

**PATIENT PACKAGE INSERT IN ACCORDANCE
WITH THE PHARMACISTS' REGULATIONS
(PREPARATIONS) – 1986**

The medicine is dispensed with
a doctor's prescription only

RILUTEK 50 mg Film Coated Tablets

SANOFI 

Active ingredient: Riluzole 50 mg

Inactive and allergenic ingredients in the preparation:
see section 6 “Further information” and section 2
“Important information regarding some of the
ingredients of the medicine”.

**Read the leaflet carefully in its entirety before using
the medicine.**

Keep this leaflet; you may need to read it again.

This leaflet contains concise information about the
medicine.

If you have further questions, refer to the doctor or
pharmacist.

This medicine has been prescribed for you. Do not pass
it on to others. It may harm them even if it seems to
you that their medical condition is similar.

The preparation is intended for the treatment of adults,
from the age of 18 years.

1. WHAT IS THE MEDICINE INTENDED FOR?

Treatment of ALS (amyotrophic lateral sclerosis) –
a motor neuron disease.

ALS is a type of neuronal disease that impairs transfer of
information from the nerves to the muscles, which leads
to weakness, muscle impairment and paralysis.

Therapeutic group: The substance acts on the central
nervous system.

2. BEFORE USING THE MEDICINE

Do not use the medicine if:

You are sensitive (allergic) to the active ingredient or
any of the additional ingredients contained in the
medicine (see section 6 “Further Information”).
You are suffering from a **liver disease** or from an
increase in enzymes associated with liver function
(transaminases).
You are pregnant or breastfeeding.

Special warnings regarding use of the medicine

Do not use the medicine without consulting a doctor
before commencing treatment:

- If you are suffering from liver problems: yellowing
of the skin or the white part of the eyes (jaundice),
itching all over the body, nausea or vomiting.
- If your kidneys are not functioning well.
- If you have a fever: it can be due to low white blood
cell count, which can cause an increased risk of
infection.

Children and adolescents

If you are under 18 years of age, use of Rilutek is not
recommended since there is no information in this
population.

Tests and follow-up

During the treatment period with this medicine,
perform liver function tests.

If, while taking the medicine, you suffer from a
disease accompanied by fever, inform the doctor
immediately.

Drug interactions

**If you are taking, or have recently taken, other
medicines, including non-prescription medicines
and nutritional supplements, tell the doctor or
pharmacist.** Especially if you are taking:

CYP1A2 enzyme inhibitors such as: caffeine, diclofenac,
diazepam, nicergoline, clomipramine, imipramine,
fluvoxamine, phenacetin, theophylline, amitriptyline
and quinolones.

CYP1A2 enzyme inducers such as: cigarette smoke,
charcoal-broiled food, rifampicin, omeprazole.

Pregnancy and breastfeeding

Do not use the medicine if you are pregnant, think you
are pregnant or are breastfeeding.

If you are planning to breastfeed, consult the doctor
before commencing use of Rilutek.

Driving and operating machinery

You can drive or operate machinery unless you feel
dizzy after taking the medicine.

**Important information regarding some of the
ingredients of the medicine**

Rilutek contains sodium. This medicine contains less
than 1 mmol (23 mg) sodium per tablet; this means it
is essentially “sodium-free”.

3. HOW SHOULD YOU USE THE MEDICINE?

Always use this preparation according to the doctor's
instructions. Check with the doctor or pharmacist if you
are uncertain about the dosage and treatment regimen
of the preparation.

The dosage and treatment regimen will be determined
by the doctor only.

The usual dosage is generally one tablet, twice a day
(every 12 hours).

Do not exceed the recommended dose.

Do not chew! Swallow the medicine with water.

The manufacturer has no information regarding the
possibility of halving, pulverizing or crushing the
tablets.

It is recommended to take the medicine one hour
before, or two hours after, a meal.

If you accidentally took a higher dosage or if a child has
accidentally swallowed the medicine, refer immediately
to a doctor or proceed to a hospital emergency room and
bring the package of the medicine with you.

If you forgot to take the medicine

If you forgot to take this medicine at the scheduled
time, skip the forgotten dose and take the next dose
on time.

Do not take a double dose to compensate for the
forgotten dose.

Adhere to the treatment as recommended by the doctor.
Even if there is an improvement in your health, do not
stop treatment with the medicine without consulting
the doctor.

**Do not take medicines in the dark! Check the label
and the dose each time you take the medicine. Wear
glasses if you need them.**

**If you have further questions regarding use of the
medicine, consult the doctor or pharmacist.**

4. SIDE EFFECTS

As with any medicine, use of Rilutek may cause side
effects in some users.

Do not be alarmed by the list of side effects. You may
not suffer from any of them.

Important

Refer to a doctor immediately

- If you have a **fever**, since Rilutek can cause a decrease
in the white blood cell count. The doctor may send
you for blood tests to check the number of white
blood cells in your blood, which are important in
fighting infection.
- Upon onset of the following effects: yellowing of the
skin or white part of the eyes (jaundice), itching all
over the body, nausea or vomiting; these can be signs
of **liver disease**. The doctor may routinely send you
for blood tests during treatment with the medicine.
- If you suffer from a cough or difficulties in breathing –
this may be a sign of lung disease.

Other side effects

Very common side effects (can affect more than one
in ten people):

Tiredness, nausea, increased liver enzyme levels.

Common side effects (can affect up to 1 in 10 people):
Dizziness, sleepiness, headache, numbness/tingling in
the mouth, rapid pulse, abdominal pain, vomiting,
diarrhea, pain.

Uncommon side effects (can affect up to 1 in 100
people):

Anemia, allergic reactions, inflammation of the
pancreas.

**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 with the doctor.**

Side effects can be reported to the Ministry of Health
by clicking on the link “Report Side Effects of Drug
Treatment” found on the Ministry of Health homepage
(www.health.gov.il) that directs you to the online form
for reporting side effects, or by entering the link:
<https://sideeffects.health.gov.il/>

5. HOW SHOULD THE MEDICINE BE STORED?

Avoid poisoning! This medicine, and any other medicine,
should be kept in a safe place out of the reach and sight
of children and/or infants in order to avoid poisoning.
Do not induce vomiting unless explicitly instructed to
do so by the doctor.

Do not use the medicine after the expiry date (exp. date)
that appears on the package. The expiry date refers to
the last day of that month.

Storage conditions:

There are no special storage conditions, but it is
recommended to store in a cool and dry place.

Store in the original package.

6. FURTHER INFORMATION

**In addition to the active ingredient, the medicine
also contains:**

Anhydrous dibasic calcium phosphate,
Microcrystalline cellulose, Croscarmellose sodium,
Hydroxypropylmethylcellulose (Hypromellose),
Magnesium stearate, Colloidal silica anhydrous,
Polyethylene glycol 6000 (macrogol 6000), Titanium
dioxide.

**What the medicine looks like and the contents of
the package:**

A package of 56 white tablets, with “RPR 202” engraved
on them.

This leaflet does not contain all the information about
your medicine. If you have any questions or are not sure
about anything, please refer to your doctor.

Registration Holder and Importer and address: sanofi-
aventis Israel Ltd., 10 Beni Gaon St., Netanya 4250499.

Revised in February 2021 according to MOH guidelines.

Registration number of the medicine in the National
Drug Registry of the Ministry of Health: 106-26-29046