Overview

Useful For
Trough level quantitation for evaluation of patients with loss of response to infliximab and infliximab-dyyb

Profile Information

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<th>Reporting Name</th>
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<tr>
<td>INFX</td>
<td>Infliximab, S</td>
<td>No</td>
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Reflex Tests

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<tr>
<td>INXAB</td>
<td>Infliximab Ab, S</td>
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Testing Algorithm
Infliximab will be performed by liquid chromatography-tandem mass spectrometry (LC-MS/MS) on all samples. When Infliximab results are below 5.1 mcg/mL, testing for antibodies to infliximab will be performed at an additional charge.

Method Name
INFXR, INFX: Selective Reaction Monitoring Liquid Chromatography-Tandem Mass Spectrometry (LC-MS/MS)

INXAB: Electrochemiluminescent Bridging Immunoassay with Acid Dissociation

NY State Available
Yes

Specimen

Specimen Type
Serum Red

Specimen Required

Patient Preparation:

1. Draw blood immediately before next scheduled dose (trough specimen).

2. **For 12 hours before specimen collection do not** take multivitamins or dietary supplements containing biotin (vitamin B7), which is commonly found in hair, skin, and nail supplements and multivitamins.

Container/Tube: Red top (serum ger/SST are not acceptable)

Specimen Volume: 1 mL
Collection Instructions: Centrifuge within 2 hours of collection.

Forms
If not ordering electronically, complete, print, and send a Gastroenterology and Hepatology Client Test Request (T728) with the specimen.

Specimen Minimum Volume
0.5 mL

Reject Due To

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<td>Gross lipemia</td>
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<tr>
<td>Gross icterus</td>
<td>Reject</td>
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Specimen Stability Information

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<tbody>
<tr>
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<tr>
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Clinical and Interpretive

Clinical Information

Infliximab (Remicade, Renflexis, Inflectra) is a chimeric immunoglobulin (IgG1 kappa) targeting tumor necrosis factor-alpha (TNF-a), and it is currently FDA-approved for the treatment of multiple inflammatory conditions. Infliximab binds to soluble TNF-a and transmembrane homotrimers, which are found on the surface of macrophages and T-cells, with similar affinity. Infliximab has the ability to mediate complement-dependent cytotoxicity and antibody-dependent cell-mediated cytotoxicity, which leads to the lysis of target cells.

Infliximab pharmacokinetic properties may vary with disease and clearance is affected by concomitant use of immunosuppressants, high concentrations of TNF-a and C-reactive proteins,(1,2) low albumin concentrations, high body mass index, and presence of antibodies to infliximab (ATI), also known as human antichimeric antibodies (HACA).(3) Males seem to clear infliximab faster than females.(3)

Several studies have demonstrated that infliximab quantitation in the setting of loss of response to therapy can aid in patient management, as trough concentrations defined as therapeutic have been associated with superior clinical response and improved prognosis.(4-6)

Evaluation of infliximab concentrations may be of value for all inflammatory diseases for which it is prescribed. Primary indications for testing of infliximab include loss of response, partial response on initiation of therapy, autoimmune or hypersensitivity reactions, primary nonresponse, reintroduction after drug holiday, endoscopic/computed tomography enterography recurrence (in inflammatory bowel disease), and acute infusion reactions.

Measurement of infliximab concentrations is indicated at trough, immediately prior to the next scheduled infusion. Low trough concentrations may be correlated with loss of response to infliximab. Assessment of antibodies to
Infliximab is suggested when infliximab quantitation at trough is 5.0 mcg/mL or less. Infliximab concentrations tend to stabilize after 14 weeks (approximately 100 days). Quantitation of peak infliximab concentrations is strongly discouraged.

The ATI assay has been verified to analyze infliximab and infliximab-dyyb (Inflectra, Pfizer Inc) with no analytical differences between the 2 drugs quantitation. Inflectra has the same primary amino acid sequence as Remicade and Renflexis. Therefore, “infliximab” will be used to refer to both the reference product and the biosimilar product interchangeably.

A biosimilar product is a biological product that is approved based on showing that it is highly similar to an FDA-approved biological product, known as the reference product. No clinically meaningful differences in terms of safety and effectiveness from the reference product are present. Only minor differences in clinically inactive components are allowable in biosimilar products. In contrast to generic medications, a prescription of biosimilars needs to come from the ordering physician and not the dispensing pharmacy (pharmacies cannot substitute a biosimilar for another medication; a separate prescription is required).

**Reference Values**

**INFLIXIMAB QUANTITATION:**

Limit of quantitation is 1.0 mcg/mL. Therapeutic ranges are disease specific.

Pediatric reference ranges are not established.

