Contractor Information

Contractor Name
First Coast Service Options, Inc.

Contractor Number
09102

Contractor Type
MAC - Part B

LCD Information

LCD ID Number
L29271

LCD Title
Rituximab (Rituxan®)

Contractor's Determination Number
J9310

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CMS National Coverage Policy
Language quoted from CMS National Coverage Determinations (NCDs) and coverage provisions in interpretive manuals are italicized throughout the Local Coverage Determination (LCD). NCDs and coverage provisions in interpretive manuals are not subject to the LCD Review Process (42 CFR 405.860[b] and 42 CFR 426 [Subpart D]). In addition, an administrative law judge may not review an NCD. See §1869(f)(1)(A)(i) of the Social Security Act.

Unless otherwise specified, italicized text represents quotation from one or more of the following CMS sources:

CMS Manual System, Pub. 100-02, Medicare Benefit Policy, Chapter 15, Section 50
CMS Manual System, Pub. 100-04, Medicare Claims Processing Manual, Chapter 17, Section 40
CMS Manual System, Pub. 100-08, Medicare Program Integrity, Chapter 13, Section 13.1.3

Primary Geographic Jurisdiction
Florida
Oversight Region
Region I

Original Determination Effective Date
For services performed on or after 02/02/2009

Original Determination Ending Date

Revision Effective Date

Revision Ending Date

Indications and Limitations of Coverage and/or Medical Necessity

Rituximab (Rituxan®)–J9310

Rituxan (Rituxan®) is a monoclonal antibody that targets a specific protein, known as CD20, on the surface of immune cells known as B-cells. Rituxan binds to CD20 and is believed to work with the body’s own immune system to attack and kill the marked B-cells.

Rituximab is FDA approved for the treatment of the following indications:

Non-Hodgkins Lymphoma (NHL)

- Patients with relapsed or refractory, low-grade or follicular, CD20-positive, B-cell, NHL as a single agent.

- Previously untreated diffuse large B-cell, CD20-positive, NHL in combination with CHOP or other anthracycline-based chemotherapy regimens.

- Previously untreated follicular, CD20-positive, B-cell NHL in combination with CVP chemotherapy.

- Non-progressing (including stable disease), low-grade, CD20-positive, B-cell NHL, as a single agent after first-line treatment with CVP chemotherapy.

Rheumatoid Arthritis (RA)

- In combination with methotrexate to reduce signs and symptoms and to slow the progression of structural damage in adult patients with moderately-to severely-active rheumatoid arthritis who have had an inadequate response to one or more TNF antagonist therapies.

Medicare will consider the use of Rituximab as medically reasonable and necessary for the FDA approved uses as well as the following off-labeled indications:
- Low grade or follicular CD20-positive, B-cell non-Hodgkin’s lymphomas (re-induction treatment appropriate for responders and patients with stable disease)

- Intermediate and high grade NHL when used as a single agent, in combination with a CHOP (Cyclophosphamide, Doxorubicin, Vincristine, and Prednisone) chemotherapy regimen, or in combination with other agents active in the disease

- Chronic lymphocytic leukemia

- Immune or idiopathic thrombocytopenia purpura

- Evans’ syndrome

- Waldenstrom’s Macroglobulinemia

- For the treatment of refractory thrombotic thrombocytopenic purpura (TTP) for patients who do not respond to plasmapheresis.

- Autoimmune hemolytic anemia

- Rituximab is covered for those patients with autoimmune hemolytic anemia condition that is refractory to conventional treatment (e.g., corticosteroid treatment and splenectomy).

Coding Information

Bill Type Codes:

Contractors may specify Bill Types to help providers identify those Bill Types typically used to report this service. Absence of a Bill Type does not guarantee that the policy does not apply to that Bill Type. Complete absence of all Bill Types indicates that coverage is not influenced by Bill Type and the policy should be assumed to apply equally to all claims.

Revenue Codes:

Contractors may specify Revenue Codes to help providers identify those Revenue Codes typically used to report this service. In most instances Revenue Codes are purely advisory; unless specified in the policy services reported under other Revenue Codes are equally subject to this coverage determination. Complete absence of all Revenue Codes indicates that coverage is not influenced by Revenue Code and the policy should be assumed to apply equally to all Revenue Codes.

