#### **Applies to:**

#### Aetna plans

#### Innovation Health® plans

Health benefits and health insurance plans offered and/or underwritten by the following:

Allina Health and Aetna Health Insurance Company (Allina Health | Aetna)

Banner Health and Aetna Health Insurance Company and/or Banner Health and Aetna Health Plan Inc.

(Banner|Aetna)

Sutter Health and Aetna Administrative Services LLC (Sutter Health | Aetna)

Texas Health + Aetna Health Plan Inc. and Texas Health + Aetna Health Insurance Company (Texas Health Aetna)



Aetna is the brand name used for products and services provided by one or more of the Aetna group of subsidiary companies, including Aetna Life Insurance Company and its affiliates (Aetna). Aetna provides certain management services on behalf of its affiliates.

#### **About this form**

You cannot use this form to initiate a precertification request. To initiate a request, call our Precertification Department or you can submit your request electronically.

This form will help you supply the right information with your precertification request. Failure to complete this form and submit all medical records we are requesting may result in the delay of review or denial of coverage.

#### How to fill out this form

As the patient's attending physician, you must complete all sections of the form. You can use this form with all Aetna health plans, including Aetna's Medicare Advantage plans. You can also use this form with health plans for which Aetna provides certain management services.

#### When you're done

Once you've filled out the form, submit it and all requested medical documentation to our Precertification Department by:

- We prefer you submit precertification requests electronically. Use our provider portal on Availity® to also upload clinical documentation, check statuses, and make changes to existing requests. Register today at <a href="mailto:availity.com/aetnaproviders">availity.com/aetnaproviders</a> or learn more about Availity at <a href="mailto:awww.availity.com/aetnatraining">awww.availity.com/aetnatraining</a>.
- Send your information by confidential fax to: **Precertification** Commercial and Medicare using FaxHub: **1-833-596-0339** 
  - The fax number above (FaxHub) is for clinical information only. Please send specific information that supports your medical necessity review. Please continue to send all other information (claims etc) to appropriate fax numbers.
- Mail your information to: PO Box 14079
   Lexington, KY 40512-4079

### What happens next?

Once we receive the requested documentation, we'll perform a clinical review. Then we'll make a coverage determination and let you know our decision. Your administrative reference number will be on the electronic precertification response.

### How we make coverage determinations

If you request precertification for a Medicare Advantage member, we use CMS benefit policies, including national coverage determinations (NCD) and local coverage determinations (LCD) when available, to make our coverage determinations. If there isn't an available NCD or LCD to review, then we'll use the Clinical Policy Bulletin referenced below to make the determination.

For all other members, we encourage you to review **Clinical Policy Bulletin #837: Shoulder Arthroplasty and Arthrodesis**, before you complete this form.

You can find the Clinical Policy Bulletins and Precertification Lists by visiting the website on the back of the member's ID card.

#### **Questions?**

If you have questions about how to fill out the form or our precertification process, call us at:

- HMO plans: 1-800-624-0756 (TTY: 711)
- Traditional plans: 1-888-632-3862 (TTY: 711)
- Medicare plans: <u>1-800-624-0756</u> (TTY: <u>711</u>)

