



Herpes Simplex Virus Typing

IVD For In Vitro Diagnostic Use

INTENDED USE

ImmunoDOT Herpes Simplex Virus (HSV) Typing test is an enzyme immunoassay (EIA) detecting HSV or glycoprotein G (gG) type specific IgG antibodies. The test detects the presence or absence of past HSV exposure and specifically determines whether past infection(s) is due to HSV Type 1, Type 2 or both Type 1 and 2.

SUMMARY AND EXPLANATION

There are two herpes antigenic types (1,2). "Definitive diagnosis of genital herpes infections is fundamental to the management of patients and the development of strategies to prevent transmission to partners and neonates" (3). Such diagnosis has proven inaccurate when based solely on clinical history and impression (4). Instead, virus, antigen or nucleic acid detection and classification is used for patients presenting with lesions or type-specific serological tests may be used when lesions are absent.

For type specific serology, either western blot (5,6,7) or assays or type specific protein (8,9) is used. Acceptable type specific classification is not possible using whole virus lysate, the commonly used antigen of early HSV serology kits. The most commonly used type specific protein is glycoprotein G. ImmunoDOT HSV Typing test uses HSV gG type 1 and type 2 recombinant proteins.

ASSAY PRINCIPLE

The assay uses an enzyme-linked immunoassay (EIA) dot technique for the detection of antibodies to HSV-1 gG, HSV-1 virus lysate proteins, HSV-2 gG or HSV-2 virus lysate proteins. An assay strip is incubated with dilute patient serum, allowing patient antibodies reactive with the test antigens to bind to the solid support membrane. In the second stage, the reaction is enhanced by removal of non-specifically bound materials. During the third stage, alkaline phosphatase-conjugated antihuman antibodies are allowed to react with bound patient antibodies. Finally, the strip is transferred to enzyme substrate reagent which reacts with bound alkaline phosphatase to produce an easily seen, distinct dot.

REAGENTS

Assay Strip. Antigens in order beginning with the window next to the label: Positive reagent control (human serum); [Total 1] HSV-1, [Total 2] HSV-2 lysate, [Specific 1] HSV-1 gG, [Specific 2] HSV-2 gG and Negative reagent control.

Diluent (#1) Consists of buffered diluent containing protein stabilizers with <0.1% NaN₃.

Enhancer (#2) Consists of sodium chloride with <0.1% NaN₃.

Conjugate (#3) Consists of alkaline phosphatase conjugated goat antihuman IgG (heavy chain specific) in buffered diluent with <0.1% NaN₃.

Developer (#4) Consists of 5-bromo-4-chloro-3-indolyl phosphate and p-nitro blue tetrazolium chloride in buffered diluent with <0.1% NaN₃.

HSV-1 Positive Control Consists of HSV-1 IgG positive human serum that is HSV-2 negative containing protein stabilizers and <0.1% NaN₃.

HSV-2 Positive Control Consists of HSV-2 IgG positive human serum that is HSV-1 negative containing protein stabilizers and <0.1% NaN₃.

Negative Control Consists of HSV-1 and HSV-2 negative human serum containing protein stabilizers and <0.1% NaN₃.

Warnings and Precautions

For In Vitro Diagnostic Use: These reagents have been optimized for use as a system. Do not interchange assay strips. Dilution or adulteration of these reagents may also affect the performance of the test. Do not use kit if evidence of microbial contamination is present. Do not use any kits beyond the stated expiration date. Analytic quality deionized or distilled water must be used as Clarifier. Close adherence to the test procedure

will assure optimal performance. Do not shorten or lengthen stated incubation times since this may result in poor assay performance.

Some assay components contain sodium azide (NaN₃) that may react with lead and copper plumbing to form highly explosive metal azides. On disposal, flush with a large volume of water to prevent azide buildup.

Warning - Potential Biohazardous Material: Human sera used in the preparation of this product were tested and found non-reactive for hepatitis B surface antigen and for antibodies to HIV-1, HIV-2, and hepatitis C virus. Because no test method can offer complete assurance that infectious agents are absent, handle reagents and patient samples as if capable of transmitting infectious disease.

Storage

Store reagents and assay strips at 2-8°C. Reagents must be at room temperature (15-30°C) before use. Avoid contamination of reagents.

SPECIMEN COLLECTION AND HANDLING

ImmunoDOT Test is performed on serum. The test requires 50 µL of serum. Lipemic or hemolyzed serum has not been shown to be an acceptable specimen.

Store samples at room temperature for no longer than eight hours. If the assay will not be completed within eight hours, refrigerate the sample at 2-10°C. If the assay or shipment of the samples will not be completed within 48 hours, freeze at -20°C.

