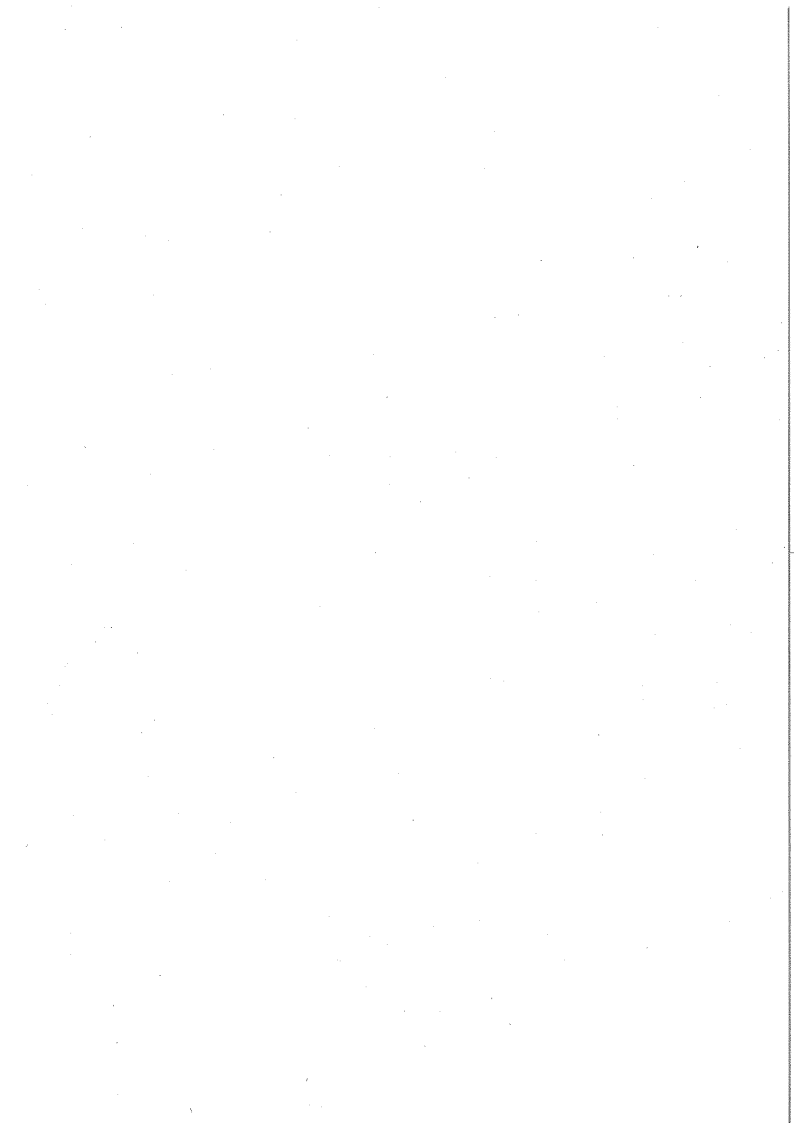
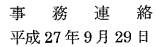
日本一般用医薬品連合会 御中

厚生労働省医薬食品局審査管理課

かぜ薬等の製造販売承認基準の英訳について

標記について、別添写しのとおり、各都道府県衛生主管部(局)薬務主管課宛て通知しましたので、御了知の上、貴会会員に対し周知方御配慮お願いします。







各都道府県衛生主管部(局)薬務主管課 御中

厚生労働省医薬食品局審査管理課

かぜ薬等の製造販売承認基準の英訳について

一般用医薬品のうち、下記のかぜ薬等の製造販売の承認基準(通知)については、別添のとおり、当該基準の英訳を作成したのでお知らせいたします。

記

別添	通知名	発出年月日等
1 .	かぜ薬の製造販売承認基準について	平成 27 年 3 月 25 日付け薬食発 0325 第 28 号
2	解熱鎮痛薬の製造販売承認基準について	平成 27 年 3 月 25 日付け薬食発 0325 第 30 号
3	鎮咳去痰薬の製造販売承認基準について	平成 27 年 3 月 25 日付け薬食発 0325 第 26 号
4	鼻炎用内服薬の製造販売承認基準について	平成 27 年 3 月 25 日付け薬食発 0325 第 23 号
5	胃腸薬製造(輸入)承認基準について	昭和 55 年 4 月 22 日付け薬発第 520 号
6	瀉下薬製造(輸入)承認基準について	昭和 57 年 5 月 17 日付け薬発第 463 号
7	鎮暈薬製造(輸入)承認基準について	昭和 59 年 6 月 1 日付け薬発第 381 号
8	眼科用薬製造(輸入)承認基準について	昭和61年7月29日付け薬発第623号
9	ビタミン主薬製剤製造(輸入) 承認基準につ	昭和63年2月1日付け薬発第90号
	いて	
1 0	浣腸薬製造(輸入)承認基準について	昭和63年2月1日付け薬発第94号
1 1	駆虫薬製造 (輸入) 承認基準について	平成元年3月28日付け薬発第300号
12	鼻炎用点鼻薬製造(輸入)承認基準について	平成3年2月1日付け薬発第109号
1 3	外用痔疾用薬製造(輸入)承認基準等につい	平成7年3月22日付け薬発第277号
	7	
1 4	みずむし・たむし用薬製造(輸入)承認基準	平成 10 年 5 月 15 日付け薬発第 447 号
	等について	
1 5	鎮痒消炎薬の製造販売承認基準について	平成23年11月1日付け薬発第1号
L	L	



Provisional Translation from Japanese Original

Mar 25, 2015 Notification PB No.28

The Standards for Marketing Approval of Cold Remedies

1. Scope of Cold Remedies

The scope of either medicines subject to these standards covers all oral medicines intended for use in treating cold symptoms (Kampo medicine* formulas are not covered).

*Kampo medicine is traditional Japanese medicine.

2. Approval Standards

The approval standards for cold remedies are as follows. For either medicines not conforming to these standards, efficacy and safety data and reasons justifying the combination should be submitted; the preparation in question will be reviewed based on these data.

(1) Types of Active Ingredients

a. The types of active ingredients that may be combined are shown in Table 1.

b. At least 1 of the active ingredients from Group 1 or 2 in Column I of Table 1 must be included. However, in the case of formulas consisting of crude drugs only, Earthworm (Lumbricus) from Column XVI of Table 1 should be combined instead of them.

2. Active ingredients from different columns of Table 1 may be combined with each

other, unless otherwise stipulated.

d. Active ingredients from Column VIII of Table 1 may be combined only in formulas that contain active ingredients from Column II of the table.

e. Up to 3 of the active ingredients from Group 1 in Column I of Table 1 can be

combined.

f. When the active ingredients from Column II, III, IV, V, VI, VIII, IX, or X or the Kampo medicine formulas from Column XVII of Table 1 are combined, one ingredient can be used from each Column. However, the active ingredients from Groups 2 and 3 in Column VI of Table 1 may be combined at the same time.

g. When the active ingredients from Group 2 in Column I of Table 1 are combined, they should not be combined simultaneously with the active ingredients from

Group 1 or 3 in the same column.

h. When the active ingredients from Group 2 from Column I of Table 1 are combined, they should not be combined simultaneously with the active ingredients from Group 2 in Column III, Group 3 in Column VI, from Column VII, Column XIII or Column XIV, Earthworm from Column XVII or the Kampo medicine formulas from Column XVII.

i. When the active ingredients from Group 3 in Column I of Table 1 are combined, they should be combined simultaneously with acetaminophen from Group 1 in the same column, and should not be combined simultaneously with other active

ingredients from the same column.

j. When the active ingredients from Group 3 in Column I of Table 1 are combined,

they should not be combined simultaneously with the active ingredients from Group 3 in Column II, Group 2 in Column III, from Column VI, Column XIII or the active ingredients from Column XIV, Earthworm from Column XVI, or the Kampo medicine formulas from Column XVII.

k. When the active ingredients from Group 2 in Column II of Table 1 are combined, they should not be combined simultaneously with the active ingredients from

Column XIV or the Kampo medicine formulas from Column XVII.

 When the active ingredients from Group 3 in Column II of Table 1 are combined, they should not be combined simultaneously with the active ingredients from Group 3 in Column I or from Column XIV or the Kampo medicine formulas from Column XVII.

- m. When the active ingredients from Group 2 in Column III of Table 1 are combined, they should not be combined simultaneously with the active ingredients from Group 2 in Column I, Group 3 in Column I, from Column IV, Column VIII, Column IX, Column XIII, Column XIV or Column XV, or Kakkontokakikyo from Column XVII.
- n. When the active ingredients from Group 2 in Column VI of Table 1 are combined, they should not be combined simultaneously with the active ingredients from Group 3 in Column I, from Column VIII, Column XIII, Column XIV or the Kampo medicine formulas from Column XVII.
- o. When the active ingredients from Group 3 in Column VI of Table 1 are combined, they should not be combined simultaneously with the active ingredients from Group 2 in Column I, Group 3 in Column I, from Column VIII, Column XIII, Column XIV or the Kampo medicine formulas from Column XVII.
- p. When the active ingredients from Column VII of Table 1 are combined, they should not be combined simultaneously with the active ingredients from Group 2 in Column I or from Column VIII or the Kampo medicine formulas from Column XVII.
- q. When the active ingredients from Column VIII of Table 1 are combined, they should not be combined simultaneously with the active ingredients from Group 2 in Column III, Group 2 and Group 3 in Column VI, from Column VII, Column XIII or Column XIV or the Kampo medicine formulas from Column XVII.
- r. When the active ingredients from Column IX of Table 1 are combined, they should not be combined simultaneously with the active ingredients from Group 2 in Column III, from Column XIII or Column XIV or the Kampo medicine formulas from Column XVII.
- s. Combinations of glycyrrhizinic acid and its salts from Column IX of Table 1 and Glycyrrhiza from Column XV are not acceptable.
- t. Combinations of Ephedra herb or Kampo medicine formulas containing Ephedra herb or their extracts and the active ingredients from Group V of Table 1 are not acceptable.
- u. Combinations between the Kampo medicine formulas from Column XVII of Table 1 and the active ingredients from Column XIII, XIV, XV or XVI are not acceptable.
- v. Apart from Kososan formula, Kampo medicine or non-Kampo crude drug medicines must be in the extract form when used in combinations.
- w. The crude drugs used in the Kampo medicine formulas from Column XVII of Table 1 and their combination ratios must be as specified in Table 2.

(2)Quantities of Active Ingredients

a. The maximum daily dose of each of the active ingredients is that specified in Table 1, unless otherwise specified. However, when the active ingredients from Column V or XIII in Table 1 are combined with the ingredients in Column X, the sum of the values obtained by dividing the amounts of each of the active ingredients by their respective maximum daily doses should not exceed 2/3rd.

b. When 2 or more of the active ingredients from Group 1 in Column I of Table 1 are combined or when 2 or more of the active ingredients from Column XIII, XIV, or XV are combined, the sum of the values obtained by dividing the amounts of each of the active ingredients by their respective maximum daily doses should not exceed 1.

c. When the active ingredients from Group 1 in Column I of Table 1 are combined with Earthworm, Kakkonto formula, Maoto formula, or Kakkontokakikyo, the sum of the values obtained by dividing the amounts of the active ingredients or the formulations combined by their respective maximum daily doses should not exceed 1.

d. When used in combinations, the amounts of the Kampo medicine formulas from Column XVII of Table 1 must not be less than 1/5th and not more than half of the maximum daily dose.

e. The lower limit of the amounts of each of the active ingredients should be half of the maximum daily dose, unless otherwise specified.

f. When 2 or more of the active ingredients from Group 1 in Column I of Table 1 are combined, the lower limit of the amounts should be 1/5th of the maximum daily dose for each active ingredient, and the sum of the values obtained by dividing the amounts of each of the active ingredients by their respective maximum daily doses should be not less than half.

g. When used in combinations, the lower limit of the amounts of the active ingredients from Columns X and XII of Table 1 is 1/5th of the maximum daily dose.

h. When used in combinations, the lower limit of the amounts of glycyrrhizinic acid and its salts from columns IX of Table 1 and the active ingredients from Columns XIII, XIV, XV, and XVI is 1/10th of the respective maximum daily doses. However, in the case of combination with Earthworm as described in (1) b, the maximum daily dose from Column XVI should be combined.

i. In cases where indications for treatment of coughing and sputum are based only on the active ingredients from Columns XIII, XIV, or XV of Table 1, when used in combinations, the lower limits of the active ingredients from Columns XIII, XIV, or XV should be half of the respective maximum daily doses. However, in cases where 2 or more of the crude drugs from Column XV are combined, the lower limit should be 1/5th of the respective maximum daily doses, and the sum of the values obtained by dividing the amounts of each of the active ingredients by their respective maximum daily dose should be not less than half.

j. The daily dose of the active ingredients from Group 2 in Column I of Table 1 should be limited to 450 mg.

k. The daily dose of the active ingredients from Group 3 in Column I of Table 1 should be limited to 300 mg, and the amount of acetaminophen from Column 1 in the same column, which is combined simultaneously, should be limited to 450 mg

1. The daily dose of the active ingredients from Group 2 in Column II of Table 1 should be limited to 1 mg as clemastine.

m. The daily dose of the active ingredients from Group 3 in Column II of Table 1 should be limited to 4 mg.

n. The daily dose of the active ingredients from Group 2 in Column III of Table 1 should be limited to 30 mg.

o. The daily dose of the active ingredients from Group 3 in Column VI of Table 1 should be limited to 750 mg.

(3)Dosage Forms

The dosage forms are tablets, capsules, pills, granules, powders, and syrups.

(4)Dosage and Administration

- a. Except for syrups, cold remedies are to be taken by oral administration 3 times a day within 30 minute after a meal. Syrups are to be taken, in principle, after every meal, However, if required, they can also be taken before going to bed. If it is absolutely necessary, they can be taken approximately every 4 hours up to a maximum of 6 times a day.
- b. For hard capsules, soft capsules larger than 6 mm in diameter, pills, and tablets, dosage for children under 5 years of age is not approved. Even for capsules smaller than 6 mm in diameter, dosage for children under 3 years of age is not approved.
- c. For tablets 6 mm in diameter or less, dosage for children under 3 years of age is not approved.
- d. For other dosage forms, dosage for infants under 3 months of age is not approved.
- e. For children under the age of 15 years, the maximum daily doses acceptable are the values obtained by multiplying the amount of the active ingredient given in 2 (2) by the coefficients for each age group in Table 3, unless otherwise specified. The maximum single dose of syrups is calculated by using the range of coefficients, and dissolving or suspending 1/6th of the calculated value in water to make less than 10 mL in each case.
- f. For formulas containing aspirin, aspirin aluminum, and sasapyrine from Group 1 in Column I, the active ingredients from Group 2 in Column I, promethazine methylenedisalicylate from Group 1 in Column II, or the active ingredients from Group 3 in Column II, dosage for children under 15 years of age is not approved.
- g. For formulas containing the active ingredients from Group 3 in Column VI, dosage for children under 8 years of age is not approved.
- h. For formulas containing the active ingredients from Group 3 in Column I or Group 2 in Column II or transxamic acid from Column IX, dosage for children under 5 years of age is not approved.
- i. For formulas containing the active ingredients from Group 2 in Column III, dosage for children under 3 years of age is not approved.
- j. For formulas containing tranexamic acid from Column IX of Table 1 with dosage for children under 15 years of age, the maximum daily dose is 420 mg. The maximum daily dose for children under 15 years of age is the amount obtained by multiplying the maximum daily dose (420 mg) in Table 1 by the coefficient corresponding to the respective age group in Table 3.

(5)Indications

Relief of various symptoms of a common cold: running nose, stuffy nose, sneezing, sore throat, cough, phlegm (sputum), chills (feeling cold due to fever), fever, headache, joint pain, and muscle pain.

However, when any single type of the active ingredients listed in the right column of the following table is not included, the indications in the left column of the table cannot be claimed.

Left column	Right column
Runny nose, stuffy nose, sneezing	Ingredients from Column II of Table 1
Cough	Ingredients from Columns III, IV, V, XIII, or XIV of Table 1
Phlegm (sputum)	Tipepidine citrate or tipepidine hibenzate from Column III of Table 1 or the ingredients from Columns V, VI, VII, XIII, or XV

(6)Packaging Units

For syrups, the maximum volume of the containers is a 2-day supply at the maximum daily dosage for children aged 6 years.

Table 1

Active ingredients and Maximum Daily Doses

ř		ive ingredients and Maximum Daily Doses	
Cate	egory	Name of active ingredient	Maximum daily dose (mg)
Group 1 Column I		Aspirin Aspirin aluminum Acetaminophen Ethenzamide Sasapyrine Salicylamide Lactylphenetidine	1500 2000 900 1500 1500 3000 600
	Group 2	Ibuprofen	450
:	Group 3	Isopropylantipyrine	300
Column II	Group 1	Isothipendyl hydrochloride Difeterol hydrochloride Tripelenamine hydrochloride Thonzylamine hydrochloride Fenethazine hydrochloride Methodilazine hydrochloride Chlorpheniramine maleate d-Chlorpheniramine maleate Carbinoxamine diphenyldisulfonate Diphenylpyraline hydrochloride Diphenylpyraline teoclate Diphenhydramine salicylate Alimemazine tartrate Diphenhydramine tannate Triprolidine hydrochloride Mebhydrolin napadisilate Promethazine methylenedisalicylate Carbinoxamine maleate Difeterol phosphate	7 90 100 50 50 8 7.5 3.5 7.5 4 4.5 75 75 5 75 4 150 40 7.5
	Group 2	Clemastine fumarate	1 [as clemastine]
	Group 3	Mequitazine	4
Column III	Group 1	Alloclamide hydrochloride Tipepidine citrate Cloperastine hydrochloride Chloperastine phendizoate Codeine phosphate Dihydrocodeine phosphate Dibunate sodium Tipepidine hibenzate Dextromethorphan hydrobromide Dextromethorphan phenolphthalinate Carbetapentane citrate	75 60 48 84 48 24 90 75 48 72 48
·.	Group 2	Dimemorfan phosphate	30
Colun	nn IV	Noscapine Noscapine hydrochloride	48 48

Colu	mn V	dl-Methylephedrine hydrochloride dl-Methylephedrine saccharinate	60	
Column	Group 1	Guaifenesin Potassium guaiacolsulfonate Potassium cresolsulphonate	250 250 250 (135)	
VI	Group 2	Bromhexine hydrochloride	12 (8)	
	Group 3	L-carbocysteine	750	
Colum	n VII	Ethyl L-cysteine hydrochloride	300	
Colum	nn VIII	Belladonna total alkaloid Isopropamide iodide extract	0.3 (0.12) 6 (1.5)	
Colu	Glycyrrhizinic acid and its salts mn IX Tranexamic acid		39 [as glycyrrhizinic acid] 750	
Colu	mn X	Caffeine and sodium benzoate Caffeine hydrate Anhydrous caffeine	(280) 300 150 150	
		Vitamin B ₁ , its derivatives, and their salts Vitamin B ₂ , its derivatives, and their		
Colu	mn XI	salts Vitamin C, its derivatives, and their	(2) 500	
	salts Hesperidin, its derivatives, and their salts		(50) 90 (18)	

; -

	Glycine	900
	Magnesium silicate	3000
	Synthetic aluminum silicate	3000
	Synthetic hydrotalcite	4000
	Magnesium oxide	500
	Dihyrdoxyaluminum and aminoacetate	1500
	(aluminum glycinate)	
]	Aluminum hydroxide gel	1000
· ·	(as dried aluminum hydroxide gel)	
	Dried aluminum hydroxide gel	1000
	Aluminum hydroxide Sodium hydrogen	900
Column XII	carbonate	
Column XII	coprecipitate	
1 . /	Aluminum hydroxide Magnesium	3000
	carbonate	
	mixed dried gel	
	Aluminum hydroxide-Magnesium	1500
_	carbonate-	
	Calcium carbonate coprecipitate	
	Magnesium hydroxide-Aluminum	1800
, i	potassium sulfate	
	coprecipitation product	
	Magnesium carbonate	2000
	Magnesium aluminometasilicate	1500

(Note) A numerical value within parentheses is the lower limit of amounts for combination.

