

MEDICARE FORM

Viscosupplementation Injectable Medication Precertification Request

Page 1 of 2

(All fields must be completed and legible for precertification review.)

For Ohio MMP:

FAX: 1-855-734-9389 **PHONE:** 1-855-364-0974

For other lines of business:

Please use other form.

Note: Single injection: Durolane and Gel-One are non-preferred. Monovisc and Synvisc-One are preferred. Multiinjection: Euflexxa, Gelsyn-3, GenVisc, Hyalgan, Hymovis, Supartz FX, Trivisc, and Visco-3 are non-preferred. Orthovisc and Synvisc are preferred.

Please indicate: Start of treatment: Start of	date	☐ Continuation of therapy (Request Additional Series Below)			
Precertification Requested By:		Phone: Fax:			
A. PATIENT INFORMATION					
First Name: Last Name:					
Address:	Cit	ty: State: ZIP:			
Home Phone:	Work Phone:	Cell Phone:			
DOB: Allergies:		Email:			
Current Weight: lbs or k	gs Height:	inches or cms			
B. INSURANCE INFORMATION					
Aetna Member ID #:	Does patient have other	ner coverage?			
Group #:	If yes, provide ID#:	Carrier Name:			
Insured:	Insured:	Insured:			
C. PRESCRIBER INFORMATION					
First Name:	Last Name:	(Check One): ☐ M.D. ☐ D.O. ☐ N.P. ☐ P.A.			
Address:		City: State: ZIP:			
Phone: Fax:	St Lic #:	NPI #: DEA #: UPIN:			
Provider Email:	Office Contact Name:	Phone:			
D. DISPENSING PROVIDER/ADMINISTRATION IN	FORMATION				
Agency Name: Administration code(s) (CPT): Address: City: Phone: Fax: TIN: PIN: NPI: E. PRODUCT INFORMATION Request is for: Gelsyn-3 (sodium hyaluronate) Hymovis (high molecular weight viscoelastic hymosomy of the sodium hyaluronate) Synvisc (hylan G-F 20) Synvisc-One (hylan G-Synvisc) Trilure	nate) ☐ Durolane (hyaluror 850 (sodium hyaluronate) ☐ hyaluronan) ☐ Orthovisc (hi lan G-F 20) ☐ TriVisc (sodiu on (1% sodium hyaluronate)	Dispensing Provider/Pharmacy: ☐ Outpatient Dialysis Center ☐ Physician's Office ☐ Retail Pharmacy ☐ Specialty Pharmacy ☐ Mail Order ☐ Other: Name: ☐ Address: City: ☐ State: ☐ ZIP: Phone: ☐ Fax: ☐ TIN: NPI: ☐ PIN: ☐ NPI: Cacid) ☐ Gel-One (cross-linked hyaluronate) ☐ Supartz FX (sodium hyaluronate) Hyalgan (sodium hyaluronate) ☐ Monovisc (sodium hyaluronate) In hyaluronate) ☐ Visco-3 (sodium hyaluronate)			
Dose:	Frequency:	HCPCS Code:			
F. DIAGNOSIS INFORMATION – Please indicate primary ICD Code and specify any other where applicable. Primary ICD Code: Other ICD Code:					
G. CLINICAL INFORMATION – Required clinical information must be completed in its entirety for all precertification requests.					
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For All Requests (includes Medicare patient requests, clinical documentation required for all requests): Note: Single injection products: Durolane and Gel-One are non-preferred. The preferred products are Monovisc and Synvisc-One.					
Multi injection products: Euflexxa, Gelsyn-3, GenVisc, Hyalgan, Hymovis, Supartz FX, TriVisc and Visco-3 are non-preferred. The preferred products are Orthovisc and Synvisc. Yes No Has the patient had prior therapy with the requested viscosupplementation product within the last 365 days? Yes No Has the patient had a trial and failure, intolerance, or contraindication to any of the following (select all that apply) Monovisc Orthovisc Synvisc Synvisc-One					

