PATIENT LEAFLET IN ACCORDANCE WITH THE PHARMACISTS' REGULATIONS (PREPARATIONS) – 1986

This medicine is dispensed without a doctor's prescription

Tiptipot Ferripel-3

Oral drops 50 mg/ml

Each ml (20 drops) contains: Iron (as Iron III Hydroxide Polymaltose Complex) 50 mg

Each drop contains: 2.5 mg iron. Inactive ingredients and allergens in the medicine - see section 6 "Additional information" and section 2 "Important information about some ingredients of the medicine".

Read the entire leaflet carefully before using the medicine. This leaflet contains concise information about the medicine. If you have any other questions, refer to the doctor or the pharmacist.

Use the medicine according to the instructions in the dosage section of this leaflet.

Consult the pharmacist if you have further questions. Refer to the doctor if signs of the ailment (symptoms) worsen or do not improve after three weeks.

1. What is the medicine intended for?

Prevention and treatment of anemia caused by iron deficiency.

Therapeutic class: Iron replacement preparations.

Iron is an essential constituent of hemoglobin, myoglobin and iron-containing enzymes. Iron deficiency may cause the following general symptoms: excessive fatigue, impaired mental function, irritability, anxiety, headaches, loss of appetite, weakening of the immune system, conspicuous paleness, cracks in

the corners of the mouth, dry skin, split hairs and brittle fingernails.

2. Before treatment with the medicine:

☑ Do not use this medicine:

- If you are sensitive (allergic) to the active ingredient or any of the additional components the medicine contains.
- If you have an excess of iron in your body (e.g., an excess of iron caused by a rare disease of iron build-up, which may lead to accumulation of iron in the tissues).
- If you have problems with efficient utilization of iron (e.g., when anemia is caused by insufficient utilization of iron).
- In states of anemia that are not caused by iron deficiency (e.g., anemia caused by increased hemoglobin breakdown or by vitamin B12 deficiency).

■ Special warnings regarding the use of the medicine

Before treatment with Tiptipot Ferripel-3, inform the doctor if:

- · You have an infection or a tumor.
- You have recently been treated or you might be treated in the future with injected iron preparations. Using this type of iron preparations during treatment with this medicine is not recommended
- · You have other diseases or allergies
- You have received blood transfusions, since there is a risk for an excess of iron due to receiving additional iron.

Tests and follow-up:

Before starting to use the medicine, the doctor will refer you for a blood test to check your blood iron and hemoglobin levels. If your symptoms are not caused by iron deficiency, this medicine will not be effective for you.

During treatment with this medicine the doctor will carry out periodic examinations, and may also refer you for blood tests. This referral is normal and should not concern you.

■ Drug-drug interactions:

If you are taking or have recently taken other medicines, including non-prescription medicines and dietary supplements, tell the doctor or the pharmacist.

■ Use of the medicine and food:

The medicine should be taken during or right after a meal. This medicine may be mixed with fruit or vegetable juice or with baby food. The slight change in color does not impair the effect or taste.

■ Pregnancy, breastfeeding and fertility:

No adverse effects of Tiptipot Ferripel-3 have been observed on the fetus or on women during pregnancy.

It is unknown whether iron passes into breastmilk.

If you are pregnant, trying to become pregnant or breastfeeding, consult the doctor before using the medicine.

■ Driving and operating machinery:
Tiptipot Ferripel-3 does not affect your ability to drive and/or operate machinery.

☐ Important information about some ingredients of the medicine:

The medicine contains sodium methyl hydroxybenzoate and sodium propyl hydroxybenzoate. These may cause allergic reactions, which may also occur after some time.

The medicine contains sucrose. If you have an intolerance to certain sugars, speak to your doctor before taking this medicine.

Sucrose may harm your teeth.



3. How should you use the medicine?

Check with the doctor or pharmacist if you are uncertain about the dosage and how to use the medicine.

