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MiSeq FGx[™] Forensic Genomics System

Solve more cases and generate more leads with the power and accuracy of Illumina next-generation sequencing.

- Highlights -

- Fully Validated, Sample-to-Answer Solution All-inclusive system ties library preparation, sequencing, data analysis, and reporting into a single integrated workflow
- Simple, Streamlined Workflow
 Interrogate 200 genetic markers in a single, streamlined workflow eliminating the need for multiple STR kits
- Access a Wider Range of Informative SNPs Biogeographical ancestry and phenotypic-informative SNPs provide valuable information not widely available with current technology
- Superior Analysis of Challenging Samples Advanced capacity to handle challenging samples such as complex mixtures or degraded DNA

Introduction

The MiSeq FGx[™] Forensic Genomics System is the first fully validated¹ sequencing system specifically designed for use in forensic genomics applications (Figure 1). With the high resolution and unmatched accuracy of next-generation sequencing (NGS), the MiSeq FGx System can transform the most fragile, degraded, or mixed samples into powerful results.

A Complete Sample-to-Answer Solution

The MiSeq FGx Forensic Genomics System delivers a complete DNA-to-Data workflow for the analysis of forensic DNA samples. The solution begins with the ForenSeq™ DNA Signature Prep Kit, which includes all reagents required to prepare the DNA library* for sequencing (Figure 2). The MiSeq ForenSeq Sequencing Kit provides sequencing reagents, an RFID labeled reagent cartridge, and wash solution, which are then loaded onto the MiSeq FGx instrument along with the sequencing-ready DNA libraries. The intuitive touch screen interface provides simple, step-by-step guidance through each stage of the sequencing run including library and reagent loading, run configuration, and run monitoring. ForenSeq Universal Analysis Software delivers a powerful suite of forensic analysis capabilities including automatic detection of mixed DNA samples, generation of population statistics, and sample comparison. The software also enables estimation of visible traits and biogeographical ancestry markers that can provide crucial investigative leads in "no suspect" cases.

Simple, Streamlined Workflow

With current forensic capillary electrophoresis (CE) methods, limited DNA quantities may require forensic analysts to choose between available fragment length-based short tandem repeat (STR) tests.





These trade-offs can result in lengthy, complex decision trees as technical limitations are weighed against potentially informative genetic data.

The MiSeq FGx System eliminates the need to choose between relevant STR tests by providing all currently available STR typing tests in a single workflow (Figure 3). One run with the MiSeq FGx System simultaneously interrogates ~200 genetic markers, including a combination of autosomal, X-, and Y-STRs. Furthermore, the low DNA input requirement of 1 ng enables testing of a broad range of samples where DNA quantity is limited.

Access a Wider Range of Informative SNPs

Another casework challenge occurs when a complete autosomal STR profile is established, but no suspect is available for direct comparison, or no hits are found in national or local criminal databases. In addition to providing all autosomal STR markers currently utilized around the world for casework and criminal DNA databasing, the ForenSeq DNA Signature Prep Kit also contains marker sets not routinely available with traditional CE methods. These include a dense set of identity informative single nucleotide polymorphisms (iiSNPs)²⁻³, which are informative for source attribution, as well as biogeographical ancestry-informative SNPs (aiSNPs)⁴, and phenotypic-informative SNPs (piSNPs)⁵, which provide estimates of eye color (blue, intermediate, brown) and hair color (brown, red, black, blond). AiSNPs and piSNPs can be critical in generating tactical investigative leads from "no suspect" cases that may otherwise have reached dead ends.

Superior Analysis of Challenging Samples

The challenges presented by degraded DNA, low quantity DNA, or complex DNA mixtures can potentially complicate or derail an investigation. The MiSeqFGx System offers an enhanced capacity to analyze these types of challenging samples.

When analyzing partially or highly degraded DNA, the SNPs in the ForenSeq DNA Signature Prep Kit are targeted by amplicon sizes ≤ 125 bp, making them extremely well-suited for analysis of degraded DNA⁶. With complex DNA mixtures, the large number of markers included in the kit— many of which are highly polymorphic— enable an improved ability to detect low-level minor components that may go undetected by traditional methods⁶.

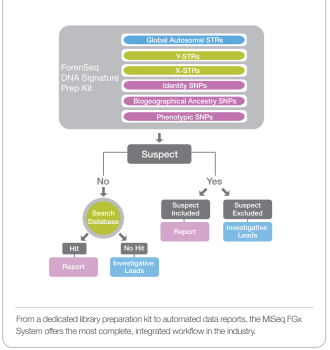
These capabilities, coupled with the inherent sensitivity of Illumina sequencing by synthesis (SBS) chemistry and the low DNA input requirement, provide superior resolution of a broad range of cases compared to conventional STR and CE analysis.

Easy, Optimized Software Solution

The MiSeq FGx System solution includes ForenSeq Universal Analysis Software: a fully optimized, software solution designed for forensic genomics applications (Figure 4). The software features a comprehensive suite of library management and analytical tools. These include data visualization at library and locus levels, STR and SNP allele and genotype calling, visualization of intra-STR sequence variation, statistical estimates of combined genotype frequency, statistical estimates of biogeographical ancestry and phenotype, quality flagging, and automated report generation. ForenSeq Universal Analysis Software is delivered pre-installed on a dedicated, standalone server. The server is sold separately to maximize flexibility for each laboratory.

