KioMedine^{vs}one composition

- Primary component: 20mg/ml animal-free KiOmedine® CM Chitosan • Other ingredients: 35mg/ml sorbitol, phosphate-buffered water
- for injection
- Pack contents: pre-filled 3ml sterile glass syringe in blister pack Indication: one single-shot intraarticular injection for the symptomatic
- treatment of knee osteoarthritis (OA)



- Clinical data from a randomized controlled trial in patients with OA of the knee have shown that a single intraarticular injection of KioMedinevsone provided significant improvement in pain, stiffness and functionality up to 6 months*.
- In clinical investigation, the safety profile of a repeated injection of KioMedine^{vs}one into the knee was not altered after an interval of 3 months.

Dosage and administration

- Remove synovial fluid before injecting KioMedine^{vs}one.
- Strict aseptic injection technique should be employed during the administration of KioMedinevsone.
- Do not use disinfectants containing quaternary ammonium salts for skin preparation as KioMedine^{vs}one may precipitate in their presence.
- Use a luer needle of appropriate size, i.e. 20G to 23G, and suitable length for injecting KioMedine^{vs}one.

Contraindications

- For intraarticular use only. Do not inject KioMedine^{vs}one in patients who have: - a known allergy or hypersensitivity to any of the product components especially edible mushrooms.
- infections or skin disease at or around the injection site.
- severe inflammation, synovitis or inflammatory arthritis of the knee joint.
- a history of autoimmune and crystal diseases, evidence of lymphatic or venous stasis or serious blood disorders.

Precaution

- To date no data is available on potential interactions of KioMedine^{vs}one injected concomitantly with other intraarticular treatments.
- KioMedinevsone should not be used in presence of suspicious inflammatory fluid.

KioMedine



Adverse events

- Potential adverse events may occur after intraarticular injections. Injection of KioMedinevsone can cause temporary joint pain, joint effusion, joint swelling, joint stiffness, joint warmth, injection site pain or synovitis of the treated joint. Rare cases of acute synovial inflammation characterized by painful effusion of the knee, and possibly low-grade fever, have been reported following an intraarticular injection of KioMedinevsone Analysis of synovial fluid reveals aseptic fluid with no crystals. These local reactions respond well to rest, cold application, oral painkillers, non-steroidal anti-inflammatory drugs (NSAIDs) and/or arthrocentesis, and may not affect the clinical benefit of the treatment.
- Intraarticular infections did not occur in the clinical trial of KioMedine^{vs}one.

Information for patients

- Before injecting KioMedine^{vs}one, please inform your patient about its composition, performance, contraindications and adverse events:
- As a precautionary measure, the patient should avoid any intense physical or excessive weight-bearing activities for 48 hours after the injection of KioMedine^{vs}one. The patient should be advised to progressively use the treated knee and perform regular physical exercise
- Transient local reactions, such as joint pain, effusion, swelling or stiffness, have been reported following the injection of KioMedine^{vs}one. These symptoms are common post-injection complications of intraarticular injections in the target population and can be managed with rest, cold application or/and pressure bandage and may not affect the clinical benefit of the treatment.
- When transient local reactions occur, the patient should be advised to take oral painkillers (paracetamol) or NSAID without delay.

For more information, please refer to the instructions for use provided with the package unit and on our website: www.kiomedine-one.com



About KiOmed Pharma

Capitalizing on a history of innovation and expertise in exclusive natural chitosan chemistry, KiOmed Pharma develops a unique pipeline of medical devices that addresses unmet medical needs in high-impact pathologies and major social burdens such as invalidating osteoarthritis, skin aging and ophthalmology.

For safety concerns and incident reports please contact our medical device vigilance department, immediately (contact-rheumatology@kiomedpharma.com)

References

- lusive license to produce chitosan from mushrooms, for applications in medicine/pharmaceuticals: patent family WO03/068824
- 2. Patent claiming KiOmedine® chitosan derivatives and products for various clinical indications, filed by KiOmed: WO2019/105719
- 3. Douette P, Chausson M, Théatre E, Hermitte L. Biological evaluation of KIO014 for viscosupplementation E-DND-KI0014-TD017, Sept 2019.
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- ABTS assay: Miller NJ, Rice-Evans C, Davies MJ A new method for measuring antioxidant activity. Biochem Soc Trans 1 May 1993; 21(2):95S. PMID: 8359548.
- Bentin J, Emans P, Skaliczki G, Haverkamp D, A single injection of animal-free CA-Chitosan provides long-lasting reduction of osteoarthritic symptoms The APROOVE clinical study. Poster WCO-ESCEO 2020.

We cannot guarantee that all of the information provided is accurate or complete, nor can we guarantee that it will be suitable for a user's individual circumstances. This brochure is therefore provided for information purposes only. *CE mark obtained for 3-month follow-up, Notified Body conformity assessment of 6-month clinical data ongoing.

Product code: UDI-DI : 05404023514004 0 - 25 C⁴ CE STERILE SYRINGE 0344

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soft implant

FOR SUSTAINABLE OSTEOARTHRITIS PAIN **RELIEF AND OUTSTANDING RESPONSE RATE**

KioMedine

KioMedine





MMMM



KioMedine^{vs}one is the new generation single shot injection intended for the symptomatic treatment of knee osteoarthritis based on world-first exclusive animal-free KiOmedine® CM Chitosan.



KiOmedine® CM Chitosan is a patented technology^{1,2} made in Belgium, resulting from years of research and innovation. Thanks to a unique structure that differs from hyaluronic acid and natural chitosan, KioMedine^{vs}one has an exclusive dual mechanism of action to tackle OA pain and other symptoms.

Scientific evidence for improved lubrication and high free radical scavenging

Enhancement of lubrication and mobility⁴

Lubrication capacity demonstrated by a low coefficient of friction (COF) and knee mobility recovery after a lesion assessed using both in vitro and ex vivo models.

Thanks to its lubrication properties, KioMedinevsone offers improved joint friction reduction and provides better knee mobility with long-lasting relief6









Scavenging property⁵

As a natural free radical scavenger, **KioMedine**^{vs}one shows capacity that is roughly five times higher (in vitro) than an unprecedented free radical scavenging capacity, which that of existing HA technologies. This is potentially linked protects the endogenous hyaluronic acid in synovial fluid. to specific interactions and the attraction between CM-This protection can slow down the irreversible degradation chitosan molecules and Reactive Oxygen Species (ROS). caused by oxidative stress, which occurs naturally during Furthermore, KiOmedine® CM Chitosan has enabled the development of OA.

KioMedine^{vs}**one** contributes significantly to the protection of endogenous synovial components and has a scavenging





Proven safety and effectiveness in OA knee pain reduction after a single injection of **KioMedine^{vs}one**

In a randomized single-blind APROOVE clinical trial, the safety and efficacy of a single shot of KioMedine^{vs}one have been proven after injection in 70 patients6.

No serious or unexpected adverse events or patient withdrawals related to the safety of the treatment were reported. Treatment Related Adverse Events (TRAEs) were transient post-injection local effects that are commonly observed after IA injection of single shot products and were consistent in nature (e.g. pain, swelling), intensity and incidence with effects reported after injection of crosslinked HA-based products^o.

Reference	APROOVE - stage 2 (63 patients)
Arthralgia	25.4%(16)
Joint effusion	6.3% (4)
Joint swelling	6.3% (4)
Musculoskeletal stiffness	1
Synovitis/Arthritis	4.8% (3)

Effective up to 6 months

Sustainable **OA** pain relief



86% of patient

the development of a resorbable implant with unique degradation resistance and up to 1-month joint residence time with no need for crosslinking^{3,5}.

