Rituximab for the Treatment of Rheumatoid Arthritis

Description of Procedure or Service

Rituximab is a chimeric murine-human monoclonal antibody directed against the CD20 surface antigen, which is expressed on pre-B and mature B lymphocytes. Rituximab induces lysis of normal and malignant CD20-expressing B cells; possible mechanisms of cell lysis include complement-dependent cytotoxicity and antibody-dependent cell-mediated cytotoxicity. B cells are thought to play a role in the pathogenesis of rheumatoid arthritis and other autoimmune diseases by producing autoantibodies and proinflammatory cytokines and by activating T lymphocytes. Rituximab reduces the number of B cells in the peripheral blood and in lymphoid tissues, thereby interrupting pathogenic processes of autoimmune diseases. Rituximab is administered by intravenous infusion.

Tumor necrosis factor inhibiting drugs such as Remicade® (infliximab), Enbrel® (etanercept), or Humira® (adalimumab) are considered first line treatment before considering the use of Rituxan® (rituximab). Patients will sometimes become intolerant or unresponsive to TNF inhibiting therapy and for these; Rituxan® (rituximab) may be considered.

Note***This policy only applies to Rituximab (Rituxan®) when used for the treatment of Rheumatoid Arthritis.

For oncologic uses of Rituximab, please see BCBSNC Corporate Medical Policy, Monoclonal Antibodies for Non-Hodgkin Lymphoma and Acute Myeloid Leukemia In the Non-Hematopoietic Stem Cell Transplant Setting

Related Policies:
Abatacept (Orencia)
Golimumab (Simponi Aria)
Infliximab (Remicade)
Tocilizumab (Actemra)

***Note: This Medical Policy is complex and technical. For questions concerning the technical language and/or specific clinical indications for its use, please consult your physician.

Policy

BCBSNC will provide coverage for rituximab (Rituxan®) for the treatment of rheumatoid arthritis when it is determined to be medically necessary because the medical criteria and guidelines shown below are met.
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Benefits Application

This medical policy relates only to the services or supplies described herein. Please refer to the Member's Benefit Booklet for availability of benefits. Member's benefits may vary according to benefit design; therefore member benefit language should be reviewed before applying the terms of this medical policy.

Rituximab may be subject to prior review requirements.

When Rituximab for the Treatment of Rheumatoid Arthritis is covered

Rituximab (Rituxan®) for the treatment of adults with rheumatoid arthritis may be considered medically necessary under the following conditions:

1. The patient has moderate to severe rheumatoid arthritis (eg, > 8 swollen and > 8 tender joints); and
2. Rituximab is administered in combination with methotrexate; and
3. Either:
   a. patient has had an inadequate response to one or more tumor necrosis factor (TNF) inhibitors; or
   b. Patient has had an inadequate response to methotrexate or other conventional synthetic disease-modifying anti-rheumatic drug (DMARD) and is not suitable for treatment with TNF inhibitors due to a recent (within 5 years) history of:
      • a lymphoma or other malignancy,
      • latent tuberculosis and contraindications to chemoprophylaxis, or
      • previous demyelinating disease.

When Rituximab for the Treatment of Rheumatoid Arthritis is not covered

1. Rituximab for the treatment of rheumatoid arthritis is considered not medically necessary when the criteria stated above are not met.
2. Rituxan® is considered not medically necessary when used in combination with TNF-inhibiting drugs.

Policy Guidelines

The U.S. Food and Drug Administration has approved rituximab for individuals who have moderately to severely active rheumatoid arthritis and inadequate response to 1 or more standard agents (eg, tumor necrosis factor inhibitors, inadequate response to methotrexate or other conventional synthetic disease-modifying anti-rheumatic drug) who receive rituximab and methotrexate, the evidence is sufficient to determine that the technology results in a meaningful improvement in the net health outcome. Rituximab should be administered by a healthcare professional with appropriate medical support to manage severe and potentially fatal infusion reactions.

Rituximab carries the following black box warnings:
   • Fatal infusion reactions within 24 hours of rituximab infusion; approximately 80% of fatal reactions occurred with first infusion.
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- Severe mucocutaneous reactions, some with fatal outcomes
- Hepatitis B virus reactivation, in some cases resulting in fulminant hepatitis, hepatic failure, and death
- Progressive multifocal leukoencephalopathy resulting in death.

Labelled warnings and precautions include:
- Tumor lysis syndrome (for patients with hematologic malignancies)
- Infections
- Cardiac arrhythmias and angina
- Bowel obstruction and perforation
- Live virus vaccines: Do not administer live virus vaccines before or during rituximab therapy.
- Cytopenias

Rituximab is pregnancy category C: There are no adequate and well-controlled studies for rituximab during pregnancy. Individuals of childbearing potential should use effective contraception while receiving rituximab and for 12 months after treatment. Rituximab may be used during pregnancy only if potential benefit justifies potential risks to the fetus.

