

FREQUENTLY ASKED QUESTIONS



1. What is EUFLEXXA?

EUFLEXXA is a hyaluronic acid (HA) that's injected directly into the knee by your healthcare provider for the treatment of osteoarthritis (OA) knee pain. It is sometimes called a "gel shot for the knee."

If simple pain medications aren't giving you the relief you need, talk to your healthcare provider about EUFLEXXA.

2. What should I know about EUFLEXXA?

EUFLEXXA closely resembles healthy HA in the knee. It's created with a precise, multi-step cleansing and filtering process that results in a highly purified viscous fluid. It's also animal free.

3. How do I know if EUFLEXXA is a good option for me?

If you experience OA knee pain while walking, climbing stairs, sitting, standing, or even resting at night, and simple pain medications and/or physical therapy or exercise no longer have the same effect, then speak with your healthcare provider to see if EUFLEXXA may be right for you.

4. How is EUFLEXXA given?

To get the maximum and long-lasting pain relief of EUFLEXXA, your doctor should give you 3 injections of EUFLEXXA. Each injection should be 1 week apart. You may feel some relief from your OA knee pain after the first shot, but a clinical study showed that most patients experience further pain relief with each additional injection in the series.*



SEE WHAT THE #1 PRESCRIBED HA CAN DO FOR YOU: TALK TO YOUR HEALTHCARE PROVIDER OR VISIT EUFLEXXA.COM TODAY

*Based on a 12-week, multicenter, prospective, randomized, controlled, double-blind study conducted in adult patients with OA knee pain.

INDICATION

EUFLEXXA (1% sodium hyaluronate) is used to relieve knee pain due to osteoarthritis. It is used for patients who do not get enough relief from simple pain medications such as acetaminophen or from exercise and physical therapy.

EUFLEXXA is only for injection into the knee, performed by a doctor or other qualified healthcare professional.

IMPORTANT SAFETY INFORMATION

- Do not take this product if you have had any previous allergic reaction to EUFLEXXA or hyaluronan products.

Please see Important Safety Information throughout and Full Prescribing Information on pages 5 and 6.

FREQUENTLY ASKED QUESTIONS (CONTINUED)



5. After receiving EUFLEXXA injections, how long can I expect my pain relief to last?

When you get all 3 injections, clinical studies show that EUFLEXXA may help provide pain relief for up to 6 months.* Each injection counts when it comes to helping keep your knees relieved from pain and stiffness. Be sure to get all 3 injections to get the full benefit of your treatment.

6. What were the clinical study results for patients who have taken EUFLEXXA?

EUFLEXXA was shown to be comparable to another HA product in a 12-week head-to-head clinical study that included 321 patients.† 62% of the patients in the study experienced pain improvement with EUFLEXXA ($P < 0.0001$) vs 55% with the other product ($P < 0.0001$).‡ After 3 injections, patients who received EUFLEXXA experienced significantly less pain associated with 5 basic functions (climbing stairs, walking, standing, sitting, and resting during the night). At the end of the 12-week study, 2 out of 3 (63%) patients who received EUFLEXXA were pain free, and 8 out of 10 (81%) patients treated with EUFLEXXA reported that they were satisfied with the reduction in OA knee pain.§



*As shown in a 26-week, randomized, double-blind, saline-controlled study of 588 patients (ITT population) with OA knee pain.

†Questions based on the Western Ontario and McMaster Universities Osteoarthritis Index (WOMAC). WOMAC is an internationally recognized and widely used set of standardized questionnaires that allows healthcare providers to assess the condition of patients with osteoarthritis of the knee and hip. Measures include pain, stiffness, and physical functioning of the joints.

‡Improvements from baseline were statistically significant for both treatment groups.

§In the 12-week study, side effects caused by EUFLEXXA were joint pain (11/160), increase in blood pressure (3/160), joint swelling (3/160), feeling of sickness of injection (3/160), tingling (2/160), back pain (1/160), joint effusion (1/160), nausea (1/160), skin irritation (1/160), and tenderness in study knee (1/160).