| Section 1: Provide the following general information Typed responses are preferred. If the responses cannot be typed, they should be printed clearly.                          |                              |
|--|------------------------------|
| Member name:   | Reference number (required): |
| Member Phone Number:   |                              |
| Member ID:   | Member date of birth:        |
| Requesting provider/facility/vendor name:  |                              |
| Requesting provider/facility/vendor NPI:   |                              |
| Requesting provider/facility/vendor phone number: 1  |                              |
| Requesting provider/facility/vendor fax number: 1-   | -                            |
| Assistant/co-surgeon name (if applicable):   | TIN:                         |
| Physical Therapist Name:   |                              |
| Physical Therapist Phone Number:   |                              |
| Physical Therapist Fax Number:   |                              |
| Has the procedure been scheduled?  |                              |
| If yes, what is the date of service:   |                              |
| Which shoulder will surgery be performed on?   |                              |
| Left Right Please submit a separate form for each shoulder.  |                              |
| Section 2: Total Sh  | noulder Arthroplasty         |
| Reason for surgery (Diagnosis):  |                              |
| (Select all that apply)  |                              |
| Osteoarthritis  Rheumatoid arthritis   |                              |
| Avascular necrosis   |                              |
| Post-traumatic arthritis   |                              |
| Malunion fracture of the proximal humerus  |                              |
| Fracture of proximal humerus   |                              |
| Malignancy of the scapula, proximal humerus, shoulder joint or adjacent soft tissues by imaging  |                              |
| Nonunion/failure of a previous proximal humeral fracture surgery (shown by imaging)  |                              |
| Does the member have any of the following contraindicati   | ons?                         |
| (Select all that apply)  |                              |
| Active infection of the joint, or active systemic bacteremia, that has not been totally eradicated   |                              |
| Active skin infection (other than recurrent cutaneous staph infections) or open wound within the planned surgical site of the shoulder   |                              |
| Allergy to components of the implant (such as cobalt, chro   | mium, alumina)               |
| Rapidly progressive neurologic disease   |                              |
| Osseous abnormalities that cannot be optimally managed and which would increase the likelihood of a poor surgical outcome (i.e., inadequate bone stock to support the implant) |                              |
| ☐ None of the above  |                              |

Continued

| Member ID:   | Reference number (required):   |
|--|--|
| Section 2: Total Shoulder Arthroplasty (continued)   |  |
| Shoulder replacement system  Will a custom total shoulder implant be utilized?  Yes  No  Computer (robotic) assisted musculoskeletal surgical navigation  Will computer (robotic) assisted musculoskeletal surgical navigation be utilized?  Yes  No   |  |
| Radiographic evidence of the following (Select all that app   Irregular joint surfaces   Glenoid sclerosis   Malunion of fracture (proximal humerus)   Avascular necrosis of the humeral head with collapse   Osteophyte changes   Flattened glenoid   Cystic changes in the humeral head   Joint space narrowing of the shoulder join   Fracture of proximal humerus   Nonunion/failure of a previous proximal humeral fracture su   Malignancy of the scapula, proximal humerus, shoulder join | rgery  |
| On exam, what is the ROM (range of motion) flexion/abduction/rotation?  Normal or Mild Limitation Significant Limitation   |  |
| How much does this limit the member's daily activities?  Mildly Moderately Severely  |  |
| What degree of pain is the member having?  Mild Moderate Severe  |  |
| Has the member experienced this degree of pain for 6 mon   | ths or longer?  Yes No   |
| Does the member have Glenoid bony erosion with posterior or anterior subluxation (Walch Classification B2 glenoid)?  Yes No  |  |
| Has the member attempted and failed at least 12 weeks not Yes No   | n-surgical treatment in the past 12 months?                            |
| Which of these treatments have been attempted in the past (Select all that apply)  NSAIDS Formal Physical Therapy: Duration (weeks): Date Activity Modification Joint injection For rheumatoid arthritis only: Anti-cytokine agents (e.g., azathioprine, cyclosporine, gold salts, hydroxychloroquine,   | es to and from: etanercept, infliximab) and non-biologic DMARDs (e.g., |