Crude drugs and Kampo medicine formulas

Oraco arago a	da Kampo medicine formulas		••
		Maximum (g	
Classification	Name of crude drug or Kampo medicine formula	Extract (converted to the amount of	Powder
		crude drug or preparation)	·
Column XIII	Ephedra Herb	4	
Column XIV	Nandina Fruit	10	_
-	Cherry Bark	4	—
	Polygala Root	5	
	Glycyrrhiza	5	1.5
	Platycodon Root	4	2
Column XV	Plantago Seed	5	
	Plantago Herb	10	_
	Lycoris Radiata Bulb	0.8	-
	Senega	4	1.5
	Fritillaria Bulb	2.5	1.5

		Maximum (g)	
Classification	Name of crude drug or Kampo medicine formula	Extract (converted to the amount of crude drug or preparation)	Powder
Column XVI	Fennel Phellodendron Bark Coptis Rhizome Zedoary German Chamomile Flower Cinnamon Bark Gentian Oriental Bezoar Animal gall (including Bear Bile) Adenophora Root Ginger Atractylodes Lancea Rhizome Clove Citrus Unshiu Peel Atractylodes Rhizome Earthworm (Lumbricus) Panax Japonicus Rhizome Ginseng	3 3 3 10 5 0.5 - 0.5 5 3 5 2 5 5 3 6 6	- 3 1.5 3 - 1 0.5 0.02 0.5 2.5 1 2 0.5 3 2 2 3
	Kakkonto Kakkontokakikyo	25 29	- -
	Keishito Kososan	15 11	- 6
Column XVII	Saikokeishito Shosaikoto Shoseiryuto	24 24 24	_ _ _
	Bakumondoto Hangekovokuto	30 16	
	Maoto	13	· –

(Note) Powder combinations will not be accepted where no maximum daily dose is given in the powder column.

Table 2

Table 2											
Name of	Kampo medicine formula	Kakkonto	Kakkontokakikyo	Keishito	Kososan	Saikokeishito	Shosaikoto	Shoseiryuto	Bakumondoto	Hangekovokuto	Maoto
	Scutellaria Root		;			2	3				
	Pueraria Root	8	. 8								
70	Glycyrrhiza	· 2	2	2	1	2	2	2	2		2
tios	Platycodon root		4			·					
ra	Apricot Kernel										4
loi	Cinnamon Bark	-3	3	4		3		3			3
nat	Cyperus Rhizome				. 4						
abi:	Brown Rice						- `		10	•	
9	Magnolia Bark									3	
pu	Schisandra Fruit							3			
් ශ ශ	Bupleurum Root		·			5	7				
rug	Asiasarum Root							3	1.		
G. G	Peony Root	3	3	4		- 3		3			\exists
l dd	Ginger	_1	1	1	1	1	1	2	-	1	
t 2	Perilla Herb				2					2	
nen	Jujube	4	4	4.		2	3		3		
Component crude drugs and combination ratios	Citrus Unshiu Peel]			3						
l m	Ginseng			•		2	3		2	-	
Ο.	Ophiopogon Tuber								8		.
•	Pinellia Tuber					. 4	5	5	5	5	
	Poria Sclerotium									5	
<u></u>	Ephedra Herb	4	4	_ [3	. I		4

Table 3

Age coefficients

Age group	Coefficient	
15 years of age and over	1	
11 to under 15 years of age	2/3	
7 to under 11 years of age	1/2	
3 to under 7 years of age	1/3	
1 to under 3 years of age	1/4	
6 months to under 1 year of age	1/5	
3 months to under 6 months of age	1/6	

Provisional Translation from Japanese Original

Mar 25, 2015 Notification PB No.30

The Standards for Marketing Approval of Antipyretic Analgesics

1. Scope of Antipyretic Analgesics

The scope of formulas subject to these standards covers oral medicines intended for the relief of pain or fever (cold remedies, formulations based on Kampo medicine* formulas and those consisting of crude drugs only are not covered).

*Kampo medicine is traditional Japanese medicine.

2. Approval Standards

The approval standards for antipyretic analgesics are as follows. For remedies deviating from these standards, efficacy and safety data and reasons justifying the combination should be submitted; the preparation in question will be reviewed based on these data.

(1) Types of Active Ingredients

a. The types of active ingredients that may be combined are shown in Table 1.

b. Either one of the active ingredients from Group 1, Group2, and Group3 in Column I of Table 1 must be included.

c. Active ingredients from different columns of Table 1 may be combined with each other, unless otherwise stipulated.

d. Up to 3 of the active ingredients from Group 1 or 2 in Column I of Table 1 can be combined.

e. When the active ingredients from Group 3 in Column I of Table 1 are combined, they should not be combined simultaneously with the active ingredients from the same column. However, this rule does not apply when they are combined simultaneously with either one of acetaminophen from Group 1 of the same column, ethenzamide in Group 2, and the active ingredients from Group 4.

f. When the active ingredients from Group 3 in Column 1 of Table 1 are combined or when they are combined simultaneously with either one of acetaminophen in Group 1 and ethenzamide in Group 2 in the same column, the active ingredients from Columns II, III, IV, V, VI, VIII, and IX can be combined. However, when the active ingredients from Group 3 in Column I of Table 1 are combined at the maximum single dose, none of the other ingredients should be combined.

g. When the active ingredients from Group 4 in Column I of Table 1 are combined, they should be combined simultaneously with either one of acetaminophen from Group 1, ethenzamide from Group 2 and the active ingredients from Group 3 in the same column, and should not be combined simultaneously with other active ingredients from Groups 1 and 2 in the same column.

h. When the active ingredients from Group 4 in Column I of the Table 1 are combined simultaneously with acetaminophen from Group 1, ethenzamide from Group 2 and the active ingredients from Group 3 in the same column, the active ingredients from Columns II, IV, V, VI, VIII, and IX can be combined.

i. When the active ingredients from Column II or IV of Table 1 are combined, only one ingredient can be used from the same column.

(2) Quantities of Active Ingredients

- a. The maximum daily dose of each active ingredient should be the dose specified in Table 1, unless otherwise specified.
- b. The lower limit of the single dose for the individual active ingredients in Groups 1 or 2 in Column 1 of Table 1 is half of the maximum single dose. When 2 or more of the active ingredients from Groups 1 and 2 in Column 1 are combined, the lower limit of the daily dose should be 1/5th of the maximum daily dose or half of the maximum single dose, whichever is lower.
- c. The lower limit of the daily dose for the active ingredients from Column II or IV of Table 1 is 1/5th of the maximum daily dose or half of the maximum single dose, whichever is lower.
- d. When used in combinations, the lower limit of the daily amounts of the active ingredients from Column VI of Table 1 is 1/5 of the maximum daily dose. However, if the medicine is taken up to twice a day, the lower limit for the single dose is 1/15th of the maximum daily dose.
- e. When 2 or more of the active ingredients from Groups 1 and 2 in Column I of Table 1 are combined, the sum of the values obtained by dividing the combined amounts of each of the active ingredients by their respective maximum daily doses (the dose within parenthesis for acetaminophen) should not exceed the combination coefficients shown in Table 2, and it must be more than half of the respective coefficient.
- f. In the case where 2 or more active ingredients from Group 1 or 2 in Column I of Table 1 are combined, the sum of the values obtained by dividing the amounts of each of the active ingredients in the combination by their respective maximum daily doses should not exceed 1.
- g. When the active ingredients from Group 1 or 2 in Column I of Table 1 are combined with the active ingredients from column VII, the stipulation in 2 (2) e will apply.
- h. The lower limit of the daily dose for the active ingredients from Columns VII, VIII, or IX of Table 1 should be 1/10th of the maximum daily dose.
- i. When only the active ingredients from Group 3 among the active ingredients from Column I of Table 1 are combined, the maximum single dose is either 200 mg or 150 mg. In the case where a single dose of 200 mg is combined, the maximum daily dose is 400 mg.
- j. When the active ingredients from Group 3 in Column I of Table 1 are combined simultaneously with acetaminophen from Group 1 in the same column or ethenzamide from Group 2 in the same column, combinations of doses should be limited to those shown in Table 3.
- k. When the active ingredients from Group 4 in Column I of Table 1 are combined simultaneously with acetaminophen from Group 1 in the same column, ethenzamide from Group 2 in the same column, or the active ingredients from Group 3 in the same column, combinations of doses should be limited to those shown in Table 4.

(3)Dosage Forms

The dosage forms should be tablets, capsules, pills, granules, and powders.

(4)Dosage and Administration

- A. The following stipulations have been made.
 - a. Once a day administration

Take the medicine not more than once a day. If possible, avoid taking the medicine on an empty stomach.

b. Twice a day administration

Take the medicine not more than twice a day with an interval of at least 6 hours between doses. If possible, avoid taking the medicine on an empty stomach.

c. Three times a day administration

Take the medicine not more than 3 times a day with an interval of at least 4 hours between doses. If possible, avoid taking the medicine on an empty stomach

- B. Dosages for infants under 3 months of age are not approved.
- C. For formulas containing aspirin, aspirin aluminum, sasapyrine, and sodium salicylate from Group 2 in Column I of the Table 1, the active ingredients from Group 3 in Column 1, or the active ingredients from Group 4 in Column I, dosage for children under 15 years of age is not approved.
- D. For formulas containing the active ingredients from Column III of Table 1, dosage for children under 5 years of age is not approved.
- E. For hard capsules, soft capsules larger than 6 mm in diameter, pills, and tablets, dosage for children under 5 years of age is not approved.
- F. For soft capsules smaller than 6 mm in diameter, pills, and tablets, dosage for children under 3 years of age is not approved.
- G. For children under the age of 15 years, the maximum daily doses acceptable are the values obtained by multiplying the amount of the active ingredient given in 2 (2) by the coefficients for each age group in Table 5.
- H. For formulas containing the active ingredients from Column III of Table 1 with dosage for children under 15 years of age, the maximum single dose is 140 mg and the maximum daily dose is 420 mg. The maximum daily dose for children under 15 years of age is the amount obtained by multiplying the maximum daily dose (420 mg) in Table 1 by the coefficient corresponding to the respective age group in Table 5.

(5) Indications

The indications should be within the following scope.

- 1) Relief of headache, toothache, pain after tooth extraction, sore throat (throat pain), earache, joint pain, neuralgia, lumbago, muscular pain, pain due to stiff shoulders, contusion pain, bone fracture pain, pain associated with sprain (sprain pain), painful menses (menstrual pain), and traumatic pain
- 2) Relief of fever at the time of chills (feeling cold due to fever) and fever

Table 1

Active Ingredients and Maximum Single and Daily Doses

<u> </u>		ingredients and maximum Sing	Ţ- <u> </u>	
	. *		Maximum	Maximum
Cate	gory	Active ingredient	single dose	daily dose
	T		(mg)	(mg)
		Acetaminophen	300	900
	Group 1			(1500)*
		Lactylphenetidine	200	600
		Aspirin	750	1500
		Aspirin aluminum	1000	2000
	Group 2	Ethenzamide	500	1500
		Sasapyrine	500	1500
		Salicylamide	1000	3000
Column I	, ,	Sodium salicylate	1000	3000
Column			-	
	C 9	173		•
	Group 3	Ibuprofen	200	450
				,
-				
	Group 4	Isopropylantipyrine	150	450
		25 op 1 op 3 tall talp 3 tall tall	100	490
	· ·		60	180
		Allylisopropylacetylurea	200	600
Colur	nn II	Bromvalerylurea	·	
		Diomitaiciyittea		
Colum	ın III	Tranexamic acid	250	750
Oolun	*** ***	Tranexamic acid	(93.4)**	(280)**
				,
		O-55.	150	300
•		Caffeine and sodium	120	250
Colum	in IV	benzoate	120	250
		Caffeine hydrate	120	200
	<u> </u>	Anhydrous caffeine		
		Vitamin B ₁ , its derivatives,		25
	,	and their salts		(1)**
		Vitamin B ₂ , its derivatives,		12
Colun	ın V	and their salts		(2)**
	,	Vitamin C, its derivatives,	İ	500
		and their salts		(50)**
-		Hesperidin, its derivatives,		90
		and their salts		(18)**

	Glycine		900
	Magnesium silicate	,	3000
	Synthetic aluminum silicate	,	3000
	Synthetic hydrotalcite		4000
	Magnesium oxide		500
	Dihyrdoxyaluminum and		1500
	aminoacetate		
	Aluminum hydroxide gel (as	,	1000
	dried aluminum hydroxide		
	gel)	·	
	Dried aluminum hydroxide		1000
	gel		-
	Aluminum		900
	hydroxide-Sodium hydrogen		
*Column VI	carbonate coprecipitate	•	
	Aluminum		3000
	hydroxide-Magnesium		
	carbonate mixed dried gel		,
	Aluminum		1500
est"	hydroxide-Magnesium		
	carbonate-Calcium carbonate		
	coprecipitate		
	Magnesium		1800
4	hydroxide-Aluminum		·
	potassium sulfate		
	coprecipitation product		
	Magnesium carbonate	, .	2000
•	Magnesium	* · · · · · · ·	1500
	aluminometasilicate		(

^{*} The figure in parentheses is used when the maximum daily dose of each active ingredient is calculated as specified in 2 (2) e.

** The figures in parentheses are the lower limits of the amounts in a combination.

(Crude drugs)

		Maximum daily dose (g)		
Category	Active ingredient	Extract (converted to the crude drug amount)	Powder	
Column VII	Earthworm(Lumbricus)	3	2	
Column VIII	Japanese Valerian Glycyrrhiza Cinnamon Bark Peony Root Mountan Bark	6 5 5 5 6	2 1.5 1 2 2	
Column IX	Japanese Zanthoxylum Peel Ginger Citrus Unshiu Peel	3 5	1 1 3	

Table 2

Combination Coefficient for Combining 2 or More of Active Ingredients from

Group 1 or 2 in Column I

Administration Number of active ingredients combined	Three times daily	Twice daily	Once daily
Two active ingredients	34/30	32/30	18/30
Three active ingredients	38/30	36/30	19/30

Table 3

Combination Patterns for Combining Active Ingredients from Group 3 in Column I and Active Ingredients from Group 1 or 2 in Column I

(daily dose, -: combination not acceptable)

Gr	oup 3 in Column I	450mg	432mg	390mg
Group 1 in Column I	Acetaminophen	195mg	_	390mg
Group 2 in Column I	Ethenzamide	•	252mg	•

Table 4

Combination Patterns for Combining Active Ingredients from Group 4 in Column I and Active Ingredients from Group 1, 2 or 3 in Column I

(daily dose, : combination not acceptable)

Gr	oup 4 in Column I	450mg	450mg	300mg
Group 1 in Column I	Acetaminophen	750mg	-	-
Group 2 in Column I	Ethenzamide	-	750mg	<u>.</u>
Group 3 in Column I	Ibuprofen	-	•	100mg

Table 5

Range of Age Coefficients

Age group	Coefficient
15 years of age and over	1
11 to under 15 years of age	2/3
7 to under 11 years of age	1/2
3 to under 7 years of age	1/3
1 to under 3 years of age	1/4
6 months to under 1 year of age	1/5
3 to under 6 months of age	1/6

Provisional Translation from Japanese Original

Mar 25, 2015 Notification PB No.26

The Standards for Marketing Approval of Antitussives and Expectorants

1. Scope of Antitussives and Expectorants

The scope of remedies subject to these standards covers oral remedies (including troches and drops) intended for use as antitussives and expectorants. However, remedies based on Kampo medicine* formulas and non-Kampo crude drug remedies consisting of crude drug only are not covered.

*Kampo medicine is traditional Japanese medicine.

2. Approval Standards

The approval standards for antitussives and expectorants are as follows. For remedies not conforming to these standards, efficacy and safety data and reasons justifying the combination should be submitted; the preparation in question will be reviewed based on these data.

(1) Types of Active Ingredients

a. Table 1 lists the active ingredients that may be used. The types of active ingredients that may be used in troches and drops are limited to those marked by △ in Table 1. The active ingredients from Column X should only be combined for troches and drops.

b. One ingredient from Columns I, II, III, XII, or XIII of Table 1 must be included. However, cases where only the active ingredients from Groups 2 and 3 in Column VI of the same table are combined simultaneously are excluded.

c. Active ingredients from different columns of Table 1 may be combined with each other, unless otherwise stipulated.

d. Active ingredients from Group IX of Table 1 may be combined only in remedies that contain active ingredients from Column I or VIII in this table.

e. In Columns I to III and Columns V to X of Table 1, only 1 ingredient from each group may be used.

However, cases where only the active ingredients from Groups 2 and 3 in Column VI of the same table are combined simultaneously are excluded.

- f. Active ingredients from Column XII of Table 1 should not be combined simultaneously with the active ingredients from Column II or V of the same table.
- g. Active ingredients from Group 2 in Column I of Table 1 should not be combined simultaneously with the active ingredients from Columns III, IV, V, XII, XIII, or XIV.

h. Active ingredients from Column IV of Table 1 should not be combined simultaneously with the active ingredients from Group 2 in Column I, or from Columns V, XII, or XIII.

i. Active ingredients from Group 2 in Column VI of Table 1 should not be combined simultaneously with the active ingredients from Column V, XII, or XIII of the same table.

j. Active ingredients from Group 3 in Column VI of Table 1 should not be

- combined simultaneously with the active ingredients from Column V, XII, or XIII of the same table.
- k. Active ingredients from Group 2 in Column VIII of Table 1 should not be combined simultaneously with the active ingredients from Column V or XIII of the same table.

(2) Quantities of Active Ingredients

- a. The maximum single dose and maximum daily dose of each active ingredient in Table 1 should be the doses specified in the same table, unless otherwise specified.
- b. When the active ingredients from Column IX are combined with those from Column II, V, or XII of Table 1 are combined, the maximum single and daily doses of the ingredients in Column IX should be half of the amounts specified in Table 1.
- c. When 2 or more of the active ingredients from Columns II and V of Table 1 are combined or when 2 or more of the active ingredients from Column XII, XIII, or XIV are combined, the sum of the values obtained by dividing the amounts of each of the active ingredients by their respective maximum daily doses should not exceed 1.
- d. The lower limit of the combined amounts of each active ingredient in Table 1 should be half of the maximum single or daily dose, unless otherwise specified. However, for the active ingredients from Column IX, the limit should be 1/5th.
- e. When the active ingredients from Group 2, Column VI of Table 1 are combined simultaneously with only the active ingredients from Group 3 in the same column, the single dose should be 4 mg and the daily dose should be limited to 12 mg.
- f. The single dose of the active ingredients from Group 3 in Column VI of Table 1 should be limited to 250 mg and the daily dose should be limited to 750 mg.
- g. The single dose of the active ingredients from Group 2 in Column VIII of Table 1 should be 0.334 mg as clemastine and the daily dose should be limited to 1 mg as clemastine.
- h. In the case of troches and drops containing Group I ingredients from Column X of Table 1 and having a dosage regimen for children, the coefficients given in Table 2 should not be used to calculate the combined amount of the ingredients from Column X.
- In the case of troches and drops to be taken 5 to 6 times per day, the lower limits of the combined amounts of each active ingredient should be half of the maximum daily dose.
- j. When the active ingredients from Column II of Table 1 are combined simultaneously with the active ingredients from Column V, the lower limits of the combined amounts should be as follows.
- When the active ingredients from Column II of Table 1 are indicated for "cough," "cough associated with wheezing (wheezy, whistling)," or "sputum," the lower limit of the amounts of the ingredients in Column V should be 1/5th of the maximum single and daily doses.
- When other ingredients with an indication of "coughing" are combined, the lower limits of the amounts of ingredients from both Column II and V should be 1/5th of the respective maximum single and daily doses. However, in the case of proportional combinations, lower limits should be such that the sum of the values obtained by dividing the amount of each active ingredient by its maximum daily dose equals half.
- When the active ingredients from Column V of Table 1 are indicated for "cough associated with wheezing (wheezy, whistling)" or "sputum," the lower limit of the amounts of the ingredients in Column II should be 1/5th of the maximum single and daily doses.

k. When used in combinations, the lower limit of the daily amounts of the active ingredients from Column XI of Table 1 is 1/5 of the maximum daily dose.

The lower limits of the amounts of crude drugs should be 1/10th of the
maximum daily dose. However, when the indications approved for a particular
crude drug are claimed, the lower limit should be half of the maximum daily
dose.

(3)Dosage Forms

The dosage forms are tablets, capsules, pills, granules, powders, troches, drops, and oral solutions (with the exception of elixirs; hereinafter the same should apply), and syrups.

(4)Dosage and Administration

a. The dosage is "3 to 4 times a day," and the timing of doses or intervals between

o doses must also be indicated.

However, as for troches, drops, and oral solutions, and syrups, the dosage may be up to 6 doses per day. For dosages of 5 to 6 doses a day, troches and drops should be taken at intervals of at least 2 hours and oral solutions and syrups at intervals of about 4 hours, in principle.

o. The dosage for troches and drops should be allowed to dissolve slowly in the

mouth without chewing.

c. For hard capsules, troches, syrups, and soft capsules larger than 6 mm in diameter, pills, and tablets, dosage for children under 5 years of age is not approved. Even for capsules smaller than 6 mm in diameter, dosage for children under 3 years of age is not approved.

d. Dosages for infants under 3 months of age are not approved.

e. For remedies containing promethazine hydrochloride or promethazine methylene disalycilate from Group 1 in Column VIII of Table 1, dosage for children under 15 years of age is not approved.