PROCEDURE

Materials Provided

Assay Strips	Reaction Vessels
Diluent (#1)	Package Insert
Enhancer (#2)	HSV-1 Positive Control
Conjugate (#3)	HSV-2 Positive Control
Developer (#4)	Negative Control

Materials Required But Not Provided

- GenBio Workstation
- Specimen collection apparatus
- Timer
- Analytic quality distilled or deionized water to be used as Clarifier
- Pipets
- Absorbent toweling to blot dry the assay strip

Set-Up

1. Turn on Workstation and adjust to appropriate temperature if necessary. Refer to Workstation Instructions.
2. Remove 4 Reaction Vessels per test from the product box and insert into appropriate slots in Workstation. For the large Workstation, add water up to the fill line of the Clarifier Vessel provided. For the small Workstation, use an appropriate container and sufficient water to cover all reactive windows of the assay strip.
3. Place 2 mL Diluent (#1) in Reaction Vessel #1; 2 mL Enhancer (#2) in Reaction Vessel #2; 2 mL Conjugate (#3) in Reaction Vessel #3; and 2 mL Developer (#4) in Reaction Vessel #4.
4. Appropriately label the Assay Strips.
5. If the large Workstation is used, insert the label end of the assay strip into the Strip Holder, one per groove, taking care not to touch the assay windows.

Assay Procedure

1. Add 50 µL serum to Reaction Vessel #1.
2. Prewet Assay Strip by immersing in Clarifier for 30-60 seconds.
3. Using several (5-10) quick up and down motions with the Assay Strip, mix thoroughly in Reaction Vessel #1. Let stand for 60-90 minutes.
4. Remove Assay Strip from Reaction Vessel and swish in the Clarifier. Use a swift back and forth motion for 5-10 seconds allowing for optimal washing of the Assay Strip's membrane windows.
5. Place Assay Strip into Reaction Vessel #2. Mix thoroughly with several (5-10) quick up and down motions. Let stand for 5 minutes.

6. Remove Assay Strip from Reaction Vessel #2 and swish in Clarifier as described (step #4).
7. Place Assay Strip into Reaction Vessel #3. Mix thoroughly with several (5-10) quick up and down motions. Let stand for 30-40 minutes.
8. Remove Assay Strip from Reaction Vessel #3 and swish in Clarifier as described (step #4). DO NOT remove the Assay Strip from the Clarifier.
9. Allow the Assay Strip to stand in the Clarifier for 5 minutes.
10. Remove Assay Strip from Clarifier and place into Reaction Vessel #4. Mix thoroughly with several (5-10) quick up and down motions. Let stand for 5 minutes.
11. Remove Assay Strip from Reaction Vessel #4 and swish in Clarifier as described (step #4).
12. Blot and allow Assay Strip to dry. It is imperative that tests of borderline specimens be interpreted after the Assay Strip has been allowed to dry. **A false positive dot may be identified if the assay strip is not dry when interpreted.**

such as western blot may be used. However, it is also possible that no type specific antibody is present and therefore typing is not possible.

- As with other serological tests, negative results do not rule out the diagnosis of herpes simplex disease.
- A single positive result only indicates previous immunologic exposure. Other methods such as nucleic acid, antigen or viral culture are required to establish current infection.
- It is unlikely that patterns other than presented will occur. Repeat the testing and if the pattern repeats, contact the manufacturer.

Reading the Assay Strip

Positive	A dot with an EASILY SEEN , distinct border is visible in the center of the window. The outer perimeter of the window must be white to pale gray.
Negative	If no dot is seen or a dot is difficult to see, interpret it as negative. Each window of the assay strip is read independently.

In order to minimize the possibility of "over interpreting" positive test results it is recommended that during initial validation of the assay (as may be required by the laboratory by regulation), the laboratory test a series of presumptive negative samples and each technician interpret the assay strips in a blinded fashion. Please call GenBio Technical Service for further clarification. (To report results, refer to Interpretation Section)

Quality Control

The assay's reagent temperature is between 42-48°C. Due to heat transfer loss, the Workstation temperature is set higher. The appropriate Workstation temperature setting is listed in the Workstation's package insert. (Contact Technical Services for additional guidance if an alternate heat source is used.)

NCCLS C24-A may be consulted for guidance on appropriate quality control practices. These should be tested according to guidelines or requirements of local, state, and/or federal regulations or accrediting organizations. Unless otherwise required, it is recommended that control sera be tested upon receipt of a kit. If the control is not reactive, results should not be reported and GenBio Technical Service should be contacted before the kit is used again.

The kit uses reagent controls to assure performance each time a test is performed. The Positive Control window (well #1) contains human serum and tests reagent reactivity. It must be reactive but the intensity must not be used as a calibrator. As a negative reagent control check, the backgrounds around dots and the bottom window (#6) must be white. If the positive is not reactive or the negative is reactive, do not interpret the assay strip.

INTERPRETATION

<u>Interpretation</u>	<u>Total 1</u>	<u>Total 2</u>	<u>Specific 1</u>	<u>Specific 2</u>
Negative	-	-	-	-
HSV Type 1	+	+	+	-
HSV Type 2	+	+	-	+
HSV Type 1&2	+	+	+	+
HSV Positive	+	+	-	-
Type Unknown	+	+	-	-

- All results from this and other serologies must be correlated with clinical history, epidemiological data and other data available to the attending physician in evaluating the patient.
- It is expected that some patient samples have both HSV Total 1 and 2 positive and that both Specific 1 and 2 negative. This indicates either a recent infection or that the patient never formed type specific antibody. If a recent infection, isolation, antigen or nucleic acid detection methods are indicated. Otherwise, another method

LIMITATIONS

The performance of this assay has not been established for ruling out diseases with similar symptoms, e.g., *Candida albicans*, Bacterioides species, *G. vaginalis*, Mobiluncus species. Instead, also use culture or other appropriate methods.

