), AE information (ADR_INFO_REPORT), seriousness of AEs (SERIOUS_ADR), reporter information (REPORTOR_ADR), primary causality assessment (ASSESSMENT_ADR), and medical history (HIST_ADR).

AEs were coded according to the World Health Organization Adverse Reaction Terminology (WHO-ART) version 092, and drugs were coded using the Anatomical Therapeutic Chemical (ATC) classification system. Causality in the 'ASSESSMENT_ADR' table was judged by the reporter as 'certain', 'probable', 'possible', 'unlikely', 'unclassified', 'unassessable', or 'not applicable'. As the reporter can report AEs without causality assessment, causality information can be omitted. Except for follow-up reports, only first reports were extracted, and drugs and AEs were paired.

Disproportionality analysis. Disproportionality analysis is a method of detecting the AE signals of a specific drug. We constructed a 2 × 2 table, which has rows with rebamipide and all other drugs and columns with specific AEs and all other AEs (Table 4).

From the table, we calculated three signal indicators, i.e., proportional reporting ratio (PRR), reporting odds ratio (ROR), and information component (IC) (Table 5). All three indicators must meet the criteria to be signals of rebamipide. The lower limit of the 95% confidence interval (CI) of the IC was calculated according to a previous study.

Signal detection. Signal detection proceeded in two different ways (Fig. 2). In the first method, only drug–AE pairs classified as certain, probable, or possible from causality assessment were used in disproportionality analysis. In this case, AEs that met the criteria of the PRR, ROR, and IC were classified as signals.

However, rebamipide may not be considered as a causative drug because it is generally known as a relatively safe drug. In addition, causality assessment is not necessary for the KAERS, and some AE reports had no causality assessment. Therefore, in the second method, disproportionality analysis was performed regardless of causality information. AEs not included on the drug label were reviewed with co-administered drugs in the 'DRUG_INFO_ADR' table and medical history in the 'HIST_ADR' table to determine whether they were adverse effects caused by rebamipide. This process was conducted by an expert with more than 10 years of experience in clinical pharmacy.

AEs not caused by the action of drugs, such as prescription errors, were excluded. Drug labels in Korea, Japan, and China were reviewed to identify signals that have been listed. The ADR information listed on the labels of the three countries was the same.

### Table 4. 2 × 2 table for disproportionality analysis of rebamipide. (A) Rebamipide–specific AE pairs. (B) Rebamipide–all other AE pairs. (C) Other drug–specific AE pairs. (D) Other drug–all other AE pairs.

|          | Specific AEs | All other AEs |
|----------|-------------|--------------|
| Rebamipide | A           | B            |
| All other drugs | C          | D            |

### Table 5. Definition and criteria of signal detection for each indicator. IC information component, PRR proportional reporting ratio, ROR reporting odds ratio.

| Indicator | Definition | Criteria of signal detection |
|-----------|------------|-----------------------------|
| PRR       | (A/(A + B))/(C/(C + D)) | PRR ≥ 2, chi-squared ≥ 4, and A ≥ 3 |
| ROR       | (A/B)/(C/D) | ROR ≥ 2, chi-squared ≥ 4, and A ≥ 3 |
| IC        | Log2(P(AE, drug)/P(AE) × P(drug)) | Under limit of 95% confidence interval ≥ 0 |

### Figure 2. Process of signal detection.
AEs frequently reported in the elderly population. Information on the patient’s age was recorded in ‘ADR_REPORT_BASIC’ table. Elderly population was defined as over 65 years of age at the time of AEs. Drug-AE pairs classified as certain, probable, or possible from causality assessment and reported among patients over 65 years of age were analyzed in this subgroup analysis. After constructing a 2 × 2 table, the ROR and 95% CI were calculated. We searched for AEs for which the lower limit of the 95% CI was greater than 1. The seriousness of frequent AEs was reviewed. Subgroup analysis was performed by limiting the control drugs to H2RAs (ATC A02BA) and PPIs (ATC A02BC). Additionally, we calculated the median time to AEs by subtracting the date of occurrence of AEs and the start date of taking rebamipide.

Data availability
KIDS-KD is available at the Korea Institute of Drug Safety & Risk Management (Ministry of Food and Drug Safety) website (https://open.drugsafe.or.kr/original/invitation.jsp). The authors do not have any special access privileges to this data. The datasets used and/or analyzed during this study are not publicly available due to privacy or ethical restrictions.

