

ARTICLE	PATCH FIT Lady
MANUFACTURER	D.Fenstec s.r.l.
TYPE OF PRODUCT	Analgesic patch with the ability to relieve the symptoms of pain, recovering the function that has suffered injury.
PRODUCT DESCRIPTION	The system basically consists of two elements: a non-woven polypropylene fabric and an acrylic adhesive. The colours used in the formula are enhanced with a certain percentage of titanium metal dioxides with different particles sizes called AT5.05: this powder reflects the far infrared waves emitted by the body - as shown by tests conducted by the Italian national research council (CNR).
CLINICAL SPECIFICATIONS	The patches promote pain relief. It is therefore clinically indicated for all cases where there is a need to eliminate toxins that have built up in the muscles: one of the causes of pain.
MATERIALS AND PROCESSING TECHNIQUES	The materials used in the device are: <ul style="list-style-type: none"> • 100% Polypropylene non-woven fabric • Acrylic adhesive mass • Ink colours • A mix of titanium dioxides of different grain sizes AT5.05 Sealed inside proper blister FIT personalized box
AVAILABLE SIZES	1 kit with: 1 back patch 10 x 4,5cm + 2 patches for the ovaries 4,5 x 3cm <ul style="list-style-type: none"> • Envelope 2 kit (FIT LADY 2K) – 9,9 x 17,5cm • Envelope with hanghole 2 kit (FIT LADY 2K) – 9,9 x 20cm
PATENT AND TRADEMARK	Product protected by industrial secret - FIT Trademark


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Company with **ISO 9001** and **ISO 13485 MD** certified quality management system

APPLICATION	Apply the two round patches to the ovary area and the shaped patch to the lumbar region corresponding with the L3 vertebra.
WARNINGS	The plasters are not to be used on wounds or on broken or reddened skin. If you suffer from specific blood circulation (especially to do with microcirculation) or muscular problems, seek the advice of your GP before applying the plasters.
CLINICAL CASE STUDIES AND FOLLOW-UP	Experimental trials and the relative publications, carried out on our device as well as on other devices based on the same principle, have shown that this phenomenon – although with a certain degree of subjectivity as subtle energy is involved – does, in actual fact, bring about benefits now acknowledged by various doctors during their clinical practices. The key ingredient - as demonstrated by a number of tests carried out using thermal imaging cameras - can result in easier blood flow in the body's smallest vessels, which has the ability to soothe pain and consequently restore muscle function.
SPECIFIC INSTRUMENTATION	There is no instrumentation required for patient application.
DURATION	5 days, including showers and swimming pool
EXPIRY	5 years
BAR CODE	2 kit 
CLASSIFICATION	CE Class I Medical Device
DATE OF LAST UPDATE	09/2019

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