For remedies containing the active ingredients from Group 3 in Column VI of

Table 1, dosage for children under 8 years of age is not approved.

g. For remedies containing the active ingredients from Column IV of Table 1 or the active ingredients from Group 2 in Column VIII, dosage for children under 5 years of age is not approved.

h. For remedies containing the active ingredients from Group 2 in Column I of

Table 1, dosage for children under 3 years of age is not approved.

i. The maximum daily dose for children under 15 years of age is the amount obtained by multiplying the maximum daily dose in Table 1 by the coefficient corresponding to the respective age group in Table 2, unless otherwise specified.

The maximum single dose of the active ingredients in oral solutions and syrups is 1/6th of the maximum daily dose (for children under 15 years of age, the maximum daily dose according to i. above), and the maximum single dose

is 10 mL, unless otherwise specified.

k. For remedies containing the active ingredients from Group 2, Column I of Table 1 with dosage for children under 15 years of age, the maximum single dose is 10 mg and the maximum daily dose is 30 mg. The maximum daily dose for children under 15 years of age is the amount obtained by multiplying the maximum daily dose (30 mg) by the coefficient corresponding to the respective age group in Table 2.

For remedies containing the active ingredients from Column IV of Table 1
with dosage for children under 15 years of age, the maximum single dose is
140 mg and the maximum daily dose is 420 mg. The maximum daily dose for

children under 15 years of age is the amount obtained by multiplying the maximum daily dose (420 mg) by the coefficient corresponding to the respective age group in Table 2.

(5)Indications

- a. The indications include "cough, cough associated with wheezing (wheezy, whistling), and sputum." However, for indications in the left column of the following table to be claimed, at least 1 of the ingredients from the corresponding right column must be included.
- b. When the active ingredients from Column IV of Table 1 are combined, the indications are "cough or sputum associated with sore throat." However, they should be combined concomitantly with any ingredient with indications of "cough" and "sputum" from the left column of the next table.
- c. When only the active ingredients from Group 2 and Group 3 in Column VI of Table 1 are combined concomitantly, the indications are "sputum and cough with sputum".
- d. For troches and drops, in addition to the above indications, the following may also be given hourse voice due to throat inflammation, rough throat, throat discomfort, sore throat, and swollen throat.

Left column	Right column
Cough	Ingredients from Columns I, II, III, XII, or XIII of Table 1
Cough associated with wheezing (wheezy, whistling)	Ingredients from Column II, V, or XII in Table 1, except for cases in which an ingredient from Column I of Table 1 is also combined.
Phlegm (sputum)	Tipepidine citrate or tipepidine hibenzate from Group 1 in Column I of Table 1 or the ingredients from Columns II, V, VI, VII, XII, or XIV
Cough associated with sore throat and sputum	Ingredients from Column IV of Table 1, only when combined concomitantly with any ingredient with indications of "cough" and "sputum."
Sputum and cough with sputum	Only when combined concomitantly with only the ingredients from Group 2 and Group 3 in Column VI of Table 1.

(6) Packaging Units

The maximum volume of containers for oral solutions and syrups is a 4-day supply at the maximum daily dose for adults (15 years of age and older).

Table 1

Active Ingredients and Maximum Single and Daily Doses

	1100110	Ingioutilib and manifest and	Maximum	Maximum
Categ	rorv	Name of active ingredient	single dose	daily dose
Cauce	CIJ		(mg)	(mg)
		Alloclamide hydrochloride	25	75
		Tipepidine citrate	20	- 60
	•	Cloperastine hydrochloride	20	60
		Chloperastine phendizoate	35	105
		Codeine phosphate	20	60
		Dihydrocodeine phosphate	10	30
	Group1	Dibunate sodium	30	90
Column I	_		25	- 75
Columni		Tipepidine hibenzate	20	60
		Dextromethorphan hydrobromide	30	90
r F	•	ΔDextromethorphan	30	90
÷ E		phenolphthalinate		co
·		Carbetapentane citrate	20	60
	G	Dimemorfan phosphate	15	60
	Group2		(10)	(30)
2' '		Trimethoquinol hydrochloride	2	6
	•	\(\Delta dl\)-Methylephedrine hydrochloride	25	75
Colum	ın II	I-Methylephedrine hydrochloride	25	. 75
		Methoxyphenamine hydrochloride	50	150
			20	60
Colum	ın III	ΔNoscapine	20	60
		Noscapine hydrochloride	250	750
Colum	n ÍV	Tranexamic acid	(70)	(280)
Colum				
191.33	_	Aminophylline	100	300
Colun	37	Diprophylline	100	300
Colun	an v	Theophylline	200	600
		Proxyphylline	70	210
		Foeniculated ammonia spirit	2mL	-
		(as 1 ingredient)		
		Ammonium chloride	300	900
	Group 1	ΔGuaifenesin	100	300 .
Column	•	ΔPotassium guaiacolsulfonate	90	270
VI		ΔPotassium cresolsulphonate	90	270
vr		I-Menthol	`-	90
	a •	Bromhexine hydrochloride	4	12
,	Group 2		(2)	(8)
f	Group 3	L-carbocysteine	250	750
	·	Ethyl L-cysteine hydrochloride	100	300
Colum	n VII	Methyl L-cysteine hydrochloride	100	300
	AL 7 AL	1 ITTOMES IN OUTSING ITS GEOGRAPHIC	1 ~~~	~~~

Alimemazine tartrate		T		<u>.</u>	
Iproheptine hydrochloride			Alimemazine tartrate	2.5	
Difeterol hydrochloride					
Tripelenamine hydrochloride	1				4
Thonzylamine hydrochloride 20 60 Fenethazine hydrochloride 30 90 Chlorpheniramine maleate 4 12 d-Chlorpheniramine maleate 2 6 Carbinoxamine 4 12 d-Chlorpheniydisulfonate Diphenyldisulfonate 30 90 Diphenyldyramine hydrochloride 30 90 Diphenhydramine salicylate 40 120 Diphenhydramine salicylate 45 135 Triprolidine hydrochloride 2 6 Promethazine tannate 45 135 Triprolidine hydrochloride 5 15 Promethazine methylene 6 18 disalycilate Carbinoxamine maleate 4 12 Difeterol phosphate 30 90 90 40 40 40 40 40 4			Difeterol hydrochloride	•	4
Fenethazine hydrochloride	-				li .
Chlorpheniramine maleate	1			1	
Group1					1
Calumn Column Column Column VIII Diphenylpyraline hydrochloride Diphenylpyraline teoclate 3 9 Diphenylpyraline teoclate 3 9 Diphenylpyraline salicylate 40 120 Diphenhydramine salicylate 40 120 Diphenhydramine tannate 50 150 Fenethazine tannate 45 135 Triprolidine hydrochloride 2 6 Promethazine hydrochloride 5 15 Promethazine methylene 6 18 disalycilate Carbinoxamine maleate 4 12 Difeterol phosphate 30 90 O Diphenylpyraline teoclate Carbinoxamine maleate 4 12 Difeterol phosphate 30 90 O O O O O O O O O					
Column VIII	-			1	
Column VIII	,			4	12
Column VIII				_	
VIII	a i	Group1			
Diphenhydramine salicylate	1				ž.
Diphenhydramine tannate	VIII		Diphenhydramine hydrochloride	E .	1
Fenethazine tannate					· ·
Triprolidine hydrochloride 2 6 Promethazine hydrochloride 5 15 15 15 Promethazine methylene 6 18 disalycilate Carbinoxamine maleate 4 12 Difeterol phosphate 30 90 90 90 100 300				1	1
Promethazine hydrochloride 5 15 18 18 18 18 18 18	,			1	
Promethazine methylene disalycilate Carbinoxamine maleate A 12					
disalycilate					
Carbinoxamine maleate 30 90				6	18
Difeterol phosphate 30 90					
Group2 Clemastine fumarate					
Group2 Clemastine fumarate [as clemastine]			Direterol phosphate		
Caffeine and sodium benzoate 100 300		Creare	Clemastine fumarate		-
Column IX		GroupZ	Ciomastine iumatate		
Column IX		l	C-86-1-1-1-1-1		
Anhydrous caffeine	Column	~ TV			
AChlorhexidine hydrochloride	Columi	11 IA		1	
Column X	<u> </u>				300-
ADequalinium chloride	Colum	n X			.
Glycine 900 Magnesium silicate 3000 Synthetic aluminum silicate 4000 Synthetic hydrotalcite 4000 Magnesium oxide 500 Dihyrdoxyaluminum and aminoacetate Aluminum hydroxide gel 1000 (as dried aluminum hydroxide gel) Dried aluminum hydroxide gel 1000 Aluminum hydroxide Sodium 900 hydrogen carbonate coprecipitate Aluminum hydroxide gel Aluminum hydroxide gel Aluminum hydroxide gel 1500 150	Colum	ш Л.			-
Magnesium silicate Synthetic aluminum silicate Synthetic hydrotalcite Magnesium oxide Dihyrdoxyaluminum and aminoacetate Aluminum hydroxide gel (as dried aluminum hydroxide gel) Dried aluminum hydroxide gel Aluminum hydroxide-Sodium hydrogen carbonate coprecipitate Aluminum hydroxide-Magnesium carbonate mixed dried gel Aluminum hydroxide-Magnesium carbonate-Calcium carbonate coprecipitate Magnesium hydroxide-Aluminum potassium sulfate coprecipitation 1800				0.25	
Synthetic aluminum silicate Synthetic hydrotalcite Magnesium oxide Dihyrdoxyaluminum and aminoacetate Aluminum hydroxide gel (as dried aluminum hydroxide gel) Dried aluminum hydroxide gel Aluminum hydroxide-Sodium hydrogen carbonate coprecipitate Aluminum hydroxide-Magnesium carbonate mixed dried gel Aluminum hydroxide-Magnesium carbonate-Calcium carbonate coprecipitate Magnesium hydroxide-Aluminum potassium sulfate coprecipitation					
Synthetic hydrotalcite Magnesium oxide Dihyrdoxyaluminum and aminoacetate Aluminum hydroxide gel (as dried aluminum hydroxide gel) Dried aluminum hydroxide gel Aluminum hydroxide Sodium hydrogen carbonate coprecipitate Aluminum hydroxide-Magnesium carbonate mixed dried gel Aluminum hydroxide-Magnesium carbonate-Calcium carbonate coprecipitate Magnesium hydroxide-Aluminum potassium sulfate coprecipitation		,			
Magnesium oxide Dihyrdoxyaluminum and aminoacetate Aluminum hydroxide gel (as dried aluminum hydroxide gel) Dried aluminum hydroxide gel Aluminum hydroxide Sodium hydrogen carbonate coprecipitate Aluminum hydroxide Magnesium carbonate mixed dried gel Aluminum hydroxide Magnesium carbonate Calcium carbonate coprecipitate Magnesium hydroxide-Aluminum potassium sulfate coprecipitation					i i
Column XI Dihyrdoxyaluminum and aminoacetate Aluminum hydroxide gel (as dried aluminum hydroxide gel) Dried aluminum hydroxide gel (as dried aluminum hydroxide gel) Aluminum hydroxide Sodium (as dried aluminum hydroxide Sodium (as dried gel) Aluminum hydroxide Magnesium (arbonate mixed dried gel) Aluminum hydroxide Magnesium (arbonate coprecipitate) Aluminum hydroxide Magnesium (arbonate coprecipitate) Magnesium hydroxide Aluminum (arbonate) potassium sulfate coprecipitation					
aminoacetate Aluminum hydroxide gel (as dried aluminum hydroxide gel) Dried aluminum hydroxide gel Aluminum hydroxide-Sodium hydrogen carbonate coprecipitate Aluminum hydroxide-Magnesium carbonate mixed dried gel Aluminum hydroxide-Magnesium carbonate-Calcium carbonate coprecipitate Magnesium hydroxide-Aluminum potassium sulfate coprecipitation					
Aluminum hydroxide gel (as dried aluminum hydroxide gel) Dried aluminum hydroxide gel Aluminum hydroxide Sodium hydrogen carbonate coprecipitate Aluminum hydroxide Magnesium carbonate mixed dried gel Aluminum hydroxide Magnesium carbonate Calcium carbonate coprecipitate Magnesium hydroxide-Aluminum potassium sulfate coprecipitation					1900
Column XI (as dried aluminum hydroxide gel) Dried aluminum hydroxide gel Aluminum hydroxide-Sodium hydrogen carbonate coprecipitate Aluminum hydroxide-Magnesium carbonate mixed dried gel Aluminum hydroxide-Magnesium carbonate-Calcium carbonate coprecipitate Magnesium hydroxide-Aluminum potassium sulfate coprecipitation (as dried aluminum hydroxide gel) 1000 900 1500 1500 1500					1000
Column XI Dried aluminum hydroxide gel Aluminum hydroxide-Sodium hydrogen carbonate coprecipitate Aluminum hydroxide-Magnesium carbonate mixed dried gel Aluminum hydroxide-Magnesium carbonate-Calcium carbonate coprecipitate Magnesium hydroxide-Aluminum potassium sulfate coprecipitation 1000 900 3000 1500 1500 1500					1000
Column XI Aluminum hydroxide-Sodium hydrogen carbonate coprecipitate Aluminum hydroxide-Magnesium carbonate mixed dried gel Aluminum hydroxide-Magnesium carbonate-Calcium carbonate coprecipitate Magnesium hydroxide-Aluminum potassium sulfate coprecipitation Aluminum hydroxide-Magnesium 1500 1800					1000
Column XI hydrogen carbonate coprecipitate Aluminum hydroxide-Magnesium carbonate mixed dried gel Aluminum hydroxide-Magnesium carbonate-Calcium carbonate coprecipitate Magnesium hydroxide-Aluminum potassium sulfate coprecipitation Aluminum hydroxide-Aluminum potassium sulfate coprecipitation					
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carbonate mixed dried gel Aluminum hydroxide Magnesium carbonate Calcium carbonate coprecipitate Magnesium hydroxide Aluminum potassium sulfate coprecipitation					2000
Aluminum hydroxide-Magnesium carbonate-Calcium carbonate coprecipitate Magnesium hydroxide-Aluminum potassium sulfate coprecipitation					3000
carbonate-Calcium carbonate coprecipitate Magnesium hydroxide-Aluminum potassium sulfate coprecipitation	,			-	1500
coprecipitate Magnesium hydroxide Aluminum potassium sulfate coprecipitation				1	1900
Magnesium hydroxide-Aluminum 1800 potassium sulfate coprecipitation					4
potassium sulfate coprecipitation	,				1000
				į	1900
product					
	I	i	oroquer :		
				.	9000
Magnesium aluminometasilicate 1500			Magnesium carbonate Magnesium aluminometasilicate		2000

(Crude drugs)

		Maximum d	aily dose (g)
		Extract	
Category	Name of crude drug or Kampo	(converted to	Powder
	medicine formula	the crude drug	
		amount)	
Column XII	Ephedra Herb	4	-
Column XIII	Nandina Fruit	10	
	Cherry Bark	4	-
	Polygala Root	5	• •
	Glycyrrhiza	5	1.5
	Platycodon Root	4	2
	Apricot Kernel	4	. *
Column XIV	Plantago Seed	5 5	-
	Plantago Herb	10	-
• • • • • • • • • • • • • • • • • • •	Lycoris Radiata Bulb	0.8	-
	Senega	4	1.5
	Ipecac	0.05	0.05
	Fritillaria Bulb	2.5	1.5
	Gambir	-	2
	Fennel	3	
	Scutellaria Root	6	.3
	Trichosanthes Seed	2	- .
	Cinnamon Bark	5	1
	Oriental Bezoar	.	0.02
· ·	Schisandra Fruit	5	*,
* * * * * * * * * * * * * * * * * * *	Asiasarum Root	3.	•
	Aster Root	5	-
Column XV	Musk	-	0.01
•	Adenophora Root	5	2.5
	Ginger	3	1
	Mulberry Bark	5	-
	Perilla Herb	2	-
	Panax Japonicus Rhizome	6	3
	Citrus Unshiu Peel	5	3
	Ginseng	6	3
	Ophiopogon Tuber	10	•
* .	Pinellia Tuber	5	-

(Note) A numerical value within parentheses is the lower limit of amounts for combination.

Table 2

Range of Age Coefficients

Age	Coefficient
15 years of age and older	1
11 to under 15 years of age	2/3
8 to under 11 years of age	1/2
5 to under 8 years of age	1/3
3 to under 5 years of age	1/4
1 to under 3 years of age	1/5
3 months to under 1 year of age	1/10

Provisional Translation from Japanese Original

Mar 25, 2015 Notification PB No.23

The Standards for Marketing Approval of Oral Remedies for Rhinitis

1. Scope of Oral Remedies for Rhinitis

The scope of remedies subject to these standards covers oral medicines (with the exception of cold remedies, anti-allergic agents, remedies based on Kampo medicine* formulas) formulated with the intent of relieving symptoms of rhinitis.

*Kampo medicine is traditional Japanese medicine.

2. Approval Standards

The approval standards for oral remedies for rhinitis are as follows. For remedies not conforming to these standards, data concerning the efficacy and safety and reasons justifying the combination should be submitted; the preparation in question will be reviewed based on these data.

(1) Types of Active Ingredients

a. Table 1 shows the types of active ingredients that may be used.

b. The active ingredients that must be used are those listed in Column I of Table 1.

c. Active ingredients from different columns of Table 1 may be combined with each other, unless otherwise stipulated.

d. When active ingredients from Column I, Column III, Column IV, or Column V are to be combined, only 1 ingredient from each column may be used.

e. When active ingredients from Column II of Table 1 are combined, up to 2 active ingredients from Group 1 may be used, but only 1 from Group 2 may be used. However, the combination of dl-methylephedrine hydrochloride and l-methylephedrine hydrochloride or that of pseudoephedrine hydrochloride and pseudoephedrine sulfate is not permitted.

f. When the active ingredients from Group 2 in Column I of Table 1 are combined, only formulas other than oral solutions and syrups can be used. They should not be combined concomitantly with the active ingredients from Column VI.

(2)Quantities of Active Ingredients

a. The maximum daily doses of individual active ingredients should be those given in Table 1, unless otherwise indicated. The maximum single dose is 1/3rd of the maximum daily dose.

However, the maximum single dose of oral solutions and syrups is 1/6th of the maximum daily dose.

b. When active ingredients from Column V of Table 1 are combined with those of Group 1 in Column II, the maximum daily dose of ingredients from Column V should be half of those specified in Table 1.

c. When 2 or more active ingredients from Column II of Table 1 are combined, the sum of the values obtained by dividing the amount of each active ingredient by the respective maximum daily dose should not exceed 2.

d. The lower limit of the daily dose for each active ingredient from Column I of Table 1 is half of its maximum daily dose.

. The lower limit of the daily dose for each active ingredient from Columns II, III,

and V of Table 1 is 1/5th of its maximum daily dose.

f. The lower limit of the daily dose for each active ingredient from Columns IV and VI of Table 1 is 1/10th of its maximum daily dose.

g. The daily dose of the active ingredients from Group 2 in Column I of Table 1 should be limited to 4 mg.

(3)Dosage Forms

The dosage forms are capsules, granules, pills, powders, tablets, oral solutions (with the exception of elixirs; hereinafter the same should apply), and syrups.

(4)Dosage and Administration

a. Dosage and administration are to be 3 times a day, in principle. The times of administration and intervals between them should be clearly indicated, but intervals between doses should be 4 or more hours. For oral solutions and syrups, taking them up to 6 times a day is acceptable, but when dosing is 6 times a day, each dose is to be taken at approximately 4-hour intervals, in principle.

b. Dosage for infants less than 3 months of age is not approved.

c. For formulas containing promethazine hydrochloride or promethazine methylenedisalicylate from Group 1 in Column I of Table 1 and the active ingredients from Group 2 in Column I, dosage for children under 15 years of age is not approved.

d. For formulas containing pseudoephedrine hydrochloride or pseudoephedrine sulfate from Group 1 in Column II of Table 1, dosage for children under 3 years

of age is not approved.

e. For hard capsules, and soft capsules, pills, and tablets larger than 6 mm in diameter, dosage for children under 5 years of age is not approved.

For soft capsules, pills, and tablets of a diameter of 6 mm or less, dosage for

children under 3 years of age is not approved.

g. The maximum daily dose for children under 15 years of age is that obtained by multiplying the maximum daily doses listed in Table 1 by the coefficient for the respective age groups in Table 2.

h. The maximum single dose for oral solutions and syrups is 10 mL.

(5)Indications

The indications are to be within the following scope:

Relief of the following symptoms due to acute rhinitis, allergic rhinitis or sinusitis; sneezing, runny nose (excessive nasal discharge), stuffy nose, watery eyes, sore throat, dull headache (heaviness in the head).

(6) Packaging Units

The maximum volume of containers for oral solutions and syrups is a 4-day supply at the maximum daily dose.

