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Introduction

Description and Indication

SynoJoynt sodium hyaluronate solution is a sterile, nonpyrogenic, clear, viscoelastic solution of hyaluronan contained in a single use, prefilled syringe. It is a viscous solution of sodium hyaluronate in buffered physiological sodium chloride. Sodium hyaluronate is a high molecular weight fraction (approximately 2.5 Da × 10^6 Da) of a natural, complex sugar polymer consisting of the repeating disaccharide units Na glucuronate and N-acetylglucosamine.

SynoJoynt sodium hyaluronate is indicated for the treatment of pain from osteoarthritis (OA) of the knee in patients who have failed to respond adequately to conservative, nonpharmacologic therapy or simple analgesics (eg, acetaminophen).

Please see full prescribing information at ArthrexRSP.com.

Dosage and Administration

Each prefilled syringe of SynoJoynt solution contains 20 mg sodium hyaluronate, 17 mg sodium chloride, 0.8 mg disodium hydrogen phosphate heptahydrate, 0.06 mg sodium dihydrogen phosphate, monohydrate, and QS to 2 mL of water for injection. SynoJoynt sodium hyaluronate is intended to be injected into the knee joint and is administered as a regimen of 3 intra-articular injections given 1 week apart, for a total of 3 injections.

Using This Reimbursement Guide

This guide is provided solely for information purposes and designed for health care professionals for general coding and claims information related to SynoJoynt sodium hyaluronate. There are many factors that affect how payors will cover and pay for SynoJoynt sodium hyaluronate, including the site of service where it is administered, what type of health insurance the patient has, and the type of benefits the payor offers. This guide contains the following information:

- Coding for SynoJoynt sodium hyaluronate by site of service, including coding for the diagnosis and administration procedure
- Contact information for the Arthrex Reimbursement Support (RSP) Program
- Prior authorization checklist
- Sample claim forms that illustrate the key components that may be required by a payor when completing a claim for SynoJoynt sodium hyaluronate
- Tips for submitting clean claims and strategies to appeal denied claims

Arthrex RSP is available to support healthcare providers with coding, coverage, and reimbursement questions to help facilitate appropriate patient access to Arthrex biologic products.

Need assistance?
Visit ArthrexRSP.com or call the Arthrex RSP hotline at (844) 604-6359 between 9:00 AM and 7:00 PM ET, Monday through Friday.
Overview of Arthrex Reimbursement Support Program

Coverage and coding for SynoJoynt™ 1% sodium hyaluronate solution may vary depending on the patient’s type of health insurance and the site of service where the product is administered (ie, physician office, hospital outpatient department, or ambulatory surgical center). It will be important to conduct a benefit investigation for each patient in order to verify the following:

- Coverage and utilization restrictions, such as prior authorization, for SynoJoynt sodium hyaluronate
- Patient copayment or coinsurance for SynoJoynt sodium hyaluronate and administration services
- Appropriate coding for SynoJoynt sodium hyaluronate
- Provider’s network status with plan

Upon request, Arthrex RSP will provide prior authorization support by submitting, if possible, any of the information available for a verbal prior authorization if the payer will accept it from the Arthrex RSP hotline.

In some circumstances, upon request Arthrex RSP may provide prior authorization support by furnishing information available for a verbal prior authorization. Arthrex RSP offers reimbursement information to practices, ambulatory surgical centers, and hospital providers. Reimbursement counselors are available to support health care professionals relating to the following:

- Benefit verification for medical benefits
- Prior authorization support
- Claims management information
- Appropriate coding and billing support
- Appeals information

Disclaimer

Information described in the SynoJoynt sodium hyaluronate reimbursement guide is intended solely for use as a resource tool to assist physician office and ambulatory surgical center billing staff regarding potential reimbursement challenges. It is the sole responsibility of the treating health care professional to diagnose and treat the patient, and to confirm coverage, coding, and claim submission guidance with the patient’s health insurance plan to ensure claims are accurate, complete, and supported documentation in the patient’s medical record. Any determination regarding if and how to seek reimbursement should be made only by the appropriate members of the staff, in consultation with the physician, and in consideration of the procedure performed or therapy provided to a specific patient. Arthrex Inc. does not recommend or endorse the use of any particular diagnosis or procedure code(s) and makes no determination if or how reimbursement may be available. Of important note, reimbursement codes and payment, as well as health policy and legislation, are subject to continual change; information contained in this version of the SynoJoynt sodium hyaluronate reimbursement guide is current as of January 2022.

The information provided in this handout was obtained from many sources and is subject to change without notice as a result of changes in reimbursement laws, regulations, rules, and policies. All content is informational only, general in nature, and does not cover all situations or all payors’ rules and policies. This content is not intended to instruct medical providers on how to use or bill for health care procedures, including new technologies outside of Medicare national guidelines. A determination of medical necessity is a prerequisite that we assume will have been made prior to assigning codes or requesting payments. Medical providers should consult with appropriate payors, including Medicare fiscal intermediaries and carriers, for specific information on proper coding, billing, and payment levels for health care procedures. It is the sole responsibility of the medical provider to determine the appropriate coding.

