INTENDED USE
The INFINITI System Assay for Factor II & Factor V is an in vitro diagnostic device that consists of reagents and instrumentation which includes polymerase chain reaction (PCR) primers, hybridization matrices, a thermal cycler, an imager, and software for detection and genotyping of Factor II (Prothrombin) G20210A and Factor V Leiden G1691A point mutations in DNA obtained from human blood samples. The INFINITI System Assay for Factor II & Factor V is a qualitative assay for use in clinical laboratories upon prescription by the attending physician.

The INFINITI System Assay for detection and genotyping of Factor II & Factor V is indicated for use as an aid to diagnosis in the evaluation of patients with suspected thrombophilia.

BACKGROUND INFORMATION
Genetic risk factors are involved in the predisposition of individuals to venous thrombosis \(^{(1)}\). The most common mutation associated with inherited thrombosis is Factor V Leiden G1691A mutation and results in resistance to activated protein C. This mutation has a relatively high prevalence in the general population (about 5% in Caucasians), and accounts for 85% to 95% of activated protein C resistant cases.\(^{(2)}\) Factor V Leiden G1691A mutation renders an enhanced risk of venous thrombosis, with odds ratios (ORs) of 3 to 8 in heterozygotes and 30 to 140 in homozygotes.\(^{(3)}\)

The second most common mutation associated with hereditary thrombosis is the G20210A mutation in the prothrombin (Factor II) gene. This is associated with increased plasma prothrombin levels and is present in 1% to 2% of the general population.\(^{(2)}\) Heterozygote carriers of the prothrombin G20210A mutation have an estimated 3 to 8-fold increased risk for venous thrombosis. Homozygote carriers are very rare and information on the increased risk for thrombosis associated with the homozygous mutation is very limited.

Heterozygosity for both Factor V Leiden G1691A and Factor II (Prothrombin) G20210A mutations results in earlier onset of thrombosis and greater severity as compared to the presence of either mutation alone in the heterozygous state.

TEST PRINCIPLE/ASSAY OVERVIEW
The INFINITI System Assay for Factor II & Factor V is designed to simultaneously detect mutations of two genes: Factor II (G20210A) and Factor V (G1691A). The assay protocol is based on five major processes:

- (a) DNA extraction from human blood samples
- (b) PCR amplification of purified DNA
- (c) Fluorescent label incorporation using analyte specific primer extension (ASPE)
- (d) Hybridization of the ASPE primers to a microarray followed by washing
- (e) Scanning of the microarray
- (f) Signal detection and analysis (determination of the Factor II and Factor V genotypes)

Steps (c) through (f) are automated by the INFINITI Analyzer

A schematic overview of the assay is shown below.
DEVICE DESCRIPTION
The INFINITI System Assay for Factor II & Factor V is an in vitro diagnostic device which utilizes AutoGenomics’ proprietary film-based microarray technology combined with process automation, reagent management and software technology for the detection and genotyping of the Factor II (Prothrombin) G20210A and the Factor V Leiden G1691A mutations from deoxyribonucleic acid (DNA) obtained from human whole peripheral blood samples.

The INFINITI System Assay for Factor II & Factor V is comprised of the BioFilmChip™ Microarray, the Intellipac™ Reagent Module, and the INFINITI Analyzer with the Qmatic™ Operating Software.

The BioFilmChip Microarray consists of a polyester film coated with proprietary multi-layer components designed for DNA analysis. The layers have been designed to provide a versatile surface to enhance test performance. The microarrays are designed to be assay specific. The INFINITI System Assay for Factor II & Factor V uses a microarray chip (L-Chip) which contains unused Capture Probes which could potentially be used for certain specific assays. Therefore, multiple assays can be developed using the same microarray.

The Intellipac Reagent Module which acts as a communication link contains four reservoirs that house the test reagents and has an integrated memory chip. Reagent information such as lot number, expiration date, and volume usage are archived in the memory chip and appear on the worklist (run report).

The INFINITI Analyzer is an instrument used for clinical multiplex systems intended to measure and sort multiple signals from a clinical sample. The INFINITI Analyzer is designed to measure fluorescence signals of labeled DNA target hybridized to BioFilmChip microarrays. The INFINITI Analyzer automates the Factor II and Factor V assays and integrates all the discrete processes of sample (PCR amplicon) handling, reagent management, hybridization, detection, and results analysis. The assays are processed automatically and the spots are read by the built-in confocal microscope. Results are analyzed and presented in numerical and graphical format.