Cate	egory	Active i	ngredient	Maximum daily dose
		Alimemazine tartra	te	5mg
		Isothipendyl hydroc		
·		Iproheptine hydrock	alorido	12mg
		Difeterol hydrochlor		150mg
				90mg
		Tripelenamine hydr		100mg
•	1	Thonzylamine hydro		50mg
		Methodilazine hydro	ochloride	8mg
		Chlorpheniramine n		12mg
		d-Chlorpheniramine	maleate	6mg
	Group1	Carbinoxamine diph	enyldisulfonate	7.5mg
Column I		Diphenylpyraline hy	drochloride	12mg
		Diphenylpyraline te	oclate	4.5mg
		Diphenhydramine h	vdrochloride	75mg
		Diphenhydramine sa		
i i		Diphenhydramine ta		75mg
	,			75mg
		Triprolidine hydroch		6mg
		Promethazine hydro		15mg
		Promethazine methy	zlenedisalicylate	40mg
		Carbinoxamine male	eate	16mg
	Group2	Mequitazine		4mg
		Phenylephrine hydro	ochloride .	30mg
		Pseudoephedrine hy	drochloride	180mg
]	Pseudoephedrine sul	fate	180mg
	Group 1	dl-Methylephedrine		110mg
	1	l-Methylephedrine h		110mg
		Methoxyphenamine		150mg
Column II				
Column				as total
	<i>'</i>	-		alkaloids
		Datura Extract		0.6mg
	Group 2	Belladonna (Total) A	lkaloids	0.6mg
	Group B	Belladonna Extract		60mg
	·	Isopropamide iodide	extract	7.5mg
		Scopolia Extract		60mg
		Bromelain		Jung
Colum	ın III	Lysozyme chloride		120,000 Units
ÖÖTÜ	111 111	Lysozyme chtoride		90 mg (potency)
		Glycyrrhizinic acid ar	nd its salts	as
	Group 1		•	glycyrrhizinic
	Group r		•	acid
			•	200mg
Column IV			Extract	
			(converted to the	n ,
	Group 2	Glycyrrhiza	E .	Powder
			crude drug amount)	
1		0.00	5g	1.5g
		Caffeine and sodium	penzoate	300mg
Colun	nn V	Caffeine and sodium l Caffeine hydrate Anhydrous caffeine	penzoate	300mg 300mg

		Extract (converted to the crude drug amount)	Powder
•	Schizonepeta Spike	3g	-
Colores VI	Asiasarum Root	3g	- ,
Column VI	Ginger	3g	1 g
	Magnolia Flower	3g	-
	Peucedanum Root	3g	-
	Angelica Dahurica	3g	1g
	Root		•

Table 2

Range of ages and coefficients

Age	Coefficient
15 years of age and over	1
11 to under 15 years of age	2/3
7 to under 11 years of age	1/2
3 to under 7 years of age	1/3
1 to under 3 years of age	1/4
6 months to under 1 year of age	1/5
3 months to under 6 months of age	1/6

Provisional Translation from Japanese Original

Apr 22, 1980 Notification PFSB No.520 Final revision Mar 28, 1986

The Standards for Marketing Approval of Gastrointestinal Medicines

1. Scope of Gastrointestinal Medicines

The scope of preparations subject to these standards covers all medicines for oral use formulated with the intent of relieving symptoms of gastrointestinal diseases (evacuants and Kampo medicine* formulas are not covered).

*Kampo medicine is traditional Japanese medicine.

2. Approval Standards

The approval standards for gastrointestinal medicines are as follows. For preparations not conforming to these standards, efficacy and safety data and reasons justifying the combination should be submitted; the preparation in question will be reviewed based on these data.

(1)Types of Active Ingredients

(a) The types of active ingredients that may be used are shown in Table 1.

(b) Preparations mainly containing active ingredients from Column I, II, III, or IV can be mutually combined with other active ingredients from Columns I, II, III, and IV as well as the active ingredients from Columns V (limited to those with a "Δ" mark in Groups 3, 4, and 5), VII, and VIII. However, notwithstanding the above rules, preparations having their main active ingredients only from Column I cannot include the following active ingredients: those in Group 2 of Column IV or those with a "Δ" mark in Group 5 of Column V. Preparations mainly containing active ingredients only from

Column IV cannot include the active ingredients from Column VII.

Preparations mainly containing active ingredients from Column V of Table 1 can include the active ingredients from Column I, II, III, IV, or VI (limited to

Scopolia Extract in Group 1 and ingredients in Group 4).

(d) Preparations mainly containing active ingredients from Column VI of Table 1 can include the active ingredients from Column I (except Group 3), II, III, or V (limited to Groups 3 and 4).

However, preparations mainly containing active ingredients from Group 1 of Column VI cannot include the active ingredients from Column II (limited to Nux Vomica Extract in Group 1 or ingredients in Group 3). When the active ingredients from Column VI (except for Group 4) are used in combination, they should be limited to 1 type from each group.

(e) When the active ingredients from Column VII (except for Group 9) of Table 1 are used in combination, they should be limited to 1 type from each group.

(f) The active ingredients from Column I (excluding Group 3) and Group 2 of Column II cannot be combined in the same preparation.

(g) When the same active ingredient appears in at least 2 columns of Table 1, it

should not be duplicated in the formula.

(h) Berberine chloride and berberine tannate in Group 1 of Column V must not be combined with Coptis Rhizome or Phellodendron Bark in Group 1 of Column II or Group 5 of Column V of Table 1. Glycyrrhizinic acid, its salts, and glycyrrhiza extracts in Group 3 of Column VII cannot be combined with Glycyrrhiza in Group 9 of Column VII.

(i) The vitamins given in the Appendix may be combined with the active ingredients listed in Table 1 as long as there is good reason for their

combination and the effect is mild.

(2)Quantities of Active Ingredients

(a) The maximum daily doses of the active ingredients listed in Table 1 (except for those in Group 1 of Column III and Group 1 of Column IV) should correspond to data in Table 1. The maximum single dose should be 1/3rd of the maximum daily dose.

(b) When not less than 2 active ingredients in Group 1 or Group 2 of Column I listed in Table 1 are combined, the sum of the values obtained by dividing the amount of each active ingredient by its respective maximum daily dose should

not exceed 2.

(d)

(c) When at least 2 active ingredients in Group 2 or Group 3 of Column II are combined, or when at least 2 active ingredients in Group 2 of Column III or at least 2 active ingredients in Group 1, 2, 3, or 4 of Column V of Table 1 are included, the sum of the values obtained by dividing the amount of each active ingredient by its respective maximum daily dose should not exceed 1 for any group.

When the crude drugs marked with "*" in Group 1 of Column II in Table 1 are combined in preparations for which the main active ingredient comes from Column I, the daily dose of the crude drug concerned should not be more than

1/10th of the maximum daily dose shown in Table 1.

(e) When preparations whose main active ingredients are from Groups 1 and 2 of Column I and which are tested for acid-neutralizing capacity or pH by the methods specified elsewhere, the acid-neutralizing capacity of the daily dose of the preparation should not be less than 150 mL when expressed as the amount of 0.1N hydrochloric acid consumed, and the pH of the preparation should not be less than 3.5.

The acid-neutralizing capacity of a single dose of the preparation should be not

less than 50 mL.

(f) In preparations mainly containing active ingredients from Group 1 of Column III of Table 1, the digestive activity of the digestive enzymes included in a single dose of the preparation should not be less than the minimum daily unit for at least 1 of the following: starch saccharifying activity, starch dextrinizing activity, starch liquefying activity, protein digesting activity, fat digesting activity, fibrin saccharifying activity, or fibrin disintegrating activity specified in Group 1 of Column III.

The minimum unit for a single dose shall be 1/3rd of the minimum daily unit.

(g) For preparations mainly containing active ingredients from Group 1 of Column IV in Table 1, the minimum daily dose of the active ingredient concerned should be the amount shown in Table 1, and the minimum single dose should be 1/3rd of the minimum daily dose.

(3)Dosage Form

The dosage forms should be capsules, granules, pills, fine granules, powders, electuaries, tablets, infusions, decoctions, or liquids for oral use (limited to mildly

acting preparations mainly containing ingredients from Column I or II).

(4)Dosage and Administration

- (a) In principle, dosage and administration should be 3 times a day.
 Oral liquids mainly containing ingredients from Column I or II, or preparations mainly containing ingredients from Column V or VI listed in Table 1 can be
 taken 1 to 3 times a day, and if they are taken not less than 2 times a day, the interval between doses must not be less than 4 hours.
- (b) For infusions and decoctions, the method of preparation at the time of use should be indicated.
- (c) The time of administration (such as before or after meals, between meals) and the administration interval should be indicated.
- (d) Dosage in infants less than 3 months of age is not approved.
- (e) For capsules, pills, or tablets larger than 6 mm in diameter, dosage in children less than 5 years of age is not approved.
- (f) For pills or tablets smaller than 6 mm in diameter, dosage in children less than 3 years of age is not approved.
- (g) The maximum daily dose for children less than 15 years of age should be obtained by multiplying the maximum daily doses listed in Table 1 by the values given in the coefficient column for the corresponding age ranges stated in Table 2.
- (h) The minimum daily doses specified in (2) (e) and (2) (f) should be multiplied by the values given in the coefficient column for the corresponding age ranges in Table 2 to obtain the minimum daily dose for children less than 15 years of age. However, the minimum daily doses specified in (2) (g) should be applied irrespective of age.

(5)Indications

- (a) The range of indications for preparations mainly containing active ingredients from the columns of Table 1 (except Columns VII and VIII) is shown in Table 3. When active ingredients from at least 2 of Columns I, II, III, and IV are used as the main ingredients, the indications should cover all of those in the columns concerned.
 - The indications in Column III of Table 3 can be claimed for preparations whose main active ingredients are from Group 1 in Column III, only if the minimum daily units of at least 1 of the following are achieved: starch saccharifying activity, starch dextrinizing activity, starch liquefying activity, protein digestive activity, and fat digestive activity.
- (b) For preparations claiming the indications mentioned in Column V or VI of Table 3, the indications listed in the other columns of the same table should not be claimed.
- (c) Notwithstanding the above standards, the indications in Column I of Table 3 cannot be claimed in cases where Nux Vomica Extract in Group 1 of Column II is included in preparations containing active ingredients from Column I in Table 1.
 - In addition, the indications in Column I of Table 3 cannot be claimed for preparations containing active ingredients only from Group 3 of Column I in Table 1.

(Table 1)

(Table 1) Classification		Active ingredient	Maximum daily dose	
		Dried aluminum hydroxide gel	3 g	
		Magnesium aluminosilicate	4 g	
		Magnesium silicate	6 g	
		Synthetic aluminum silicate	10 g	
	,	Synthetic hydrotalcite	4 g	
		Magnesium oxide	1 g	
ı		Magnesium hydroxide aluminum hydroxide co-precipitate	4 g	
,,,		Aluminum hydroxide gel	30 mL	
<u>.</u>	•		(1.2 g as aluminum oxide)	
		Aluminum hydroxide sodium bicarbonate co-precipitate	2 g	
	Group 1	Dried mixed aluminum hydroxide and magnesium carbonate gel	3 g	
H		Aluminum hydroxide magnesium carbonate calcium carbonate co-precipitate	4 g	
шш		Magnesium hydroxide	2.4 g	
Column I	,	Sodium bicarbonate	5 g	
Ö		Magnesium carbonate	2 g	
· •		Precipitated calcium carbonate	3 g	
		Magnesium aluminometasilicate	4 g	
		Anhydrous dibasic calcium phosphate	2.4 g	
		Dibasic calcium phosphate	3 g	
		Cuttlefish Bone	3 g	
		Abalone Shell	3 g	
		Oyster Shell	3 g	
	2	Aminoacetic acid	0.9 g	
	Group	Dihydroxyaluminum aminoacetate	3 g	
	Gre			
	Group 3	Scopolia Extract	30 mg	

	Classification		Maximum daily dose (g)						Maximum daily dose	
			Active ingredient	Extract (converted to crude drug	Powder	Classification		Active ingredient	Extract (converted to crude drug	Powder
		T	Aniseed	amount)		<u> </u>		O':	amount)	
			Amiseed	3	1			Citrus	5	3
		1.	Aloe		0.15	'		Unshiu Peel *Capsicum		
-			Fennel	3	1			Bitter	5	0.1
					-			Orange Peel	9	3
			Turmeric	6	2			Animal bile (including	_	0.5
ı			I make a Deci					Bear Bile)	1	
			Lindera Root	5	1			Picrasma Wood	5	0.5
			Isodon Herb	10	3			Nutmeg	3	1
			Scutellaria Root	6	3			Ginseng	, 6	- 3
			Phellodendron	. 3						
			Bark	3	3			Mentha Herb (including	3	1
- [Coptis	3	1.5			peppermint)		-
			Rhizome					Long pepper	2	0.5
			Processed Garlic Bulb	-].	0.2			Atractylodes	5	2
			Zedoary	3	3			Rhizome		
			Pogostemon	8	3	•		Hop Strobile Nux Vomica	3	1
			Herb	Ĭ	"			Extract		0.03
			Calamus Root	6	2	•		Menyanthes	4	1.3
	н.		Processed	3	1	<u>.</u> .		trifolia herb	. 4	1.5
	g	O.	Ginger	1		Д	1			
l	Column II	Group 1	Orange Fruit	5	2	Column II	Group	Saussurea	3	. 1
	ပိ	ජි	Immature	_		200	Q.	Root		
			Orange	5	2	_		Bitter	3	1
		•	Cinnamon	5	1			Cardamon Japanese		[
,			Bark	3	1	:		Japanese Gentian	15	0.5
l		-	Gentian	1.5	0.5			Alpinia	3	1
			Red Ginseng	6	3			Officinarum		1
	[ĺ			Rhizome		.
			Magnolia Bark	5	1.5			Fennel Oil	0.0	8
			Euodia Fruit	3	1			Cinnamon Oil	0.0	3
			*Pepper	5	1.5		L	Ginger Oil	0.03	3
	1		Calumba	5	1.5	ŀ		Cardamon	0.03	
				ļ		1		Oil	0.00	
	1		Condurango	9	3			Clove Oil	0.02	2
	ŀ		*Japanese Zanthoxylum	3	1			Bitter	0.03	3
			Peel		. [Orange Peel	•	
			Resurrection	6	2	İ		Oil Mentha Oil	0.03	
		1	Lily Rhizome Perilla Fruit						•	
			Amomum	6 3	3			Lemon Oil	0.03	
			Seed	1	1		j	Menthol	0.18	}
			Ginger	3	1		1	d/Menthol	0.18	i

				·	•			·	
		Cardamon	3	1			-		
		Immature	5	3			.	•	ł
		Citrus Unshiu				C3	Betaine hydrochloride	0.6	
		Peel	6	. 2		Group	L-Glutamic	•	
		Acorus Gramineus	0	. 4		}ro	acid	1.8	
		Rhizome					hydrochloride		•
		Centaury	2	0.7					
		Herb							
		Swertia Herb	1.5	0.05			Carnitine	0.6	
		Atractylodes	5	2		က	chloride	Vio	
		Lancea				Group			
		Rhizome				Gr	Bethanechol		ŀ
		Perilla Herb	2 3	1		_	chloride	0.045	
		Star Anise Rhubarb	0.2	0.1					1 .
		Panax	6	3			.		
		Japonicus	"			ıp 4	[n.,	10	
		Rhizome				Group	Dried yeast	10	
ľ	1	Clove	2	0.5	-	0	.		

Class	ification	Active ingredient	Minimum daily unit	Note 1)
		Starch digestive enzymes	Starch saccharifying activity:	250 unit
			Starch dextrinizing activity	210 units
	-		Starch liquefying activity:	360 units
	Group	Protein digestive enzymes	Proteolytic activity:	1,500 units
	Grö	Fat digestive enzymes	Fat digestive activity:	100 units
H	-	Fibrin digestive enzymes	Fibrin saccharifying activity:	13 units
Column III			Fibrin disintegrating activity:	25 units
		Active ingredient	Maximum daily dose	(g)
		Ursodesoxycholic acid	0.06	
	7	Oxycholanates	0.15	
	dn	Cholic acid	0.9	
٠	Group	Gall powder	1.5	
		Gall extract (powder)	0.5	
	[Dehydrocholic acid	0.5	
		Animal bile (including Bear Bile)	0.5	

Note 1) Methods for measuring the digestive activity of each digestive enzyme are specified separately.

		Active ingredient	Minimum	daily dose
-	Group 1	Live bacteria for intestinal regulation	1 × 10 ⁶	
			Maximum d	aily dose (g)
Column IV	1p 2		Extract (converted to crude drug amount)	Powder
	Group	Mallotus Bark	. 5	1.5
	0	Gambir	-	. 2
		Processed Mume	10	3 -
-		Cassia Seed	10	3
		Geranium Herb	10	3

Classif	cation	Active ingredient	Maximum da	aily dose (g)
.		Acrinol	0.3	-,
İ	. :	Berberine chloride	0.3	
	-	Guaiacol	0.6	
	ďn	Creosote	0.5	
	Group 1	Phenyl salicylate	1	
,		Guaiacol carbonate	1.2	
		Berberine tannate	0.3	
-		Bismuth subsalicylate	3	
		Bismuth subnitrate	2	
	2	Bismuth subcarbonate	3	
	Group 2	Bismuth subgallate	2	
	Ę,	Tannic acid	1.2	
	,	Albumin tannate	. 4	
-	28	Methylene thymol tannin	. 2	
	23.	Kaolin	10	
	ന	Natural aluminum silicate	. 10	•
Λ	Group	Aluminum hydroxynaphthoate	0.9	
uu	용	Pectin	0.6	
Column V	Ů	Medicinal carbon	5	•
ပိ	4	Precipitated calcium carbonate	3	
1.54	đn	Calcium lactate	5	
	Group	Dibasic calcium phosphate	. 3	
			Extract (g)	
		. •	(converted to	Powder (g)
			crude drug	1 OWACL (G/
			amount)	
		△ Gambir	-	2
		△ Processed Mume	10	3
		Phellodendron Bark	9	3
	ī.	Coptis Rhizome	3	1.5
	Group 5	Sophora Root	3 ⁻ 10	1.5 3
	Ä	△ Geranium Herb Rhus Javanica Nutgall	-	3
		△ Crataegus Fruit	8	3
		Swertia Herb	-	0.9
1 .		Myrica Rubra Bark	5	2

Classific	cation	Active ingredient	Maximum	ı daily dose
		Oxyphencyclimine hydrochloride	7	mg
Ì		Dicyclomine hydrochloride	30 mg	
		Methixene hydrochloride	l .	75 mg
		Scopolamine hydrobromide		3 mg
1		Atropine methylbromide	1	ng
	·	Anisotropine methylbromide	30 1	*
	Group	Scopolamine methylbromide	•	3 mg
'	Ç,	IHyoscyamine methylbromide	,	25 mg
l		Methylbenactyzium bromide	30 r	•
]		Belladonna extract	60 i	•
-		Isopropamide iodide	F	mg
		Diphenylpiperidinomethyldioxolane iodide	60 r	
₽ I		Scopolia Extract	60 r	- ·
a L		Scopolia Rhizome (Total) Alkaloid citrates	. 1 r	
Column VI	Group 2	Papaverine hydrochloride	90 n	
	Group 3	Ethyl aminobenzoate	0.6	mg
	-		Extract (g) (converted to crude drug amount)	Powder (g)
[.	4	Corydalis Tuber	5	1.5
	Group 4	Glycyrrhiza	5	1.5
.	ž.	Magnolia Bark	5	1.5
,		Peony Root	5	2

Classifi	cation	Active ingredient	Maximum daily dose (g)
	Group 1	Sodium azulene sulfonate	0.006
	Group 2	Aldioxa	0.3
	Group 3	Glycyrrhizinic acid, its salts, and glycyrrhiza extracts	(as glycyrrhizinic acid) 0.2
-	Group 4	L-Glutamine	2
n VII	Group 5	Potassium copper chlorophyllin Sodium copper chlorophyllin	0.2 0.2
Column VII	Group 6	Histidine monohydrochloride	0.18
	Group 7	Pepsin decomposition products of pig stomach wall Acid hydrolysis products of pig stomach wall	0.3 0.3
	Group 8	Methylmethioninesulfonium chloride	0.15
	Group 9		Extract (g) (converted to Powder crude drug (g) amount)
'	Gro	Mallotus Bark Corydalis Tuber Glycyrrhiza	5 1.5 5 1.5 5 1.5

Dimethylpolysiloxane 0.18 g	olumn VIII	Dimethylpolysiloxane	0.18 g	
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(Table 2)

Age coefficients

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Age	Coefficients
15 years of age or over 11 to under 15 years of age 8 to under 11 years of age 5 to under 8 years of age 3 to under 5 years of age 1 to under 3 years of age 3 months to under 1 year of age	1 2/3 1/2 1/3 1/4 1/5

(Table 3)

Main ingredient	Indications
Column I	Hyperacidity, heartburn, feeling of discomfort in the stomach, feeling of fullness in the stomach, constricted feeling in the stomach (stomach heaviness), heaviness in the stomach, heaviness in the chest, belching (burping), nausea (retching, stomach retching, retching due to hangovers and overdrinking, sick feeling, and feeling of sickness), vomiting, excessive drinking (overdrinking), and stomachache
Column II	Loss of appetite (anorexia), feeling of fullness in the stomach and abdomen, indigestion, weak stomach, excessive eating (overeating), excessive drinking (overdrinking), heartburn, constricted feeling in the stomach (stomach heaviness), heaviness in the chest, nausea (retching, stomach retching, retching due to hangovers and overdrinking, sick feeling, and feeling of sickness), and vomiting
Column III	For promoting digestion, indigestion, loss of appetite (anorexia), excessive eating (overeating), constricted feeling in the stomach (stomach heaviness), heaviness in the chest, and feeling of fullness in the stomach and abdomen due to indigestion
Column IV	Intestinal regulation (regulation of stool), feeling of fullness in the abdomen, soft stool, and constipation
Column V	Diarrhea, diarrhea due to indigestion, food poisoning, vomiting and purging, water poisoning, loose bowels, soft stool, and diarrhea accompanied by abdominal pain ^{Note 1)}
Column VI	Stomachache, abdominal pain, gripping pain (colic, spasms), hyperacidity, and heartburn

Note 1) Only when scopolia extract in Group 1 of Column VI is included.