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References
1. Ivashkin, V. T., Trukhmanov, A. S. & Gonik, M. I. Rebamipide using in gastrooesophageal reflux disease treatment. Ter. Arkh. 92(4), 98–104 (2020).
2. Arakawa, T. et al. 15th anniversary of rebamipide: Looking ahead to the new mechanisms and new applications. Dig. Dis. Sci. 50(Suppl 1), S3–S11 (2005).
3. Dailypharm. [2020 Prescription Drug Market] Rebamipide. http://www.dailypharm.com/Users/News/NewsView.html?DAILY PHARM_MOBILE=ok&ID=273120?DAILYPHARM_MOBILE=ok. Accessed 12 April 2022 (2021).
4. Yang, B. R., Lee, J. Y. & Kim, M. G. The risk of pulmonary adverse drug reactions of rebamipide and other drugs for acid-related diseases: An analysis of national pharmacovigilance database in South Korea. J. Dig. Dis. 23, 118–123 (2021).
5. Korean Statistical Information Service. Status of Health Insurance Benefits of Frequent Diseases by Disease Subclassification of the Elderly Aged 65 or Older. https://kosis.kr/statHtml/statHtml.do?orgId=350&tblId=DT_35001_A667412&vw_cd=MT_ZTITLE&list_id=350_35001_66&scrid=SA&seqNo=6&lang_mode=ko&obj_var_id=itmit&itm_id=8&com_path=K1&path=%25EB%25B3%25B4%25EA%25B1%2525. Accessed 1 March 2022 (2021).
6. Mes, M. L., Fixen, D. R. & Linnebur, S. A. Adverse effects of proton-pump inhibitor use in older adults: A review of the evidence. Ther. Adv. Drug Saf. 8(9), 273–297 (2017).
7. By the American Geriatrics Society Beers Criteria Update Expert Panel. American Geriatrics Society 2019 Updated AGS Beers Criteria(R) for potentially inappropriate medication use in older adults. J. Am. Geriatr. Soc. 67(4), 674–694 (2019).
8. Pitts, P. J., Louet, H. L., Moride, Y. & Conti, R. M. 21st century pharmacovigilance: Efforts, roles, and responsibilities. Lancet Oncol. 17(11), e486–e492 (2016).
9. Edwards, I. R. & Aronson, J. K. Adverse drug reactions: Definitions, diagnosis, and management. Lancet 356(9237), 1255–1259 (2000).
10. Shin, H., Cha, J., Lee, Y., Kim, J. Y. & Lee, S. Real-world data-based adverse drug reactions detection from the Korea Adverse Event Reporting System database with electronic health records-based detection algorithm. Health Inform. J. 27(3), 14604582211033014 (2021).
11. Yu, J., Zeng, Y., Dai, Y., Zhou, Z., Yu, X. & Shi, D. Hepatotoxicity-related adverse effects of proton pump inhibitors: A cross-sectional study of signal mining and analysis of the FDA Adverse Event Report System Database. Front. Med. 8, 648164 (2021).
12. Imai, T., Hazama, K., Kosuge, Y., Suzuki, S. & Ootsuka, S. Preventive effect of rebamipide on NSAID-induced lower gastrointestinal tract injury using FAERS and JADER. Sci. Rep. 12(1), 2631 (2022).
13. Garcia, A. K. A., Fonseca, L. F., Furuya, R. K., Rabelo, P. D. & Rossetto, E. G. Effect of chewing gum on thirst: An integrative review. Rev. Bras. Enferm. 72(2), 484–493 (2019).
14. Yacoub, M. R. et al. Drug induced exfoliative dermatitis: State of the art. Clin. Mol. Allergy 14, 9 (2016).
15. de Groot, A. C. Systemic allergic dermatitis (systemic contact dermatitis) from pharmaceutical drugs: A review. Contact Dermatitis 86, 145–164 (2021).
16. Vinholi, P. J. The role of platelets in bleeding in patients with thrombocytopenia and hematological disease. Clin. Chem. Lab. Med. 57(12), 1808–1817 (2019).
17. Padala, P. R. et al. Selective serotonin reuptake inhibitors-associated apathy syndrome: A cross sectional study. Medicine 99(33), e21497 (2020).
18. Panicker, J. N., Fowler, C. J. & Kessler, T. M. Lower urinary tract dysfunction in the neurological patient: Clinical assessment and management. Lancet Neurol. 14(7), 720–732 (2015).
19. Tang, I. T., Shepp, D. H. Herpes simplex virus infection in cancer patients: prevention and treatment. Oncology (Williston Park) 6(7), 101–106, 109; discussion 9–10 (1992).
20. Ljungman, P. Herpes infections in immunocompromised patients: Problems and therapeutic interventions. Ann. Med. 25(4), 329–333 (1993).
21. Nasit, C. C. Medication-induced infantile nephritis in the 21st century. Adv. Chronic Kidney Dis. 24(2), 72–79 (2017).
22. Kanwar, A., Thaden, J. J. & Nkomo, V. T. Management of patients with aortic valve stenosis. Mayo Clin. Proc. 93(4), 488–508 (2018).
23. Qian, Y. et al. Observational and genetic evidence highlight the association of human sleep behaviors with the incidence of fracture. Ann. Med. 53(4), 1339 (2021).
24. Oades, R. D. et al. Assessment of the effects of antihistamine drugs on mood, sleep quality, sleepiness, and dream anxiety. Int. J. Psychiatry Clin. Pract. 18(3), 161–168 (2014).
25. Biedry, L., Lien, K. W. & Biggelsen, S. Efficacy and tolerability of newer antihistamines in the treatment of allergic conjunctivitis. Drugs 65(5), 215–228 (2005).
26. Park, K. et al. Signal detection of imipenem compared to other drugs from Korea Adverse Event Reporting System Database. Yonsei Med. J. 58(3), 564–569 (2017).
30. Wilson, A. M., Thabane, L. & Holbrook, A. Application of data mining techniques in pharmacovigilance. *Br. J. Clin. Pharmacol.* 57(2), 127–134 (2004).
31. Gould, A. L. Practical pharmacovigilance analysis strategies. *Pharmacoepidemiol. Drug Saf.* 12(7), 559–574 (2003).
32. Kim, J., Oh, J. M., Son, S. H. & Shin, W. G. Use of information component (IC) and relative risk (RR) for signal detection of drug interactions of clopidogrel: Data-mining study using Health Insurance Review & Assessment Service (HIRA) claims database. *Korean J. Clin. Pharm.* 21(2), 90–99 (2011).

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Competing interests
The authors declare no competing interests.

Additional information
Correspondence and requests for materials should be addressed to M.G.K.

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