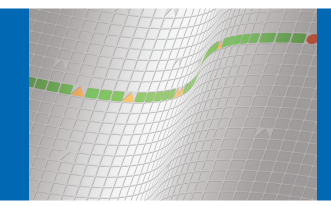


# GeneMapper<sup>®</sup> *ID-X* Software Version 1.0

**Note:** To improve the clarity of graphics in this PDF file, use the zoom tool to increase magnification to 150% or greater.





Software

Version 1.0

GeneMapper<sup>®</sup> ID-X

Getting Started Setting Up the Software Creating a Project,

Analyzing, and Reviewing Analysis Workflow Summaries

Manually Reviewing and Interpreting Data

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Performing Quality Control of Sample Results

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# Preface

## How to Use This Guide

Purpose of This Guide	<ul> <li>The <i>GeneMapper</i><sup>®</sup> <i>ID-X Software Version 1.0 Getting Started Guide</i> explains how to perform AmpFℓSTR<sup>®</sup> kit data analysis using the GeneMapper<sup>®</sup> <i>ID-X</i> Software Version 1.0. This guide functions as both:</li> <li>A tutorial, using example experimental data provided with the GeneMapper <i>ID-X</i> Software.</li> <li>A guide for your own experiments.</li> <li>In addition, this guide introduces you to the features of the GeneMapper<sup>®</sup> <i>ID-X</i> Software Version 1.0.</li> </ul>
Audience	This guide is written for forensic analysts who perform $AmpF\ell STR^{\circledast}$ kit data analysis using the GeneMapper <i>ID-X</i> Software.
Assumptions	This guide assumes that you have:
	<ul> <li>Installed GeneMapper <i>ID-X</i> Software version 1.0 as described in the <i>GeneMapper<sup>®</sup> ID-X Software Version 1.0 Installation</i> <i>Guide.</i></li> <li>Used AmpF<i>l</i>STR<sup>®</sup> amplification kit data for human</li> </ul>
	<ul> <li>identification (HID) applications.</li> <li>Developed a working knowledge of the Microsoft<sup>®</sup> Windows<sup>®</sup></li> </ul>
	• Developed a working knowledge of the Microsoft® Windows® XP operating system.
Text Conventions	This guide uses the following conventions:
	• Bold text indicates user action. For example:
	<ul> <li>Type 0, then press Enter for each of the remaining fields.</li> <li><i>Italic</i> text indicates new or important words and is also used for emphasis. For example:</li> </ul>
	Before analyzing, <i>always</i> prepare fresh matrix.

• A right arrow symbol ( ▶ ) separates successive commands you select from a drop-down or shortcut menu. For example:

Select File ▶ Open ▶ Spot Set.

Right-click the sample row, then select **View Filter** > **View All Runs**.

# User Attention<br/>WordsTwo user attention words appear in Applied Biosystems user<br/>documentation. Each word implies a particular level of observation<br/>or action as described below:

**Note:** – Provides information that may be of interest or help but is not critical to the use of the product.

**IMPORTANT!** – Provides information that is necessary for proper instrument or software operation, accurate chemistry kit use, or safe use of a chemical.

Examples of the user attention words appear below:

**Note:** Each registered user has his or her own set of preferences. When you set these options, if affects only the user currently logged in.

**IMPORTANT!** To verify your client connection to the database, you need a valid user ID and password.

## How to Obtain More Information

Related Documentation

The following related documents are shipped with the system:

- GeneMapper<sup>®</sup> ID-X Software Version 1.0 Installation Guide Provides procedures for installing version 1.0 of the GeneMapper<sup>®</sup> ID-X Software.
- *GeneMapper*<sup>®</sup> *ID-X Software Version 1.0 Administrator's Guide* – Provides procedures for creating user accounts, user groups, and security groups; configuring the audit trail and E-signature tools; and maintaining version 1.0 of the GeneMapper<sup>®</sup> *ID-X* Software.
- *GeneMapper*<sup>®</sup> *ID-X Software Version 1.0 Help* Contains context-sensitive help for all screens, and provides procedures and background information needed to use the software.
- *GeneMapper*<sup>®</sup> *ID-X Software Version 1.0 Quick Reference Guide* – Provides abbreviated procedures for analyzing, viewing, and interpreting data using GeneMapper<sup>®</sup> *ID-X* Software.
- *GeneMapper*<sup>®</sup> *ID-X Software Version 1.0 Reference Guide* Describes peak detection, sizing, and genotyping algorithms, and the GeneMapper<sup>®</sup> *ID-X* Software quality value system.