(Appendix)

1. Vitamins that can be included in preparations mainly containing active ingredients from Column II or III are indicated below, together with their maximum daily doses.

Ingredient	Maximum daily dose
Vitamin B ₁ , its derivatives, and their salts	25 mg

2. Vitamins that can be included in preparations mainly containing active ingredients from Column IV are listed below, together with their maximum daily doses.

Ingredient	Maximum daily dose
Nicotinamide Calcium panthothenate Biotin Vitamin B ₁ , its derivatives, and their salts Vitamin B ₂ , its derivatives, and their salts Vitamin B ₆ , its derivatives, and their salts Vitamin C, its derivatives, and their salts	5 mg 30 mg 25 μg 25 mg 12 mg 50 mg 500 mg

However, the combination of biotin and nicotinamide is permitted only when including live lactic acid bacteria or lactic acid producing bacteria for intestinal regulation.

3. Vitamins that can be included in preparations mainly containing active ingredients from Column V are listed below together with their maximum daily doses.

	Ingredient	Maximum daily dose
,	s derivatives, and their salts s derivatives, and their salts	25 mg 12 mg

Provisional Translation from Japanese Original

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The Standards for Marketing Approval of Laxatives

Scope of Laxatives

The scope of preparations subject to these standards covers oral medicines intended for the relief of the symptoms of constipation or the elimination of intestinal contents (except for preparations covered by the Standards for Marketing Approval of gastrointestinal medicines and Kampo medicine* formulas.

* Kampo medicine is traditional Japanese medicine.

2. Approval Standards

The approval standards for laxatives are as follows.

For preparations not conforming to these standards, concerning the efficacy and safety data and reasons justifying the combination should be submitted; the preparation in question will be reviewed based on these data.

Types of Active Ingredients

- The types of active ingredients that may be used in laxatives are shown in Tables 1 and 2.
- At least 1 of the active ingredients in Table 1 must be used.
- Preparations mainly containing the active ingredients from Group I, II, III, or IV in Column A of Table 1 may be made by mutual combination of the active ingredients in these 4 groups, and may also include the active ingredients in Table 2.
- When active ingredients from Group I, Group II, or Group III in Column A of Table 1 are combined, only 1 ingredient from each group should be used. When active ingredients from Group IV are used, up to 4 active ingredients from this group may be included.

However, when active ingredients from 2 or more groups, among Groups I, II, III, and IV, are combined, up to 4 active ingredients from Column A of Table 1

(except Group V) may be combined.

- The following combinations are not permitted among the active ingredients of Group IV in Column A of Table 1: Aloes with aloin, Cascara sagrada bark with casanthranol, Pharbitis seeds with Pharbitis seed resin, Senna or Senna fruit with sennoside or sennosides A and B, and Jalap tuber with Jalap resin.
- For preparations mainly containing the active ingredients from Group V of Column A in Table 1, combinations with the other active ingredients in these standards are not permitted.
- When the active ingredients from Column B of Table 1 are used as a main ingredient, only 1 active ingredient can be used in a preparation and none of the other active ingredients covered by these standards should be combined.
- When the active ingredients from Column I or II of Table 2 are combined, up to 4 active ingredients in the same column may be used. When active ingredients in both Columns I and II of Table 2 are combined, up

to 5 of the active ingredients from the whole table may be used.

(i) Other than the active ingredients in Tables 1 and 2, vitamins in the Appendix may be included if there is a sound basis for their combination and the effect is mild.

(2) Quantities of Active Ingredients

(a) The maximum single and daily doses of the active ingredients from Column A of Table 1 are as indicated in the table.

b) The maximum single doses of the active ingredients from Column B of Table 1

are as indicated in the table.

(c) The maximum daily dose of each of the active ingredients from Column I (except live bacteria for intestinal regulation) and Column II of Table 2 are as given in the table. The maximum single dose should be 1/3rd of the maximum daily dose.

(d). When 2 or more of the active ingredients from Column A of Table 1 are combined, the sum of the values obtained by dividing the amounts of each of the active ingredients by their respective maximum daily doses should not

exceed 2.

(e) When 2 or more of the active ingredients from either Column I or Column II of Table 2 are combined, the sum of the values obtained by dividing the amounts of each of the active ingredients by their respective maximum daily doses should not exceed 2 in each column.

(f) The minimum daily dose of live bacteria for intestinal regulation from Column I of Table 2 is as given in the same group, and the minimum single dose should

be 1/3rd of the minimum daily dose.

(3) Dosage Forms

The dosage forms are capsules, granules, pills, fine granules, powders, lingual tablets (limited to preparations mainly containing the active ingredients from Group V of Column A of Table 1), tablets, infusions, decoctions, chocolate preparations and liquids for oral use (limited to syrups and preparations mainly containing the active ingredients from Group I of Column A or those from Column B of Table 1).

(4) Dosage and Administration

(a) Preparations should, in principle, be taken by oral administration 1 to 3 times daily, and the administration times and intervals must be clearly indicated. When the preparation is taken twice a day or more, the interval between doses must be not less than 4 hours. However, preparations mainly containing the active ingredients from Column

B of Table 1 should be taken not more than once a day, to be taken when

required.

(b) For preparations mainly containing the active ingredients from Column A of Table 1, the dosage range for different degrees of constipation must be indicated.

Since there are individual differences with respect to the degree of constipation, it must be stated that the minimum dose should be taken initially and then the dose should be gradually increased (or decreased) depending on the condition of relief.

(c) In principle, dosage for children under 3 years of age is not permitted.

(d) Regardless of the rules described in (a), (b), or (c), preparations mainly containing the active ingredients from Group V of Column A in Table 1 will be approved only for small children and infants. Entries for dosage and

administration should be made in accordance with Table 5.

(e) In the case of infusions and decoctions, the method of preparation at the time of use should be clearly indicated.

For capsules, and pills and tablets larger than 6 mm in diameter, dosage for

children under 5 years of age is not approved.

(g) The maximum single and daily doses for those under 15 years of age are the values obtained by multiplying the coefficients corresponding to the respective age groups in Table 3 by the maximum single and daily doses shown in Tables 1 and 2.

However, the minimum daily dose of live bacteria for intestinal regulation from Column I of Table 2 should be applied irrespective of age.

(5) Indications

(a) The indications for preparations mainly containing the active ingredients from Column A of Table 1 are shown from Column I of Table 4. However, the indications for preparations mainly containing the active ingredients from Group V of Column A in Table 1 are as specified in Table 5.

b) The indications for preparations mainly containing the active ingredients from

Column B of Table 1 are as specified from Column II of Table 4.

(6) Packaging Units

The maximum volume of syrup containers is a 2-day supply at the maximum daily dose for adults (15 years of age and over).

<u>Table</u>	1					
Classification		Active ingredients	Maximum single dose		Maximum daily dose	
		(g)			(g)	
		Magnesium oxide	0.7 (2)			2
	Group I	Magnesium hydroxide	0.7 (2.	1)		2.1
	on	Magnesium carbonate	2.7			. 8
n 7	Ġ	Sodium sulfate	5			15
Column A		Magnesium sulfate	.5			15
- <u>-</u> [5]		Carboxymethylcellulose calcium	2			6
۱	Group II	Carboxymethylcellulose sodium	2	•		6
	, on	Plantago ovata coating (Ispaghula	3.5			10.5
	ਲੋਂ .	husk)				
	Ħ					
	Group III	Sodium dioctyl-sulfosuccinate	0.067 (0.3	12)		0,2
	o	Domain dioctyr suriosuccinate	0.001 (0.	,		3,2
	වි					
	**.		,			
	.*	Aloin	0.02			0.06
		Sulfur	0.5			1.5
		Casanthranol	0.067 (0.			0.2
		Sennoside (as sennosides A and B)	0.016 (0.			0.048
	, I	Sennosides A and B	0.016 (0.0			0.048
	Inc	Bisacodyl	0.007 (0.0			. 0.02
	Group IV			Extract (g)	1 .	Extract (g)
			Powder	(converted	Powder	(converted
			(g)	to crude	(g)	to crude
		`	(6)	drug	8.	- drug
				amount)		amount)
1	,	Aloes .	0.25	0.25	0.75	0.75
			(0.38)	(0.38)		,
		Rose fruit	0.67	1.7	2	5
		Cascara sagrada bark		1 (1.5)		3 .
		Pharbitis seed	0.1	-	0.3	-
-		Pharbitis seed resin	0.05		0.15	-
		Senna	0.5	2	1.5	6
			(0.75)	(3)		
		Senna fruit	0.5	. –	1.5	
		•	(0.75)			
Í		Rhubarb	1 (1.5)	1.4 (2)	3	4
		Frangula bark	_	1 (1.5)	-	. 3
		Jalap root	0.1		0.3	_
		Jalap resin	0.05	_	0.15	<u> </u>
	>			•		
	Group V			11 8		
	ő	Malt extract	As per Ta	ante o		
	J					
		1				
Column B	·		-			
		tic castor oil		20 mL		_
nlc nlc	Castor	oil ·		$20 \mathrm{mL}$		_
ರ		ϵ				
	<u> </u>		1			

(Note) Figures in parentheses are the maximum single dose applicable when the dosage is once or twice a day.

Table 2

Classification	Active ingredient		num daily ose (g)
-	Ursodeoxycholic acid	<u> </u>	0.06
	Oxycolanate	•	0.15
	Dried yeast		10
	Cholic acid		0.9
•	Dimethylpolysiloxane		0.18
	Live bacteria for		1×10° (*)
	intestinal regulation		1×10- (-)
	Sodium bicarbonate		3
	Dehydrocholic acid		_
	Denyarochoric aciti.		0.5
		İ	Extract
		Powder	(g)
			(converted
		(g)	to crude
			drug
	Linseed	2	amount)
	Japanese valerian	2	_
Column I	Glycyrrhiza	•	
	Cassia seed	1.5	5
		3	10
	Smilax rhizome	1.5	5
	Gardenia fruit	1	3
	Rehmannia root	1.5	5
	Peony root	2	. 5
•	Houttuynia herb	5	15
·	Cimicifuga rhizome	1	3
	Cnidium rhizome	1.5	5
	Jujube	1.5	5
	Bile extract (powder)	0.	5
	Japanese angelica	1.5	5
	root		
	Animal bile	0.5	
	Moutan bark	1.3	4
	Hemp fruit	5	_
	Coix seed	6	20

(*) Minimum daily dose

		Maximum daily dose (g)		
	1		Extract (g)	
Classification	Active ingredient	1	1 . ~	
Cidobilication	Mostve mgredient	Powder	(converted	
		(g)	drug	
		1	amount)	
	Fennel	0.5	1.5	
	Plectranthus	1.5	5	
	herb		"	
	Scutellaria root	1.5	3	
	Phellodendron -	1.5	1.5	
	Bark			
	Coptis Rhizome	0.75	1.5	
	Zeodary	1.5	1.5	
	Calamus Root	1	3	
	Immature orange	1.	2.5	
	Cinnamon Bark	0.5	2.5	
	Gentian	0.25	0.75	
	Magnolia bark	0.75	2.5	
	Condurango	1.5	4.5	
	Resurrection Lily	1	3	
	Rhizome			
	Ginger	0.5	1.5	
Column II	Swertia herb	0.025	0.75	
Column	Atractylodes	1	2.5	
	Lancea Rhizome			
	Perilla Herb	0.5	1	
	Citrus Unshiu	1.5	2.5	
	Peel			
	Bitter orange peel	1.5	2.5	
	Ginseng	1.5	3	
	Mentha herb	0.5	1.5	
	Mentha oil	0.0	15	
	Atractylodes	1	2.5	
	rhizome			
	Nux vomica	0.0	15	
	extract			
	dl-Menthol	0.09		
į	I-Menthol .	0.09		
	Saussurea root	0.5	1.5	
	Japanese gentian	0.25	0.75	

Table 3

Age coefficient

,	
Age	Coefficient
15 years of age and over	`1
11 to under 15 years of age	2/3
7 to under 11 years of age	1/2
3 to under 7 years of age	1/3

Table 4

`	Indications
Column I	O Constipation O Relief of the following symptoms due to constipation: dull headache, hot flush, skin roughness, eruption, loss of appetite (anorexia), fullness in the abdomen, abnormal fermentation in the intestines, and hemorrhoids
Column II	O Rapid excretion of intestinal contents (food poisoning, etc.)

Table 5

Dosage and administration (maximum single dose)	Indications
1 to under 3 years of age: 15 g/dose 6 months to under 1 year of age: 9 g/dose Under 6 months of age: 9 g/dose Take orally up to 3 times a day in each case	Constipation in infants and small children

Appendix

Ingredients	Maximum daily dose
Vitamin B ₁ , its derivatives, and their salts	25 mg
Vitamin B ₆	50 mg
Nicotinamide	5 mg
Calcium panthothenate	30 mg

(Note) Nicotinamide is to be combined only when lactic acid bacteria or lactic acid producing bacteria are used as live bacteria for intestinal regulation.

Provisional Translation from Japanese Original

Jun 1, 1984 Notification PB No.381

The Standards for Marketing Approval of Antivertigo Medicines

1. Scope of Antivertigo Medicines

The scope of preparations subject to these standards covers oral medicines (Kampo medicine* formulas are not covered) intended to prevent or relieve symptoms associated with motion sickness, such as dizziness, nausea, and headaches.

*Kampo medicine is traditional Japanese medicine.

2. Approval Standards

The approval standards for antivertigo medicines intended to prevent or relieve symptoms associated with motion sickness (hereinafter referred to as motion sickness drugs) are as follows.

For motion sickness drugs and antivertigo medicines other than motion sickness drugs not conforming to these standards, efficacy and safety data and reasons justifying the combination should be submitted; the preparation in question will be reviewed based on these data.

(1) Types of Active Ingredients

(a) The types of active ingredients that may be combined are shown in Table 1.

(b) At least one ingredient from either Column I or Group 1 of Column II of Table 1 must be combined.

(c) Though the active ingredients in Column I, II, III, IV, V, VI, or VII of Table 1 may all be mutually combined, the types of active ingredients that may be combined in oral liquid preparations should be those in Column I, Group 1 of Column II, Column V, and Column VII.

(d) Up to 2 ingredients from each of Column I or V in Table 1 may be included (however, only 1 ingredient from each of Group 1 or 2 of Column V may be combined).

One active ingredient each from Column II, III, IV, VI, or VII may be included.

(e) Other than the active ingredients in Table 1, vitamins listed in the Appendix may be included if there is a sound basis for their combination and the effect is mild.

(2) Quantities of Active Ingredients

(a) Table 1 shows the maximum single and daily doses for each of the active ingredients listed.

(b) When 1 active ingredient listed in either Column I or Group 1 of Column II of Table 1 is used, the lower limit of the single dose of each active ingredient

should be half of the maximum single dose.

(c) When 2 of the active ingredients in Column I of Table 1 are used, the lower limit of the single dose of each active ingredient should be 1/5th of the maximum single dose. In addition, the sum of the values obtained by dividing the amounts of each active ingredient by their respective maximum single dose should be not less than 0.5 and not more than 1.

When active ingredients in Column I or Group 1 of Column II of Table are combined mutually, the lower limit of the single dose of each active ingredient should be 1/5th of the maximum single dose. Further, the sum of the values obtained by dividing the amounts of each active ingredient by their respective maximum single dose should be not less than 0.5 and not more than 2.

The lower limit of the single dose of each active ingredient in Group 2 or 3 of Column II, Column IV, Column V, or Column VI of Table 1 should

be 1/5th of the maximum single dose.

When 2 ingredients from Column V of Table 1 are combined, the sum of the values obtained by dividing the amounts of each active ingredient by their respective maximum single dose should not exceed 1.

The lower limit of the single dose of each active ingredient in Column VII of

Table 1 should be 1/10th of the maximum single dose.

The maximum daily dose of each active ingredient listed in the Appendix is as specified in the table.

(3) Dosage Form

The dosage forms are capsules, granules, pills, fine granules, powders, tablets (including chewable tablets), and oral liquids.

(4) Dosage and Administration

Dosage is by oral administration from 1 to 3 times a day (with the exception of 1 to 4 times a day for single active ingredient preparations containing dimenhydrinate). The time of administration and intervals between doses should be clearly indicated. For medicines designed to be taken twice a day or more, the interval between doses must be at least 4 hours.

In principle, dosage for children under 3 years of age is not approved. In the case of preparations containing ethyl aminobenzoate, dosage is not approved for children under 6 years of age, and as for preparations containing promethazine hydrochloride or promethazine methylene disalicylate, dosage

for those under 15 years of age is not approved.

For capsules, and pills and tablets larger than 6 mm in diameter, dosage for

children under 5 years of age is not approved.

The maximum single and daily doses for children under 15 years of age is obtained by multiplying the maximum single and daily doses given in Table 1 by the coefficient for each age group given in Table 2.

The method of administration must be clearly indicated for chewable tablets.

Indications

The indications are "prevention and relief of dizziness, nausea, and headache associated with motion sickness."

Packaging Units

In principle, the volume of containers for oral liquids should be the amount for a single dose and should not exceed 30 mL.

Table 1

Table 1 Column			Active ingredient	Maximum single dose (mg)	Maximum daily dose (mg)
Column I		Ι	Difenidol hydrochloride	25	75
			Diphenylpyraline hydrochloride	4	12
			Diphenhydramine hydrochloride	50	150
			Promethazine hydrochloride	25	50
	•		Meclizine hydrochloride	50	75
			Diphenhydramine salicylate	60	180
			Dimenhydrinate	50	200
			Diphenhydramine tannate	150	450
			Fenethazine tannate	30	90
			Diphenylpyraline teoclate	3	
			Diphenhydramine fumarate		9
			Promethazine methylenedisalicylate	60	180
			dl-Chlorpheniramine maleate	30	60
				4	12
			d Chlorpheniramine maleate	2	6
		<u>. </u>	Pheniramine maleate	30	90
		Group 1	Scopolamine hydrobromide	0.25	0.50
			Oxyphencyclimine hydrochloride	2.34	. 7
			Dicyclomine hydrochloride	10	30
`			Methixene hydrochloride	2.92	8.75
			Atropine methylbromide	2	. 6
	1:	Π .	Anisotropine methylbromide	10	30
Column II		Group II	Scopolamine methylbromide	1.6	4.8
lan l	١,		Hyoscyamine methylbromide	0.75	2.25
රි	`	0	Metylbenactyzium bromide	10	30
			Belladonna extract	20	60
			Isopropamide iodide	2.5	7.5
			Diphenylpiperidinomethyldioxolan iodide	20	60
	_		Scopolia extract	20	60
		Group III	Papaverine hydrochloride	30	. 90
Colı	ımn I	II	Ethyl aminobenzoate	100	300
			Cerium oxalate	100	300
			Ethyl p-piperidinoacetylaminobenzoate	200	600
Coh	ımn F	V	Allylisopropylacetylurea	60	180
	T		Bromovalerylurea	200	600
Column V	l dr		Caffeine	50	150
	ű		Caffeine citrate	100	300
	ļ		Anhydrous caffeine	50	150
-	Πď	ļ	Aminophylline	. 100	300
Group II			Diprophylline	100	300
			Theophylline	100	300
	mn V		Sodium bicarbonate	1,000	3,000
Column VII		/II	Mentha oil	5	15
,		ŀ	dl-Menthol	30	90
`			I-Menthol	30	90

Table 2

Age	Coefficient	
15 years old and over	. 1	
11 years old Under 15	2/3	
7 years old Under 11	1/2	
3 years old Under 7	1/3	

Appendix

Ingredients	Maximum daily dose (mg)
Vitamin B ₁ , its derivatives, and their salts	25
Vitamin B2, its derivatives, and their salts	12
Vitamin B ₆ , its derivatives, and their salts	50
Nicotinamide	60
Calcium panthothenate	30

Provisional Translation from Japanese Original

Jul 29, 1986 Notification PB No.623

The Standards for Marketing Approval of Ophthalmic Medicines

1. Scope of Ophthalmic Medicines

The scope of preparations subject to these standards covers medicines to be applied to the mucous membrane of the eyes to treat symptoms of eye diseases and those to be used when inserting contact lenses.

