Overview

Useful For
Risk assessment for development of hepatocellular carcinoma in patients with chronic liver disease

Highlights
This test includes a GALAD (gender, age, alpha-fetoprotein [AFP]-L3, AFP, des-gamma-carboxy prothrombin [DCP]) model score calculation to calculate the probability of hepatocellular carcinoma (HCC). The GALAD model has been demonstrated to have higher diagnostic accuracy for the detection of HCC when compared to the use AFP, AFP-L3, and DCP markers alone or in combination. The performance of the GALAD score has also been reported to be superior to ultrasound for HCC detection.

Profile Information

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Method Name
Isotachophoresis with Laser-Induced Fluorescence

NY State Available
Yes

Specimen

Specimen Type
Serum

Advisory Information
GALAD score testing (this test) should not be performed for pregnant patients as alpha-fetoprotein results are elevated during pregnancy.

Necessary Information
Gender and age are required.

Specimen Required

Container/Tube:

Preferred: Red top

Acceptable: Serum gel
**Specimen Volume:** 0.5 mL

**Collection Instructions:** Centrifuge and aliquot serum.

**Specimen Minimum Volume**
0.25 mL

**Reject Due To**

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**Specimen Stability Information**

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**Clinical and Interpretive**

**Clinical Information**

Worldwide, hepatocellular carcinoma (HCC) is the third leading cause of cancer-related death.(1) While HCC can be treated effectively in its early stages, most patients are not diagnosed until they are symptomatic and at higher grades and stages, which are less responsive to therapies. Alpha-fetoprotein (AFP) is the standard serum tumor marker utilized in the evaluation of suspected HCC. However, increased serum concentrations of AFP might be found in chronic hepatitis and liver cirrhosis, as well as in other tumor types (eg, germ cell tumors)(2), decreasing the specificity of AFP testing for HCC. Furthermore, AFP is not expressed at high levels in all HCC patients, resulting in decreased sensitivity, especially in potentially curable small tumors.

**AFP-L3:**

AFP is differentially glycosylated in several hepatic diseases. For example, UDP-alpha-(1→6)-fucosyltransferase is differentially expressed in hepatocytes following malignant transformation.(3) This enzyme incorporates fucose residues on the carbohydrate chains of AFP. Different glycosylated forms of AFP can be recognized following electrophoresis by reaction with different carbohydrate-binding plant lectins. The fucosylated form of serum AFP that is most closely associated with HCC is recognized by a lectin from the common lentil (*Lens culinaris*). This is designated as AFP-L3 (third electrophoretic form of lentil lectin-reactive AFP). AFP-L3 is most useful in the differential diagnosis of individuals with total serum AFP of 200 ng/mL or less, which may result from a variety of benign pathologies, such as chronic liver diseases.

**Des-gamma-carboxy prothrombin (DCP):**

(DCP, also known as the protein induced by vitamin K absence or antagonist II (PIVKA-II), is an abnormal form of the coagulation protein, prothrombin. DCP is a nonfunctional prothrombin resulting from a lack of carboxylation of 10 glutamic acid residues in the N-terminal portion of the molecule. In normal liver, prothrombin undergoes post-translational carboxylation before release into the peripheral blood. The carboxylation converts specific amino-terminal glutamic acid residues to gamma-carboxyglutamic acid. The vitamin K dependent carboxylase responsible
for the carboxylation is absent in many HCC cells, and an abnormal prothrombin with all or some of unconverted glutamic acid is secreted. Therefore, this noncarboxylated form (DCP) has been used as an HCC biomarker.

DCP is considered a complementary biomarker to AFP and AFP-L3 for assessing the risk of developing HCC. The elevation of both AFP-L3 and DCP indicate progression of HCC, albeit they reflect different features of the progression. In a prospective study of patients in the United States with an established diagnosis of HCC, the sensitivities for AFP, AFP-L3, and DCP were 68%, 62%, and 73%, respectively. When the 3 markers were combined, the sensitivity was 86%. In another study, DCP levels were shown to correlate with tumor size and metastatic HCC. In this study, compared to AFP and AFP-L3, DCP had the highest sensitivity (87%) and the highest positive predictive value (87%) in patients with HCC due to chronic hepatitis B and C infections. A number of studies have shown that elevated serum DCP is significantly related to portal vein invasion or intrahepatic metastasis, which significantly affect prognosis for patients with HCC.