The generally accepted dosage is: Adults and children over the age of 12 years:

For prevention of anemia: 15-30 drops per day.

For treatment of anemia: 45-60 drops per day.

Dosage for children:

For prevention of anemia: 1 mg iron/kg body weight per day.

Do not exceed a dosage of 15 mg per day in infants up to 2 years of age. In preterm infants: 2.5-5 mg (1-2 drops)/kg body weight per day.

In accordance with the Mother and Child Health Clinic recommendations, use the following table, which indicates the dosage based on the child's age.

Child's age	Required dose in drops	Required dose in mg per day
From 4 months	3 drops	About 7 mg
From 6 months	6 drops	15 mg

For treatment of anemia: up to 6 mg iron (2 drops)/kg body weight per day. The daily dose may be taken as a single dose or in divided doses.

Do not exceed the recommended dose. Method of use:

Hold the bottle vertically upside-down. A drop will promptly form at the tip of the bottle's dropper. If no drop is forming, tap the bottle gently until a drop forms. Do not shake the bottle.

Do not pour the medicine directly from the bottle into the baby's mouth! If you took an overdose or by mistake a child swallowed this medicine, go immediately to the doctor or the emergency room of the hospital and take the package of the medicine with you.

Do not take medicines in the dark!

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Check the label and the dose
every time you take the medicine. Wear
glasses if you need them.

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If you have any other questions
regarding use of the medicine, consult
the doctor or the pharmacist.

4. Side effects

abdominal pain.

As with any medicine, using Tiptipot
Ferripel-3 may cause side effects in some
users. Do not be alarmed when reading
the list of side effects. You may not
experience any of them.

Very common side effects - side effects that occur in more than one out of ten users:

Stool discoloration due to iron excretion. This phenomenon is harmless.

Common side effects - side effects that occur in 1-10 out of 100 users: Nausea, constipation, diarrhea and

Uncommon side effects - side effects that occur in 1-10 out of 1,000 users: Vomiting, teeth discoloration, gastritis,

Vomiting, teeth discoloration, gastritis, itching, rash, hives, redness in the skin, headache.

Rare side effects - side effects that occur in 1-10 out of 10,000 users: Muscle cramps and pain.

The abovementioned side effects are usually transient.

If a side effect occurs, if one of the side effects worsens, or if you suffer from a side effect not mentioned in this leaflet, consult your doctor.

Reporting side effects:

Side effects may be reported to the Ministry of Health by clicking on the link "Report side effects due to medicinal treatment" found on the Ministry of Health website homepage (www.health.gov.il), which will direct you to the online form for

reporting side effects, or by clicking on the following link:

https://sideeffects.health.gov.il/

5. How to store the medicine?

Avoid poisoning! This medicine and any other medicine must be kept in a closed place out of the reach and sight of children and/or infants to avoid poisoning. Do not induce vomiting without an explicit instruction from the doctor.

Do not use the medicine after the expiry date (exp. date) appearing on the package and the bottle.

The expiry date refers to the last day of that month.

Storage:

Store in a dark place, at a temperature lower than 25°C.

After first opening, the product may be

After first opening, the product may be used for 3 months.

6. Additional information In addition to the active ingredient, the medicine also contains:

Sucrose, Cream Essence, Sodium Methyl Hydroxybenzoate, Sodium Propyl Hydroxybenzoate, Sodium Hydroxide, Dilute Hydrochloric Acid, Purified Water What does the medicine look like and what are the contents of the package: A glass bottle containing 15 ml of darkbrown solution.

Manufacturer/license holder and address: CTS Chemical Industries Ltd., 3 Hakidma st., Kirvat Malachi.

This leaflet was reviewed and approved by the Ministry of Health in 10/2012 and has been updated in accordance with the Ministry of Health instructions in 12/2020. Registration number of the medicine in the national drug registry of the Ministry of Health: 1074628828-00





