# Rheumatoid Arthritis (RA) Measures Group

### PQRI Data Collection Sheet\*

PURI Data Collection Silect					
				Male ☐ Female	
Patient's Name Practice Medical Record Nur	Practice Medical Record Number (MRN)		Birth Date (mm/dd/yyyy)		
National Describer Identifier (NDI)			Data of Engagnetor		
National Provider Identifier (NPI)			Date of Encounter		
Step 1 Preliminary reporting requirements					
You must identify your intent to report the RA Measures Gro patient claim (G8490: I intend to report the RA Measures Groove than one claim.					
Step 2 Determine patient eligibility (Codes determining a patient's eligibility code(s) identified in Step 3 below.)	ility mus	st be rep	ported on the <b>same claim</b> as th	e quality	
	Yes	No			
Patient is aged 18 years and older on date of encounter.			Refer to date of birth listed above or o	n claim form.	
Patient has a line item diagnosis of RA.			714.0, 714.1, 714.2, 714.81		
There is a CPT E/M Service Code for an office visit/ consultation, home visit, or for work related/medical disability evaluation.			99201, 99202, 99203, 99204, 99205, 99212, 99213, 99214, 99215, 99241, 99242, 99243, 99244, 99245, 99341, 99342, 99343, 99344, 99345, 99349, 99350, 99455, 99456		
If <b>No</b> is checked for any of the above, STOP. This patient is no not report a CPT category II code or G-code.	not eligible	for report	ing on this measures group.		
Step 3 Complete individual measures					
Assessment and Classification of Disease Activity			Report one code for assessment/classification of disease activity OR one code for NOT assessed.		
<ul> <li>PQRI Measure #177</li> <li>reporting frequency: disease activity must be assessed, classified and reported once during the calendar year</li> <li>a standardized, systematic approach for evaluating the level of disease activity must be utilized. Standardized descriptive or numeric scales and/or composite indexes include but are not limited to: DAS28, SDAI, CDAI, RADAI, RAPID</li> </ul>			Disease activity assessed and classified as low	□ 3470F	
			Disease activity assessed and classified as moderate	□ 3471F	
			Disease activity assessed and classified as high	□ 3472F	
			OR		
			Disease activity NOT assessed	□ 3470F–8P	
Assessment of Functional Status		Report one code for assessment of functional status OR one code for NOT assessed.			
PQRI Measure #178					
<ul> <li>reporting frequency: functional status must be assessed an during the calendar year</li> </ul>	nd reported	d once			
<ul> <li>a standardized descriptive or numeric scale, standardized questionnaire, or notation of assessment of the impact of RA on patient activities of daily living must be utilized. Examples include but are not limited to: Health Assessment Questionnaire (HAQ), Modified HAQ, HAQ-2; American College of Rheumatology's Classification of Functional Status in RA</li> </ul>			Functional status assessed	□ 1170F	

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□ 1170F–8P

Functional status NOT assessed

<sup>\*</sup>For additional information on the PQRI program and reporting on measures groups, please visit the CMS Web site at http://www.cms.hhs.gov/pqri.