2. Approval Standards

The approval standards for ophthalmic medicines are as follows.

For preparations not conforming to these standards, efficacy and safety data and reasons justifying the combination should be submitted; the preparation in question will be reviewed based on these data.

(1) Types of Active Ingredients

- (a) Active ingredients that may be used in ophthalmic medicines are listed in Table I.
- (b) At least 1 active ingredient from Column A, B, C, or D; Group 1, 2, or 3 of Column E; Column F, G, or H; Group 1 of Column I; or Column J in Table I must be used.
- (c) Preparations mainly containing the active ingredients in Column A, B, C, or D; Group 1, 2, or 3 of Column E; or Group 1 of Column F (hereinafter referred to as "ordinary eye drops") in Table I may be formulated through the mutual combination of any of the active ingredients in these columns and groups, and may also include the active ingredients in Group 4, 5, or 6 of Column E or those in Group 2 or 3 of Column F in Table I.
- (d) Preparations mainly containing active ingredients in Column G (hereinafter referred to as "antibacterial eye drops") in Table I may include up to 3 active ingredients from Column A, B, C, D, E, or F.
- (e) Preparations mainly containing active ingredients in Groups 2 or 3 of Column F or those in Column H of Table I (hereinafter referred to as "artificial tears") may be formulated through the mutual combination of any of the active ingredients in Group 2 or 3 of Column F or those in Column H, and may also include the active ingredients in Group 1 of Column F or those in Column I.
- (f) Preparations mainly containing active ingredients in Group 1 of Column I (hereinafter referred to as "contact lens insertion preparations") of Table I may also include active ingredients in Column F or H or those in Group 2 of Column I.
- (g) Preparations mainly containing active ingredients in Column C, D, H, or J, listed in Table I, are used for washing the eyes and are referred to as "eyewashes." Those mainly containing active ingredients from Column C or D may be formulated by combining any of the active ingredients from Column C or D, and may also include active ingredients from Column E or F. Preparations mainly containing active ingredients from Column H or J of

Table I can include only 1 active ingredient from Column H or J, and no other active ingredients mentioned in these standards should be used.

(h) When the active ingredients from Column A, D, or G of Table I are combined,

only 1 ingredient from each column may be used.

(i) When the active ingredients from Column C, E, or F of Table I are combined, up to 3 ingredients from each column may be used, but only 1 from each group is permitted.

(2) Quantities of Active Ingredients

(a) The maximum concentrations of the active ingredients from Column A, B, C, D, E, F, or G; Group 1 of Column I; or Column J should be those given in mentioned in Table I.

However, in the case of eyewashes, the maximum concentrations of the active ingredients in Columns C, D, E, and F should be 1/10th of the maximum

concentrations mentioned in Table I.

(b) When 2 or more of the active ingredients from any 1 of Column C, E, or F of Table I are combined, the sum of the values obtained by dividing the concentration of each active ingredient by its respective maximum concentration should not exceed 2.

However, in the case of eyewashes, the maximum concentration stipulated in

(2) (a) shall apply.

- (c) In the case of ordinary eye drops, when only 1 active ingredient from Column A, B, C, or D; Group 1, 2, or 3 of Column E; or Group 1 of Column F of Table I is included, the minimum concentration of the ingredients should be half of the maximum concentration. When 2 or more of these active ingredients are combined, the minimum concentration of each shall be 1/5 of the maximum concentration.
- (d) In the case of antibacterial eye drops, when active ingredients in Column G of Table I are included, the minimum concentration of these active ingredients should be half of the maximum concentration. When active ingredients from Column A, B, C, or D; Group 1, 2, or 3 of Column E; or Group 1 of Column F are included, their minimum concentrations should be 1/5 of the maximum concentration.
- (e) In the case of artificial tears, when active ingredients listed in Column F or Group 1 of Column I in Table I are used, their minimum concentrations should be 1/10th the maximum concentration. pH values must be in the range of 5.5 to 8.0, and specific osmotic pressures (specific osmotic pressures with respect to physiological saline) must be in the range of 0.85 to 1.55 when pH and osmotic pressures are measured by the methods specified elsewhere.

(f) For contact lens insertion preparations, when 1 active ingredient from Group 1 of Column I in Table I is used, the minimum concentration should be half of the maximum concentration. When 2 active ingredients are included, their minimum concentrations should be 1/5th of the maximum concentration. When active ingredients in Column F are combined, their minimum concentrations should be 1/10th of the maximum concentration.

(g) In the case of eyewashes, when active ingredients from Column C, D, or J of Table I are combined, the minimum concentration should be 1/5th of the maximum concentration specified in (2) (a). When active ingredients in Column E or F are used, the minimum concentration should be 1/10th of the maximum concentration specified in (2) (a). pH values must be in the range of 5.5 to 8.0, and specific osmotic pressures (specific osmotic pressure with respect to physiological saline) must be in the range of 0.60 to 1.55 when pH and osmotic pressures are measured by the methods specified elsewhere.

- (h) Unless otherwise specified, when active ingredients in Groups 4, 5, and 6 of Column E, or Groups 2 and 3 of Column F in Table I are combined, the minimum concentration should be 1/10th of the maximum concentration.
- (3) Dosage Form

The dosage form shall be ophthalmic solutions (eye drops and eyewashes).

(4) Dosage and Administration

- (a) Ordinary eye drops, antibacterial eye drops, and artificial tears are to be administered 3 to 6 times a day.
- (b) For contact lens insertion preparations, the detailed method of use should be stated.
- (c) Eyewashes are to be used 3 to 6 times a day to wash the eyes.

(5) Indications

(a) The range of indications for ordinary eye drops is shown in Table II-1. However, for indications in the upper column of the following table to be claimed, at least 1 of the ingredients from the columns listed in the corresponding lower column must be included.

Upper column	Lower column
Conjunctival congestion	Columns A, C, and D
Inflammation of eyes (snow blindness), blepharitis	Columns C and D and
(inflammation of the eyelids), and itchy eyes due to	Group 1 of Column E
ultraviolet light and other rays	<u> </u>

- (b) The range of indications for antibacterial eye drops is shown in Table II-2.
- (c) The range of indications for artificial tears is shown in Table II-3.

 However, "treatment of feeling of discomfort when inserting soft contact lenses" cannot be claimed when the effect is brought about due to the effect of ingredients on the lenses, such as adsorption on the lenses.
- (d) The range of indications for contact lens insertion preparations is shown in Table II-4.
 - However, "ease of insertion of soft contact lenses" cannot be claimed when the effect is brought about due to the effect of ingredients on the lenses, such as adsorption on the lenses.
- (e) The range of indications for eyewashes is shown in Table II-5.

(6) Packaging Units

- (a) The maximum volume of containers for ordinary eye drops, antibacterial eye drops, and artificial tears is 20 mL.
- (b) The maximum volume of containers for contact lens insertion preparations is 100 mL.
- (c) The maximum volume of containers for eyewashes is 500 mL.

Table I	•		
Column	Group	Active ingredient	Maximum concentration (%)
A		Epinephrine	0.003
		Epinephrine hydrochloride	0.003 (as epinephrine)
		Ephedrine hydrochloride	0.1
		Terahydrozoline hydrochloride	0.05
•		Naphazoline hydrochloride	0.003
-		Naphazoline nitrate	0.003
		Phenylephrine hydrochloride	0.1
`	,	dl-Methylephedrine hydrochloride	0.1
В		Neostigimine methylsulfate	0.005
C	1	e-Aminocaproic acid	5
	2	Allantoin	0.3
	3.	Berberine chloride	0.025
		Berberine sulfate	0.025
	4	Sodium azulene sulfonate	0.02
•	5	Dipotassium glycyrrhizinate	0.25
, .	6	Zinc sulfate	0.25
. '		Zinc lactate	0.25
	7	Lysozyme chloride	0.5 (potency)
D		Diphenhydramine hydrochloride	0.05
		Chlorpheniramine maleate	0.03
E	1	Sodium flavine adenine dinucleotide	0.05
,	2	Cyanocobalamin	0.02
	3	Retinol acetate	50,000 units/100 mL
	-	Retinol palmitate	50,000 units/100 mL
	4	Pyridoxine hydrochloride	0.1
	5	Panthenol	0.1
		Calcium pantothenate	0.1
		Sodium pantothenate	0.1
* /	6	Tocopherol acetate	0.05
F	1	Potassium L-aspartate	1
		Magnesium L-aspartate	1
		Mixture of magnesium L-aspartate and	2
		potassium L-aspartate (equal mixture)	1
	2	Aminoethyl sulfonic acid	1
	3	Sodium chondroitin sulfate	0.5

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G		Sulfamethoxazole	4
		Sodium sulfamethoxazole	4
	,	Sulfisoxazole	4
		Sodium sulfisomidine	5
H		Potassium chloride	
		Calcium chloride	
		Sodium chloride	_
		Sodium bicarbonate	_
		Sodium carbonate	_
		Dried sodium carbonate	
		Magnesium sulfate	,
		Sodium hydrogen phosphate	-
		Monobasic sodium phosphate	_
		Monobasic potassium phosphate	
I	1	Polyvinyl alcohol	2
		Polyvinylpyrrolidone	2.5
	2	Hydroxyethl cellulose	_
		Hydroxypropylmethyl cellulose	_
		Glucose	. –
	,	Methylcellulose	-
J		Alkylpolyaminoethylglycine	0.1
		Boric acid	2

Table II

l (general ophthalmic drops)	Eyestrain, redness of the conjunctiva, prevention of eye troubles (after swimming, or to wash out sweat or dust etc.), ophthalmia by ultraviolet rays etc. (snow blindness etc.), blepharitis (running eye), foreign body feeling by contact lenses, itchy eyes, blurred vision (eye mucus)
2 (antibiotic ophthalmic drops)	Conjunctivitis (pink-eye), chalazia, blepharitis (running eye), itchy eyes
3 (Artificial tears)	Eyestrain, prevention of dry-eyes, foreign-body feeling by contact lenses, blurred vision (eye mucus)
4 (eye-lotions for contact lenses)	Help to wear hard contact lenses or soft contact lenses
5 (eye washes)	Irrigation of eyes, prevention of eye troubles (after swimming, or to wash out sweat or dust etc.)

Provisional Translation from Japanese Original

Feb 1, 1985 Notification PB No.90 Final revision Mar 22, 1995

The Standards for Marketing Approval of Vitamin Preparations

Scope of Vitamin Preparations

Vitamin Preparations, as defined here, are oral vitamin preparations which contain one or more vitamins for the purpose of alleviating symptoms against which such a vitamin should be effective or for vitamin supplementation.

2. Standards

The following standards shall be applied to Vitamin Preparations.

For vitamin preparations which do not conform to these standards, the submission of documents regarding the efficacy, safety, and the basis for combination shall be required for review.

(I)Types of Active Ingredients

- A) The types of active ingredients which may be combined in vitamin preparations are listed in the attached Table 1.
- B) For preparations mainly consisting of the active ingredients listed in Column I of the attached Table 1 (hereinafter referred to as Vitamin A preparations), those mainly consisting of the active ingredients in Group 1 may include the active ingredients listed in Column II or IV of the same Table and those mainly consisting of the active ingredients in Group 2 may include the active ingredients in Group 1 of Column I, Column III, IV, or VIII.
- C) Preparations mainly consisting of the active ingredients listed in Column II of the attached Table 1 (hereinafter referred to as Vitamin D preparations) may include the active ingredients listed in Group 1 of Column I, Column III, VIII, or Group 7 of Column X of the same Table.
- D) Preparations mainly consisting of the active ingredients listed in Column III of the attached Table 1 (hereinafter referred to as Vitamin E preparations) may include the active ingredients listed in Column IV, Group 2 of Column V, Column VI, VIII, Group 1 or 2 of Column IX, Group 2, 3, 6, or 9 of Column X, or Group 1 or 2 of Column XI of the same Table.
- E) Preparations mainly consisting of the active ingredients listed in Column IV of the attached Table 1 (hereinafter referred to as Vitamin B₁ preparations) may include the active ingredients listed in Column III, V, VI, VII, Group 1 or 2 of Column IX, Group 1, 6, or 9 of Column X, or Group 1 of Column XI of the same Table.
- F) Preparations mainly consisting of the active ingredients listed in Column V of the attached Table 1 (hereinafter referred to as Vitamin B₂ preparations) may include the active ingredients listed in Column IV, VI, VIII, IX, Group 4, 5, 6, or 8 of Column X, or Group 3 of Column XI of the same Table.
- G) Preparations mainly consisting of the active ingredients listed in Column VI of the attached Table 1 (hereinafter referred to as Vitamin B₆ preparations) may include the active ingredients listed in Column III, IV, V, VII, VIII, IX, Group 4, 5, 6, or 8 of Column X, or Group 3 of Column XI of the same Table.
- H) Preparations mainly consisting of the active ingredients listed in Column VIII of the attached Table 1 (hereinafter referred to as Vitamin C preparations) may include the active ingredients listed in Column III, V, VI, IX, or Group 4, 5, or 8 of Column X of the same Table.
- I) Preparations mainly consisting of the active ingredients in Group 1 of Column I and Column II of the attached Table 1 (hereinafter referred to as Vitamin A and D preparations) may include the active ingredients listed in Column III, IV, VIII, or Group 7 of Column X of the same Table.
- J) Preparations mainly consisting of the active ingredients listed in Columns V and VI of the attached Table 1 (hereinafter referred to as Vitamin B₂ and B₆ preparations) may include the

active ingredients listed in Column VIII, IX, Group 4, 5, or 8 of Column X, or Group 3 of Column XI of the same Table.

K) Preparations mainly consisting of the active ingredients listed in Columns III and VIII of the attached Table 1 (hereinafter referred to as Vitamin E and C preparations) may include the active ingredients listed in Group 2 of Column V, Column VI, Group 1 or 2 of Column IX, or Group 3 of Column X of the same Table.

L) Preparations mainly consisting of the active ingredients listed in Columns IV, VI, and VII of the attached Table 1 (hereinafter referred to as Vitamin B₁, B₆ and B₁₂ preparations) may include the active ingredients listed in Column III, Group 1 or 2 of Column IX, or Group 6 of Column X of the same Table.

M) If active ingredients from Column II, III, IV, V, VI, or VII of the attached Table 1 are combined, only one active ingredient from each column may be used.

N) If active ingredients from Column VIII of the attached Table 1 are combined, no more than 2 active ingredients from the column may be used.

O) If active ingredients from Column I, IX, or Group 4 or 8 of Column X of the attached Table 1 are combined, only one active ingredient from each column or group may be used.

(2)Quantities of active ingredients

A) When the active ingredients in the attached Table 1 are used as the main ingredients of vitamin preparations, the maximum daily dose, minimum daily dose, maximum single dose, and minimum single dose shall be those given in Section A of the Table.

B) When the active ingredients in the attached Table 1 in vitamin preparations are used as active ingredients other than the main vitamins, the maximum daily dose, minimum daily dose, and maximum single dose shall be those given in Section B of the Table.

C) When 2 of the active ingredients in Column I or VIII of the attached Table 1 are combined or when 2 or more of the active ingredients in Group 7 of Column X are combined, the sum of the values obtained by dividing the amounts of each active ingredient used by their respective maximum daily dose shall not exceed one, or the sum of the values obtained by dividing the amounts of each active ingredient used by their respective minimum daily dose should be at least one.

(3)Dosage forms

The dosage forms of vitamin preparations shall be capsules, granules, pills, powders, electuaries, tablets, jelly type drops, or oral liquids.

(4)Dosage and administration

A) In principle, the dosage of vitamin preparations shall not exceed 3 doses a day.

B) Dosage and administration suggesting that the preparations may be given to infants less than 3 months of age are not permitted.

C) Hard capsules and soft capsules, pills or tablets over 6 mm in diameter intended to be taken by children less than 5 years old are not permitted.

Soft capsules, pills or tablets not more than 6 mm in diameter intended to be taken by children less than 3 years old are not permitted.

E) The maximum and minimum daily and single doses for people under 15 years of age shall be calculated by multiplying the maximum and minimum daily and single doses shown in the attached Table 1 by the values specified in the Coefficient column for the corresponding age ranges in the attached Table 2.

(5) Indications

The indications of vitamin preparations should be within the scope of the attached Table 3.

Attached Table 1

		·····	т		r		
				A,	1	3	_
Cla icat	ssif ion	Active ingredient	Maximum daily dose	Minimum daily dose	Maximum daily dose	Minimum daily dose	Remarks
		Retinol acetate	4,000I.U.	2,000I.U.	2,000I.U.	500I.U.	as vitamin A
П	Group	Retinol palmitate	4,000I.U.	2,000I.U.	2,000I.U.	500I.U.	as vitamin A
Column I	S.	Vitamin A oil	4,000I.U.	2,000I.U.	2,000I.U.	500I.U.	as vitamin A
oln	63	Cod liver oil	4,000I.U.	2,000I.U.	2,000I.U.	500I.U.	as vitamin A
Ç	Group	Strong cod liver oil	4,000I.U.	2,000I.U.	2,000I.U.	500I.U.	as vitamin A
II a	۲.	Ergocalciferol	400I.U.	200I.U.	2001.U.	50I.U.	as vitamin D
Column II		Cholecalciferol	400I.U.	200I.U.	200I.U.	50I,U.	as vitamin D
		d-a-Tocopherol succinate	300mg (100mg)	100mg (50mg)	100mg	10mg	
	``	dl-a-Tocopherol succinate	300mg (100mg)	100mg (50mg)	100mg	10mg	
Ħ		dl a-Tocopherol calcium succinate	300mg (100mg)	100mg (50mg)	100mg	10mg	as dl'a tocopherol succinate
Column III		d-α-Tocopherol acetate	300mg (100mg)	100mg (50mg)	100mg	10mg	
S		dl u Tocopherol acetate	300mg (100mg)	100mg (50mg)	100mg	10mg	
		d α -Tocopherol	300mg (100mg)	100mg (50mg)	100mg	10mg	
		dl a Tocopherol	300mg (100mg)	100mg (50mg)	100mg	10mg	
		Thiamine hydrochloride	30mg (10mg)	1mg (1mg)	25mg (10mg)	lmg	
		Thiamine nitrate	30mg (10mg)	1mg (1mg)	25mg (10mg)	1mg	
	Group 1	Bisthiamine nitrate	30mg (10mg)	1mg (1mg)	25mg (10mg)	lmg	as thiamine disulfide
	Ğ	Thiamine disulfide	30mg (10mg)	lmg (1mg)	25mg (10mg)	1mg	
N		Thiamine dicetylsulfate	30mg (10mg)	lmg (lmg)	25mg (10mg)	1mg	as thiamine nitrate or thiamine hydrochloride
Column IV		Dicethiamine hydrochloride	100mg	5mg	25mg	lmg	as thiamine hydrochloride
ŏ	1 .	Fursultiamine hydrochloride	100mg	5mg	25mg	lmg	as fursultiamine
	•	Octotiamine	100mg	5mg	25mg	lmg	
	63	Cycothiamine	100mg	5mg	25mg	lmg	
	dn	Bisibuthiamine	100mg	5mg	25mg	lmg	
	Group	Bisbentiamine	100mg	5mg	25mg	1mg	as thiamine hydrochloride
		Fursultiamine	100mg	5mg	25mg	1mg	_
		Prosultiamine	100mg	5mg	25mg	1mg	· · · · · · · · · · · · · · · · · · ·
		Benfotiamine	100mg	5mg	25mg	lmg	as thiamine hydrochloride
	_	Flavin adenine dinucleotide sodium	45mg	5mg	12mg	2mg	as flavin adenine dinucleotide
Column V	Group	Riboflavin	30mg	2mg	12mg	2mg	
l m	١	Riboflavin sodium phosphate	30mg	2mg	12mg	2mg	as riboflavin
රී	Group 2	Riboflavin butyrate	20mg	5mg	12mg	2mg	

F		Pyridoxine hydrochloride	100mg	10mg	50mg	5mg	
N G					33.28	Jung	1
Column VI		Pyridoxal phosphate	60mg	10mg	50mg	5mg	
		Hydroxocobalamin hydrochloride	1,500µg	60µg	60µg	lug	as hydroxocobalamin
Column VII		Hydroxocobalamin acetate	1,500pg	60µg	60µg	lµg	as hydroxocobalamin
Coln		Cyanocobalamin	1,500µg	60µg	60µg	1µg	
		Hydroxocobalamin	1,500µg	60µg	60µg	1µg	
MIII		Ascorbic acid	2,000mg	50mg	500mg	50mg	
Column VIII		Calcium ascorbate	2,000mg	50mg	500mg	50mg	as ascorbic acid
S		Sodium ascorbate	.2,000mg	50mg	500mg	50mg	as ascorbic acid
	ıp 3	Nicotinic acid		/	60mg	12mg	
	Group	Nicotinamide			60mg	12mg	-
Column IX	22	Panthenol			30mg	5mg	-
H H	Group	Calcium pantothenate	· /	<i>/</i> .	30mg	5mg	
\[\bar{1}{2}\]	G	Sodium pantothenate		. •	30mg	5mg	
1	Group 3	Biotin		•	500µg	10µg	
	Group 1	Mixture of potassium aspartate and magnesium aspartate (equal mixture)			400mg	200mg	
	Group 2	Inositol hexanicotinate			400mg	80mg	
	Group 3	Ursodeoxycholic acid			60mg	10mg	
	Group 4	L-Cysteine hydrochloride		/ 1	160mg	30mg	
		L-Cysteine		/ [160mg	30mg	
mn X	Group 5	Orotic acid	/	/ [200mg	60mg	
Colu	Group 6	y Oryzanol			10mg	5mg	
		Calcium glycerophosphate		, [300mg	30mg	as calcium
	[~	Calcium gluconate	/	-	300mg	30mg	as calcium
	a l	Precipitated calcium carbonate Calcium lactate		-	300mg	30mg	as calcium
.	Group	Anhydrous dibasic calcium	- /		300mg	. 30mg	as calcium
		phosphate	, / -		300mg	30mg	as calcium
	_	Dibasic calcium phosphate	/	.]_	300mg	30mg	as calcium
	Group 8	Glucuronolactone	/-		1,000mg	200mg	
		Glucuronamide	<i></i>		1,000mg	200mg	
	Group 9	Sodium chondroitin sulfate	/		900mg	180mg	

	Group 1	Processed Gar	lic Bulb	200mg	20mg	
XI	Group 2	Ginseng	Extract (Crude drug conversion value)	.3g	0.6g	
Column	පි		Powder	1.5g	0.3g	
ပိ	Group 3	Coix seeds	Extract (Crude drug conversion value)	10g	1g	-
	ď		Powder	3g _,	0.3g	

(Note) The figures in parentheses in the maximum daily dose or minimum daily dose columns indicate the maximum or minimum single dose, respectively.