Portable document format (PDF) versions of this guide and the other documents listed above are also available on the *GeneMapper*<sup>®</sup> *ID-X Software Version 1.0 Documentation CD*.

**Note:** To open the user documentation included on the *GeneMapper*<sup>®</sup> *ID-X Software Version 1.0 Documentation CD*, use the Adobe<sup>®</sup> Acrobat<sup>®</sup> Reader<sup>®</sup> software available from **www.adobe.com**.

**Note:** For additional documentation, see "How to Obtain Support" on page xiii.

### Obtaining Information from the Help System

The GeneMapper<sup>®</sup> ID-X Software has a Help system that describes how to use each feature of the user interface. Access the Help system by doing one of the following:

- Click 🕐 in the toolbar of the Project window
- Select Help > Contents and Index
- Press F1

You can use the Help system to find topics of interest by:

- Reviewing the table of contents
- Searching for a specific topic
- Searching an alphabetized index

You can also access PDF versions of all documents in the GeneMapper<sup>®</sup> ID-X Software document set from the Help system.

#### **Send Us Your Comments** Applied Biosystems welcomes your comments and suggestions for improving its user documents. You can e-mail your comments to:

### techpubs@appliedbiosystems.com

**IMPORTANT!** The e-mail address above is only for submitting comments and suggestions relating to documentation. To order documents, download PDF files, or for help with a technical question, go to http://www.appliedbiosystems.com, then click the link for Support. (See "How to Obtain Support" on page xiii).

## How to Obtain Support

For HID support, you can send an e-mail to **HIDTechSupport@appliedbiosystems.com** or call **888-821-4443** option **1**.

For HID support outside North America, contact your local support office.

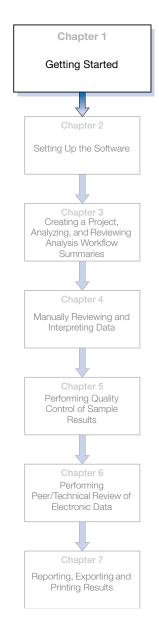
For the latest services and support information for all locations, go to **http://www.appliedbiosystems.com**, then click the link for **Support**.

At the Support page, you can:

- Access worldwide telephone and fax numbers to contact Applied Biosystems Technical Support and Sales facilities
- Search through frequently asked questions (FAQs)
- Submit a question directly to Technical Support
- Order Applied Biosystems user documents, MSDSs, certificates of analysis, and other related documents
- Download PDF documents
- Obtain information about customer training
- · Download software updates and patches

Preface How to Obtain Support

# Getting Started 1



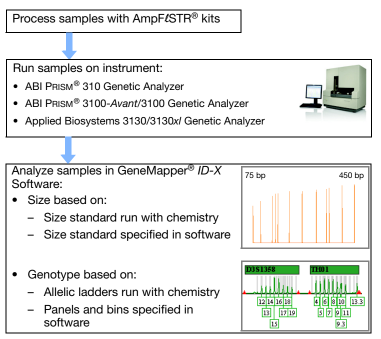
This chapter covers:

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GeneMapper $^{\ensuremath{\mathbb{R}}}$ ID-X Software and Analysis Workflows $\hdots$ . $\hdots$ 6
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# Overview of the GeneMapper<sup>®</sup> *ID-X* Software Version 1.0

GeneMapper<sup>®</sup> *ID-X* Software Version 1.0 is an automated genotyping software solution for all human identification (HID) data analysis needs, including forensic casework, databasing, and paternity testing. The workflow for performing AmpFℓSTR<sup>®</sup> kit data analysis with GeneMapper *ID-X* Software is shown below.



**Note:** GeneMapper<sup>®</sup> *ID-X* Software has undergone a verification process defined by Applied Biosystems. However, human identification laboratories analyzing forensic, paternity, databasing and single-source samples that choose to use GeneMapper *ID-X* Software for data analysis should perform their own appropriate validation studies.



