ForenSeq™ Universal Analysis Software

Interrogate the broadest range of forensically relevant loci with a simple user interface and powerful analysis algorithms.

**Highlights**

- **Complete, Sample-to-Answer Workflow**
  Manage the sequencing workflow from run set-up to data analysis and report generation

- **Perform Automated Data Visualization and Reporting with Simple Graphical User Interface**
  Compare samples automatically, generate population statistics, and view data through an intuitive user interface

- **Generate Investigative Leads Through Analysis Software Advances**
  Estimate visible traits such as hair color and eye color as well as biogeographical ancestry

**Introduction**

Illumina ForenSeq Universal Analysis Software is part of the MiSeq FGx™ Forensic Genomics System—a fully validated next-generation sequencing (NGS) system specifically designed for use in forensic genomics applications. The MiSeq FGx System and ForenSeq Universal Analysis Software enable the simultaneous analysis of approximately 200 forensically relevant single nucleotide polymorphisms (SNPs) and short tandem repeats (STRs)—including marker sets not routinely available with traditional methods (Table 1).

To maximize flexibility and scalability of the MiSeq FGx System for each laboratory, the ForenSeq Universal Analysis Software is sold separately on a standalone, dedicated server.

**Complete, Sample-to-Answer Workflow**

The MiSeq FGx Forensic Genomics System uses a pipeline of software applications to perform sequencing runs and complete data analysis (Figure 1). ForenSeq Universal Analysis Software guides the sequencing workflow from run-setup and sample data entry to the final stages of data analysis and report generation, MiSeq FGx Control Software (MCS), installed on the MiSeq FGx instrument control computer, captures flow cell images, operates the flow cell stage, and controls reagent delivery and temperature. During the run, Real-Time Analysis (RTA) software, performs image analysis, base calling, and assigns base-by-base quality scores. ForenSeq Universal Analysis Software then initiates the final stages of analysis, including demultiplexing, sequence alignment, allele calling, genotyping, and reporting.

**Table 1: Simultaneous Analysis of Forensically Relevant Loci with ForenSeq Universal Analysis Software**

<table>
<thead>
<tr>
<th>Feature</th>
<th>Markers$^b$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Global Autosomal STRs</td>
<td>27</td>
</tr>
<tr>
<td>Y-STRs</td>
<td>24</td>
</tr>
<tr>
<td>X-STRs</td>
<td>7</td>
</tr>
<tr>
<td>Identity SNPs</td>
<td>94</td>
</tr>
<tr>
<td>Phenotypic SNPs</td>
<td>22</td>
</tr>
<tr>
<td>Biogeographical Ancestry SNPs</td>
<td>56</td>
</tr>
<tr>
<td>Total Number of Loci$^b$</td>
<td>&gt; 200</td>
</tr>
</tbody>
</table>

$^a$ SNP and STR chromosome locations can be found in the ForenSeq DNA Signature Prep Kit User Guide (support.illumina.com/downloads/forenseq-dna-signature-prep-guide-15049528.html).

$^b$ Over 200 markers analyzed when running primer set B.
Perform Automated Data Visualization and Reporting with Simple Graphical User Interface

Illumina ForenSeq Universal Analysis Software provides an easy data analysis workflow and user-friendly viewing and reporting features. The graphical user interface has a simple, intuitive design that can be viewed from any laboratory computer through the Google Chrome web browser. The Run Overview screen features easy run-setup, sample information, and index tracking options (Figure 2). Within the analysis software, sequencing run data can be associated with specific, user-defined projects and results can be viewed by run or by project in the Projects screen.

At run completion, ForenSeq Universal Analysis Software offers data visualization through several types of display screens. The Sample Details screen provides a summarized view of SNP and STR locus intensity scatter plots. Below the intensity charts, individual allele calls are displayed, along with convenient icons displayed when specific quality flags are triggered (Figure 3). To view more indepth reports on specific loci, drill down using the Locus Detail pop-up screens. The Locus Detail screens display intensity charts, the base-by-base DNA target sequence, stutter, and more (Figure 4). The analysis software also offers a full suite of run metrics and sample quality controls that can be evaluated after the run. Quality metrics and sample quality controls are displayed with color indicators allowing forensic analysts to quickly scan the results and determine whether controls fall within the recommended ranges.