99999 Not Applicable

CPT/HCPCS Codes
ICD-9 Codes that Support Medical Necessity

200.00 - 200.88  RETICULOSARCOMA UNSPECIFIED SITE - OTHER NAMED VARIANTS OF LYMPHOSARCOMA AND RETICULOSARCOMA INVOLVING LYMPH NODES OF MULTIPLE SITES

202.00 - 202.98  NODULAR LYMPHOMA UNSPECIFIED SITE - OTHER AND UNSPECIFIED MALIGNANT NEOPLASMS OF LYMPHOID AND HISTIOCYTIC TISSUE INVOLVING LYMPH NODES OF MULTIPLE SITES

204.10 - 204.12  CHRONIC LYMPHOID LEUKEMIA, WITHOUT MENTION OF HAVING ACHIEVED REMISSION - CHRONIC LYMPHOID LEUKEMIA, IN RELAPSE

273.3  MACROGLOBULINEMIA

283.0  AUTOIMMUNE HEMOLYTIC ANEMIAS

287.31  IMMUNE THROMBOCYTOPENIC PURPURA

287.32  EVANS' SYNDROME

446.6  THROMBOTIC MICROANGIOPATHY

714.0  RHEUMATOID ARTHRITIS

714.1  FELTY'S SYNDROME

714.2  OTHER RHEUMATOID ARTHRITIS WITH VISCERAL OR SYSTEMIC INVOLVEMENT

Diagnoses that Support Medical Necessity
See ICD-9 Codes that Support Medical Necessity

ICD-9 Codes that DO NOT Support Medical Necessity
All other diagnosis codes not listed as covered in the “ICD-9 Codes that Support Medical Necessity” section of this LCD.

XX000  Not Applicable

ICD-9 Codes that DO NOT Support Medical Necessity Asterisk Explanation
Diagnoses that DO NOT Support Medical Necessity
All other diagnoses not listed as covered in the “ICD-9 Codes that Support Medical Necessity” section of this LCD.

General Information

Documentation Requirements
Medical record documentation maintained by the ordering/referring physician must substantiate the medical need for the use of these chemotherapy drugs by clearly indicating the condition for which these drugs are being used. This might include the type of cancer, staging, if applicable, prior therapy and the patient’s response to that therapy. This documentation is usually found in the history and physical or in the office/progress notes.

If the provider of the service is other than the ordering/referring physician, that provider must maintain copies of the ordering/referring physician’s order for the chemotherapy drug. The physician must state the clinical indication/medical need for using the chemotherapy drug in the order.

For patients receiving Rituximab, an explanation of lymphoma type and previous treatment(s) should be maintained in the patient’s medical record.

For the off-label indication of autoimmune hemolytic anemia, in addition to the above documentation requirements, the following documentation must be supported in the medical record: Hgb and Hct, reticulocyte count, bilirubin, hepatoglobulin, indirect and direct hepatoglobulintests), patients subjective complaints.

Appendices

Utilization Guidelines
It is expected that these services would be performed as indicated by current medical literature and/or standards of practice. When services are performed in excess of established parameters, they may be subject to review for medical necessity.

Dosage for the off label indication of autoimmune hemolytic anemia is as follows: 375 mg/m2 once weekly for four consecutive weeks.

See the Food and Drug Administration (FDA) drug label for recommended dosages for specific FDA indications.

Rituximab (Rituxan®) is supplied as a 100mg/10mL and 500mg/50mL solution in a single-use vial.

Sources of Information and Basis for Decision


U.S. Food and Drug Administration, Department of Health and Human Services, CDER web site updates, January 2008.


Advisory Committee Meeting Notes

This Local Coverage Determination (LCD) does not reflect the sole opinion of the contractor or Contractor Medical Director. Although the final decision rests with the contractor, this LCD was developed in cooperation with advisory groups, which includes representatives from numerous societies.

Start Date of Comment Period

End Date of Comment Period

Start Date of Notice Period
12/04/2008

Revision History Number
Original
LCR B2009-
December 2008 Bulletin

This LCD consolidates and replaces all previous policies and publications on this subject by the carrier predecessors of First Coast Service Options, Inc. (Triple S and FCSO).

For Florida (00590) this LCD (L29271) replaces LCD L25077 as the policy in notice. This document (L29271) is effective on 02/02/2009.

Reason for Change

Last Reviewed On Date

Related Documents
This LCD has no Related Documents.

LCD Attachments
There are no attachments for this LCD.

All Versions
Updated on 11/30/2008 with effective dates 02/02/2009 - N/A