The INFINITI Analyzer has two main components: pipetting and optics modules. A variety of electronic components inside the instrument are used for its operation. These include multiple stepper motors, heating and cooling devices, a barcode reader, a photomultiplier tube, and a camera all connected to USB ports.
- Pipetting Module - The pipetting module performs all the operations related to dispensing and aspiration of reagent and processing the amplified sample to be dispensed on the microarray. When the sample has been processed and hybridized to the microarray, it is transferred to the optics module for measurement of fluorescence signal.
• Optics Module - The optics module is a lightproof assembly comprised of a 3-axis stage; camera, lasers, and a photo multiplier tube (PMT). It is the enclosed casement into which the microarray is transported automatically prior to being processed on the stringency station. The optics’ stage follows X-Y-Z motions that can be stepped at a very precise rate (2.0 micron per step). Using excitation wavelengths of a 760nm laser diode, the camera takes a 1.2x1.2mm picture for each registration spot of a fluorescent dye. Analyses of these pictures allow the location of three registration spots to be determined. With respect to the position of the three registration spots, coordinates of all the bio-spots can be located. While scanning, the stage moves along the Z-axis to focus the chip and the X and Y-axes to locate the individual spots on the microarray.

Instructions on how to use the INFINITI Analyzer are provided in the INFINITI Analyzer Operator’s Manual.

The INFINITI Analyzer hardware is controlled by the Qmatic™ operating software, which is installed with-in the on-board computer and utilizes a LCD screen display. The INFINITI Analyzer modules are controlled by multitasking real time software. The Qmatic™ operating software has a schedule manager that is capable of controlling all operations of the INFINITI Analyzer such as assay protocol, fluid handling, robotics, optical detection and result analysis. Results are available for review via the LCD screen. Management reports include results in numerical and graphical format. The operator can also print the displayed results in tabular form (printer not included with INFINITI Analyzer).

WARNINGS AND PRECAUTIONS

Handling Requirements
• **For in vitro diagnostic use. To be used by qualified laboratory personnel.**
• This test is to be used only with whole blood collected in EDTA. Do not freeze/thaw blood samples. Specimens should be assayed as soon as possible.
• Do not use Heparin with this procedure; Heparin might interfere with the PCR.
• To minimize the risk of cross contamination, sample preparation, PCR reaction set up and PCR product analysis should be performed according to approved guidelines such as CLSI (Molecular Diagnostic Methods for Genetic Diseases: Approved Guideline).
• All patient specimens are potentially hazardous and care should be taken when handling materials of human origin. No test method can offer complete assurance that HCV, HIV or other infectious agents are absent.
  
  Follow the CLSI Guidelines (Molecular Diagnostics Methods for Infectious Diseases; Approved Guidelines; MM3-A).
• Do not pool/mix reagents from different lots.
• Do not use a kit or reagent past its expiration date.
• Store kits and reagents according to the product label.

Laboratory Procedures
• Follow normal precautions for handling laboratory reagents. Do not mix reagents from different containers or from different lots.
• Follow safe laboratory procedures: do not pipette by mouth; wear protective clothing (e.g., disposable gloves laboratory coats) and eye protection; do not eat, drink or smoke in the laboratory work areas; wash hands thoroughly after handling samples and reagents.

Waste Handling
• Dispose of unused reagents, specimens and waste according to applicable country, federal, state and local regulations
• Material Safety data Sheets (MSDS) are available upon request from AutoGenomics Customer Service
Sample Preparation
- Refer to the safety instructions in the package insert provided with the DNA extraction kit used.
- The PCR product can not be stored prior to loading it onto the microarray. Use immediately.

INFINITI Analyzer
- **Read the INFINITI Analyzer Operator’s Manual before operating the instrument.** Pay particular attention to “Notes”.
- Follow the Caution and Safety Warning in the Operator’s Manual.
- Refer to the Installation Requirements Section when installing the instrument.
- Refer to the Errors Section when errors are encountered while operating the instrument.
- Refer to the Help Section when problems are encountered.