| Member ID: Reference number (required):  |  |
|--|--|
| Section 3: Reverse Total Shoulder Arthroplasty   |  |
| Reason for Surgery (Diagnosis):  |  |
| (Select all that apply)  |  |
| Massive rotator cuff tears with pseudo-paralysis and without osteoarthritis  |  |
| Deficient rotator cuff with glenohumeral arthropathy   |  |
| Failed hemiarthroplasty  |  |
| Failed total shoulder arthroplasty with failed rotator cuff that is non-repairable   |  |
| Proximal humeral fractures that are not repairable or cannot be reconstructed with other techniques and are associated with a deficient rotator cuff                           |  |
| Reconstruction after a tumor resection   |  |
| Does the member have any of the following contraindications?   |  |
| (Select all that apply)  |  |
| Active infection of the joint, or active systemic bacteremia, that has not been totally eradicated   |  |
| Active skin infection (other than recurrent cutaneous staph infections) or open wound within the planned surgical site of the shoulder   |  |
| Allergy to components of the implant (such as cobalt, chromium, alumina)   |  |
| Rapidly progressive neurologic disease   |  |
| Osseous abnormalities that cannot be optimally managed and which would increase the likelihood of a poor surgical  |  |
| outcome (i.e., inadequate bone stock to support the implant)  None of the above  |  |
|  |  |
| Shoulder replacement system  Will a quetom total aboutdor implant be utilized?  Vee No.  |  |
| Will a custom total shoulder implant be utilized?    Yes    No   |  |
| Computer (robotic) assisted musculoskeletal surgical navigation  |  |
| Will computer (robotic) assisted musculoskeletal surgical navigation be utilized?   Yes   No   |  |
| Radiographic evidence of the following (Select all that apply)?  |  |
| MRI Massive Rotator Cuff Tear  |  |
| MRI Rotator Cuff Tear  |  |
| ☐ Irregular joint surfaces   |  |
| Glenoid sclerosis  |  |
| ☐ Osteophyte changes   |  |
| Flattened glenoid  |  |
| Cystic changes in the humeral head   |  |
| Joint space narrowing of shoulder joint  |  |
| Failed total shoulder arthroplasty with failed rotator cuff that is non-repairable  Shoulder fracture that is not repairable or cannot be reconstructed with other techniques. |  |
| Shoulder fracture that is not repairable or cannot be reconstructed with other techniques  Need for reconstruction after a tumor resection                                     |  |
| Failed hemiarthroplasty  |  |

| Member ID:   | Reference number (required):                     |
|--|--|
| Section 3: Reverse Total Shoulder Arthroplasty (continued)   |  |
| On exam, what is the ROM (range of motion) flexion/abduc  Normal or Mild Limitation  Significant Limitation  | tion/rotation?                                   |
| How much does this limit the member's daily activities?  Mildly Moderately Severely  |  |
| What degree of pain is the member having?  Mild Moderate Severe  |  |
| Has the member experienced this degree of pain for 6 mon   | ths or longer?  Yes No                           |
| Does the member have Glenoid bony erosion with posterior or anterior subluxation (Walch Classification B2 glenoid)?  Yes No  |  |
| Does the member have avascular necrosis of the humeral head with collapse in the presence of severe osteoarthritis of the shoulder?  Yes No  |  |
| Has the member attempted and failed at least 12 weeks non-surgical treatment in the past 12 months?  |  |
| Yes No   | is surgicular dedutions in the past 12 months.   |
| Which of these treatments have been attempted in the past  | t year?  |
| (Select all that apply)  | •  |
| NSAIDS   |  |
| Formal Physical Therapy: Duration (weeks): Date  | tes to and from:                                 |
| Activity Modification  |  |
| ☐ Joint injection  |  |
| For rheumatoid arthritis only: Anti-cytokine agents (e.g., etanercept, infliximab) and non-biologic DMARDs (e.g., azathioprine, cyclosporine, gold salts, hydroxychloroquine, leflunomide, methotrexate, or sulfasalazine) |  |
| Section 4: Total Shoulder Revision Arthroplasty  |  |
| Reason for surgery (Diagnosis)   |  |
| (Select all that apply)  |  |
| Fracture or mechanical failure of 1 or more components of the prosthesis or worn or dislocated plastic insert  |  |
| ☐ Displaced periprosthetic fracture  |  |
| Progressive or substantial periprosthetic bone loss  |  |
| Migration of the humeral head  |  |
| Confirmed peri-prosthetic infection by gram stain and culture  |  |
| Instability or dislocation of the glenoid or humeral components  |  |
| Aseptic loosening of one or more prosthetic components   |  |
| Bearing surface wear leading to symptomatic synovitis  |  |
| Persistent shoulder pain of unknown etiology that has not re   | esponded to non-surgical care for six (6) months |