This handout’s information represents no promise or guarantee concerning coverage, coding, billing, and payment levels. Arthrex specifically disclaims liability or responsibility for the results or consequences of any actions taken in reliance on information in this handout or through the Arthrex Reimbursement Support Program. The SynoJoynt sodium hyaluronate reimbursement guide does not constitute legal, coding, coverage, reimbursement, business, clinical, or other advice and no warranty regarding completeness or accuracy is implied.
Overview of Arthrex Reimbursement Support Program

Arthrex RSP assists with the provision of information to health care professionals in order to appropriately expedite patient access to care. Many reimbursement research requests can be completed in 1 to 2 business days from the time complete information is submitted to the program.

It is helpful to have the following information available when calling the Arthrex RSP to speak with a reimbursement counselor:

- Physician’s name, address, phone number, and provider number (NPI, TID, etc)
- Patient’s name, date of birth, address, and Social Security number
- Insurance company name, phone number, and fax number
- Name of policy holder
- Policy identification and group numbers
- Diagnosis
- Site of care
- Office contact name and phone number

In addition to reimbursement assistance, Arthrex RSP can provide additional resources that may include the following:

- Patient case management information
- Product ordering management

In order to use Arthrex RSP, health care professionals are asked to fill out and sign a benefit verification request form. You can obtain the form by contacting Arthrex RSP, accessing it at www.ArthrexRSP.com, or requesting one from your Arthrex sales representative.

Need assistance?
Visit ArthrexRSP.com or call the Arthrex RSP hotline (844) 604-6359 between 9:00 AM and 7:00 PM ET, Monday through Friday.
Coding and Associated Services for SynoJoynt™ Sodium Hyaluronate

Many payors recognize Healthcare Common Procedure Coding System (HCPCS) Level II national codes to report products (drugs and medical devices), supplies, and services not included in the Current Procedural Terminology (CPT) code.

Payors may accept the following HCPCS code for SynoJoynt sodium hyaluronate:

<table>
<thead>
<tr>
<th>HCPCS Code</th>
<th>Description</th>
<th>Billing Units</th>
<th>Site of Service</th>
<th>Claim Form (Location)</th>
<th>Payor Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>J7331</td>
<td>Hyaluronan or derivative, SynoJoynt, for intra-articular injection, 1 mg</td>
<td>20 (1 mg = 1 billing unit; each syringe = 20 billing units)</td>
<td>Physician office</td>
<td>CMS-1500 (Box 24D)</td>
<td>All</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Hospital outpatient</td>
<td>CMS-1450 (Field 44)</td>
<td>All</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Ambulatory surgical center</td>
<td>CMS-1450 (Field 44)</td>
<td>All</td>
</tr>
</tbody>
</table>

SynoJoynt sodium hyaluronate is supplied in a 3 mL single-use syringe containing 2 mL of product
- Each milliliter contains 10 mg sodium hyaluronate
- 2 mL contains 20 mg sodium hyaluronate
- Administration of SynoJoynt sodium hyaluronate does not vary by patient

National Health-Related Items Code

For devices such as SynoJoynt sodium hyaluronate, the manufacturer may adopt a unique, 3-segment number, known as the National Health-Related Items Code (NHRIC). Proper billing—especially to Medicare, Medicaid, or via electronic data interchange—may require the NHRIC be submitted in the 11-digit numeric 5-4-2 format (eg, 82197-0721-16). Hyphens should not be used when entering the actual data on your claim. For example:

<table>
<thead>
<tr>
<th>Eleven-Digit Example for SynoJoynt Sodium Hyaluronate</th>
<th>Reporting on CMS Claim Forms</th>
</tr>
</thead>
<tbody>
<tr>
<td>82197-0721-16</td>
<td>82197072116</td>
</tr>
</tbody>
</table>


Coding for Administration Services

CPT codes may be used to identify professional services (eg, administration procedure) provided in the physician office.