STORAGE / STABILITY
- **BioFilmChip Microarray:** 12 months at Room Temperature (25°C to 30°C)
- **Intellipac Reagent:** 12 months Refrigerated (2°C to 8°C)
  - **Note:** Do not use after Intellipac has been opened for 30 days.
- **Amplification Mix:** 18 months Frozen (-30°C to -15°C)
  - **Note:** Specific product expiration date is printed on the product label

SPECIMEN COLLECTION AND STABILITY
- Peripheral blood drawn in an EDTA (purple-top) tube.
- Do not freeze/thaw blood samples. Specimens should be assayed as soon as possible.

MATERIALS PROVIDED (EACH PACKAGE IS SUFFICIENT FOR 48 TESTS)
- **AutoGenomics Product Number A1-105** FII-FV Magazine – BioFilmChip™ Microarray (with specific capture probes); 4 magazines per package
- **AutoGenomics Product Number A1-205** FII-FV Intellipac™ Reagent Management Module: 24 tests per module; 2 modules per package. Each Intellipac module contains
  - 1.4 ml ASPE master mix composed of
    - d[AGT]TP Mix
    - Cy5 –dCTP
    - ASPE Primer Mix
    - Extension Buffer
    - 2.6ml Hybridization Buffer
    - SSC
    - Sodium Azide
    - Hybridization Positive Control (Pos C-PTAG10, Cy5)
- **AutoGenomics Product Number A1-305** FII-FV Amp Mix. Each package contains 4 x 250µl vials of Amp Mix and 1 x 100µl vial of Taq DNA Polymerase
  - **AMP Mix contains**
    - d[AGT]TP Mix
    - dCTP
    - PCR Primer Mix
    - MgCl₂
    - PCR Buffer
- **AutoGenomics Product Number 12-001:** Wash buffer
REAGENTS REQUIRED BUT NOT PROVIDED BY AUTOGENOMICS

- DNA Extraction Kits - The INFINITI System Assay for Factor II & Factor V can detect Factor II (Prothrombin) G20210A and Factor V Leiden G1691A mutations using genomic DNA isolated from blood with sufficient purity, i.e., with the ratio of absorbance at 260nm to absorbance at 280nm of 1.7 to 2.0. Any DNA extraction method that meets this specification may be used. The INFINITI System Assay for Factor II & Factor V has been tested with several commercially available kits and the user can contact AutoGenomics for further information.
- Distilled Water (DNase and RNAse free)

EQUIPMENT
The following equipment is required but not provided with the assay reagents

- AutoGenomics Product Number 10-001: INFINITITM Analyzer with Qmatic operating software.
- AutoGenomics Product Number 11-001: INFINITITM PipetteTips
- AutoGenomics Product Number 11-003: INFINITITM 24 well plate/lids
- Pipettors
- Mini Centrifuge
- Pipette tips
- Microfuge tube Racks
- Thermocycler
- Vortex
- 0.2 ml thin wall tubes for PCR
- 0.5 ml microcentrifuge tubes
- 24-Well Plate (24 WP)

ASSAY PROCEDURE
DNA Extraction
Follow the instructions provided with the DNA extraction kit used.

PCR Reaction
Note: The PCR product cannot be stored prior to loading it onto the microarray. Use immediately.

1. Thaw and bring to room temperature the tube of Amp Mix and keep Taq DNA polymerase on ice.
2. Vortex the Amp Mix tube 2 to 5 sec to mix reagents and centrifuge briefly to bring the content to the bottom of the tubes. Include one negative control in every PCR reaction set up.

Amp Mix 17.75 µl
Taq DNA Polymerase 0.25 µl
Template DNA (25 ng/µl) 2.0 µl

Total 20.0 µl

For the PCR negative control, add 1µl of DNase free water.
3. Vortex tubes for 2 to 5 seconds to mix and then centrifuge briefly.
4. Place tubes in the thermocycler and cycle using the following conditions:

<table>
<thead>
<tr>
<th>Step No.</th>
<th>Temperature °C</th>
<th>Time</th>
<th>No. of Cycles</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>95</td>
<td>4 minutes</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>95</td>
<td>15 seconds</td>
<td>40</td>
</tr>
<tr>
<td></td>
<td>56</td>
<td>15 seconds</td>
<td></td>
</tr>
<tr>
<td></td>
<td>72</td>
<td>15 seconds</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>4</td>
<td>Hold</td>
<td></td>
</tr>
</tbody>
</table>

<table>
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<th>Step No.</th>
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<td>95</td>
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<td></td>
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<td>15 seconds</td>
<td></td>
</tr>
<tr>
<td></td>
<td>72</td>
<td>15 seconds</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>4</td>
<td>Hold</td>
<td></td>
</tr>
</tbody>
</table>
Note:
Setting PCR reaction in 3 X 8 microwell is recommended to minimize sample handling in subsequent steps.