Continued on next page



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB		
C. CLINICAL INFORMATION (confinued)	Required clinical information mu	at he completed in its antiroty for all	proportification requests		
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests. Please explain if there are any other medical reason(s) that the patient cannot use any of the following (select all that apply)					
☐ Monovisc ☐ Orthovisc ☐ Synvisc-One					
	ontod symptomatic astocarthritis	(OA) of the tibiofomeral articulation	of the knee?		
☐ Yes ☐ No Does the patient have documented symptomatic osteoarthritis (OA) of the tibiofemoral articulation of the knee? Which knee will the viscosupplement be used? ☐ Left knee ☐ Right knee ☐ Both knees					
☐ Yes ☐ No Is there radiologic evidence of osteoarthritis (OA) of the knee?					
☐ Yes ☐ No Is the patient symptomatic?☐ ☐ Yes ☐ No Is the patient symptomatic?☐ ☐ Yes ☐ No Is the patient symptoms of osteoarthritis (OA) does the patient have? (Check ALL that apply)					
 ☐ Knee Pain ☐ Bony enlargement ☐ Bony tenderness ☐ Crepitus (noisy, grating sound) on active motion 					
☐ Erythrocyte sedimentation rate (ESR) less than 40 mm/hr ☐ Less than 30 minutes of morning stiffness					
 No palpable warmth of synovium ☐ Over 50 years of age Rheumatoid factor less than 1:40 titer (agglutination method) 					
Synovial fluid signs (clear fluid of normal viscosity and WBC less than 2000/mm3)					
Which of the following radiologic findings support the clinical diagnosis of osteoarthritis (OA)?					
Please select: Joint space narrowing Subchondral sclerosis Osteophytes and sub-chondral cysts No Does the patient have knee pain that interferes with functional activities (e.g. ambulation or prolonged standing)?					
Yes No Can the knee pain be attributed to any other forms of joint disease (other than osteoarthritis)?					
Yes No Has the patient completed conservative therapy in each joint to be treated with viscosupplementation? Yes No Is the patient unable to tolerate conservative therapy because of adverse side effects?					
Please indicate which of the following conservative therapies the patient completed:					
☐ Physical therapy ☐ Acetaminophen ☐ Topical capsaicin cream ☐ NSAID's, Specify:					
☐ Yes ☐ No Has the conservative treatment resulted in functional improvement after therapy?					
☐ Yes ☐ No Has the patient failed to adequately respond to aspiration and injection of intra-articular steroids?					
Yes No Are there any contraindications to the patient receiving viscosupplementation injections (e.g. active joint infection, bleeding disorder or skin infections at the injection site)?					
Yes No Is the patient scheduled to undergo a total knee replacement within 6 months of starting viscosupplementation treatment?					
Yes No Will the drug requested be used concomitantly with any of the following? Please select: With intra-articular anesthetics With intra-articular corticosteroids With intra-articular platelet rich plasma					
☐ With intra-articular mannitol/sorbitol ☐ With intra-articular mesenchymal stem cells ☐ With another viscosupplement					
Yes No Does the patient have morning stiffness of less than 30 minutes in duration?					
☐ Yes ☐ No Does the patient have crepitus on motion of the knee?					
For All Additional Series Requests (clinical documentation required for all requests): What product did the patient last receive?					
Enter date of last injection from prior series://					
Yes No Have at least six months elapsed since the last injection in the prior series?					
Yes No Has the patient had a documented reduction in the dose of NSAID's, other anti-inflammatories, or other analgesics during the 6-month period following the previous injection series?					
		nflammatories, or other analgesics	for a comorbid medical condition in addition		
to OA of the knee? If yes, please identify the comorbid medical condition:					
☐ Yes ☐ No ☐ N/A Was there a reduction in the number of intra-articular steroid injections or aspirations during the 6-month period following the series?☐ Yes ☐ No Is there objective documentation to support significant improvement of functional capacity as a result of previous injection series?					
Yes No Is there objective documentation to support significant improvement in pain as a result of previous injections?					
H. ACKNOWLEDGEMENT					
Request Completed By (Signature Requi	red):		Date:/		
Any person who knowingly files a request fany insurance company by providing mater insurance act, which is a crime and subjects	ially false information or conce	als material information for the p			

The plan may request additional information or clarification, if needed, to evaluate requests.