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Genome AnalyzerIIx System

The most proven, widely adopted next-generation sequencing platform

Highlights

- Broadest Applications Flexibility Study the genome, epigenome, and transcriptome
- Broadest Spectrum of Genomic Variation Characterize genomic variants with short- and long-insert paired-end reads with insert sizes from 200 bp to 5 kb
- Unmatched Combination of Read Length and Number of Reads Achieve 2 × 150 bp reads and > 640 million reads per flow cell
- High Data Quality Generation

Yield greater than 85% of bases higher than Q30 at 2 x 50 b

A Revolution in Genomics

Illumina's Genome Analyzer*IIx* is a proven platform for genetic analysis and functional genomics, and has transformed the way experiments are developed and executed (Figure 1). Massively parallel sequenc- ing technology leverages clonal cluster formation and proprietary reversible terminator chemistry to dramatically improve the speed, and reduce the cost, of large-scale sequencing.

Broadest Applications Flexibility

The Genome Analyzer *IIx* supports a wide range of applications, includ- ing whole-genome and candidate region resequencing, transcriptome analysis, small RNA discovery, methylation profiling, and genome-wide protein-nucleic acid interaction analysis.

Simple, Fast, and Automated

The Genome Analyzer IIx system offers the simplest and- fastest work- flow for a broad range of high-throughput sequencing applications. Sample libraries are prepared in just a few hours with ready-to-use kits. Clonal clusters are automatically generated on Illumina Genome Analyzer IIx flow cells using the cBot cluster generation system. In less than four hours, up to 12 multiplexed samples can be isothermally amplified in each channel of the eight-channel flow cell

Illumina sequencing technology provides an easy-to-use protocol that does not require emulsion PCR. This allows for a self-contained system that minimizes handling errors and contamination concerns, eliminating the need for robotics or clean rooms. The system is de-signed to fit in any lab, from individual research labs to core labs and genome centers. The streamlined workflow

of the Genome Analyzer *IIx* system generates meaningful data quickly and efficiently, while reduc- ing project time lines and costs (Figure 2).

TruSeqTM Technology

The TruSeq family of reagents represents the latest advancement of Illumina's sequencing by synthesis (SBS) chemistry. From sample prep through DNA sequencing, TruSeq technology enables Illumina sequencing to deliver the most accurate data across a broad range of applications.

SBS chemistry enables massively parallel sequencing of millions of fragments using a proprietary reversible terminator-based method that detects single bases as they are incorporated into growing DNA strands. A fluorescently-labeled terminator is imaged as each dNTP is added and then cleaved to allow incorporation of the next base. Since all four reversible terminator-bound dNTPs are present during each sequencing cycle, natural competition minimizes incorporation bias. Base calls are made directly from signal intensity measurements during each cycle, which greatly reduces raw error rates compared to other technologies. The end result is highly accurate base-by-base sequencing that eliminates sequence-context specific errors, enabling robust base calling across the genome, including repetitive sequence regions and within homopolymers.

Illumina sequencing delivers the most accurate human genome at any level of coverage. The highest yield of error free reads and most base calls above Q30 provide researchers the highest confidence in their data integrity to draw sound biological conclusions.

Single- or Paired-End Read Support

The Genome Analyzer IIx system supports sequencing of both single- read and paired-end libraries. It is the only platform available that offers a short-insert paired-end capability for high-resolution sequencing as well as long-insert paired-end reads for efficient sequence assembly, de novo sequencing, and large-scale structural variation detection. The TruSeq library construction protocol minimizes the time from sample isolation to usable results. Single-read or short-insert paired-end sample preparation of genomic DNA takes as few as two hours (5 minutes of hands-on time) using Nextera Library Prep Kits. The combination of short inserts and 2×150 bp or longer reads increases the ability to align and sample the genome, expanding the Genome Analyzer's utility for other applications.

Low Input Requirements

The Genome Analyzer *IIx* system requires sample inputs as low as 100 ng, enabling a host of applications where sample is limited (e.g., immunoprecipitates, laser-dissected materials, and small model systems).