| Member ID:   | Reference number (required):     |
|--|----------------------------------|
| Section 4: Total Shoulder Revision Arthroplasty (continued)  |                                  |
| If etiology unknown, which of these treatments have been   | attempted in the past 12 months? |
| (Select all that apply)  |                                  |
| ☐ NSAIDS   |                                  |
|  | Dates to and from:               |
| Activity Modification  |                                  |
| Assistive device (for example, sling)  |                                  |
| None of the above  |                                  |
| Does the member have any of the following contraindications?   |                                  |
| (Select all that apply)  |                                  |
| Active infection of the joint, or active systemic bacteremia, that has not been totally eradicated   |                                  |
| Active skin infection (other than recurrent cutaneous staph infections) or open wound within the planned surgical site of the shoulder   |                                  |
| Allergy to components of the implant (such as cobalt, ch   | romium, alumina)                 |
| Rapidly progressive neurologic disease   |                                  |
| Osseous abnormalities that cannot be optimally managed and which would increase the likelihood of a poor surgical outcome (i.e., inadequate bone stock to support the implant) |                                  |
| None of the above  |                                  |
| Shoulder replacement system  |                                  |
| Will a custom total shoulder implant be utilized?  | No                               |
| Computer (robotic) assisted musculoskeletal surgical navigation  |                                  |
| Will computer (robotic) assisted musculoskeletal surgical navigation be utilized?   Yes   No   |                                  |
| How much does this limit the member's daily activities?  |                                  |
| Mildly Moderately Severely   |                                  |
| What degree of pain is the member having?  |                                  |
| ☐ Mild ☐ Moderate ☐ Severe   |                                  |
| Has the member experienced this degree of pain for 6 mor   | nths or longer?  Yes No          |

| Member ID:  | Reference number (required): |
|---|------------------------------|
| Section 5: Shoulder Hemiarthroplasty  |                              |
| Reason for surgery (Diagnosis)  (Select all that apply)  Osteoarthritis  Rheumatoid arthritis  Avascular necrosis  Post-traumatic arthritis  Malunion of fracture (proximal humerus)  Arthritic conditions in which the glenoid bone stock is inated Rotator cuff tear arthropathy  Fracture of proximal humerus  Nonunion/failure of a previous proximal humeral fracture  |                              |
| Does the member have any of the following contraindication  |                              |
| <ul> <li>(Select all that apply)</li> <li>Active infection of the joint, or active systemic bacteremia, that has not been totally eradicated</li> <li>Active skin infection (other than recurrent cutaneous staph infections) or open wound within the planned surgical site of the shoulder</li> <li>Allergy to components of the implant (such as cobalt, chromium, alumina)</li> <li>Rapidly progressive neurologic disease/paralytic disorder of the shoulder</li> <li>Osseous abnormalities that cannot be optimally managed and which would increase the likelihood of a poor surgical outcome (i.e., inadequate bone stock to support the implant)</li> <li>None of the above</li> </ul> |                              |
| Shoulder replacement system  Will a custom total shoulder implant be utilized?  Yes  I  | No                           |
| Computer (robotic) assisted musculoskeletal surgical navigation  Will computer (robotic) assisted musculoskeletal surgical navigation be utilized?  Yes No  |                              |
| Radiographic evidence of the following (Select all that app    Irregular joint surfaces   Glenoid sclerosis   Malunion of a fracture (proximal humerus)   Avascular necrosis of the humeral head with collapse   Rotator cuff tear arthropathy   Osteophyte changes   Flattened glenoid   Cystic changes in the humeral head   Joint space narrowing of shoulder joint   Fracture of proximal humerus   Nonunion/failure of a previous proximal humeral fracture  |                              |