Attached Table 2

Age	Co	efficient
15 years old and over	1	(1)
11 years old Under 15	2/3	(2/3)
7 years old-Under 11	1/2	(2/3)
3 years old-Under 7	1/3	(1/2)
1 year old-Under 3	1/4	(1/2)
6 months Under 1	1/5	(1/2)
3 months Under 6 months	1/6	(1/2)

(Note) The coefficients in parentheses are used for the active ingredients in Columns I and II for vitamins A, D, and A and D preparations.

Attached Table 3

	Preparations	Indications		
preparations	Preparations with Group 1 ingredients	Relief of the following symptoms: dryness of the eyes Night blindness (nyctalopia) Supplementation of Vitamin A in the following cases: during pregnancy and lactation, decreased strength during and after illness, and for growing children		
Vitamin A pr	Preparations with Group 2 ingredients	Relief of the following symptoms: dryness of the eyes Night blindness (nyctalopia) Supplementation of Vitamin A and D in the following cases: during pregnancy and lactation, decreased strength during and after illness, and for growing children and the elderly		
Vitamin D preparations		To treat bone and teeth developmental defects Prevention of rickets Supplementation of Vitamin D in the following cases: during pregnancy and lactation, and for growing children and the elderly		

Preparations	Indications
Vitamin E preparations	Relief of the following symptoms due to peripheral circulatory disturbances: stiffness in the shoulder and neck, numbness/chills in the limbs and chilblains
	Relief of the following symptoms in the climacterium: stiffness in the shoulder and neck, chills, numbness in the limbs and hot flashes, irregular menstruation
	(A physician or pharmacist should be consulted if there is no improvement after about one month of administration)
	Supplementation of Vitamin E in the following case: for the elderly
Vitamin B ₁ preparations	Relief of the following symptoms: neuralgia, muscle and joint pain (lumbago, stiff shoulder, frozen shoulder), numbness in the limbs, constipation, and eye strain
proparations .	Beriberi
	(A physician or pharmacist should be consulted if there is no improvement after about one month of administration)
	Supplementation of Vitamin B ₁ in the following cases: physical fatigue, during
	pregnancy and lactation, decreased strength during and after illness
Vitamin B ₂	Relief of the following symptoms: angular stomatitis, canker sores, stomatitis,
preparations	glossitis, eczema, dermatitis, rash, sores, acne, skin roughness, rosacea, congestion of the eye, and itchy eyes
•	(A physician or pharmacist should be consulted if there is no improvement after about one month of administration)
	Supplementation of Vitamin B ₂ in the following cases: physical fatigue, during pregnancy and lactation, and decreased strength during and after illness
Vitamin B ₆	Relief of the following symptoms: angular stomatitis, canker sores, stomatitis,
preparations	glossitis, eczema, dermatitis, rash, sores, acne, skin roughness, and numbness in the limbs
	(A physician or pharmacist should be consulted if there is no improvement after about one month of administration)
	Supplementation of Vitamin Be in the following cases: during pregnancy and
	lactation, and decreased strength during and after illness
Vitamin C	Relief of the following symptoms: spots, freckles, and pigmentation due to
preparations	sunlight/rash Proportion of blooding in the following and the following sunlight for the following sun
	Prevention of bleeding in the following cases: bleeding of the gums and nose bleeds (A physician, pharmacist, or dentist should be consulted if there is no
	improvement after about one month of administration) Supplementation of Vitamin C in the following cases: physical fatigue, during
	pregnancy and lactation, decreased strength during and after illness, and for the
	elderly
Vitamin A and D	Relief of the following symptoms: dryness of the eyes
preparations	Bone and teeth developmental defects
•	Night blindness (nyctalopia)
	Prevention of rickets
	Supplementation of Vitamin A and D in the following cases: during pregnancy and
	lactation, decreased strength during and after illness, and for growing children and the elderly
Vitamin B ₂ and B ₆	Relief of the following symptoms: angular stomatitis, canker sores, stomatitis,
preparations	glossitis, eczema, dermatitis, rash, sores, acne, and skin roughness
	(A physician or pharmacist should be consulted if there is no improvement after
	about one month of administration) Supplementation of Vitamin B ₂ and B ₆ in the following cases: physical fatigue,
	during pregnancy and lactation, and decreased strength during and after illness

Preparations	Indications
Vitamin E and C preparations	Relief of the following symptoms due to peripheral circulatory disturbances: stiffness in the shoulder and neck, numbness/chills in the limbs and chilblains Relief of the following symptoms: spots, freckles, and pigmentation due to sunlight/rash Prevention of bleeding in the following cases: bleeding of the gums and nose bleeds (A physician, pharmacist, or dentist should be consulted if there is no improvement after about one month of administration) Supplementation of Vitamin E and C in the following cases: physical fatigue, decreased strength during and after illness, and for the elderly
Vitamin B1, B6, and	
B ₁₂ preparations	shoulder, frozen shoulder), numbness in the limbs, and eye strain
- · -	(A physician or pharmacist should be consulted if there is no improvement after
	about one month of administration)
1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1	Supplementation of Vitamin B ₁ , B ₆ , and B ₁₂ in the following cases: physical fatigue,
	during pregnancy and lactation, and decreased strength during and after illness

Provisional Translation from Japanese Original

Feb 1, 1988 Notification PB No.94 Final revision May 15, 1998

The Standards for Marketing Approval of Enemas

1. Scope of Enemas

The scope of preparations subject to these standards covers medicines for rectal application formulated with the intent of treating constipation.

2. Approval Standards

The approval standards for enemas are as follows.

For preparations not conforming to these standards, efficacy and safety data and reasons justifying the combination should be submitted; the preparation in question will be reviewed based on these data.

(1) Types of Active Ingredients

- (a) The types of active ingredients the may be used are those listed in Table 1 for liquid preparations and those listed in Table 2 for suppositories.
- (b) The active ingredients that must be included are those from Column I of Table 1 and Column I or II of Table 2.
- (c) The active ingredients from Column II of Table 1 can be combined with the active ingredients from Column I.
- (d) The active ingredients from Columns I and II of Table 2 may not be used in the same preparation.

(2) Quantities of Active Ingredients

- (a) The maximum and minimum single doses of the active ingredients in Tables 1 and 2 are those specified in the respective tables.
- (b) The concentration of glycerin in Column I of Table 1 for liquid preparations is 42% to 50%.
- (3) Dosage Form

The dosage forms are liquids and suppositories.

- (4) Dosage and Administration
 - (a) Liquid preparations
 - [1] When dilution is required, water should be added so that the concentration of glycerin reaches 42% to 50%.
 - [2] When no effect is obtained by intra rectal administration of a single dose of the preparation, administer the same amount again.
 - (b) Suppositories

If no effect is obtained by the insertion of a single suppository, insert 1 more. In the case of suppositories containing ingredients from Column II of Table 2, the daily dose is limited to 0.02 g.

(c) Dosages for children under 3 years of age is not approved.

- (d) For children under 12 years of age, the single dose of the active ingredients in Table 1 is that obtained by multiplying the single doses listed in the table by the coefficient for the corresponding age range in Table 3. The single dose of the active ingredients from Column I of Table 2 is that obtained by multiplying the single doses listed in the table by the coefficient in Table 4. The single dose of the active ingredients from Column II of Table 2 is that obtained by multiplying the single doses listed in the table by the coefficient in Table 5.
- (5) Indications
 The indication is limited to constipation.

Table 1

Liquids-

Column	Active	Single dose (g)		
	ingredient	Minimum	Maximum	
I	Glycerin	12	18	
II	D-Sorbitol		10	

Table 2

Suppositories

Column	Active	Single dose (g)		
	ingredient	Minimum	Maximum	
I	Glycerin	1.5	2.5	
II	Bisacodyl	0.005	0.01	

Table 3

Age	Coefficient		
12 years of age or over	1		
6 to under 12 years of age	2/3		
1 to under 6 years of age	1/3		
Under 1 year of age	1/6		

Table 4

Age	Coefficient		
12 years of age or over	1		
3 to under 12 years of age	2/3		

Table 5

Age	Coefficient		
12 years of age or over	1		
6 to under 12 years of age	1/2		
3 to under 6 years of age	1/5		

Provisional Translation from Japanese Original

Mar 28, 1989 Notification PB No.300 Final revision May 15, 1998

The Standards for Marketing Approval of Anthelmintics

1. Scope of Anthelmintics

The scope of preparations subject to these standards covers all oral preparations intended to eradicate parasites (Kampo medicine* formulas are not covered).

* Kampo medicine is traditional Japanese medicine.

2. Approval Standards

The approval standards for anthelmintics are as follows.

For preparations not conforming to these standards, efficacy and safety data and reasons justifying the combination should be submitted; the preparation in question will be reviewed based on these data.

(1) Types of Active Ingredients

(a) The types of active ingredients that may be used are shown in Table 1.

- (b) One or more of the active ingredients from Column A of Table 1 must be included.
- (c) Preparations mainly containing active ingredients from Group 1 of Column A in Table 1 may include active ingredients from Column B or C.

(d) Preparations mainly containing active ingredients from Group 2a of Column A in Table 1 may include active ingredients from Column B.

- (e) Preparations mainly containing active ingredients from Group 2b of Column A in Table 1 may include active ingredients from Group 2 of Column B, or Column D. However, the active ingredient from Group 2 of Column D may be included only when an active ingredient from Group 2 of Column B is also included.
- (f) Preparations mainly containing active ingredients from Group 3 of Column A or Group 4 of Column A in Table 1 may not include any other active ingredient.
- (g) Preparations mainly containing active ingredients from Groups 1 and 2 of Column A, those mainly containing active ingredients from Groups 1 and 3 of Column A, and those mainly containing active ingredients from Groups 1, 2, and 3 of Column A in Table 1 may also include active ingredients from Column B or C.
- (h) In the case of Columns B and C in Table 1, only 1 active ingredient from each column may be used in the preparation.
- (i) Only 1 active ingredient from Group 2 of Column A in Table 1 may be included from this group.

(2) Quantities of Active Ingredients

- (a) The maximum daily dose of each of the active ingredients in Table 1 is the amount shown in this table.
- (b) When an active ingredient from Group 1 of Column A in Table 1 is combined

with another active ingredient from Column A, or when active ingredients from Group 1 of Column B in Table 1 are combined, the lower limit of the daily dose is half of the maximum daily dose.

(c) When an active ingredient from Group 2 of Column A in Table 1 is combined with another active ingredient from Column A, the lower limit of the daily dose

is 1/4th of the maximum daily dose.

(d) When an active ingredient from Group 3 of Column A in Table 1 is combined with another active ingredient from Column A, the lower limit of the daily dose is 3/4 of the maximum daily dose.

(e) The lower limit of the daily dose of the active ingredients from Group 4 of

Column A in Table 1 is 2/5th of the maximum daily dose.

(f) The lower limit of the daily dose of the active ingredients from Group 2 of Column B, and Column D of Table 1 is 1/10th of the maximum daily dose.

g) The lower limit of the daily dose of the active ingredients from Column C of

Table 1 is 1/5th of the maximum daily dose.

- (h) When 2 or more of the active ingredients from Column A of Table 1 are combined, the lower limit of the daily dose of each active ingredient is 1/5th of the maximum daily dose, and the sum of the values obtained by dividing the amount of each active ingredient combined by its maximum daily dose must be at least half, and should not exceed 2/3.
 - However, when 2 or more of the active ingredients only from Group 3 of Column A are combined, the sum of the values obtained by dividing the amount of each active ingredient combined by its maximum daily dose should be at least 3/4 and not exceed 1.
- (i) When 2 or more of the active ingredients from Group 1 of Column D in Table 1 are combined, the sum of the values obtained by dividing the amount of each active ingredient combined by its maximum daily dose should not exceed 1.

3). Dosage Form

The dosage forms are capsules, granules, pills, powders, tablets, decoctions (only preparations mainly containing the active ingredients from Group 2b of Column A in Table 1), chocolate tablets, and oral liquids.

(4) Dosage and Administration

(a) Dose regimen

(i) Preparations mainly containing the active ingredients from Group 1 of Column A in Table 1

Take twice a day on an empty stomach, or take once before bed after a light evening meal and once on the following morning.

Do not take more than twice in succession.

(ii) Preparations mainly containing the active ingredients from Group 2a of Column A in Table 1

Take once or twice a day on an empty stomach.

Do not take more than twice in succession.

(iii) Preparations mainly containing the active ingredients from Group 2b of Column A in Table 1

Take once or twice a day on an empty stomach.

- (iv) Preparations mainly containing the active ingredients from Group 3 of Column A in Table 1
 - [1] For eradication of ascarids

Take once or twice a day on an empty stomach for 1 to 2 days.

Do not take for more than 2 successive days.

[2] For eradication of oxyurids

Take once or twice a day on an empty stomach for 1 week. Do not take for more than 7 successive days.

(v) Preparations mainly containing the active ingredients from Group 4 of Column A in Table 1

Take once a day.

Do not take more than twice in succession.

(vi) Preparations mainly containing the active ingredients from Groups 1 and 2 of Column A, those mainly containing the active ingredients from Groups 1 and 3 of Column A, and those mainly containing the active ingredients from Groups 1, 2, and 3 of Column A in Table 1

Take once or twice a day on an empty stomach, or take once before bed after a light evening meal and once on the following morning. Do not take more than twice in succession.

(b) For decoctions, the method of preparation at the time of use should be clearly described.

(c) Dosage for infants younger than 3 months of age is not approved.

(d) For capsules, and pills and tablets larger than 6 mm in diameter, dosage for children under 5 years of age is not approved.

(e) For pills and tablets, dosage for infants younger than 3 years of age is not approved, even if the diameter is less than 6 mm.

(f) The maximum daily doses for children under 15 years of age are the amounts obtained by multiplying the maximum daily dose in Table 1 by the coefficients for the respective age groups shown in Table 2.

(5) Indications

(i) Preparations mainly containing the active ingredients from Group 3 of Column A in Table 1

Eradication of ascarids and oxyurids

(ii) Preparations mainly containing the active ingredients from Group 4 of Column A in Table 1

Eradication of oxyurids (iii) Other preparations

Eradication of ascarids

Table 1

	le 1					
Clas	ssification Active ingredient Maximum daily dose			Remarks		
	Gro	սր 1	Santonin	200 mg		
		a	Kainic acid	20 mg		
		b	Digenea	Powder	Extract	
	Group 2		• •		(converted to	
	où.				the crude	
	Q.				drug	*
⋖					amount)	
Column A					10 g	-
[2]	Gro	up 3		For	For oxyurids	
ပြ	•			ascarids		
			Piperazine adipate	4000 mg	2000 mg	As piperazine hexahydrate
			Piperazine citrate	4000 mg	2000 mg	As piperazine hexahydrate
			Piperazine hexahydrate	4000 mg	2000 mg	-
		• '	Piperazine malate	4000 mg	2000 mg	As piperazine hexahydrate
1 1			Piperazine phosphate	4000 mg	2000 mg	As piperazine hexahydrate
	Gro	up 4	Pyrvinium pamoate	250	mg	As pyrvinium base
	Group 1		Sulfur	1000		
		•	Magnesium oxide	2000	mg	
			Dioctyl sodium		mg	
			sulfosuccinate			
			Bisacodyl	20	mg	
Column B	Group 2			Powder	Extract	
H H.		: * -			(converted to	
- lo	,		•		the crude	
					drug	· · · · ·
					amount)	
			Aloes	0.75 g	0.75 g	
			Senna Leaf	1.5 g	6 g	
			Rhubarb	. 3 g	4 g	
C	olumi	ı C	Aminoethylsulfonic acid	2000		· .
1			Bile extract (powder)	500 mg		
			Bile powder	1500 mg		
			Dehydrocholic acid	500 mg		
	Gro	up 1		Powder	Extract	
		_	•		(converted to	-
			·	,	the crude	
Ω		,			drug	
🖁					amount)	
Column D			Melia Bark	<u> </u>	10 g	· .
			Japanese Zanthoxylum		3 g	
			Peel	`		
	<u> </u>		Rangoon Creeper Fruit	_	3 g	
	Gro	up 2	Glycyrrhiza	_	3.3 g	Į.

Table 2

Age group	Coefficient
15 years of age and over	1
11 to under 15 years of age	2/3
8 to under 11 years of age	1/2
5 to under 8 years of age	1/3
3 to under 5 years of age	1/4
1 to under 3 years of age	1/5
3 months to under 1 year of age	1/7

Provisional Translation from Japanese Original

Feb 1, 1991 Notification PB No.109 Final revision Jan 19, 2012

The Standards for Marketing Approval of Nasal Drops for Rhinitis

Scope of Nasal Drops for Rhinitis

The scope of preparations subject to these standards covers intranasal medicines intended for the relief of symptoms of rhinitis.

2. Approval Standards

The approval standards for nasal drops for rhinitis are as follows.

For preparations not conforming to these standards, efficacy and safety data and reasons justifying the combination should be submitted; the preparation in question will be reviewed based on these data.

- (1) Types of Active Ingredients
 - a. The types of active ingredients that may be used are shown in Table 1.
 - b. The active ingredients that must be included are those from Column I of Table 1.
 - c. Active ingredients from different columns of Table 1 may be combined with each other.
 - d. When the active ingredients from Column I, II, III, or IV of Table 1 are combined, only 1 ingredient per column is permitted.
- (2) Quantities of Active Ingredients
 - a. The maximum concentration of each of the active ingredients is shown in Table 1.
 - b. The minimum concentration of each of the active ingredients from Column I of Table 1 is half of the respective maximum concentrations, and that of the active ingredients from the other columns is 1/5th of the respective maximum concentrations.
- (3) Dosage Form

The dosage forms are intranasally applied liquid preparations.

- (4) Dosage and Administration
 - a. Preparations are to be applied intranasally not more than 6 times a day. The application method and intervals must be clearly indicated. The application interval is to be at least 3 hours.
 - b. Dosages for infants under 2 years of age are not approved.
 - c. The maximum concentrations for children under 7 years of age are half of the maximum concentration shown in Table 1.
- (5) Indications

The indications are to be within the following scope: relief of the following

symptoms due to acute rhinitis, allergic rhinitis or sinusitis; stuffy nose, runny nose (excessive nasal discharge), sneezing, dull headache (heaviness in head).

