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Since this translation is prepared in July 2025 based on the Japanese text current at that time, the translation may not reflect the latest information, due to continuous revision of package inserts.

The latest Japanese text is available on PMDA website.

Kracie

Oct 2023 (1st version)

Standard Commodity Classification No. of Japan	
875200	
Approval No.	16100AMZ03510000
Date of Initial Marketing in Japan	Oct 1986

Kampo product

Kracie Ninjin'yoeito Extract Fine Granules

Storage : Store at room temperature
Shelf Life : 3 years

KB-108

EK-108

3. COMPOSITION AND PRODUCT DESCRIPTION

3.1 Composition

Brand name	Kracie Ninjin'yoeito Extract Fine Granules	
Active ingredients	The daily dose of this product (7.5 g) contains 6,700 mg of Ninjin'yoeito extract powder, derived from a mixture of the following crude drugs.	
	JP Ginseng	3.0 g
	JP Japanese Angelica Root	4.0 g
	JP Peony Root	2.0 g
	JP Rehmannia Root	4.0 g
	JP Atractylodes Rhizome	4.0 g
	JP Poria Sclerotium	4.0 g
	JP Cinnamon Bark	2.5 g
	JP Astragalus Root	1.5 g
	JP Citrus Unshiu Peel	2.0 g
	JP Polygala Root	2.0 g
	JP Schisandra Fruit	1.0 g
	JP Glycyrrhiza	1.0 g
Inactive ingredients	JP Magnesium Stearate, JP Light Anhydrous Silicic Acid, JP Microcrystalline Cellulose, Hydrated Silicon Dioxide	

(JP: The Japanese Pharmacopoeia)

3.2 Product Description

Dosage form	Fine granules
Color	Light brown to brown
Odor	Characteristic odor
Taste	Slightly bitter and sweet
ID code	KB-108 (3.75 g per sachet) EK-108 (2.5 g per sachet)

4. INDICATIONS

Loss of strength after illness, fatigue and anorexia, night sweats, cold hands and feet, anemia

6. DOSAGE AND ADMINISTRATION

The usual adult dosage for oral use is 7.5 g daily in two or three divided doses before or between meals.

The dosage may be adjusted according to the patient's age, body weight, and symptoms.

8. IMPORTANT PRECAUTIONS

8.1 When this product is used, the patient's "SHO" (constitution/symptoms) should be taken into consideration. The patient's progress should be carefully monitored, and if no improvement in symptoms or findings is observed, continuous administration should be avoided.

8.2 Since this product contains Glycyrrhiza, careful attention should be paid to the serum potassium level, blood pressure, etc. [See Sections 10.2, 11.1.1, 11.1.2.]

8.3 When this product is used in combination with other Kampo products, etc., attention should be paid to the duplication of the contained crude drugs.

9. PRECAUTIONS CONCERNING PATIENTS WITH SPECIFIC BACKGROUNDS

9.1 Patients with Complication or History of Diseases, etc.

9.1.1 Patients with an extremely weak gastrointestinal tract
Anorexia, epigastric distress, nausea, vomiting, abdominal pain, diarrhea, etc. may occur.

9.1.2 Patients with anorexia, nausea, or vomiting
These symptoms may be aggravated.

9.5 Pregnant Women

This product should be used in pregnant women or women who may possibly be pregnant only if the expected therapeutic benefits outweigh the possible risks associated with treatment.

9.6 Breast-feeding Women

Considering the therapeutic benefits and the benefits of breastfeeding, continuation or discontinuation of breastfeeding should be considered.

9.7 Pediatric Use

No clinical studies have been conducted in children.

9.8 Geriatric Use

Since the physiological functions are generally decreased in elderly patients, careful supervision is recommended; measures such as reducing the dose may be considered.

10. INTERACTIONS

10.2 Precautions for Co-administration (This drug should be administered with caution when co-administered with the following.)

Drugs	Signs, Symptoms, and Treatment	Mechanism and Risk Factors
Glycyrrhiza-containing preparations Shakuyakukanzoto Hochuekkito Yokukansan, etc. Preparations containing glycyrrhizic acid and its salts Monoammonium Glycyrrhizinate/Glycine/L-cysteine Monoammonium Glycyrrhizinate/Glycine/DL-Methionine combination tablets, etc. [See Sections 8.2, 11.1.1, 11.1.2]	Pseudoaldosteronism is likely to occur. As a result of hypokalaemia, myopathy is likely to occur.	Since glycyrrhizic acid has an accelerating effect on potassium excretion in the renal tubules, an acceleration of decrease in the serum potassium level has been suggested.

11. ADVERSE REACTIONS

The following adverse reactions may occur. Patients should be carefully monitored, and if any abnormalities are observed, appropriate measures such as discontinuation of administration should be taken.

11.1 Clinically Significant Adverse Reactions

11.1.1 Pseudoaldosteronism (frequency unknown)

Pseudoaldosteronism such as hypokalaemia, blood pressure increased, retention of sodium/body fluid, edema, and body weight gain may occur. Patients should be carefully monitored (e.g., measurement of serum potassium levels), and if any abnormalities are observed, administration should be discontinued, and appropriate measures such as administration of potassium preparations should be taken. [See Sections 8.2, 10.2]

11.1.2 Myopathy (frequency unknown)

Myopathy may occur as a result of hypokalaemia. Patients should be carefully monitored, and if any abnormalities such as feelings of weakness, muscle cramp in extremities, or paralysis are observed, administration should be discontinued and appropriate measures such as administration of potassium preparations should be taken. [See Sections 8.2, 10.2]

11.1.3 Hepatic impairment, jaundice (frequency unknown)

Hepatic impairment and/or jaundice with marked elevations of AST, ALT, Al-P, γ -GTP, etc. may occur.

11.2 Other Adverse Reactions

	Frequency unknown
Hypersensitivity	Rash, Redness, Pruritus, Urticaria, etc.
Gastrointestinal	Anorexia, Epigastric Distress, Nausea, Vomiting, Abdominal pain, Diarrhea, etc.

12. INFLUENCE ON LABORATORY TESTS

Administration of this product may cause an increase in blood AG (1,5-anhydro-D-glucitol).

15. OTHER PRECAUTIONS

15.1 Information Based on Clinical Use

Eczema or dermatitis may be aggravated.

20. PRECAUTIONS FOR HANDLING

20.1 To maintain the quality of the product, avoid moisture as much as possible and store in a cool place, away from direct sunlight.

20.2 Avoid moisture, especially after opening, and handle with care.

20.3 Since this product is made from crude drugs, the color of the product may vary.

22. PACKAGING

(KB-108)

3.75 g \times 28 packets [sachets]

3.75 g \times 168 packets [sachets]

(EK-108)

2.5 g \times 42 packets [sachets]

2.5 g \times 294 packets [sachets]

500 g [bottle]

26. MARKETING AUTHORIZATION HOLDER, etc.

26.1 Manufactured and Distributed by:

Kracie, Ltd.

20-20, Kaigan 3-Chome, Minato-ku, Tokyo 108-8080, Japan

26.2 Distributed by:

Kracie Pharmaceutical, Ltd.

20-20, Kaigan 3-Chome, Minato-ku, Tokyo 108-8080, Japan