IMPORTANT SAFETY INFORMATION (continued)

- You should not have EUFLEXXA injected into the knee if you have a knee joint infection or skin diseases or infections around the injection site.

Please see Important Safety Information throughout and Full Prescribing Information on pages 5 and 6.

LOOKING FOR A NEW OA PAIN RELIEF OPTION? VISIT EUFLEXXA.COM TO FIND A HEALTHCARE PROFESSIONAL NEAR YOU WHO OFFERS EUFLEXXA

FREQUENTLY ASKED QUESTIONS (CONTINUED)



7. Can both my knees be treated with EUFLEXXA at the same time?

Yes. Ask your healthcare provider what will work best for you. If you have pain in both knees, also known as bilateral OA, get 3 weekly injections in each knee to get the full benefit of EUFLEXXA.

8. Will the injections hurt?

While injection pain is individual, your physician may discuss with you and provide you with options to numb the knee before EUFLEXXA is given to offset any potential discomfort.

9. What should I do after my injections?

Avoid any major physical activity for 48 hours after your injection, including standing on your feet for over an hour. Ice your knee if you experience any swelling or pain. If you experience joint pain, back pain, limb/muscle pains or swelling, or if you have any other problems, call your healthcare provider. Regardless of whether you experience discomfort after the first injection, you should still get all 3 injections to help achieve maximum pain relief.*

10. What are the side effects of EUFLEXXA?

The most common adverse events reported in a clinical trial of EUFLEXXA were joint pain, back pain, limb pain, muscle pain, and joint swelling.

11. When should I follow up with my healthcare provider?

Follow up with your healthcare provider at 6 months[†] to discuss the result of your treatment (ie, relief from OA knee pain) and whether you would benefit from retreatment with EUFLEXXA.



DO YOU HAVE ANY MORE QUESTIONS?
EXPLORE [EUFLEXXA.COM](https://www.euflexxa.com) TO LEARN
MORE ABOUT THE TREATMENT

*Based on a 12-week, multicenter, prospective, randomized, controlled, double-blind study conducted in adult patients with OA knee pain.

[†]Extended treatment was verified in a multicenter, open-label, 26-week trial (N=433).

IMPORTANT SAFETY INFORMATION (continued)

- EUFLEXXA has not been tested in pregnant women, women who are nursing or in children less than 18 years of age. After you receive your EUFLEXXA injection you should avoid physical activities for 48 hours such as jogging, tennis, heavy lifting, or standing on your feet for a long time (more than one hour at a time).

Please see Important Safety Information throughout and Full Prescribing Information on pages 5 and 6.

FREQUENTLY ASKED QUESTIONS (CONTINUED)



12. What types of healthcare providers treat with EUFLEXXA?

Mostly specialists treat OA knee pain with EUFLEXXA. Specialists who give EUFLEXXA may include orthopedic surgeons/orthopedists, rheumatologists, sports medicine physicians, pain management specialists, physiatrists (rehabilitation physicians), nurse practitioners, and physician assistants.

13. How do I know if my insurance covers EUFLEXXA?

EUFLEXXA has broad commercial coverage and is covered under Medicare Part B. Your healthcare provider's office will verify your insurance coverage for you. You can also visit euflexxa.com for more information. If your plan doesn't cover EUFLEXXA, your physician's office can provide details on how you can buy EUFLEXXA directly.

14. Does EUFLEXXA offer a patient support program?

Yes, the EUFLEXXA patient support program provides helpful information like appointment reminders, tips for after your injections, scheduling follow-up appointments, and more. Just text **HAPPY** to **973-355-7159** from your smartphone to enroll.



