

Summary Public Assessment Report

**Fullreizn
(tadalafil)**

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(tadalafil)

Film-coated tablets, 5 mg, 10 mg and 20 mg

This is a summary of the public assessment report (PAR) for Fullreizn. It explains how Fullreizn was assessed and its authorisation recommended as well as its conditions of use. It is not intended to provide practical advice on how to use Fullreizn.

For practical information about using Fullreizn, patients should read the package leaflet or contact their doctor or pharmacist.

What is Fullreizn and what is it used for?

Fullreizn is a 'generic medicine'. This means that Fullreizn is similar to a 'reference medicine' already authorised in the European Union (EU) called Cialis. Fullreizn is a treatment for adult men with erectile dysfunction.

How does Fullreizn work?

When a man cannot get, or keep a hard, erect penis suitable for sexual activity. Fullreizn has been shown to significantly improve the ability of obtaining a hard erect penis suitable for sexual activity.

How is Fullreizn used?

The pharmaceutical form of Fullreizn is film-coated tablet.

Please read section 3 of the package leaflet for detailed information on dosing recommendations, the route of administration, and the duration of treatment.

The medicine can only be obtained with a prescription.

What benefits of Fullreizn have been shown in studies?

Because Fullreizn is a generic medicine, studies in patients have been limited to tests to determine that it is bioequivalent to the reference medicine, Cialis. Two medicines are bioequivalent when they produce the same levels of the active substance in the body.

What are the possible side effects of Fullreizn?

Because Fullreizn is a generic medicine and is bioequivalent to the reference medicine, its benefits and possible side effects are taken as being the same as the reference medicine. For the full list of restrictions, see the package leaflet.

Why is Fullreizn approved?

It was concluded that, in accordance with EU requirements, Fullreizn has been shown to have comparable quality and to be bioequivalent to the reference medicine Cialis. Therefore, the Medical Products Agency in Sweden decided that, as for Cialis, the benefits are greater than its risks and recommended that it can be approved for use.

What measures are being taken to ensure the safe and effective use of Fullreizn?

A risk management plan has been developed to ensure that Fullreizn is used as safely as possible. Based on this plan, safety information has been included in the summary of product characteristics and the package leaflet for Fullreizn, including the appropriate precautions to be followed by healthcare professionals and patients.

Known side effects are continuously monitored. Furthermore new safety signals reported by patients/healthcare professionals will be monitored/reviewed continuously as well.

Other information about Fullreizn

The marketing authorisation for Fullreizn was granted on 2017-03-30 in Sweden.

The full PAR for Fullreizn can be found on the following website:
<http://mri.medagencies.org/Human/>. For more information about treatment with Fullreizn, please read the [package leaflet](#) or contact your doctor or pharmacist.

This summary was last updated in 2017-04.