Project Detail reports and Sample Detail reports can be generated automatically within the software. Each report can be easily printed or exported in .xlsx format. Password protected user accounts ensure secure access to the MiSeq FGx sequencing instrument and to the analysis server.

Generate Investigative Leads through Analysis Software Advances

Compared to current capillary electrophoresis-based methods, Illumina ForenSeq Universal Analysis Software provides a number of data analysis advantages. These include, the capacity to interrogate a greater number of markers (Table 1) and the ability to recover the maximum amount of useful genomic information from degraded DNA, low quality DNA, or complex mixtures. Furthermore, all markers, including SNPs, autosomal, X-, and Y-STRs, can be analyzed simultaneously using a single, streamlined workflow.

Beyond analysis of autosomal, X-, and Y-STRs, the software enables the analysis of marker sets not routinely available with traditional capillary electrophoresis methods. These include a dense set of identity-informative SNPs (iSNPs), which are informative for source attribution, phenotypic-informative SNPs (pSNPs), which provide estimates of eye color (blue, intermediate, brown) and hair color (brown, red, black, blond), and biogeographical ancestry-informative SNPs (aSNPs). Biogeographical ancestry estimation is presented as
Figure 3: ForenSeq Universal Analysis Software Sample Details Screen — The Sample Details screen displays overall sample results as an intensity plot, along with detailed summary tables for all loci included in the sequencing library.

Figure 4: ForenSeq Universal Analysis Software Locus Details Screen — The pop-up Locus Detail screen shows the allele call, intensity bar graph, and stutter. View the Repeat Sequence column to see the full, base-by-base target sequence, and any intra-STR variation present in the sample.

Figure 5: ForenSeq Universal Analysis Software Estimation of Visible Traits — The ForenSeq Universal Analysis Software enables estimation of visible traits including eye and hair color. Markers for biogeographical ancestry can also be tested and reviewed.
Table 2: ForenSeq Server Components and Specifications
- 4 × Seagate 1 TB Sata hard drives
- RAID controller allowing for data redundancy and speed
- Intel 2 GHz × 64 processor with 6 cores/12 threads
- Intel socket R server board
- 32 GB DDR3 RAM
- 550 W power supply
- ULL, FCC, CE certified
- Windows Server 2012 R2 Standard with 5 CALs

Ordering Information

<table>
<thead>
<tr>
<th>Product</th>
<th>Catalog No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>ForenSeq Universal Analysis</td>
<td>SE–550–1001</td>
</tr>
<tr>
<td>Software and Server</td>
<td></td>
</tr>
</tbody>
</table>

Learn More
To learn more about ForenSeq Universal Analysis Software, visit: www.illumina.com/informatics/sequencing-microarray-data-analysis/forenseq-universal-analysis-software.ilmn
To learn about the MiSeq FGx instrument, visit: www.illumina.com/systems/miseq-fgx.ilmn
To learn about the ForenSeq DNA Signature Prep Kit, visit: www.illumina.com/products/forenseq-dna-signature-kit.ilmn
To learn about mitochondrial DNA analysis with the MiSeq FGx System, visit: www.illumina.com/applications/forensics/missing_persons_and_unidentified_human_remains.ilmn

References
1. The full MiSeq FGx System workflow is validated per the Scientific Working Group on DNA Analysis Methods (SWGDAM) guidelines (www.swgdam.org).

Summary
ForenSeq Universal Analysis Software contains comprehensive sample management and analytical capabilities, including sample and index management, application-specific workflows, data visualization at sample and locus levels, quality flags to simplify data analysis, and easily exportable reports.