ADDITIONAL SUPPORT IS JUST A STEP AWAY. VISIT EUFLEXXA.COM TO SIGN UP FOR THE HAPPY KNEES PROGRAM FOR FRIENDLY REMINDERS AND TIPS TO HELP YOU STAY ON TRACK WITH TREATMENT

IMPORTANT SAFETY INFORMATION (continued)

- The most common adverse events related to EUFLEXXA injections were joint pain, back pain, limb pain, muscle pain, and joint swelling.

You are encouraged to report negative side effects of prescription drugs to the FDA. Visit www.FDA.gov/medwatch or call 1-800-FDA-1088. You may also contact Ferring Pharmaceuticals Inc. at 1-888-FERRING.

Please see Important Safety Information throughout and Full Prescribing Information on pages 5 and 6.



EUFLEXXA® is a registered trademark of Ferring B.V.
©2019 Ferring B.V. All rights reserved. 12/19 US-EU-1900050



WHAT SHOULD I DO AFTER RECEIVING A EUFLEXXA INJECTION?

- Avoid physical activity for 48 hours following the injection to keep your knee from swelling.
Some examples of activities to avoid include:
 - Running
 - Tennis
 - Hiking
 - Jumping
 - Swimming
 - Heavy lifting (weight lifting)
 - Jogging
 - Bicycling
 - Aerobic exercise
- Do not stand on your feet for more than one hour at a time during the first 48 hours following your injection of EUFLEXXA.
- You should ask your doctor when you should begin major physical activities again.

WHEN SHOULD I CALL MY DOCTOR? (TROUBLESHOOTING)

If you experience any of the adverse effects or symptoms described earlier or if you have any other problems, you should call your doctor immediately.

WHAT OTHER NON-SURGICAL TREATMENTS ARE AVAILABLE FOR OSTEOARTHRITIS?

If you have osteoarthritis, there are other non-surgical treatment options available and these include:

- **Non-drug treatments**
 - Avoiding activities that cause pain in your knee
 - Exercise
 - Physical therapy
 - Weight loss (if overweight)
 - Removal of excess fluid from the knee
- **Drug therapy**
 - Pain medications such as acetaminophen or stronger prescription medications
 - Drugs that reduce inflammation, such as aspirin and other nonsteroidal anti-inflammatory agents (NSAIDs) such as ibuprofen and naproxen
 - Corticosteroids that are injected directly into the joint

WHAT DID CLINICAL STUDIES WITH EUFLEXXA SHOW?

A medical study involving 321 patients with knee pain due to osteoarthritis was performed in Germany. The study compared EUFLEXXA against another hyaluronan once a week for 3 weeks (control arm).

Pain, stiffness and function of the knee joint and patients' and doctors' judgment of treatment success were measured for 12 weeks. Patients were those with knee pain due to osteoarthritis who had not received pain relief with other medications. Patients experienced pain relief from EUFLEXXA injections similar to those patients in the control arm.

Another study involving 588 patients with knee pain due to osteoarthritis was conducted in the United States. Two hundred ninety three (293) patients were injected with EUFLEXXA and 295 with saline (salt water). The pain scores were used to compare the effectiveness of EUFLEXXA to saline injection: Patients were asked to rate how pain was felt on the 100 mm scale after 50 foot walk at 1, 2, 3, 6, 12, 18 and 26 weeks. EUFLEXXA group improved 25.7 mm from the baseline pain score, whereas the saline group improved 18.5 mm. There was more improvement in EUFLEXXA group than the saline group. The difference was 6.6 mm on 100 mm pain scale in favor of EUFLEXXA group. Study results showed significant improvement in osteoarthritis knee pain relief with EUFLEXXA therapy lasting up to 6 months. The study also showed that a repeated cycle of EUFLEXXA for an additional 26 weeks (1 year total) was safe.

EUFLEXXA has not been proven to relieve pain in any other joints.

WHAT ADVERSE EVENTS WERE OBSERVED IN THE CLINICAL STUDIES?

The number of subjects reporting adverse events was generally similar between the EUFLEXXA and Saline groups. Serious events were not observed in these clinical studies.