<table>
<thead>
<tr>
<th>CPT Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>20610</td>
<td>Arthrocentesis, aspiration, and/or injection, major joint or bursa (eg, shoulder, hip, knee, subacromial bursa); without ultrasound guidance</td>
</tr>
<tr>
<td>20611</td>
<td>Arthrocentesis, aspiration, and/or injection, major joint or bursa (eg, shoulder, hip, knee, subacromial bursa); with ultrasound guidance</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Modifier</th>
<th>Modifier Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>RT</td>
<td>Right side (used to identify procedures performed on the right side of the body)</td>
</tr>
<tr>
<td>LT</td>
<td>Left side (used to identify procedures performed on the left side of the body)</td>
</tr>
<tr>
<td>50</td>
<td>Bilateral procedure</td>
</tr>
<tr>
<td>EJ</td>
<td>Indicates subsequent injections of a series. Do not use for first injection of each series.</td>
</tr>
</tbody>
</table>
Coding and Associated Services for SynoJoynt™ Sodium Hyaluronate

ICD-10-CM Diagnosis Codes

International Classification of Diseases, 10th Revision, Clinical Modification (ICD-10-CM) diagnosis codes are used to report diseases and conditions. ICD-10-CM diagnosis codes identify why a patient needs treatment by documenting the medical necessity for prescribing SynoJoynt™ sodium hyaluronate. Coding to the highest level of specificity may expedite the claims adjudication process. The following ICD-10-CM diagnosis codes are consistent with the product indications. Arthrex RSP does not offer product support for off-label indications. The following diagnosis codes may be appropriate to describe patients with osteoarthritis of the knee; however, you—as the patient’s health care provider—are ultimately responsible for independently determining what code(s) are appropriate based on your clinical assessment of the patient:

<table>
<thead>
<tr>
<th>ICD-10-CM</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>M17.0</td>
<td>Bilateral primary osteoarthritis of knee</td>
</tr>
<tr>
<td>M17.10</td>
<td>Unilateral primary osteoarthritis, unspecified knee</td>
</tr>
<tr>
<td>M17.11</td>
<td>Unilateral primary osteoarthritis, right knee</td>
</tr>
<tr>
<td>M17.12</td>
<td>Unilateral primary osteoarthritis, left knee</td>
</tr>
<tr>
<td>M17.2</td>
<td>Bilateral post-traumatic osteoarthritis of knee</td>
</tr>
<tr>
<td>M17.30</td>
<td>Unilateral post-traumatic osteoarthritis, unspecified knee</td>
</tr>
<tr>
<td>M17.31</td>
<td>Unilateral post-traumatic osteoarthritis, right knee</td>
</tr>
<tr>
<td>M17.32</td>
<td>Unilateral post-traumatic osteoarthritis, left knee</td>
</tr>
<tr>
<td>M17.4</td>
<td>Other bilateral secondary osteoarthritis of knee</td>
</tr>
<tr>
<td>M17.5</td>
<td>Other unilateral secondary osteoarthritis of knee</td>
</tr>
<tr>
<td>M17.9</td>
<td>Osteoarthritis of knee, unspecified</td>
</tr>
</tbody>
</table>

Coding for SynoJoynt sodium hyaluronate may vary by payor type and plan type (ie, Medicare, private payor, Medicaid). Upon request, Arthrex RSP may conduct appropriate benefit verifications that provide coverage and coding information that is specific to your patient’s health insurance coverage. All information and submissions to CoPilot must be truthful, accurate, non-misleading, and/or of known application to the particular patient. The program is available Monday through Friday from 9:00 AM to 7:00 PM ET at (844) 604-6359.

Medicare National Average Reimbursement Rate Information*

<table>
<thead>
<tr>
<th>Site of Service</th>
<th>CPT Code</th>
<th>Website for Look-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>Physician Office</td>
<td>20610</td>
<td><a href="https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/PFSlook-up/index.html">https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/PFSlook-up/index.html</a></td>
</tr>
<tr>
<td></td>
<td>20611</td>
<td></td>
</tr>
<tr>
<td>Hospital Outpatient</td>
<td>20610</td>
<td><a href="https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/HospitalOutpatientPPS/Addendum-A-and-Addendum-B-Updates.html">https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/HospitalOutpatientPPS/Addendum-A-and-Addendum-B-Updates.html</a></td>
</tr>
<tr>
<td></td>
<td>20611</td>
<td></td>
</tr>
<tr>
<td>Ambulatory Surgical Center</td>
<td>20610</td>
<td><a href="https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/ASCPayment/11_Addenda_Updates.html">https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/ASCPayment/11_Addenda_Updates.html</a></td>
</tr>
<tr>
<td></td>
<td>20611</td>
<td></td>
</tr>
</tbody>
</table>

*Reimbursement rates for CPT codes vary by geography; consult the CMS website for regional rates applicable to the practice or contact the local Medicare administrative contractor for regional rates.
Prior Authorization Checklist

Arthrex RSP may assist you with obtaining information for prior authorization (PA) for SynoJoynt™ 1% sodium hyaluronate solution. Alternatively, the checklist below may be used to ensure that you are obtaining the necessary information from your patient’s insurer.