**INFLIXIMAB ANTIBODIES**

Absence of antibodies to infliximab (ATI) is defined as <50 U/mL

Presence of ATI is reported as positive when concentrations are > or =50 U/mL

**Interpretation**

Low trough concentrations may be correlated with loss of response to infliximab. For infliximab trough concentrations 5.0 mcg/mL or less, testing for antibodies to infliximab (ATI) is suggested.

For infliximab trough concentrations above 5.0 mcg/mL, the presence of ATI is unlikely; patients experiencing loss of response to infliximab may benefit from an increased dose or a shorter infusion interval.

Results above 35 mcg/mL are suggestive of a blood draw at a time-point in treatment other than trough.

**Cautions**

Toxicity effects other than acute hypersensitivity infusion reactions have not been described nor correlated with infliximab concentrations.

During the initial induction phase of treatment (weeks 0, 2, and 6), steady-state has not been achieved and concentrations of infliximab may vary significantly between infusions.(3)

Therapeutic concentrations of infliximab may vary according to the disease (eg, Crohn disease versus ulcerative colitis versus rheumatoid arthritis).

Samples containing more than 12.5 ng/mL biotin (vitamin B7) may interfere (in the form of depressed signal) with INXAB / Infliximab Antibodies, Serum.

For antibodies-to-infliximab (ATI), pediatric and adult reference ranges were validated, and the presence of an ATI is
established as greater than or equal to 50 U/mL by our bridging electrochemiluminescent/acid dissociation method.

The presence of endogenous infliximab is a recognized interference in most ATI methods. This assay includes an acid dissociation step, which partially mitigates this interference. Tolerance up to 12.5 mcg/mL infliximab has been documented, although this is also determined by the titer of the ATI present in the patient sample.

**Clinical Reference**


**Performance**

**Method Description**

**Infliximab Quantitation:**

This test is performed using liquid-chromatography and tandem mass spectrometry. Preanalytical sample preparation includes a trypsin digestion; 2 unique clonotypic peptides from the heavy and light chains of the infliximab chimeric structure (IgG1 kappa) are monitored. (Willrich MA, Murray DL, Barnidge DR, et al: Quantitation of infliximab using clonotypic peptides and selective reaction monitoring by LC-MS/MS. Int Immunopharmacol. 2015;28:513-520)

**Infliximab Antibodies:**

This lab developed immunoassay is designed to measure antibodies-to-infliximab (ATI) in human serum by means of electrochemiluminescence (ECL) on the MesoScale Discovery (MSD) platform. The assay uses a "bridging" format in which the ATI forms a link between biotin labeled infliximab and SULFO-Tag labeled infliximab. The biotin binds to a streptavidin (SA) coated surface and the SULFO-Tag creates a signal in the presence of a conjugate following application of an electric current. During sample preparation, serum is mixed with acetic acid to break the infliximab/ATI complex. Biotinylated and SULFO-Tagged infliximab are then added and bind with ATI that is present in the sample. After the incubation with the labeled drug, the calibrators, controls, and samples are added to a SA plate that has been blocked with a solution of bovine serum albumin (BSA). The biotinylated infliximab then binds to the SA plate. After an incubation period, the SA plate is washed and MSD read buffer is added. Immediately after the
addition of read buffer, the plate is analyzed. The read buffer provides an appropriate chemical environment for ECL when a voltage is applied to the electrodes on the plate. This voltage causes bound SULFO-Tagged infliximab to emit measureable light. The intensity of emitted light is measured and correlated to a set of standards with known concentrations of ATI by means of a 4-point logistics curve fitting method. (Willrich M, Balsanek J, Ladwig P, et al: A-374 Antibodies-to-Infliximab: Assay Development and Correlation with Infliximab Concentrations in Serum Samples of Treated Patients. AACC Annual Meeting; Chicago: AACC Press; 2014 pS110; Balsanek J, Willrich MAV, Murray DL, Snyder M: Sa1268 Clinical Development of an Electrochemiluminescent Immunoassay to Measure Antibodies-to-Infliximab. Gastroenterology. 2014;146[5]:S-248)

PDF Report

No

Day(s) and Time(s) Test Performed

INFX: Monday, Wednesday, Thursday; 4 p.m.

INXAB: Monday, Wednesday, Friday; 8 a.m.

Analytic Time

3 days

Maximum Laboratory Time

6 days

Specimen Retention Time

2 weeks

Performing Laboratory Location

Rochester

Fees and Codes

Fees

- Authorized users can sign in to Test Prices for detailed fee information.
- Clients without access to Test Prices can contact Customer Service 24 hours a day, seven days a week.
- Prospective clients should contact their Regional Manager. For assistance, contact Customer Service.

Test Classification

This test was developed and its performance characteristics determined by Mayo Clinic in a manner consistent with CLIA requirements. This test has not been cleared or approved by the U.S. Food and Drug Administration.

CPT Code Information

80230

82397-(if appropriate)

LOINC® Information

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