The following are the most common adverse events and symptoms that occurred during clinical studies of EUFLEXXA:

- Pain in the knee or at the injection site
- Stiffness, swelling or warmth in or around the knee

HOW DO I GET MORE INFORMATION ABOUT EUFLEXXA? (PATIENT ASSISTANCE)

If you have any questions or problems, talk to your doctor. If you would like more information on EUFLEXXA, please call 1-888-FERRING (1-888-337-7464) toll-free or visit www.euflexxa.com.

MANUFACTURED FOR:
FERRING PHARMACEUTICALS INC.
PARSIPPANY, NJ 07054

6309501103 Rev. 07/2016

EUFLEXXA is a registered trademark of Ferring BV

allowing only one parameter to be below 20 or above 80 at both the pre-screening visit and visit 1). For those patients who dropped out of the study before Week 12, the last evaluation was used. For those patients who requested NSAID or analgesic during the study, the last evaluation before start of NSAID/analgesic was used for the analysis. The results indicate that the effect of EUFLEXXA on pain relief was not inferior to that of a commercially available hyaluronan.

Table 6. Changes from Baseline to Last Visit in Overall Pain Score (primary end point, average of five pain scores)

	EUFLEXXA		Active Control (commercially available hyaluronan)		Standard Deviation	P value (non-inferiority)
	N	Change from Baseline (mm)	N	Change from Baseline (mm)		
ITT – patient	160	29.9	161	28.4	21	0.0032
Evaluable – patient	103	33.5	105	32.18	20	0.0083

26 Week Multicenter Clinical Trial

This was a multicenter, randomized, double-blind trial evaluating the efficacy and safety of EUFLEXXA as compared to saline comparator in subjects with chronic osteoarthritis of the knee. The intervention consisted of three weekly injections into the target knee with evaluations from baseline through Week 26 (1, 2, 3, 6, 12, 18, and 26). The primary objective was to demonstrate superiority over saline comparator from baseline to Week 26 using the pain level reported following a 50 foot walk test, measured by 100 mm visual analog scale. The following secondary endpoints were also evaluated: OARS1 responder rate at Week 12 and Week 26; WOMAC pain, disability, and joint stiffness score changes from baseline to Week 12 and 26; and change in Patient Global Assessment from baseline to Week 12 and Week 26.

Patient Population and Demographics

A total of 821 subjects were screened for the study, and 588 subjects were randomized. Approximately 88% of the randomized subjects completed the study, with similar proportions completing in each treatment group. Sixty-eight (11.6%) subjects discontinued the randomization/treatment phase prematurely: 34 (11.5%) in the saline group and 34 (11.6%) in the EUFLEXXA group. The most common reasons for discontinuation were the subject's withdrawing consent 25 (4.3%) and AEs 17 (2.9%). A total of 433 (73.6%) subjects entered the open-label extension study.

Clinical Results

Primary Endpoint

In the primary efficacy analysis, the EUFLEXXA group showed a larger mean decrease in pain scores on the 50-foot walk test from baseline to Week 26 than the saline group: -25.7 (28.85) mm versus -18.5 (32.53) mm, respectively. The group difference in least squares mean change from baseline of -6.6 mm (95% CI = -10.8 to -2.5 mm) was statistically significant (p-value = 0.002). Figure 1 depicts the adjusted mean change in pain scores on 50-foot walk test from baseline to week 26 (ITT Population).

Table 7. The Adjusted Mean Change in Pain Scores on 50-foot Walk Test from Baseline to Week 26 (ITT^a Population).