(6) Packaging Units

The maximum volume of containers for liquids is limited to 30 mL.

e l		
Classification	Active ingredient	Maximum concentration (%)
Column I	Epinephrine	0.01
45	Ephedrine hydrochloride	0.5
	Tetrahydrozoline hydrochloride	0.1
	Naphazoline hydrochloride	0.05
	Phenylephrine hydrochloride	0.5
	dl-Methylephedrine hydrochloride	0.5
$\mathcal{F}_{\mathcal{A}} = \frac{1}{2} \mathcal{F}_{\mathcal{A}}$	Tetrahydrozoline nitrate	0.1
	Naphazoline nitrate	0.05
Column II	Iproheptine hydrochloride	0.5
	Diphenhydramine hydrochloride	0.2
•	Diphenhydramine	0.2
	Chlorpheniramine maleate	0.5
Column III	Acrinol	0.05
٠,	Cetylpyridinium chloride	0.05
	Benzalkonium chloride	0.02
	Benzethonium chloride	0.02
Column IV	Lidocaine hydrochloride	0.5
	Lidocaine	0.5
Column V	Dipotassium glycyrrhizinate	0.3
	Methyl salicylate	0.05

Provisional Translation from Japanese Original

Mar 22, 1995 Notification PFSB No.277

The Standards for Marketing Approval of Antihemorrhoids (External Preparations)

1. Scope of Antihemorrhoids (External Preparations)

The scope of preparations subject to these standards covers medicines intended for the relief of hemorrhoidal symptoms in the anus and rectum (Kampo medicine* formulas and non-Kampo crude drug remedies consisting of crude drug only are not covered).

*Kampo medicine is traditional Japanese medicine.

2. Approval Standards

The approval standards for antihemorrhoids (external preparations) are as follows. For preparations deviating from these standards, efficacy and safety data and reasons justifying the combination should be submitted, and the preparation in question will be reviewed based on these data.

(1) Types of Active Ingredients

- a. The types of active ingredients that may be combined are listed in Table 1.
- b. Active ingredients that must be included are those from Column I in Table 1.
- c. Active ingredients in different columns in Table 1 may be mutually combined, unless otherwise specified elsewhere.
- d. When active ingredients from Column II, III, V, or VI are to be combined, only 1 ingredient from each column is allowed.
- e. When active ingredients from Column VIII or IX are to be combined, only 1 ingredient from the same group is allowed.
- f. It is permissible to use 2 of the active ingredients from Group 1 in Column I of Table 1, but the combination of dibucaine hydrochloride with dibucaine and the combination of lidocaine hydrochloride with lidocaine are not permitted.
- g. In Column VII of Table 1, the combination of allantoin with aluminum chlorohydroxy allantoinate, that of dried aluminum potassium sulfate with aluminum potassium sulfate, and that of purified yolk lecithin with egg yolk oil is not permitted.

(2) Quantities of Active Ingredients

- a. The maximum concentration of each of the active ingredients listed in Table 1 is given in "A" for ointments to be applied by rubbing or external liquids. The maximum single dose of each of the active ingredients is given in "B" for ointments to be applied by an applicator and for suppositories.
- b. The minimum concentration or the lowest single dose of each of the active ingredients listed in the individual columns (except for the ingredients of Group 2 in Columns VII and IX) of Table 1 is 1/5th of the corresponding maximum concentration or the maximum single dose. However, if 1 or more of the active

ingredients from Column I is used, the concentration of at least 1 active ingredient must be at least half of the maximum concentration or the maximum single dose.

c. The minimum concentration or the lowest single dose of each of the active ingredients listed in Group 2 of Columns VII and IX is 1/10th of the corresponding maximum concentration or maximum single dose.

d. When 2 active ingredients listed in Group 1 of Column I in Table 1 are combined, the sum of the values obtained by dividing the individual concentrations or doses by their respective maximum concentration or maximum single dose must not exceed 1.

(3) Dosage Form

The dosage forms should be suppositories (including soft capsules), ointments, and external liquids (including aerosols).

(4) Dosage and Administration

- a Ointments to be applied by rubbing and external liquids
 The preparations should be applied to the anal area up to 3 times a day at
 maximum. For external liquids, the method of application should be indicated
 clearly.
- b. Ointments to be applied by an applicator and suppositories
 - [1] The preparations should be applied to the anal area or the rectum 1 dose at a time, up to 3 times a day, at maximum.
 - [2] For ointments to be applied by an applicator, the method of application should be indicated clearly.
 - [3] Dosage for children younger than 7 years of age is not approved.
 - [4] The maximum single dose for those 7 to <15 years of age is half of the maximum single dose given in "B" of Table 1.

(5) Indications

The scope of indications is "Relief of pain, itching, swelling, bleeding, and erosion associated with bleeding piles (ripped piles)/blind piles, and disinfection. The indications of "erosion" and "disinfection" should be limited to ointments to be applied by rubbing and external liquids. The indications given in the upper column of the following table should be limited to cases in which 1 of the active ingredients from a group or column in the lower column of the following table is used at an amount not less than half of the maximum concentration or the maximum single dose as specified in Table 1.

Upper column	Lower column	
Itching	Group 1 of Column I, III, VI	
Swelling and bleeding	Column II, III, IV	
Erosion	Column IV	
Disinfection	Group 1 of Column V	

Table 1 Classification		Active ingredient	A Maximum concentration (%)	B Maximum single dose (mg)
Column I Group 1		Ethyl aminobenzoate	10	200
		Dibucaine hydrochloride	0.5	10
		p-Butylaminobenzoyl diethylaminoethyl hydrochloride	0.1	2
		Procaine hydrochloride	2	40
		Meprylcaine hydrochloride	0.5	10
		Lidocaine hydrochloride	3	60
-		Oxypolyethoxydodecane	3	60
		Dibucaine	0.5	10
		Mepivacaine	0.75	15
		Lidocaine	3	60
	Group 2	Scopolia Extract	5	100
Column II	-	Epinephrine solution	0.001 (as epinephrine)	_
-		Ephedrine hydrochloride	1	20
	-	Tetrahydrozoline hydrochloride	0.05	1
		Naphazoline hydrochloride	0.05	1
•		Phenylephrine hydrochloride	0.25	5
· · · · · · · · · · · · · · · · · · ·		dl-Methylephedrine hydrochloride	0.5	10
Column III		Hydrocortisone acetate	0.5	5
•		Prednisolone acetate	0.1	1
,		Hydrocortisone	0.5	5
•		Prednisolone	0.1	1
Column IV		Zinc oxide	20	400
		Tannic acid	5	100
Column V	Group 1	Acrinol	0.2	4
		Alkyl polyaminoethylglycine	0.2	4
		Isopropylmethylphenol	0.1	2
÷		Cetylpyridinium chloride	0.2	4
•		Dequalinium chloride	0.1	2
× 2		Berberine chloride	1.5	30
		Benzalkonium chloride	0.1	2
		Chlorhexidine hydrochloride	0.5	10
	· · ·	Chlorhexidine gluconate solution	1	· ·
		Cetrimide	0.125	2.5
•		Resorcin	2	40
	Group 2	Sulfadiazine	5	100
		Sulfisomidine	5	100
	÷	Sulfisomidine sodium	5	100
		Homosulfamine	5	100
Column	Group 1	Diphenylpyraline hydrochloride	0.1	2
VI		Diphenhydramine hydrochloride	1	20
		Diphenhydramine	1	20
:		Chorpheniramine maleate	0.2	4
	Group 2	Crotamiton	5	100

Column	Group 1	Allantoin	1		20	
VII		Aluminium chlorhydroxy allantoinate	1 .		20	
		Ichthammol	10		200	
		Lysozyme chloride	1.5 (potency)	30 (potency))
		Dried aluminum potassium sulfate	1.1		22	
		Glycyrrhetinic acid	1.5		30	
	,	1,4-Dimethyl-7-isopropylazulene	0.04		0.8	
		Purified yolk lecithin	5		100	
		Egg yolk oil	5		100	
		Aluminum potassium sulfate	2		40	
	Group 2	·	Extract (converted	Powder	Extract (converted	Powder
			to crude		to crude	
	- '		drug .		drug	
		Lithospermum root	amount)	2.5	amount) 50	50
		Horse Chestnut Seed	25	2.0	500	
1.00.		Witch hazel leaf	25		500	
		Processed Garlic Bulb	1	<u> </u>	20	l. <u></u>
Column	Group 1	Cod liver oil	120,000 I.U.	/100 g (as	2,400 I.U.	
VIII	Oroup 1	Cou iivoi oii	vitamin A)	. 100 B (00	(as vitamin A)	
		Strong cod liver oil	120,000 I.U.	/100 g (as	2,400 I.U.	
			vitamin A)		(as vitamin A)	
,		Retinol palmitate	120,000 I.U. vitamin A)	/100 g (as	2,400 I.U. (as vitamin A)	
		Vitamin A oil	120,000 I.U.	/100 g (as	2,400 I.U.	
		,	vitamin A)	, 200 B (40	(as vitamin A)	
	Group 2	Tocopherol acetate	3		60	
· -	-	Tocopherol	3.		60	
Column IX	Group 1	d-Camphor	1 1 0.75		20	
		dl-Camphor			20	
	Group 2	Mentha Oil			15	
		I-Menthol	0.5		10	-
٠.		dl-Menthol	0.5		10	
	Group 3	Eucalyptus Oil	0.5	-*	10	

Provisional Translation from Japanese Original

May 15, 1998 Notification PSB No.447

The Standards Marketing Approval of Athlete's Foot and Ringworm Remedies

1 Scope of Athlete's Foot and Ringworm Remedies

The scope of preparations subject to these standards covers external medicines intended for the relief of symptoms associated with athlete's foot and ringworm Kampo medicine* formulas and non-Kampo crude drug remedies consisting of crude drug only are not covered).

*Kampo medicine is traditional Japanese medicine.

2 Approval Standards

The approval standards for athlete's foot and ringworm remedies are as follows. For preparations deviating from these standards, efficacy and safety data and reasons justifying the combination should be submitted; the preparation in question will be reviewed based on these data.

(1) Types of Active Ingredients

- a. The types of active ingredients that may be combined are listed in Table 1.
- b. At least 1 of the active ingredients from either Column I (apart from the ingredients in Groups 12 and 13) or Column II of Table 1 must be combined.
- Active ingredients in different columns listed in Table 1 may be mutually combined.
- d. When active ingredients from Column V of Table 1 are to be combined with other ingredients in the same Column, the use of only 1 ingredient is allowed.
- e. Up to 3 active ingredients from Column I of Table 1 may be used. However, with the exception of undecylenic acid and zinc undecylenate in Group 1, the use of only 1 ingredient from each group is allowed. Active ingredients marked with "Δ" must not be combined with the other ingredients in this column.
- f. When active ingredients from Group 1 of Column III or Group 1 of Column IV listed in Table 1 are to be combined, the use of only 1 ingredient from the same group is allowed.
- g. Up to 3 active ingredients from Group 2 of Column III listed in Table 1 may be used. However, acetic acid should not be combined with the other ingredients in this group.
- h. In Column VI, the combination of allantoin with aldioxa and the combination of glycyrrhizinic acid or its salts with glycyrrhetinic acid are not permitted. In Column VII, the combination of d-camphor with dl-camphor and the combination of mentha oil with dl-menthol and l-menthol are not permitted.
- (2) Quantities of Active Ingredients
 - a. The maximum concentration of each of the active ingredients is shown in Table 1.
 - b. The minimum concentration of individual active ingredients listed in Column I (except for Groups 12 and 13) and Column II of Table 1 is 1/5th of the maximum

concentration (for ingredients with a concentration in parentheses, the minimum concentration is 1/5th of the one in the parentheses). In this case, the concentration of 1 or more ingredients must be at least half of the specified maximum concentration (for ingredients with concentrations in parentheses, the minimum concentration must be the one provided in parentheses).

- c. The minimum concentration of individual active ingredients listed in Groups 12 and 13 of Column I and those listed in Columns III, IV, V, VI, VII, VIII, and IX of Table 1 is 1/10th of the maximum concentration. However, in the case of benzalkonium chloride in Group 1 of Column III, the concentration must be as listed in the maximum concentration column.
- (3) Dosage Form The dosage forms are aerosols, ointments, external liquids, and external powders.
- (4) Dosage and Administration Preparations should be applied to the skin surface several times a day. The method of application should be clearly indicated.
- (5) Indications

 The indications are to be within the scope of "athlete's foot, jock itch, and ringworm."

r	ble 1	the state of the s	
Classification		Active ingredient	Maximum concentration (%)
I I	Group 1	Undecylenic acid	10
Column		Zinc undecylenate	20
්ටී		Δ Phenyl-11-iode-10-undecynoate	0.5
	Group 2	Δ Exalamide	5
	Group 3	△ Clotrimazole	1
		Δ Econazole nitrate	1
		Δ Miconazole nitrate	1
		△ Tioconazole	1
	Group 4	△ Zinc diethyldithiocarbamate	25
	Group 5	△ Ciclopirox olamine	1
	Group 6	Δ Siccanin	1 (potency)
		△ Trichomycin	15,000,000 units/100 g
		△ Pyrrolnitrin	0.5 (potency)
	Group 7	Thianthol	30
	Group 8	2,3,6 Tribromphenol caproate	2
	Group 9	Trimethylcetylammonium pentachlorophenate	2
	Group 10	△ Tolciclate	1
		Tolnaftate	2
	Group 11	Δ Haloprogin	1
	Group 12	Sulfur	10
	Group 13	Hibiscus syriacus bark (converted to the crude drug amount)	10
	Group 1	Salicylic acid	10 (2)
Column II	Group 2	Zinc oxide	60 (2)
	Group 1	Acrinol	
l un	Group 1	Alkylpolyaminoethyl glycine	0.2
Column III		Berberine benzoate	1
٥			0.5
		Isopropylmethylphenol	3
		Dequalinium chloride	0.5
		Benzalkonium chloride	0.05
		Benzethonium chloride	0.5
		Chlorhexidine hydrochloride	1 -
		Chlorhexidine gluconate solution	2.5
		Dequalinium acetate	1
		Hinokitiol	0.1
		Resorcin	5
	Group 2	Benzoic acid	12
İ		Chlorobutanol	1
		Acetic acid	2
		Phenol	2
		1 1101101	l ^z

Ν	Group 1	Diphenylpyraline hydrochloride	0.2
um	,	Diphenhydramine hydrochloride	2
Column IV		Chlorpheniramine	0.5
,		Diphenhydramine salicylate	2
		Diphenylimidazole	0.2
		Diphenhydramine	1
		Chlorpheniramine maleate	0.5
	Group 2	Crotamiton	10
Col	umn V	Ethyl aminobenzoate	6
·		Dibucaine hydrochloride	0.5
	, me	Procaine hydrochloride	2
	9-	Lidocaine hydrochloride	2.5
		Oxypolyethoxydodecane	3 .
		Dibucaine	0.5
	1. 7	Lidocaine	2.5
Ħ	Group 1	Allantoin	1
пп		Aldioxa	0.2
Column VII		Ichthammol	6
		Glycyrrhizinic acid and its salts	1
		Glycyrrhetinic acid	1
		Methyl salicylate	2.5
		Dimethyl isopropylazulene	0.04
	Group 2	Lithospermum root (converted to the crude drug amount)	6
		Japanese angelica root (converted to the crude drug amount)	6
Col	umn VII	d-Camphor	4
		dl-Camphor	4
		Thymol	2.5
		Mentha oil	0.5
		dl-Menthol	3 .
		dl-Menthol	3
		d [*] Borneol	5
Col	umn VIII	Urea	10
		Diethyl phthalate	25
Col	umn IX	Aluminum hydroxychloride	10
Column 171			_t

Provisional Translation from Japanese Original

Nov 1, 2011 Notification PFSB No.1101-1

The Standards for Marketing Approval of Antipruritic and Anti-inflammatory Drugs

1. Scope of Antipruritic and Anti-inflammatory Drugs

The scope of preparations subject to these standards covers medicines mainly containing adrenocortical hormones or antihistamines for dermal application formulated with the intent of using as antipruritic and anti-inflammatory drugs.

2. Approval Standards

The approval standards for antipruritic and anti-inflammatory drugs are as follows: For antipruritic and anti-inflammatory drugs mainly containing adrenocortical hormones or antihistamines that do not conform to these standards, efficacy and safety data and reasons justifying the combination should be submitted; the preparation in question will be reviewed based on these data.

(1) Types of Active Ingredients

- a) The active ingredients that may be combined in the preparations are shown in the Table.
- b) At least 1 ingredient from either Column I or Column II of the Table must be combined.
- c) Preparations mainly containing the active ingredients from Column I of the Table may include the active ingredients from Column II, III, IV, V, VI, VII, VIII, IX, X, or XII.
- d) Preparations mainly containing the active ingredients from Column II of the Table may include the active ingredients from Column III, IV, V, VI, VII, VIII, IX, X, XI, or XII.
- e) In the case of Column I, II, IV, V, VII, VIII, or IX in the Table, only 1 active ingredient from each column may be used in a preparation. When the active ingredient from Group 1 or 2 of Column X, or Group 1 or 3 of Column XII is combined, only 1 active ingredient from each group may be used in a preparation.
- (2) Quantities of Active Ingredients
 - a) The maximum concentration of each of the active ingredients in the Table is that shown in the table.
 - b) The minimum concentration of each of the active ingredients listed in Columns II, III, V, VI, VIII, Groups 2 and 3 of Column X, Column XI, and Group 2 of Column XII is 1/5th of the maximum concentration (for ingredients with a concentration in parentheses, the minimum concentration must be the amount shown in the parentheses). However, in the case of preparations mainly containing the active ingredients from Group 1 of Column I or Group 2 of Column II, the minimum concentration of each active ingredient must be at

- least half of the maximum concentration, and in the case of preparations mainly containing the active ingredients from Group 2 of Column I or Group 1 of Column II, the concentration is fixed to the maximum concentration.
- c) The minimum concentration of each of the active ingredients listed in Column IV, VII, or IX, Group 1 of Column X, or Groups 1 and 3 of Column XII of the Table is 1/10th of the maximum concentration (for ingredients with a concentration in parentheses, the minimum concentration must be the amount shown in the parentheses).
- (3) Dosage Form

The dosage forms are liquids for external use, sprays, ointments, creams, and gels. However, for sprays, preparations mainly containing the active ingredients listed in Column I of the Table are excluded.

(4) Dosage and Administration

The preparation should be applied to the skin surface several times a day. The method of application must be clearly indicated.

(5) Indications

The indications are shown by main ingredient in the following table.

Main ingredients	Indications
Group 1 of Column I Eczema, dermatitis, miliaria, irritated skin, itching, chilblain, insurticaria	
Group 2 of Column I	Eczema, dermatitis, miliaria, irritated skin, itching, insect bites, urticaria
Column II	Eczema, dermatitis, skin sore, miliaria, irritated skin, itching, chilblain, insect bites, urticaria

Classification		Active ingredient	Maximum concentration (%)
Column I	Group 1	Cortisone acetate	0.5
	1	Dexamethasone acetate	0.025
		Dexamethasone	0.025
		Hydrocortisone acetate	0.5
	,	Hydrocortisone	0.5
		Prednisolone acetate	0.25
		Prednisolone	0.25
	Group 2	Hydrocortisone butyrate	0.05
~ · · ·		Prednisolone valerate acetate	0.15
Column II	Group 1	Isothipendyl hydrochloride	0.75
		Chlorpheniramine	0.5
		Chlorpheniramine maleate	1
		Diphenhydramine	1
	Group 2	Diphenhydramine hydrochloride	2
Column III		Crotamiton	10
Column IV		Glycyrrhizic acid and its salts	1
*		Glycyrrhetic acid	1
Column V		Glycol salicylate	2
		Methyl salicylate	5
Column VI		Allantoin	1
Column VII		Isopropyl methylphenol	0.5
		Benzalkonium chloride	0,3
		Benzethonium chloride	0.1
Column VIII	-	Calamine	8
		Zinc oxide	37 (1.5)
Column IX		Ethyl aminobenzoate	
		Oxy polyethoxy dodecane	3
		Dibucaine	0.5
•		Dibucaine hydrochloride	0.5
		Lidocaine	2
		Lidocaine hydrochloride	2
Column X	Group1	d-Camphor de Camphor	7 (0.1)
		dl-Camphor .	7 (0.1)
-	Group 2	Mentha oil	2
		df-Menthol	5 (0.1)
		I-Menthol .	5 (0.1)
Group 3		d [*] Borneol	0.3
Column XI	_	Ammonia water	15
Column XII	Group 1	Tocopherol	2 (0.1)
		Tocopherol acetate	2 (0.1)
	Group 2	Panthenol	5
	Group 3	Vitamin A oil	500,000 I.U./100 g as vitamin A
		Retinol palmitate	500,000 I.U./100 g as vitamin A