NARON m

- ♦NARON m is an antipyretic analgesic containing acetaminophen for relief of fever and pain such as headache and menstrual pain.
- ◆It contains glycine to protect the stomach membrane and vitamins B₁ and B₂, which often become depleted during a fever.
- ◆It can be used by family members that are at least 7 years old.
- ♦ NARON m contains no components that may induce drowsiness.

Indication

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

Dosage and administration

Take the following dose with cold or lukewarm water. Avoid taking the drug on an empty stomach whenever possible. Provide a dosing interval for at least 4 hours.

Adults (15 years or over): 2 tablets per dose, 3 times daily

7 to 14 years: 1 tablet per dose, up to 3 times daily

Under 7 years: Do not take

Precaution

- · Comply with the prescribed dosage and administration instructions.
- \cdot The use of the drug in children should be supervised by a parent.

How to take out the tablets

Press the convex part of the PTP sheet containing the tablet firmly with your fingertips, break the aluminum foil on the back side, and take out the tablets. (If the tablets are inadvertently taken together with the PTP sheet containing them, it will lead to unexpected accidents, such as the sharp edge sticking into the esophageal mucosa)

Ingredient

In 2 tablets

Acetaminophen 300mg

Glycine 300mg

Thiamine nitrate (vitamin B₁) 8mg

Riboflavin (vitamin B₂) 4mg

Excipients

Anhydrous silicic acid, cellulose, magnesium aluminometasilicate, hydroxypropylcellulose, sodium starch glycolate, magnesium stearate

Precaution

- No need to worry if you have yellow urine after taking the drug because it is caused by the vitamin B₂ contained in the drug.
- Red dots may appear on the tablets due to the vitamin B₂, which is contained in the drug, but this will not alter the effect.

Precautions

When not to use the product

(If you do not follow these instructions, the current symptoms may worsen or adverse reactions/incidents are more likely to occur)

- This product should not be taken in the following persons
 - Patients who have had an allergic symptom to this drug or its ingredients.
 - Patients who have experienced asthma from taking this drug, other antipyretic analgesics, cold remedies.
- This medicine should not be taken together with any of the following medicines Other antipyretics/analgesics, cold medicine, sedatives
- Do not drink alcohol before and after taking this drug

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Do not take this medicine for a long time

Consultation

• The following persons should contact a physician, dentist, pharmacist, or registered salesperson for a consultation before administration

Patients undergoing medical treatment from a physician or dentist.

Pregnant women or women suspected of being pregnant.

The elderly.

Patients who have experienced allergic symptoms associated with drugs, etc.

Persons diagnosed as having the following.

Heart disease, kidney disease, liver disease, gastric/duodenal ulcer

• If the following symptoms are observed after taking this drug, these may be adverse reactions, so immediately discontinue the use of this drug, and show this document to your physician, pharmacist, or registered salesperson for a consultation

Skin: rash/redness, itching

Gastrointestinal system: nausea/vomiting, and loss of appetite

Neuropsychiatric system: dizziness Other: excessive temperature decrease

• The following serious symptoms may occur in rare cases. In such cases, immediately seek medical aid.

Shock (anaphylaxis): Symptoms, such as itching of skin, urticaria, hoarseness, sneezing, itchy throat, breathing

difficulties, palpitations, and clouding of consciousness may occur immediately after take.

Oculomucocutaneous syndrome (stevens-johnson syndrome), toxic epidermal necrolysis, acute generalised exanthematous pustulosis: hyperthermia, ocular hyperaemia, eye discharge, lip erosion, pain throat, widespread skin rash/redness, small pimples (small pustules) on reddened skin, general malaise, anorexia, etc. may persist or suddenly worsen.

Drug-induced hypersensitivity syndrome: Symptoms such as widespread redness of the skin, generalised rash, fever, malaise, and swelling of lymph nodes (neck, armpit, groin, etc.) may occur.

Hepatic function failure: Symptoms, such as fever, itching, rash, jaundice (yellowing of skin and white of eyes), brown urine, general malaise, loss of appetite, etc. may occur.

Kidney disorders: Symptoms, such as fever, rash, reduced urinary volume, general oedema, general malaise, arthralgia (painful joints) and diarrhea, etc. may occur.

Interstitial pneumonia: Shortness and/or difficulties of breath when go upstairs or overwork, sudden dry cough and/or fever and its continuance.

Asthma: Symptoms, such as wheezing or hissing when breathing, and difficult breathing, etc. may occur.

• If the symptoms do not resolve after taking 5 to 6 doses, discontinue the use of this drug, and bring this document to a physician, dentist, pharmacist, or registered salesperson for a consultation

Precautions for storage and handling

- · Avoid direct sunlight, and store the product in a cool place with little humidity.
- · Store the product beyond the reach of children.
- Do not transfer the drug to other containers. (It may lead to misuse or quality deterioration)
- Do not use the product past the expiration date. Even before the expiration date, use the product within 6 months after it is opened. (to assure the quality)

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- •This product is a pharmaceutical product approved under a Japanese law, the Law for Ensuring the Quality, Efficacy and Safety of Drugs and Medical Devices, with a view to its sale and use in Japan.
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