1

## Features of the GeneMapper® *ID-X* Software

	The software provides a quality value system and a set of streamlined data review tools and features for both expert system and traditional manual review workflows. Included in the system are the following features:
Analysis Requirement Check	The analysis requirement check identifies unmet requirements before analysis starts. For example, the software checks if there is a sample with Sample Type = Allelic Ladder listed in the Samples table. The system can be set up to stop analysis and display an alert if this sample type is not found.
Allelic Ladder	The allelic ladder quality assessment:
Quality Assessment	• Evaluates allelic ladders (based on system-defined allelic ladder quality requirements) before proceeding to sample analysis
	• Flags run folders without at least one passing allelic ladder. You can review allelic ladders before proceeding with analysis.
	• Automatically excludes low-quality ladders from analysis and continues analysis with passing ladders. You can optionally override the software assessment and use low-quality ladders to generate bin offsets.
Analysis	For efficient data evaluation, the analysis summary provides:
Summary	• An easy-to-view summary of analysis results
	• An overview of allelic ladder, control, and sample quality
	• A separation of passing samples from samples that do not meet one or more quality thresholds
	• Interactive links to specific categories of samples
	(passing/check/low quality, allelic ladder/control/sample)
Comprehensive Quality Value System	<b>Note:</b> For more information on the Quality Value system, see "The GeneMapper <sup>®</sup> ID-X Software Quality Value System" on page 7.

Manual Review Tools	For efficient data review in traditional manual review and expert systems workflows, the manual review tools provide:
	<ul> <li>Process quality value (PQV) flags (, , , , )</li> <li>Automatic spike labeling based on intelligent rules</li> <li>User-defined "artifact" peak labels</li> <li>Marker-specific quality value details with thresholds and observed values displayed to show deviations from thresholds and identify sources of anomalies</li> <li>Mark Sample for Deletion and Delete All Labels from Sample functions in the Samples plot to allow easy elimination of low-quality samples directly from the plot window</li> <li>Override Genotype Quality (GQ) and Composite Genotype Quality (CGQ) functions to "manually accept" genotypes at the marker and sample level</li> <li>Detailed label edit table display and visual indicators to indicate</li> </ul>
	edits (gray PQVs) for electronic peer/technical review
Quality Control	The quality control features:
	<ul> <li>Evaluate sample concordance and allele matching</li> <li>Support additional custom positive controls and allow automatic concordance checks of custom controls</li> </ul>
	• Compare samples in a project to one another to determine if they contain profiles similar to neighboring samples
	• Compare samples in a project to laboratory reference and custom control profiles using a user-defined match percent threshold
Chain-of-Custody Systems for Electronic Data	The chain-of-custody systems for electronic data can be custom-configured (or turned off) by the GeneMapper <i>ID-X</i> System administrator as needed. These systems provide:
	• Security that controls user access to software functions and data, and allows custom configuration that meets the data-sharing needs of your laboratory and limits access to data when needed
	• Auditing that tracks changes and provides audit history reports

• E-signature that requires user-authentication before changes are saved

For more information on these features, see the *GeneMapper*<sup>®</sup> *ID-X* Software Version 1.0 Administrator's Guide.

Multi-User The multi-user database environment:

Environment

- Allows multiple users to share projects in a centralized database
- Facilitates efficient data sharing between analysts for second analysis and review
- · Limits the need to import and export projects
- Allows central management of analysis settings

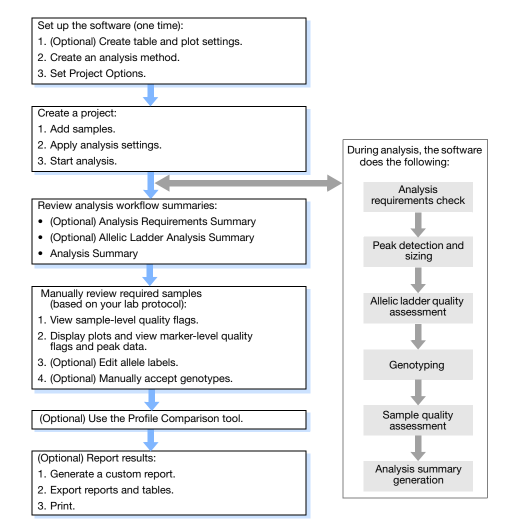
**Report Manager** For customized table-formatted reports, the Report Manager:

- · Allows you to change column names and column order
- Can export a traditional horizontal allele table for a selected group of samples for use as a genotype summary or import into LIMS or other downstream applications



# GeneMapper<sup>®</sup> *ID-X* Software and Analysis Workflows

The following flowchart summarizes the steps for performing a typical data analysis workflow using the GeneMapper<sup>®</sup> ID-XSoftware. To the left are the steps the user performs when analyzing samples and interpreting results. To the right are the software operations that occur automatically during analysis.