| Member ID:  | Reference number (required):                                |
|---|---|
| Section 5: Shoulder Hemiarthroplasty (continued)  |   |
| On exam, what is the ROM (range of motion) flexion/abduc  Normal or Mild Limitation  Significant Limitation                   | tion/rotation?  |
| How much does this limit the member's daily activities?  Mildly Moderately Severely   |   |
| What degree of pain is the member having?  ☐ Mild ☐ Moderate ☐ Severe   |   |
| Has the member experienced this degree of pain for 6 months or longer?  Yes No  |   |
| Does the member have avascular necrosis of the humeral of the shoulder?  Yes No   | nead with collapse in the presence of severe osteoarthritis |
| Has the member attempted and failed at least 12 weeks of ☐ Yes ☐ No   | non-surgical treatment in the past 12 months?               |
| Which of these treatments have been attempted in the pas-   | year?   |
| (Select all that apply)  NSAIDS   |   |
| <ul><li>☐ Formal Physical Therapy: Duration (weeks): Date</li><li>☐ Activity Modification</li><li>☐ Joint injection</li></ul> | res to and from:  |
| For rheumatoid arthritis only: Anti-cytokine agents (e.g.,  | etanercept, infliximab) and non-biologic DMARDs (e.g.,      |
| azathioprine, cyclosporine, gold salts, hydroxychloroquine,   | leflunomide, methotrexate, or sulfasalazine)                |

| Member ID:   | Reference number (required): |
|--|------------------------------|
| Section 6: Request for hospital admission pre and/or post-surgery  |                              |
| Are you requesting:  |                              |
| Section 7: Location where  | procedure will be performed  |
| Will the procedure be performed: ☐ Inpatient ☐ Outpatient  |                              |
| If procedure to be performed outpatient indicate the setting:  Outpatient hospital Ambulatory Surgical Center (free standing) Office   |                              |
| If request is for Outpatient hospital check any/all that apply:  Less than 12 years of age American Society of Anesthesiologists (ASA) Physic Danger of airway compromise Morbid obesity (BMI > 35 with comorbidities or BMI Pregnant Advanced liver disease Poorly controlled diabetes (hemoglobin A1C > 7) End stage renal disease (ESRD) with hyperkalemia Active substance use related disorders (Includes alc | > 40)                        |

| Member ID:   | Reference number (required):   |
|--|--|
| Section 7: Location where procedure will be performed (continued)  |  |
| <u> </u>   | Ongoing symptoms from previous MI<br>Symptomatic cardiac arrhythmia  |
| <u> </u>   | with: Drug Eluting Stent (DES) Bare Metal Stent placed in last year Current use of Aspirin or prescription anticoagulants                                      |
| Uncontrolled epilepsy  | Mini stroke/transient ischemic attack (TIA) Cerebral palsy Amyotrophic lateral sclerosis oral issues   |
| Respiratory conditions:  Moderate to severe obstructive sleep apnea  |  |
| Unstable respiratory status:  Poorly controlled asthma (FEV1 < 80% despite medical magnetic components)  Ventilator dependent patient  | nanagement)  |
|  | nfusion products to correct a coagulation defect nticipated need for blood or blood product transfusion istory of Disseminated Intravascular Coagulation (DIC) |
| <ul> <li>☐ Personal or family history of complication of anesthesia</li> <li>☐ History of solid organ transplant requiring anti-rejection medicatio</li> <li>☐ Other unstable or severe systemic diseases, intellectual disabilities outpatient hospital setting</li> <li>☐ This will be a prolonged surgery (&gt;3 hrs.)</li> </ul> | · /  |
| Do any of the following apply when procedure(s) to be performed at a surgical center  List specific equipment not available:  There are no participating general or specialty free-standing amb allow procedure(s) planned   | ng free-standing ambulatory surgical center or office based  |

| Member ID:  | Reference number (required): |
|---|------------------------------|
| Section 8: Provide the following documentation for your request   |                              |
| <ul> <li>Documentation of the indication for total arthroplasty, hemiarthroplasty or repeat shoulder arthroplasty</li> <li>Clinical records documenting the symptoms the patient experiencing</li> <li>Documentation of all conservative treatments, including type, duration, and outcome and</li> <li>Documentation of radiographic evidence of destructive degenerative joint disease.</li> </ul>  |                              |
| Section 9: Read this important information  |                              |
| Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive any insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent insurance act, which is a crime and subjects such person to criminal and civil penalties. |                              |
| Section 10: Sign the form   |                              |
| Just remember: You can't use this form to initiate a precertification request. To initiate a request, you can submit your request electronically or call our Precertification Department.   |                              |
| Signature of person completing form:  |                              |
| Date: / /   |                              |

Contact name of office personnel to call with questions:

Telephone number: 1-