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Assessment and Classification of Disease Prognosis	Report one code for assessment/classification of disease prognosis OR one code for NOT assessed.		
PQRI Measure #179			
<ul> <li>reporting frequency: disease prognosis must be assessed, classified and reported once during the calendar year</li> <li>classification should be based upon at a minimum the following: functional limitation, extraarticular disease (eg, vasculitis, Sjorgen's syndrome, RA lung</li> </ul>	Disease prognosis assessed and classified as good	□ 3476F	
disease, rheumatoid nodules), rheumatoid factor (RF) positivity, positive anti-cyclic citrullinated peptide (anti-CCP) antibodies (both characterized dichotomously, per CEP recommendation), and/or bony erosions by radiography  • RA patients with features of poor prognosis have active disease with high tender and swollen joint counts, often have evidence of radiographic erosions, elevated levels of RF and or anti-CCP antibodies, and an elevated erythrocyte sedimentation rate, and an elevated C-reactive protein level	Disease prognosis assessed and classified as poor	□ 3475F	
erythrocyte seannentation rate, and an elevated c-reactive protein level	OR		
	Disease prognosis NOT assessed	□ 3475F–8P	
Disease Modifying Anti-Rheumatic Drug (DMARD) Therapy	. 3	t one code for DMARD therapy OR one code for	
PQRI Measure #108  • reporting frequency: DMARD therapy must be prescribed and reported once during the calendar year	DMARD therapy prescribed, dispensed, or administered	□ 4187F	
	Not prescribed, dispensed, or administered for medical reasons <sup>†</sup>	□ 4187F–1P	
<ul> <li>biologic DMARD therapy includes adalimunab, etanercept, infliximab, abatacept, anakinra and rituximab</li> </ul>	Document reason in medical chart		
. ,	OR		
	DMARD therapy NOT prescribed, dispensed, or administered	□ 4187F–8P	
Tuberculosis (TB) Screening Prior to First Course of Biologic DMARD Therapy	Report one code for biologic DMARD therapy status. If patient is receiving a first course of biologic DMARD therapy (excluding rituximab), you will also need to report one code for TB screening or one code for NOT performed.		
	Patient not receiving a first course of biologic DMARD therapy (or biologic DMARD prescription is for rituximab)	□ 4196F	
PQRI Measure #176 • reporting frequency: TB screening must be performed and results interpreted 6 months prior to patient's first course of biologic DMARD therapy and reported once during the calendar year	Patient receiving a first course of biologic DMARD therapy AND TB screening performed and results interpreted within 6 months prior to therapy	□ 4195F AND 3455F	
<ul> <li>biologic DMARD therapy includes adalimunab, etanercept, infliximab, abatacept, anakinra (rituximab is excluded)</li> <li>patients are considered to be receiving a first course of therapy using a biologic DMARD only if they have never previously been prescribed or dispensed a biologic DMARD</li> </ul>	Patient receiving a first course of biologic DMARD therapy AND TB screening not performed or results not interpreted for medical reasons (ie, patient positive for TB and documentation of past treatment; patient has recently completed a course of anti-TB therapy)	☐ 4195F and 3455F–1P	
	Document reason in medical chart		
	OR		
	Patient receiving a first course of biologic DMARD therapy AND TB screening NOT performed or results NOT interpreted	☐ 4195F and 3455F–8P	

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Glucocorticoid Management		Report one code for glucocorticoid use (including dose) or ONE code for NOT assessed. If patient is receiving ≥ 10 mg daily prednisone (or equivalent) for longer than 6 months with improvement or no change in disease activity, you will also need to report one code for management plan OR one code for NOT documented.	
PQRI Measure #180  • reporting frequency: glucocorticoid use (and management plan, if appropriate) must be assessed and reported once during the calendar year  • 1 mg prednisone = 1 mg prednisolone; 5 mg cortisone; 4 mg hydrocortisone; 0.8 mg triamcinolone; 0.8 mg methylprednisolone; 0.15 mg dexamethasone; 0.15 mg betamethasone  • glucocorticoid management plan: attempt to taper steroids OR a new prescription for a non-glucocorticoid DMARD OR increase in dose of non-glucocorticoid DMARD dose for persistent RA disease activity at current or reduced dose	Assessment of glucocorticoid use (including dose) and management plan, if appropriate	Patient not receiving glucocorticoid therapy	□ 4192F
		Patient receiving < 10 mg daily prednisone (or equivalent) OR glucocorticoid use is for less than 6 months OR RA disease activity is worsening	□ 4193F
		Patient receiving ≥ 10 mg daily prednisone (or equivalent) for longer than 6 months with improvement or no change in disease activity AND management plan documented	□ 4194F AND 0540F
		Patient receiving ≥ 10 mg daily prednisone (or equivalent) for longer than 6 months with improvement or no change in disease activity AND management plan not documented for medical reasons (ie, glucocorticoid prescription is for a medical condition other than RA)	□ 4194F AND 0540F–1P
		Document reason in medical chart	
		<b>OR</b> (Report one of the following options)	
		Glucocorticoid use NOT assessed	
		Patient receiving ≥ 10 mg daily prednisone (or equivalent) for longer than 6 months with improvement or no change in disease activity AND management plan NOT documented	□ 4194F AND 0540F–8P

#### **Step 4 Reporting Instructions**

This measure can be reported for each eligible patient in one of two ways:

- 1. Report the corresponding CPT category II code(s) as selected above for each of the six measures in the RA Measures Group np.
- 2. If **all** quality actions for the patient have been performed for each of the six measures in the RA Measures Group, **G8499** may be reported. *Note: G8499 is not appropriate for this patient if any CPT category II codes with the 8P modifier have been selected from Step 3.*