1

# The GeneMapper<sup>®</sup> *ID-X* Software Quality Value System

Overview	The GeneMapper <sup>®</sup> <i>ID-X</i> Software quality value system:	
	• Assesses the quality of allelic ladders before analysis and does not consider low-quality allelic ladders for genotyping.	
	• Assesses the quality of data at the sample and marker level using PQVs.	
	• Can be used in an optimized and validated expert system or traditional manual review workflow to quickly identify data quality issues and aid in interpretation of samples that do not meet all thresholds.	
	• Can be used in an optimized and validated expert system workflow to quickly segregate samples that require manual review from those that do not.	
	The PQV results of the quality assessment are displayed as color-coded flags: Pass (, ), Check, (), Low Quality (). The color of the flag depends on software-specified thresholds or user-defined thresholds set in the analysis method.	
Optimizing and Validating an Expert System	Before using any software as an expert system, optimize and validate the thresholds for each AmpFlSTR <sup>®</sup> kit and instrument platform combination by processing a variety of samples that challenge each of the different quality flags.	
	<b>IMPORTANT!</b> Different kit/instrument combinations may require different thresholds.	



### **Quality Value System Checks and Assessments**

The GeneMapper *ID-X* Software quality value system performs the following checks and assessments:

- Analysis requirements checks Before analysis starts, identifies any conditions that may prevent analysis or cause unexpected results.
- Sizing quality assessment Evaluates the quality of the size standard profile in each sample.
- Allelic ladder quality assessment Evaluates allelic ladder quality. Also determines if an allelic ladder is used for creating bin offsets.
- Marker-level quality assessment Evaluates labeled peaks within each marker. Contributes to the overall genotype quality assessment.
- **Sample-level quality assessment** Evaluates the quality of the entire sample.
- Genotype quality assessment Evaluates the quality of each marker in a sample. Contributes to the overall composite genotype quality assessment.

The following sections contain a brief description of each quality value system check and assessment along with a list of each sample-level and marker-level quality value.

**Note:** For more information on the quality value system, see the *GeneMapper*<sup>®</sup> *ID-X Software Version 1.0 Reference Guide*.

### **Analysis Requirements Checks**

The analysis requirements checks are performed and results displayed either in the Samples table before analysis starts or in the Analysis Requirements Summary after analysis starts.

Acronym	Full Name	Description and Flags 📄 📥
ARNM	Analysis Requirement Not Met	<ul> <li>Indicates if all analysis requirements are met. These requirement checks are performed when analysis is started:</li> <li>Sample File Not Found</li> <li>Analysis Method Not Selected</li> <li>Analysis Method Not Found in the Database</li> <li>Panel Not Selected</li> <li>Panel Not Found in the Database</li> <li>Binset Not Selected</li> <li>Binset in Analysis Method Does Not Match Binset Selected in the Panel Manager</li> <li>Size Standard Not Selected</li> <li>Size Standard Not Found in Database</li> <li>Size Standard Not Found in Database</li> <li>Natrix Not Selected</li> <li>Matrix Not Found or Contains Invalid Data</li> <li>No Allelic Ladder Selected in Run Folder</li> <li>GMID v3.x Analysis Method Selected</li> <li>SNP Panel Selected</li> </ul>



### Sizing Quality Assessment

The quality value system evaluates the quality of the size standard profile within each sample (SQ) and allows you to flag size standards with poor peak resolution. Sizing quality assessment is displayed in the Samples table after analysis completes.

Acronym	Full Name	Description and Flags 🔳 📥 🥮	
SQ	Sizing Quality	Evaluates the similarity between the fragment pattern for the size standard dye specified in the size standard definition and the actual distribution of size standard peaks in the sample, calculates an interim SQ (a value between 0 and 1), then applies the broad peak weighting specified in the analysis method, as described in the <i>GeneMapper</i> <sup>®</sup> <i>ID-X Software Version 1.0 Reference Guide</i> .	
		<b>Note:</b> The GeneMapper <i>ID-X</i> Software does not genotype samples with SQ.	

### Allelic Ladder Quality Assessment

The quality value system performs an allelic ladder quality assessment to determine if a ladder is used in genotyping (to create bin offsets). Allelic ladder samples are analyzed before all other samples. An allelic ladder sample must have a SQ and a CGQ to be used for creating bin offsets. For an allelic ladder to have a CGQ, all the markers within the allelic ladder must pass the following rules:

Rule	Description
1	All ladder alleles specified in the panel used to analyze are detected.
2	In each marker, the peak height ratio of the first and second peak is greater than 50%. This rule eliminates allelic ladders if the stutter peak before the first true allele peak is labeled as an allele.
3	No spikes are detected above 20% (default) of the highest allele peak in the same dye color within the extended marker range.
	<b>Note:</b> Spike detection for allelic ladders is performed within each extended marker range (no gaps are present between markers; the end point of each marker is extended past the marker definition in the panel to the beginning of the next marker).
	<b>Note:</b> The Allelic Ladder Spike Cut-off value is user-definable in the Peak Quality tab of the analysis method.
4	The peak height ratio between the lowest and highest peak is equal to or greater than 15%.