To avoid potential contamination, a separate area is recommended for the PCR.

Sample Loading
Follow instructions in the INFINITI Analyzer Operator’s Manual (Part Number EM 34000).
1. Dispense 20 µl of the PCR product into the 3x8 well plates and load onto the INFINITI Analyzer.
2. Load the appropriate magazines, Intellipac, tips and wash buffer.

Operation of the INFINITI Analyzer
Please refer to the INFINITI Analyzer Operator’s Manual (Part Number EM 34000)

QUALITY CONTROL
It is recommended that a positive (heterozygous and/or homozygous for both genotypes) sample, from a cell line or patient sample, for each mutation; a negative control (a sample that does not contain the mutation of interest, i.e., a wild type sample); and a “No Template Control” (Molecular Grade water) be included with each test run. Please contact AutoGenomics for recommendations.

Note: The thermal cycler used should be regularly maintained and calibrated with an external temperature standard, according to the laboratory’s regulatory and QC requirements.

LIMITATIONS
The results obtained from this method should be used and interpreted only in the context of the overall clinical diagnosis. AutoGenomics is not responsible for any clinical decisions that are taken.

The absence of the mutations detected by this Procedure is no guarantee that other mutations are not present. Presence of other mutations is possible and is not detected by this kit.

INTERPRETATION OF RESULTS
Results from the INFINITI System Assay for Factor II & Factor V are reported to the user as a genotype “call”, indicating which genotype was detected in the sample, i.e., Wild Type, Homozygous, or Heterozygous for Factor II (Prothrombin) G20210A mutation, and Wild Type, Homozygous, or Heterozygous for Factor V Leiden G1691A mutation. In addition to the genotype “call”, the ratio and the signal (RFU) are provided for each analyte.

If the call is “Indeterminate”, repeat the sample assay and refer to the Trouble Shooting section of the INFINITI Analyzer Operator’s Manual.

If errors occur during the assay, “Test Error” message (e.g., “low DNA”) is shown. An Error Log is generated which identifies the problem. Depending on the error message/problem, the assay may have to be repeated.

DISPOSAL
Waste materials for the INFINITI System Assay for Factor II & Factor V are common waste materials generated in clinical laboratories, and should be handled/disposed of in accordance with the policies/procedures in place in the clinical laboratory.
PERFORMANCE CHARACTERISTICS

Analytical Specificity
Studies related to specificity were conducted during assay development. PCR primer specificity was determined by amplicon size on a gel and sequencing the amplicon. ASP primer specificity was determined by the correct calls made by the assay using known genomic samples. Capture probe specificity was determined by hybridizing different oligos and demonstrating that correct oligo hybridizes to the known spot.

Analytical Sensitivity
Studies demonstrated that the INFINITI System Assay for Factor II & Factor V can detect as low as 1ng DNA and as high as 300ng DNA. A minimum of 25ng DNA is recommended for this assay.

Percent Agreement
The results of the comparison studies conducted in three clinical sites comparing the INFINITI System Assay for Factor II & Factor V to FDA cleared device (predicate) demonstrated

