ATTACHMENT G: JOURNAL INFORMATION PIECE FOR NEUROLOGISTS
Important Safety Information for Neurologists About Demyelinating Disorders in Co-managing Rheumatoid Arthritis Patients Receiving ACTEMRA®

ACTEMRA® (tocilizumab) is an interleukin-6 (IL-6) receptor antagonist that has been approved by the Food and Drug Administration (FDA) for three indications:

- Adult patients with moderately to severely active rheumatoid arthritis (RA) who have had an inadequate response to one or more Disease-Modifying Anti-Rheumatic Drugs (DMARDs) with a recommended ACTEMRA dosing interval of every 4 weeks for intravenous (IV) or every other week or weekly for subcutaneous (SC) administration.

- Children 2 years of age and older with active Polyarticular Juvenile Idiopathic Arthritis (PJIA) with a recommended ACTEMRA dosing interval of every 4 weeks for IV administration.

- Children 2 years of age and older with active Systemic Juvenile Idiopathic Arthritis (SJIA) with a recommended ACTEMRA dosing interval of every 2 weeks for IV administration.

The safety and efficacy of ACTEMRA for conditions other than RA, PJIA and SJIA have not yet been established.

Neurologists co-managing RA patients should be aware of important safety information regarding treatment with ACTEMRA.

Demyelinating disorders: The impact of treatment with ACTEMRA on demyelinating disorders is not known, but multiple sclerosis and chronic inflammatory demyelinating polyneuropathy were reported rarely in clinical studies of adults with RA. Monitor patients for signs and symptoms potentially indicative of demyelinating disorders. Prescribers should exercise caution in considering the use of ACTEMRA in patients with preexisting or recent onset demyelinating disorders.

Reporting Adverse Events
It is important that you report any serious neurologic adverse event, including demyelinating disorders, that occurs in a patient being treated with ACTEMRA, even if you do not think there is a causal relationship. The information that you, as a neurologist, provide about these events may inform therapy and monitoring decisions for future patients.

Reporting and maintains patient confidentiality. Your patient’s name or contact information is not needed. HIPAA does not apply to this adverse event reporting.
You can report your cases to Genentech or directly to the FDA:
- Genentech at 1-888-835-2555
- MedWatch (FDA safety information and adverse event reporting program) at 1-800-332-1088 or online at www.fda.gov/medwatch/report.htm

Please visit www.ACTEMRA.com for Prescribing Information and Medication Guide.