Billing/Coding/Physician Documentation Information

This policy may apply to the following codes. Inclusion of a code in this section does not guarantee that it will be reimbursed. For further information on reimbursement guidelines, please see Administrative Policies on the Blue Cross Blue Shield of North Carolina web site at www.bcbsnc.com. They are listed in the Category Search on the Medical Policy search page.

Applicable codes: J9310

Diagnoses that are subject to medical necessity review: 714, 714.0, 714.1, 714.2, 714.3, 714.30, 714.31, 714.32, 714.33, 714.4, 714.8, 714.81, 714.89, 720.0

ICD-10 diagnosis codes: M05.00, M05.011, M05.012, M05.019, M05.021, M05.022, M05.029, M05.031 M05.032, M05.039, M05.041, M05.042, M05.049, M05.051, M05.052, M05.059, M05.061, M05.062, M05.10, M05.111, M05.112, M05.119, M05.319, M05.069, M05.071, M05.072, M05.079, M05.09, M05.30, M05.311, M05.312, M05.321, M05.322, M05.329, M05.331, M05.332, M05.339, M05.341, M05.342, M05.349, M05.351, M05.352, M05.359, M05.361, M05.362, M05.369, M05.371, M05.372, M05.379, M05.39, M05.60, M05.611, M05.612, M05.619, M05.621, M05.622, M05.629, M05.631, M05.632, M05.639, M05.641, M05.642, M05.649, M05.651, M05.652, M05.659, M05.661, M05.662, M05.669, M05.671, M05.672, M05.679, M05.69, M06.80, M06.811, M06.812, M06.819, M06.821, M06.822, M06.829, M06.831, M06.832, M06.839, M06.841, M06.842, M06.849, M06.851, M06.852, M06.859, M06.861, M06.862, M06.869, M06.871, M06.872, M06.879, M06.88, M06.89,M08.00, M08.09, M08.29, M08.011, M08.012, M08.019, M08.021, M08.022, M08.029, M08.031, M08.032, M08.039, M08.041, M08.042, M08.049, M08.051, M08.052, M08.059, M08.061, M08.062, M08.069, M08.071, M08.072, M08.079, M08.08, M08.20, M08.211, M08.212, M08.219, M08.221, M08.222, M08.229, M08.231, M08.232, M08.239, M08.241, M08.242, M08.249, M08.251, M08.252, M08.259, M08.261, M08.262, M08.269, M08.271, M08.272, M08.279, M08.28, M08.29,
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M08.00, M08.011, M08.012, M08.019, M08.021, M08.022, M08.029, M08.031, M08.032, M08.039, M08.041, M08.042, M08.049, M08.051, M08.052, M08.059, M08.061, M08.062, M08.069, M08.071, M08.072, M08.079, M08.09, M08.40, M08.411, M08.412, M08.419, M08.421, M08.422, M08.429, M08.431, M08.432, M08.439, M08.441, M08.442, M08.449, M08.451, M08.452, M08.459, M08.461, M08.462, M08.469, M08.471, M08.472, M08.479, M08.48, M12.00, M12.011, M12.012, M12.019, M12.021, M12.022, M12.029, M12.031, M12.032, M12.039, M12.041, M12.042, M12.049, M12.051, M12.052, M12.059, M12.061, M12.062, M12.069, M12.071, M12.072, M12.079, M12.08, M12.09, M06.4, M45.0, M45.1, M45.2, M45.3, M45.4, M45.5, M45.6, M45.7, M45.8, M45.9

BCBSNC may request medical records for determination of medical necessity. When medical records are requested, letters of support and/or explanation are often useful, but are not sufficient documentation unless all specific information needed to make a medical necessity determination is included.

Scientific Background and Reference Sources


Senior Medical Director review, 3/20/2008.


Specialty Matched Consultant Panel - 1/2010


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Policy Implementation/Update Information

4/1/08 New policy developed. Under the "When Covered" section; "Rituxan® (rituximab) may be medically necessary for the treatment of rheumatoid arthritis when the medical criteria and guidelines shown below are met: 1. The patient has moderate to severe rheumatoid arthritis; and 2. The patient has failed to respond adequately to at least one Tumor Necrosis Factor (TNF) inhibiting drug or is intolerant to all TNF-inhibiting drugs (i.e., Remicade (infliximab), Enbrel (etanercept), or Humira (adalimumab); and 3. Rituxan® (rituximab) is to be used in combination with methotrexate; and 4. The patient is 18 years old or older. 5. Continued use of Rituxan® (rituximab) can only be renewed after 6 months have passed from the last course of treatment and retreatment is necessary to control symptoms." Under the "When Not Covered" section; "1. When the criteria stated above are not met. 2. Rituxan® should not be used in combination with TNF-inhibiting drugs." Senior Medical Director review, 3/20/2008. References added. Notification given April 1, 2008. Policy effective 7/1/2008. References added.