Patient Name: ____________________________________________  DOB: ______________________

Payor Name: ____________________________________________  Phone: ____________________  Date: ____________

<table>
<thead>
<tr>
<th>Questions to Ask</th>
<th>Answers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Is a PA required?</td>
<td>Yes</td>
</tr>
<tr>
<td>What information is needed by the insurer for the PA?</td>
<td>Diagnosis</td>
</tr>
<tr>
<td>Does the patient need to have a failure, contraindication, or intolerance to the following treatment options?</td>
<td>Nonpharmacologic (eg, exercise, physical therapy, weight loss if overweight)</td>
</tr>
<tr>
<td>Does the patient need to have documented symptomatic OA of the knee?</td>
<td>Yes</td>
</tr>
<tr>
<td>Does the patient need to have tried any other medications for the condition?</td>
<td>Yes (if yes, complete below) Medication/therapy:</td>
</tr>
<tr>
<td>Does the insurer have a specific PA form?</td>
<td>Yes</td>
</tr>
<tr>
<td>If the insurer has a specific PA form, how is the form obtained? Provide website, provider portal address, and/or fax number.</td>
<td>Online:</td>
</tr>
<tr>
<td>How is the PA submitted to the insurer? Provide phone, fax, and/or portal address.</td>
<td>Phone:</td>
</tr>
<tr>
<td>Will the insurer provide a PA number to include on the claim form?</td>
<td>Yes (if yes, complete below) PA number:</td>
</tr>
<tr>
<td>How long does it take the insurer to review the PA request?</td>
<td></td>
</tr>
<tr>
<td>How long is the PA valid for SynoJoynt sodium hyaluronate?</td>
<td></td>
</tr>
</tbody>
</table>

Need assistance?
Visit ArthrexRSP.com or call the Arthrex RSP hotline (844) 604-6359 between 9:00 AM and 7:00PM ET, Monday through Friday.
This document is provided for guidance only. Please call the Arthrex RSP hotline at (844) 604-6359.

Box 21—Diagnosis: Enter the appropriate diagnosis code (e.g., ICD-10-CM: M17.12, unilateral primary osteoarthritis, left knee)

Note: Other diagnosis codes may be applicable

Box 24G—Units: Enter the appropriate number of units of service (e.g., J7331 is per 1 mg, for a syringe of SYNOJOYNT, 20 units)

Box 24D—Procedures/Services/Supplies: Enter the appropriate CPT/HCPCS codes and modifiers
- J code: J7331 for SynoJoynt sodium hyaluronate, per mg
- Administration: eg, 20610, arthrocentesis, aspiration, and/or injection, major joint or bursa, without ultrasound guidance
- Modifier: LT for left knee or RT for right knee

Box 21—ICD Indicator: Identify the type of ICD diagnosis code used (enter a “0” for ICD-10-CM)
This document is provided for guidance only. Please call the Arthrex RSP hotline at (844) 604-6359.

**Fields 42-43:** Enter the appropriate revenue code and description corresponding to the HCPCS code in Field 44
- **0636** for SynoJoynt sodium hyaluronate
- **0510** for knee joint injection administered in the outpatient clinic

*Note: Other revenue codes may apply*

**Field 44:** Enter appropriate CPT/HCPCS codes and modifiers
- **Drug:** J7331 hyaluronan or derivative, SynoJoynt sodium hyaluronate, for intra-articular injection, 1 mg
- **Administration:** 20610 for knee joint injection without ultrasound guidance
- **Modifier** LT (left knee) or RT (right knee)

**Field 46:** Enter the appropriate number of units of service
- Enter 20 units of J7331 to denote use of SynoJoynt sodium hyaluronate for 1 syringe

**Field 66:** Identify the type of ICD diagnosis code used

**Fields 67 and 67A-67Q:** Enter the appropriate diagnosis code ICD-10-CM: M17.12 for unilateral primary osteoarthritis of the left knee (specific 4th and 5th digits depend on medical record documentation)

*Note: Other diagnosis codes may apply*
Tips for Clean Claims Submissions

Reasons for denied claims may include:

- Use of incorrect codes on claim
- Missing or incorrect information on claim form (e.g., misspelled patient name)
- Provider's network status with plan
- Incorrect number of units reported
- Omission of letter of medical necessity
- Failure to obtain a PA before initiating treatment or failure to include the PA approval number on the claim form

Since payors may have different guidelines for coding and claims filing, you may consider checking with individual plans to research claims submission requirements.

It is possible that not all payors will be familiar with SynoJoynt™ 1% sodium hyaluronate solution because it is a newer product and billed with its own unique HCPCS code. Payors may need more information about a product if they are unfamiliar with it, and may request additional information about the patient's treatment or diagnosis in order to determine whether a treatment is medically necessary.

A letter of medical necessity may help to explain why SynoJoynt sodium hyaluronate is medically necessary for the patient's treatment. Claims may include supporting materials such as:

- Customized letter of medical necessity
- Invoice
- FDA approval letter
- Chart notes
- Patient medical history
- Prior therapies
- Package insert