EXPECTED RESULTS

It has been shown that gG specific antibody typically develops over the course of six months. Appearance is slower and later than antibodies to other, non-type specific proteins. Therefore, it is expected that some patients will be HSV antibody positive yet anti-gG negative.

Eighty-four sera collected from U.S. blood donors were tested using ImmunoDOT HSV Typing kit. These results are shown in Table 1.

Table 1

<u>Interpretation</u>	<u>Percent</u>
Negative	15.5%
HSV Type 1	36.9%
HSV Type 2	11.9%
HSV Type 1&2	32.1%
HSV Positive Type Unknown	3.6%

PERFORMANCE CHARACTERISTICS

Eighty-four sera collected from U.S. blood donors were tested using: ImmunoDOT HSV Typing kit [IDOT], an alternate commercial microtiter kit for specific detection of either type 1 or type 2 gG antibodies (Kit A), and two alternate commercial microtiter kits for detection of HSV lysate antibodies (Not able to specify type) [Kits B and C].

Total (Traditional Method)

Comparison between ImmunoDOT Total 1 and 2 results and two commercial microtiter kits (Kits B and C) is shown in Table 2. Agreement is 100% and therefore sensitivity and specificity are 100%.

Table 2

<u>Alternate Kit</u>	<u>ImmunoDOT</u>	
	Negative	Positive
Negative	13	0
Positive	0	71

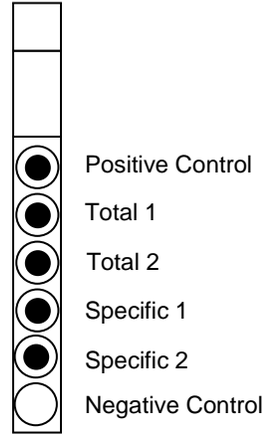
Specific (HSV Serology Type)

Comparison between ImmunoDOT Specific results and the alternate commercial type specific microtiter kit is shown in Table 3.

One sample is HSV antibody negative using kits B and C, reported negative by ImmunoDOT but reported HSV Type 2 by the Alternate gG assay. This is considered a false positive and not shown in Table 3. Agreement between the two gG assays is 90% (75/83).

Table 3

<u>ImmunoDOT</u>	<u>Alternate</u>	<u>Number</u>
Negative	Negative	15 (18%)
Type 1	Type 1	32 (39%)
Type 2	Type 2	9 (11%)
Type 1&2	Type 1&2	22 (27%)
Type 1	Negative	0
Type 1	Type 2	0
Type 1	Type 1&2	0
Type 2	Negative	0
Type 2	Type 1	0
Type 2	Type 1&2	2 (2%)
Type 1&2	Negative	0
Type 1&2	Type 1	2 (2%)
Type 1&2	Type 2	0
Negative	Type 1	1 (1%)
Negative	Type 2	0
Negative	Type 1&2	0



There is 94% (78/84) agreement between the two methods. Assuming alternate test is correct (relative performance) is listed in Table 4.

Table 4

<u>Interpretation</u>	<u>Relative Sensitivity</u>	<u>Relative Specificity</u>
Negative	100% (15/15)	100% (15/15)
Type 1	95% (56/59)	100% (15/15)
Type 2	94% (31/33)	100% (15/15)

Precision

Like any visually interpreted test, dot intensity is directly related to precision. The darkest dots are most reliable while weaker reactions (less intensity) are proportionately less reliable (equivocal or borderline).

Bibliography

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QUICK REFERENCE PROCEDURE

ImmunoDOT HSV IgG Typing

Set-Up

- Make sure Workstation is at temperature.
- Place reaction Vessels into slots in Workstation and add water to the Clarifier Vessel.
- Place 2 mL Diluent (1) in Vessel #1; 2 mL Enhancer (2) in Vessel #2; 2 mL Conjugate (3) in Vessel #3; and 2 mL Developer (4) in Vessel #4.

Procedure

- Add 50 µL serum to Vessel #1.
- **Prewet** assay strip in Clarifier for 30 - 60 seconds.
- Place strip in Vessel #1, mix, **let stand 60-90 min.**
- Remove strip, place in Clarifier, swish 5-10 sec.
- Place strip in Vessel #2, mix, **let stand 5 min.**
- Remove strip, place in Clarifier, swish 5-10 sec.
- Place strip in Vessel #3, mix, **let stand 30-40 min.**
- Remove strip, place in Clarifier, **let stand 5 min.**
- Place strip in Vessel #4, mix, **let stand 5 min.**
- Remove strip, place in Clarifier, swish, blot, dry, and read.

To place an order for ImmunoDOT products, contact your local distributor or call GenBio directly for the distributor nearest you and for additional product information.

For assistance, please call toll-free 800-288-4368.



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