Exceptional Data Quality

The MiSeq FGx System leverages Illumina SBS chemistry—the most trusted and widely adopted next-generation sequencing technology in the industry⁷. Exceptional data quality is achieved by a proprietary, reversible terminator-based method that detects single bases as they are incorporated into massively parallel DNA strands. Fluorescent



terminator dyes are imaged as each dNTP is added and then cleaved to allow incorporation of the next base. With all four reversible, terminator-bound dNTPs present during each sequencing cycle, natural competition among bases minimizes incorporation bias⁸. Base calls are made directly from signal intensity measurements during each incorporation cycle, reducing raw error rates compared to other technologies⁹⁻¹². The result is highly accurate base-by-base sequencing that virtually eliminates sequence context-specific errors, even within repetitive sequence regions or homopolymers.

Summary

The MiSeq FGx Forensic Genomics System is a fully validated sequencing system specifically designed for use in forensic genomics applications. The complete sample-to-answer system includes the ForenSeq DNA Signature Prep Kit, the MiSeq FGx Instrument, and the ForenSeq Universal Analysis Software package. With the speed and accuracy of Illumina next-generation sequencing, criminal justice has a powerful new ally.

Learn More

To learn more 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

For more on the ForenSeq Universal Analysis Software, visit: www.illumina.com/informatics/sequencing-microarray-data-analysis/ forenseq-universal-analysis-software.ilmn

Figure 3: MiSeq FGx System Investigative Workflow

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ForenSeq Universal Analysis Software features DNA library management tools, data analysis and visualization, and automated reporting. The Run Overview screen features easy data entry fields for Run Name and Run Description. Sample information, index combinations, and application options can also be imported with a tab delimited .txt file. Sample types include Known Sample, Forensic Sample, Reagent Blank, Negative Amplification Control, Positive Amplification Control, and Human Sequencing Control.

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

- The full MiSeq FGx System workflow is validated per the Scientific Working Group on DNA Analysis Methods (SWGDAM) guidelines (www.swgdam. org).
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- Illumina (2014) ForenSeq DNA Signature Prep Kit Data Sheet (www.illumina. com/products/forenseq-dna-signature-kit.ilmn).

MiSeq FGx System Performance Parameters

eature	Performance
ow Input DNA	robust performance ≥ 100 pg, optimal input 1 ng
ocus Multiplexing apability	~200 loci
ccurate Low-level	detects minor contributors
ixture Detection	at < 5% of major
ample Multiplexing apability ^a	8-96 samples
eep Coverage	14 million reads per run
hort Amplicon etection	≥ 65 bp

Join the Illumina Community

Join a worldwide community of over 60,000 scientists using Illumina technology for research studies, clinical applications, and forensic genomics analysis. Illumina schedules community events throughout the year, bringing researchers and analysts together from around the world. User group meetings, scientific symposia, community newsletters, and blog forums provide venues to discuss new research methods and establish collaborations.

Our dedicated service and support team is an integral part of the Illumina community, consisting of more than 300 people worldwide—75% of whom have advanced degrees. Illumina support begins when the sequencing system is delivered and continues through installation, setup, and training of laboratory personnel. The global, Illumina technical support team is available 24/7 to answer questions every step of the way. In addition to onsite assistance, we offer webinars, online training courses, and instructor-led training courses at the state-of-the-art Illumina campus in San Diego.

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MiSeq FGx Instrument Specifications

Instrument Configuration

RFID tracking for consumables MiSeq FGx Control Software

Instrument Control Computer (Internal)^a Base Unit: Intel Core i7-2710QE 2.10 GHz CPU Memory: 16 GB RAM Hard Drive: 750 GB Operating System: Windows 7 embedded standard

Operating Environment

Temperature: 22°C ± 3°C (66–77°F) Humidity: Non-condensing 20%-80% Altitude: Less than 2,000 m (6,500 ft) Air Quality: Pollution degree rating of II Ventilation: Maximum of 1,364 BTU/h For Indoor Use Only Light Emitting Diode (LED) 530 nm, 660 nm Dimensions W×D×H: 68.6 cm × 56.5 cm × 52.3 cm (27.0 in × 22.2 in × 20.6 in) Weight: 54.5 kg (120 lbs) Crated Weight: 90.9 kg (200 lbs) **Power Requirements** 100-240V AC @ 50/60Hz, 10A, 400W **Radio Frequency Identifier (RFID)** Frequency: 13.56 MHz Power: 100 mW

Product Safety and Compliance

NRTL certified IEC 61010-1 CE marked FCC/IC approved

a. Computer specifications are subject to change.

Ordering Information

Product	Catalog No.		
MiSeq FGx Instrument	SY-411-1001		
MiSeq ForenSeq Sequencing Kit	MS-201-1001		
ForenSeq DNA Signature Prep Kit	FC-450-1001		
ForenSeq Index Adapter Fixture	FC-451-1001		
ForenSeq Universal Analysis Software	SE-550-1001		

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