DCP can be elevated in other conditions besides HCC. Conditions such as obstructive jaundice, intrahepatic cholestasis causing chronic decrease in vitamin K, and ingestion of drugs such as warfarin or wide-spectrum antibiotics can result in high concentrations of DCP. In addition, 25% to 50% of patients with HCC will have a DCP value within the reference range. Because of this, a normal DCP value does not rule out HCC.

Gender, Age, AFP-L3, AFP, DCP (GALAD) Score and Probability of HCC:

Biomarkers of HCC include AFP, AFP-L3, and DCP. The GALAD model combines these three biomarkers with the patient’s gender and age to estimate the probability of HCC in patients with chronic liver disease based on the following equation $Z = -10.08 + 0.09 \times \text{age} + 1.67 \times \text{sex} + 2.34 \log(10)(\text{AFP}) + 0.04 \times \text{AFP-L3} + 1.33 \times \log(10)\text{(DCP)}$, where sex = 1 for males, 0 for females. The probability estimate of HCC is calculated as follows

$$\Pr(\text{HCC}) = \exp(Z)/(1 + \exp[Z]).$$

The GALAD model has been demonstrated to have higher diagnostic accuracy for the detection of HCC when compared to the use AFP, AFP-L3, and DCP markers alone or in combination. The performance of the GALAD score has also been reported to be superior to ultrasound for HCC detection.

Reference Values

TOTAL ALPHA-FETOPROTEIN (AFP):

<4.7 ng/mL

AFP L3-PERCENT:

<10%

DES-GAMMA-CARBOXY PROTHROMBIN:

<7.5 ng/mL

GAL1:

Not applicable

GAL2:

Not applicable

Interpretation
Alpha-fetoprotein (AFP)-L3 results of 10% or more are associated with a 7-fold increased risk of developing hepatocellular carcinoma (HCC). Patients with AFP-L3 levels of 10% or more should be monitored more intensely for evidence of hepatocellular carcinoma according to current practice guidelines.

Total serum AFP results above 200 ng/mL are highly suggestive of a diagnosis of HCC. In patients with liver disease, a total serum AFP level above 200 ng/mL is near 100% predictive of HCC. With decreasing total AFP levels, there is an increased likelihood that chronic liver disease, rather than HCC, is responsible for the AFP elevation.

Based on a retrospective study at Mayo Clinic, for patients with total AFP levels 200 ng/mL or less, AFP-L3 specificity approaches 100% for HCC when its percentage exceeds 35% of the total AFP.(4)

AFP concentrations over 100,000 ng/mL have been reported in normal newborns, and the values rapidly decline in the first 6 years of life.

Des-gamma-carboxy prothrombin (DCP):

In patients with an elevated DCP result (> or = 7.5 ng/mL), the risk of developing HCC is 36.5% (95% CI 23.5%-49.6%). The risk of developing HCC with a negative DCP result (<7.5 ng/mL) is 7.6% (95% CI 4.4%-10.8%).

Gender, Age, AFP-L3, AFP, DCP (GALAD) Score and Probability of HCC:

The probability of the presence of HCC is estimated from the GALAD model score. Higher GALAD model scores correlate with increased risk of HCC. The area under the curve (AUC) of a receiver operating characteristic (ROC) curve of the GALAD score was 0.95 for all HCC detection, and 0.92 for the detection of early stage HCC. Additionally, the AUC of the GALAD score (0.95) was higher than that of ultrasound alone for all HCC detection (AUC of 0.82, P < 0.01).