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Number Tested</th>
<th>Number of Correct Calls on First Run</th>
<th>Number of Invalid Calls* on First Run</th>
<th>Agreement First Run</th>
<th>Number of Correct Calls including Repeat Run</th>
<th>Number of Invalid Calls* on Repeat Run</th>
<th>Agreement After Repeat Run</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT</td>
<td>146**</td>
<td>126</td>
<td>20</td>
<td>86.3%</td>
<td>144</td>
<td>2</td>
<td>98.6%</td>
</tr>
<tr>
<td>MUT</td>
<td>22**</td>
<td>22</td>
<td>0</td>
<td>100%</td>
<td>22</td>
<td>n/a</td>
<td>100%</td>
</tr>
<tr>
<td>HET</td>
<td>40</td>
<td>39</td>
<td>1</td>
<td>97.5%</td>
<td>39</td>
<td>1</td>
<td>97.5%</td>
</tr>
<tr>
<td>Overall</td>
<td>208</td>
<td>187</td>
<td>21</td>
<td>89.9%</td>
<td>205</td>
<td>3</td>
<td>98.6%</td>
</tr>
<tr>
<td>WT</td>
<td>113</td>
<td>101</td>
<td>12</td>
<td>89.4%</td>
<td>113</td>
<td>0</td>
<td>100%</td>
</tr>
<tr>
<td>MUT</td>
<td>15</td>
<td>15</td>
<td>0</td>
<td>100%</td>
<td>15</td>
<td>n/a</td>
<td>100%</td>
</tr>
<tr>
<td>HET</td>
<td>47</td>
<td>45</td>
<td>2</td>
<td>95.7%</td>
<td>47</td>
<td>0</td>
<td>100%</td>
</tr>
<tr>
<td>Overall</td>
<td>175</td>
<td>161</td>
<td>14</td>
<td>92.0%</td>
<td>175</td>
<td>0</td>
<td>100%</td>
</tr>
</tbody>
</table>

* No discordant results. Invalid results refer to “indeterminate” results.
** Samples tested were taken from suspected thrombophilia patients except:
- 15 of the 146 WT samples (negative for Factor II G20210A mutation) were from a known genomic WT sample which was split into 15 individual samples.
- 15 of 22 Factor II G20210A mutation samples were from a known genomic sample which was split into 15 individual samples.

Precision/Reproducibility – Assay
Chip-to-chip: Using the same sample and the same INFINITI Analyzer, the assay was run in five replicates using three BioFilmChip microarrays from one lot. The CVs using average triplicate spots for each mutation ranged from 9 – 12% for wild-type present calls. This was repeated two other times, each time using a different INFINITI Analyzer. All calls were 100% correct.
Lot-to-lot: Three lots of BioFilmChip microarray were tested using the same instrument four times, each time using a different sample. Two-way ANOVA on the RFU readings did not detect lot-to-lot difference on three of the four test runs (p > 0.05), and detected lot-to-lot difference on one test run (0.05 > p > 0.01). Genotype calls were 100% correct.

Day-to-day: Known genomic sample was assayed 12 times on each of three days using one instrument. The RFU signal %CV ranged from 1.35 to 14.87 on day 1, 0.77 to 19.72 on day 2 and 0.41 to 21.2 on day 3. Genotype calls were 100% correct.

Precision/Reproducibility - Instrument

- One DNA sample was analyzed using three different INFINITI Analyzers and one lot of BioFilmChips, five times (five runs).
  
  Intra-Instrument: The %CV using a single chip five times on a single instrument ranged from 0.9 to 28.3%CV. Genotype calls were 100% reproduced within each instrument.

  Inter-Instrument: The %CV using a single chip five times on each of three instruments ranged from 0.5% to 12%CV. All genotype calls were 100% correct and reproducible.

- Three instruments were tested on three different days using a Standard (non-assay) Microarray Chip. For each instrument tested, each capture probe spot on the Standard Microarray Chip was read 24 times, then averaged, and a %CV calculated for the spot. The following lists the ranges for the %CV for the three instruments tested.

<table>
<thead>
<tr>
<th>Instrument</th>
<th>Ave %CV</th>
<th>%CV Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>4.03%</td>
<td>1.9-7.5%</td>
</tr>
<tr>
<td>2</td>
<td>3.99%</td>
<td>2.7-6.5%</td>
</tr>
<tr>
<td>3</td>
<td>3.24%</td>
<td>1.9-5.3%</td>
</tr>
</tbody>
</table>

Sample Carry-over

No carry-over was detected when a series of 300ng of a wild type sample (FV-WT; FII-WT) was followed by 10ng of a positive sample (FV-WT; FII-M), followed by 300ng of a heterozygous sample (FV-H; FII-H), followed by a “No Template Control” or water, was run six times.

Interference

Results of the interference studies demonstrate that there is no interference with the INFINITI System Assay for Factor II & Factor V from 8mg/dL bilirubin, 70mg/dL cholesterol, and 1333v/dL heparin. No studies were conducted with oral anti-coagulants; therefore, no claims are made.

REFERENCES

