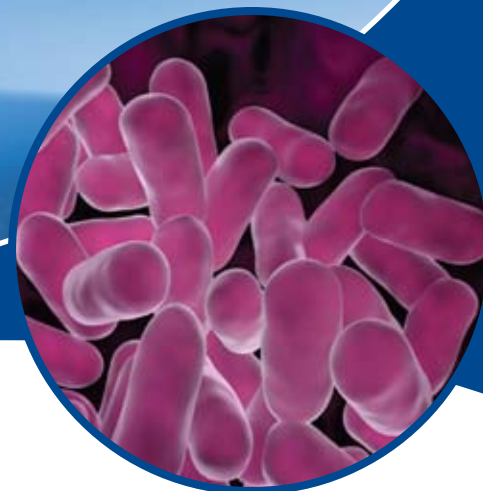


# MassARRAY® iSEQ™ – Comparative Sequence Analysis

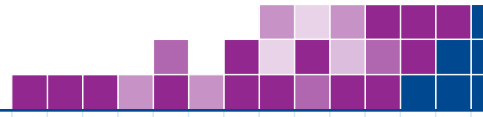


Setting a New Standard in Molecular Typing.

Accurate. Sensitive. Rapid.

SEQUENOM®  
[www.sequenom.com](http://www.sequenom.com)

CCGATGATCGACCAGTATGCGCATGATGATCGAAGTATGCGCATTATGCGCAT  
GCGCATTATGCGCATGATGATCGAAGCCGATGATCGACCAGTATGCGCATGA  
GCGCATTATGCGCGCATGATGATCGAAGTATCATGATGATCGAAGCCGATGA



## Overview

### Advancing Molecular Typing

iSEQ™ is new MassARRAY® application for automated comparative sequence analysis that combines the sensitivity of PCR with the accuracy of mass spectrometry. The combination creates a highly accurate, reproducible method for identifying and typing microbes, viruses, and other haploid organisms. The iSEQ™ solution allows you to analyze one or multiple target regions on multiple samples in a convenient homogeneous assay format, and enables automatic sample identification or sample clustering. The software package delivers automatic data analysis, customizable plate setups, data portability and database management.

## The iSEQ™ Solution for Comparative Sequence Analysis

iSEQ™ offers a combination of automation, versatility and discriminatory power that is superior to any other comparative sequence analysis system. The versatility of the iSEQ™ software and open database access enables comparative sequencing with a wide range of global reference databases. Analysis results for 384 reactions are automatically achieved in less than an hour.

### Flexible

- Scalable throughput
- Comparative sequence analysis with a wide range of reference databases
- Reads length routinely up to 800 bp

### Sensitive and Accurate

- Two levels of amplification: PCR and *in vitro* transcription
- Discriminatory power down to a single nucleotide

### Cost Effective

- 10 µl PCR reactions in 384 microplate format
- No product purification required

### Efficient

- Automated data analysis
- Ideal for a few or several target regions
- Data portability

### Rapid

- Data acquisition for 384 reactions in < 1hr
- Analyze up to 3,072 reactions / day
- Throughput up to 614,400 bp / day



## Testimonial

"In our laboratory, we use the MassARRAY identification system in several projects. This technology serves as a basis for developing new methods for rapid genotyping of hepatitis B and C viruses and quasispecies analysis of the hepatitis C virus. Currently, a new project is underway to adapt the MassARRAY system to sequencing of whole genomes of hepatitis A and B viruses."

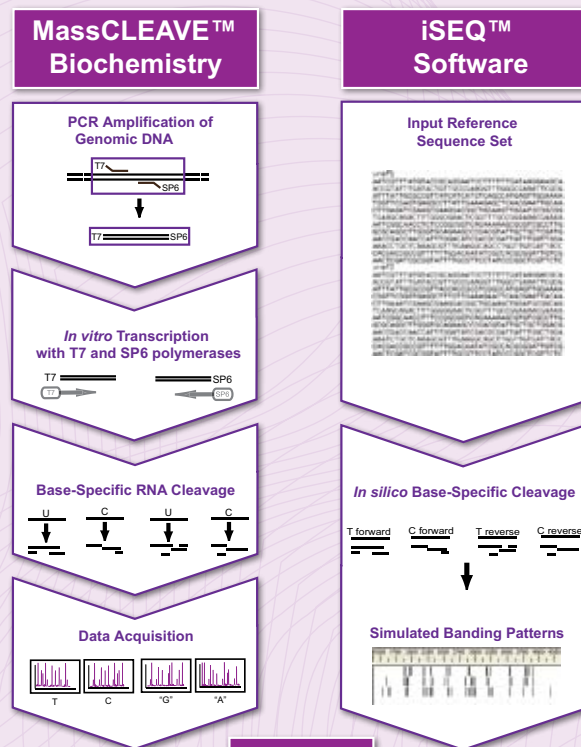
Laboratory of Molecular Epidemiology and Bioinformatics, Division of Viral Hepatitis, Centers for Disease Control and Prevention (CDC).