Genome AnalyzerIIx Performance Parameters

| Read Length | Run Time (Days) | Output (Gb) |
|---------------------------|-----------------|-------------|
| 1 × 35 bp | ~2 | 10 – 12 |
| $2 \times 50 \text{ bp}$ | ~5 | 25 – 30 |
| 2 × 75 bp | ~7 | 37.5 – 45 |
| 2 × 100 bp | ~9.5 | 54 – 60 |
| $2 \times 150 \text{ bp}$ | ~14 | 85 – 95 |

^{*}Sequencing output generated using TruSeq SBS V5 kit with PhiX library and cluster densities between 508,000-630,000 clusters/mm2 that pass filtering on a GAIIx

Throughput

Up to 6.5 Gb per day for a 2 x 100 bp run

Reads

Up to 320 million clusters passing filter and up to 640 million paired-end reads

Performance

The Genome Analyzer IIx generates a significant yield of bases greater than Q30

- Greater than 85% bases higher than Q30 at 2 x 50 bp
- Greater than 80% bases higher than Q30 at 2 x 100 bp

Data Analysis Support

The analysis software and hardware included with the Genome Analyzer*IIx* contribute to an end-to-end sequencing approach that enables researchers to rapidly move from raw data acquisition to publishable, biologically meaningful results. Illumina's Sequencing Control Software (SCS) offers real-time analysis processing that automatically produces image intensities and quality-scored base calls directly on the instrument computer. These reads can be aligned to a reference sequence and analyzed using the Pipeline analysis software. In combination with the Consensus Assessment of Sequence and Variation (CASAVA) software, GenomeStudio® data analysis software provides intuitive, graphical analysis and interaction with DNA and RNA data.

Specifications for the Genome Analyzer IIx System

| Illumina Genome Analyzer <i>IIx</i> | Illumina cBot | Illumina Paired-End Module |
|-------------------------------------|---------------------------------|-------------------------------|
| Catalog No. | | |
| SY-301-1301 | SY-301-2002 | SE-301-1002 |
| Instrument Configuration | | |
| CE Marked and ETL Listed | CE Marked and ETL Listed | CE Marked and ETL Listed |
| instrument Installation, setup, | instrument Installation, setup, | instrument Installation setup |
| and accessories Computer and | and accessories | and accessories |
| flat panel display | | |
| Data collection and analysis | | |

| software | | | |
|-----------------------------------------------------------------------------------------|------------------------------|---|--|
| Instrument Control | | | |
| Computer | | | |
| Base Unit: Intel Xeon x5560 | Embedded Mini-ITX Board | | |
| 2.8 GHz, Quad Core Memory: | with Celeron M Processor | | |
| 4 GB RAM Hard Drive: 4 × | Memory: 1 GB RAM Hard | | |
| 300 GB SCSI Operating | Drive: 80 GB Operating | | |
| System: Windows XP | System: Windows Embedded | | |
| Monitor: 19" LCD flat panel | Monitor: Integrated 8" touch | | |
| | screen | | |
| Note: The computer specifications may be regularly upgraded. Contact your local Account | | | |
| Manager for current configuration. | | | |
| Operating Environment | · | _ | |

Operating Environment

| -1 | | |
|---------------------------------------------------------|-------------------------------------------|---------------------------------------------------------|
| Temperature: $22^{\circ}\text{C} \pm 3^{\circ}\text{C}$ | Temperature: $22^{\circ}C \pm 3^{\circ}C$ | Temperature: $22^{\circ}\text{C} \pm 3^{\circ}\text{C}$ |
| Humidity: Non-Condensing | Humidity: Non-Condensing | Humidity: Non-Condensing |
| 20%–80% Altitude: Less than | 20%–80% Altitude: Less than | 20%–80% Altitude: Less than |
| 2,000 m (6,500 ft) Air | 2,000 m (6,500 ft) Air | 2,000 m (6,500 ft) Air |
| Quality: Pollution Degree | Quality: Pollution Degree | Quality: Pollution Degree |
| Rating of II Ventilation: | Rating of II For Indoor Use | Rating of II For Indoor Use |
| Maximum of 3400 BTU/h | Only | Only |
| (1000W) For Indoor Use Only | | |
| Laser | | |
| Three laser system: 660, 635, | Class 2 laser: 630–650 nm | |
| and 532 nm | | |
| Dimensions | | |
| W×D×H: $102 \text{ cm} \times 67 \text{ cm} \times$ | W×D×H:38cm×62cm×40cm | W×D×H:24cm×61cm×44cm |
| 92 cm | Weight: 34 kg | Weight: 13 kg |
| Weight: 187 kg | Crated Weight: 36 kg | Crated Weight: 34 kg |
| Crated Weight: 232 kg | | |
| Power Requirements | | |
| 100-240V AC 50/60 Hz, | 100-240V AC 50/60 Hz, 4A, | 100-240V AC 50/60 Hz, 3A |
| 20A, 900 Watts | 400 Watts | Max, 250 Watts |
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Illumina recommends an uninterruptible power supply (UPS) with an output capacity of at least 3 kVA.

Instrument Bench

Illumina recommends a movable table with locking casters capable of supporting the weight of the instrument and computers.

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