	Change from Baseline at Week 26		Difference in Changes (EUFLEXXA - Saline) from Baseline ^{b,c,d}	2-Sided 95% Lower and Upper Bound of Confidence Interval of the Difference ^d in Changes ^e	2-Sided P-Value ^e
	Saline (n=295) (SD)	EUFLEXXA (n=291) (SD)			
50-foot walk test, measured on a 100mm horizontal VAS score improvement at 26 weeks	-18.5 (32.53)	-25.7 (28.85)	-6.6 mm	-10.8, -2.5	0.002

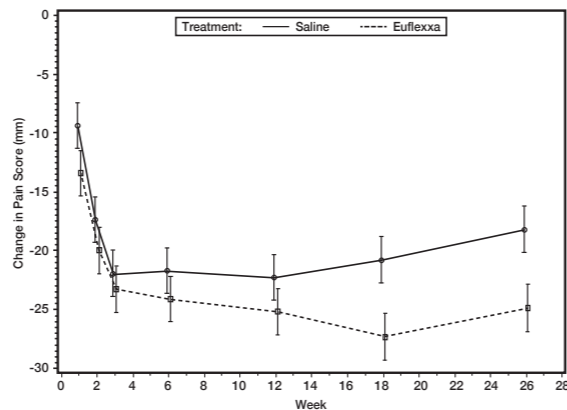
^a ITT= Intent to Treat

^b Negative (-) values favor EUFLEXXA.

^c The analysis is based on repeated measure mixed model Analysis of Covariance (ANCOVA) from baseline through 26 weeks on mean change from baseline 50-foot walk test, measured on a 100mm horizontal VAS score improvement at 26 weeks, with a weekly injection of EUFLEXXA for 3 weeks.

^d difference = least squares mean difference

Figure 1 Adjusted Mean Change in Pain Scores on 50-foot Walk Test from Baseline to Week 26 (ITT Population)



Secondary Endpoints

Table 8. OARS1 Responder Rates Using 50-foot Walk Test (ITT)

Visit Response/Statistics	Saline N=295	EUFLEXXA N=291	All Treatments N=586	Overall Comparison (2-sided 95% Lower and Upper Bound Statistics of Confidence Interval of Odds Ratio) ^c
Week 12				
No. of subjects with data	274	263	537	
Yes-n (%)	167 (60.9)	173 (65.8)	340 (63.3)	
No-n (%)	107 (39.1)	90 (34.2)	197 (36.7)	
Odds ratio ^a (95% CI)				1.3 (0.9, 1.8)
P-value				0.202
Week 26				

No. of subjects with data	264	254	518	
Yes-n (%)	155 (58.7)	169 (66.5)	324 (62.5)	
No-n (%)	109 (41.3)	85 (33.5)	194 (37.5)	
Odds ratio ^b (95%CI)				1.4 (1.0, 2.1)
P-value				0.047

OARS1 = Osteoarthritis Research Society International; ITT = intent-to-treat; N = number of subjects in a given treatment group for the population analyzed; n = number of subjects; (%) = percentage of subjects based on N; CI = confidence interval.

Note: The p-value for the odds ratio corresponds to the Wald chi-square test for EUFLEXXA versus saline with respect to OARS1 responder rates from a logistic regression adjusting for treatment group and study center.

Note: A subject was considered a responder if there was high improvement in pain or function >50% and absolute change >20 mm or improvement in at least two of the three following categories: pain >20% and absolute change >10 mm, function >20% and absolute change >10 mm, and/or Patient Global Assessment >20% and absolute change >10.

^a P_g (Log Odds Ratio) = 1.27 for 12 weeks and 1.4 for 26 weeks, based on a logistic regression model

(Log Odds Ratio) = \log [probability (responder)/probability (non-responder)]_{EUFLEXXA} / [probability (responder)/probability (non-responder)]_{saline}

^c When odds ratio > 1, [probability (responder)/probability (non-responder)]_{EUFLEXXA} > [probability (responder)/probability (non-responder)]_{saline}

Table 9. Other Secondary Endpoints at 26 Weeks for ITT (n=291)

	Change from Baseline at Week 26		The Difference ^d in Changes (EUFLEXXA - Saline) from the Baseline ^b	2-Sided Test P-Value ^a
	Saline (SD) (n=295)	EUFLEXXA (SD) (n=291)		
WOMAC C ^c (disability)	-14.6 (25.79)	-19.5 (24.68)	-4.3 mm	0.019
WOMAC B (joint stiffness)	-15.4 (29.33)	-19.6 (31.27)	-3.8 mm	0.075
WOMAC A (pain)	-16.3 (26.82)	-19.2 (26.81)	-3.3 mm	0.085
Patient Global Assessment	-17.8 (28.82)	-22 (30.38)	-4.5 mm	0.035

Note: The analysis is based on repeated measure mixed model Analysis of Covariance (ANCOVA) from baseline through 26 weeks on mean change from baseline.