## Accurate, Rapid Comparative Sequence Analysis

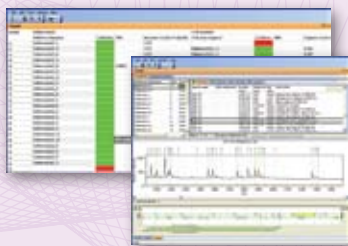
### How It Works

iSEQ™ uses reference sequences as a comparative measure to identify microbes, viruses, or other haploid organisms. The starting point of the protocol is the amplification of a target region of interest. T7- and SP6- promoter tagged primers are used to amplify the template.

After SAP treatment, *in vitro* transcription provides RNA transcripts which are base-specifically cleaved. The resulting RNA cleavage products are analyzed by matrix-assisted laser desorption ionization time-of-flight (MALDI-TOF) mass spectrometry. The iSEQ™ software automatically generates a report containing ID results by listing the best matching reference sequences and any sequence variations for each target region.



### Results



Sample Identification  
Sequence Variation Detection  
Cluster Analysis (Distance Matrix)

**Instrumentation** – Sequenom developed the MassARRAY system and SpectroCHIP® arrays to meet the requirements of moderate- to high-throughput comparative sequence analysis. Several system options are available depending on throughput and study requirements.

**Software** – The iSEQ™ software provides an advanced and versatile solution for automated data analysis:

- **Identification** – compares unknown samples to a set of reference sequences and identifies a best match
- **Sequence Variation Detection** – detects single base pair deviations and identifies new sequences
- **Clustering** – produces distance matrices that facilitate sample clustering based on homologous mass spectrometric patterns

Numerical and graphical interpretation tools are available for easy visualization and analysis. Data quality control and confidence ratings are provided for accurate identification.



## ■ MassARRAY® System

### MassARRAY® Compact System

The MassARRAY technology is trusted by the leading genetics institutions worldwide. The bench top MassARRAY Compact System is a multi-application platform that addresses the following applications:

- Comparative Sequence Analysis
- SNP Genotyping
- Quantitative Gene Expression
- Methylation Analysis

### MassARRAY® Advantage

The iSEQ™ software for the MassARRAY System addresses the need for rapid and accurate comparative sequence analysis. The combination of the iSEQ™ comparative sequence analysis and iPLEX® SNP Genotyping capabilities enable automated identification and detection of microbes and viruses for research applications, including clinical microbiology, epidemiological and surveillance monitoring, biodefense, agricultural and food science applications, as well as forensics.

MassARRAY Genotyping facilitates identification and prioritization of genetic targets within each stage of

biomedical research.

The Quantitative Gene Expression and Methylation applications allow functional analysis and target characterization.



Flexibility of Scale with  
Versatility of Application

## ■ Publications

Honisch, C., et al. (2007)

“Automated comparative sequence analysis by base-specific cleavage and mass spectrometry for nucleic acid-based microbial typing.”  
*Proc Natl Acad Sci USA* 104 (25): 10649-54.

Herring, C. D., et al. (2006)

“Comparative genome sequencing of *Escherichia coli* allows observation of bacterial evolution on a laboratory timescale.”  
*Nat Genet* 38 (12): 1406-12.

Honisch, C., et al. (2004)

“High-throughput mutation detection underlying adaptive evolution of *Escherichia coli*-K12.”  
*Genome Res* 14 (12): 2495-502.

Lefmann, M., et al. (2004)

“Novel mass spectrometry-based tool for genotypic identification of mycobacteria.”  
*J Clin Microbiol* 42 (1): 339-46.

Stanssens, P., et al. (2004)

“High-throughput MALDI-TOF discovery of genomic sequence polymorphisms.”  
*Genome Res* 14 (1): 126-33.

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The products described in the brochure are for Research Use Only, and not for use in diagnostic procedures or for other purposes.