The sensitivity and specificity performance characteristics of the GALAD score for HCC will be influenced by the selected GALAD score cut-off. For example at an optimal AUC cutoff of -0.76, the GALAD score had 91% sensitivity and 85% specificity for HCC detection. At a more specific GALAD score cutoff of 0.88, the observed sensitivity was 80% for HCC detection with an observed specificity of 97%.

The GALAD model was developed and validated in patient cohorts with a prevalence of HCC ranging from 35% to 49%. The performance of the model may be altered in populations with different HCC prevalence. In addition, the clinical performance of the GALAD score varies by etiology of HCC and therefore may be different in different regions of the world.

Cautions

Serum markers are not specific for malignancy, and values may vary by method. Do not interpret alpha-fetoprotein (AFP), AFP-L3, and des-gamma-carboxy prothrombin (DCP) levels as absolute evidence of the presence or absence of malignant disease. Results should be used in conjunction with information from the clinical evaluation of the patient, cytology, and imaging procedures.

Values obtained with different assay methods or kits cannot be used interchangeably.

Some patients who have been exposed to animal antigens, either in the environment or as part of treatment or imaging procedures, may have circulating anti-animal antibodies present. These antibodies may interfere with the assay reagents to produce unreliable results in the AFP-L3 and DCP assays.

Test results for AFP are not interpretable if the patient is pregnant.
DCP-producing tumors other than hepatocellular carcinoma can show elevated DCP values. Liver disease caused by other etiologies such as alcohol-induced liver disease, hemochromatosis, Wilson disease, autoimmune hepatitis, and steatohepatitis have not been studied with the DCP assay.

Medications containing vitamin K preparations may cause a negative bias with DCP values. Medications containing vitamin K antagonist or antibiotic may cause a positive bias with DCP values.

The total AFP and AFP-L3 test values must be obtained using the uTASWako i30 in the GALAD score calculation.

**Clinical Reference**


**Performance**

**Method Description**
Testing is performed using the uTASWako i30 instrument and the test system reagents.

Alpha-Fetoprotein (AFP) L3:

Total AFP is measured by laser-induced fluorescence with separation of the lentil lectin-reactive AFP-L3 and lectin nonreactive forms of AFP by isotachophoresis of their immune-complexes. Results are expressed as the percent ratio of AFP-L3 to total AFP. (Package insert: uTASWako i30 AFP-L3, Wako Diagnostics, Richmond, VA, 06.1.24K02)

Des-Gamma-Carboxy Prothrombin (DCP):

Sample is added to the reagent well with the fluorescent dye labeled antihuman prothrombin antibody (mouse monoclonal) to form the primary immunocomplex. The second labeled antibody solution, anion-conjugated antihuman DCP antibody (mouse monoclonal), is concentrated by isotachophoresis when voltage is applied. The concentrated anion-conjugated antibody then reacts with the primary immunocomplex to form the secondary immunocomplex. This secondary complex is further concentrated during isotachophoresis and is separated from unbound fluorescent dye-labeled antibody by capillary gel electrophoresis. The remaining dye labeled DCP is measured by laser-induced fluorescence. The concentration of DCP in the specimen is directly proportional to the amount of fluorescence. (Package insert: uTASWako i30 DCP, Wako Diagnostics, Richmond, VA. V 11.03.08K02)

Gender, Age, AFP-L3, AFP, DCP (GALAD) Model Score and Probability of Hepatocellular Carcinoma (HCC):

The GALAD model is a statistical model for estimating the likelihood of HCC in patients with chronic liver disease. The GALAD score is calculated based on gender, age, and measured concentrations of AFL-L3, AFP and DCP.

PDF Report
No

Day(s) and Time(s) Test Performed
Monday, Wednesday, Friday; 10 a.m.

Analytic Time
Same day/1 day

Maximum Laboratory Time
3 days

Specimen Retention Time
12 months

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 has been cleared, approved or is exempt by the U.S. Food and Drug Administration and is used per manufacturer's instructions. Performance characteristics were verified by Mayo Clinic in a manner consistent with CLIA requirements.

**CPT Code Information**

82107

83951

**LOINC® Information**

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