^a P-values are not adjusted for the multiplicity.

^b Negative (-) values for WOMAC C and Patient Global Assessment are in favor of EUFLEXXA.

^c The Western Ontario and McMaster Universities Arthritis Index (WOMAC) is a set of standardized questionnaires used by healthcare professionals to evaluate the condition of patients with osteoarthritis of the knee and hip. WOMAC Pain Scale is 100mm.

^d difference=least square mean difference

No significant treatment group differences were observed in the change in number of study-specific acetaminophen tablets used per week or in the proportion of subjects who were pain free at Week 26 or last visit.

DETAILED DEVICE DESCRIPTION

Each syringe of EUFLEXXA contains:

Sodium hyaluronate	20 mg
Sodium chloride	17 mg
Disodium hydrogen phosphate dodecahydrate	1.12 mg
Sodium dihydrogen phosphate dihydrate	0.1 mg
Water for injection	q.s.

INTERACTIONS

None currently known

HOW SUPPLIED

EUFLEXXA is supplied in 2.25 mL nominal volume, disposable, pre-filled glass syringes containing 2 mL of EUFLEXXA. Only the contents of the syringe are sterile. EUFLEXXA is nonpyrogenic.

This product is not made with natural rubber latex.

Product Number: 55566-4100-1

3 disposable syringes per carton

STORAGE INSTRUCTIONS

Do not use EUFLEXXA if the package is open or damaged. Store in the original package at 2°-25°C (36°-77°F). Protect from light. Do not freeze.

CAUTION

Federal law restricts this device to sale by or on the order of a physician.

DIRECTIONS FOR USE

1. Each package of EUFLEXXA is manufactured using aseptic filling techniques. Do not use if the blister package is opened or damaged.
2. Remove joint effusion, if present.
3. Peel off the blister Tyvek backing (The syringe should be used immediately after the individual syringe blister is opened).
4. While holding the blister open side down, bend the blister and allow the syringe to fall gently onto the clean surface. Alternatively, hold the blister open side up and bend back the blister until the barrel's luer end is exposed. Gripping the luer end of the barrel, remove the syringe from the blister. **Do not remove the syringe from the plunger end.**
5. Remove the tip cap from the syringe and attach an appropriately sized sterile needle, for example 17 to 21 gauge.

Attention: Do not apply pressure to the plunger rod while the needle is being affixed. Verify that the needle is properly locked to the Luer Lock Adapter (LLA). Do not overtighten the LLA; this can lead to loosening of the LLA from the barrel.

6. Apply gentle pressure to the plunger in order to expel air from the syringe needle and to verify that the syringe is operating properly.
7. The syringe is ready for use.
8. Inject intra-articularly into the knee synovial capsule using strict aseptic injection procedures. Inject the full syringe contents, 2 ml into one knee only. If treatment is being administered to both knees, use a separate syringe for each knee. Discard any unused EUFLEXXA.
9. For single use only. Do not resterilize.
10. Store at 2°-25°C (36°-77°F). Protect from light. Do not freeze. If refrigerated, remove from refrigeration at least 20-30 minutes before use.
11. A dose of 2 ml is injected intra-articularly into the affected knee at weekly intervals for three weeks, for a total of three injections.

 **euflexxa**[®]
1% SODIUM HYALURONATE

 **euflexxa**[®]
1% SODIUM HYALURONATE



6309501103



6309501103