11/3/08 Added "Note" at the end of the "Description" section indicating; "NOTE: This policy only applies to Rituximab (Rituxan®) when used for the treatment of Rheumatoid Arthritis." (btw)

3/2/10 Specialty Matched Consultant Advisory Panel review 1/5/2010. Updated "Description" section. No change to policy statement. Added wording to clarify #3 in the “When Covered” section to indicate; “Rituxan (rituximab) is to be used preferably in combination with methotrexate unless contraindicated”. References added. (btw)

5/11/10 Under the “When Covered” section, updated guideline #5 to indicate “Subsequent courses of Rituxan (rituximab) should be administered every 24 weeks or based on clinical evaluation, but not sooner than every 16 weeks.” Update due to a change in Rituxan’s FDA-approved labeling. References added. (LR)

10/26/10 Added diagnoses codes to the “Billing/Coding” section. (lpr)

3/15/11 Specialty Matched Consultant Advisory Panel review 2/2011. Under “When Covered” section, moved #5 “Subsequent courses of Rituxan (rituximab) should be administered every 24 weeks or based on clinical evaluation, but not sooner than every 16 weeks” to Policy Guidelines section since this refers to continuation of treatment and not initial approval. Under “When Not Covered” section, added phrase “not medically necessary” to statement #1 “Rituximab for the treatment of rheumatoid arthritis is considered not medically necessary when the criteria stated above are not met. (lpr)

8/16/11 In the description section, added a cross reference to the Corporate Medical Policy, Monoclonal Antibodies for Non-Hodgkin Lymphoma, including Chronic Lymphocytic, &Acute Myeloid Leukemia in the Non-Hematopoietic Stem Cell Transplant Setting. (lpr)

11/22/11 Removed the x from the ICD-9 codes 714.0, 714.4, 714.8, 720.0 in the Billing/Coding section since there are no 5th digits for these codes. (lpr)
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3/20/12  Specialty Matched Consultant Advisory Panel review meeting 2/29/2012. Under “When Not Covered” statement 2, changed to not medically necessary. No change to policy statement. (lpr)

9/4/12  Added ICD-9 code 714.89 to the Billing/Coding section for 2012 code update. (lpr)

12/1/12  "When Covered" section modified to revise statement “The patient has failed to respond adequately to at least one Tumor Necrosis Factor (TNF) inhibiting drug or is intolerant to all TNF-inhibiting drugs (i.e., Remicade® (infliximab), Enbrel® (etanercept), or Humira® (adalimumab);” to “The patient has failed to respond adequately or is intolerant to Remicade ® (infliximab).” Notice 12/1/12 effective 2/1/13.

3/12/13  Specialty Matched Consultant Advisory Panel review meeting 2/20/2013. No change to policy statement. Reference added. (lpr)

7/1/13  Added ICD-10 codes to the “Billing/Coding” section. (lpr)

10/15/13  Added trial of Simponi Aria (golimumab) to statement #2 under When Covered section. Added ICD-10 diagnosis code M05.319 to Billing/Coding section. Medical director review 10/2013. (lpr)

12/31/13  Added ICD-10 diagnosis code M08.029 to Billing/Coding section and changed M05.11 to M05.111 for 2014 code update. (lpr)

4/1/14  Specialty Matched Consultant Advisory Panel review meeting 2/25/2014. Under Description section, changed DC20 to CD20 in description of the drug; changed “bad” cells to “targeted” cells to describe B cells. Medical director review. No change to policy statement. (lpr)

7/1/14  Removed ICD-10 effective date from Billing/Coding section. (lpr)

3/10/15  Specialty matched consultant advisory panel review meeting 2/25/2015. No change to policy statement. (lpr)

4/1/16  Description section and coverage guidelines updated for clarity. No change to policy intent. References added. Specialty Matched Consultant Advisory Panel review 2/24/16. (an)

12/30/16  Minor changes to Description section. Reference updated. No change to policy statement. (an)


Medical policy is not an authorization, certification, explanation of benefits or a contract. Benefits and eligibility are determined before medical guidelines and payment guidelines are applied. Benefits are determined by the group contract and subscriber certificate that is in effect at the time services are rendered. This document is solely provided for informational purposes only and is based on research of current medical literature and review of common medical practices in the treatment and diagnosis of disease. Medical practices and knowledge are constantly changing and BCBSNC reserves the right to review and revise its medical policies periodically.