### Marker-Level Quality Assessments

Marker-level quality assessments indicate the quality of each marker in a sample and are displayed in the Genotypes table after analysis completes.

Acronym	Full Name	Description and Flags 📄 📥	
OS	Off-scale	Indicates if any fluorescence signal within the marker exceeds the detection threshold of the instrument.	
BIN	Out of Bin Allele	Indicates if labeled peaks do not fall inside bins. These peaks are labeled with OL (Off ladder).	
PHR	Peak Height Ratio	Indicates if the peak height ratio between the lowest and highest peak is less than the Min Peak Height Ratio defined in the analysis method.	
MPH	Max Peak Height	Indicates if any peak heights (in RFU) within the marker size range exceed the Max Peak Height value (in RFU) set in the analysis method.	
LPH	Low Peak Height	<ul> <li>Indicates if any peak heights (in RFU) within the marker size range are below the following thresholds set in the analysis method:</li> <li>Homozygous Min Peak Height</li> <li>Heterozygous Min Peak Height</li> </ul>	
AN	Allele Number	Indicates if the software detects no alleles, more than the Max Expected Alleles set in the analysis method (Peak Quality tab), or no X allele detected in amelogenin.	
BD	Broad Peak	Indicates if the width of any peak exceeds the Max Peak Width (half height in base pairs) defined in the analysis method (Peak Quality tab).	
CC	Control Concordance	Indicates if a positive, custom, or negative control produces the expected profile.	
SPK	Marker Spike	<ul> <li>Allelic ladders – Indicates if spikes are detected within each extended marker range (no gaps are present between markers; the end point of each marker is extended past the marker definition in the panel to the beginning of the next marker).</li> <li>Samples – Indicates if spikes are detected within a marker size range. The software uses a proprietary algorithm that detects spikes based on the peak morphology.</li> </ul>	
OVL	Overlapping Alleles	Indicates if a labeled peak (allele or artifact) falls within the size ranges of two neighboring markers.	



### **Genotype Quality Assessment**

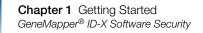
For samples, the quality value system assigns the GQ for each marker based on the individual marker quality flags. For allelic ladders, the quality value system assigns a GQ for each marker based on the allelic ladder quality requirements, as described in "Allelic Ladder Quality Assessment" on page 10. The GQ is used to determine the CGQ, and is displayed in the Genotypes table after analysis completes.

Acronym	Full Name	Description and Flags 🔳 🍐	
GQ (samples)	Genotype Quality	Indicates the genotype quality of the marker in the sample. The genotype quality for a sample marker is determined based on the presence of labeled peaks detected (after filtering) and the GQ weighting specified in the analysis method.	
		If no labeled peaks are detected (and the sample is not a negative control), the GQ is set to 0. If one or more labeled peaks are detected, the GQ is initially set to 1 with a final value determined by the GQ weighting of individual marker-level quality values as specified in the analysis method.	
GQ (allelic ladders)	Genotype Quality	Indicates the genotype quality of the marker in the allelic ladder. The genotype quality for an allelic ladder marker is determined using system-defined quality rules (as described in the <i>GeneMapper<sup>®</sup> ID-X Software Version 1.0 Reference Guide</i> ) to ensure:	
		All expected peaks are present.	
		• Peak height ratio of the first and second peak is greater than 50%.	
		<ul> <li>No spikes are present in the extended marker range (within or between markers).</li> </ul>	
		<ul> <li>The peak height ratio between the lowest and highest peak is equal to or greater than 15%.</li> </ul>	
		<b>IMPORTANT!</b> If the Allelic Ladder GQ Weighting for Spikes is set to 0 (off) in the analysis method, the GQ may be, even if spikes are present in the allelic ladder.	

### Sample-Level Quality Assessments

Sample-level quality assessments that indicate the quality of the entire sample are displayed in the Samples table after analysis completes.

Acronym	Full Name	Description and Flags 📕 📥	
SOS	Sample Off-scale	Indicates if any fluorescence signal within the analysis range exceeds the detection threshold of the instrument.	
MIX	Mixed Source	Indicates a potential mixed-source sample.	
OMR	Outside Marker Range	Indicates if labeled peaks are detected between two marker size ranges defined in the panel.	
SSPK	Sample Spike	<ul> <li>Allelic ladders – Indicates if spikes are detected within the sizing range.</li> <li>Samples – Indicates if spikes are detected within or between two</li> </ul>	
		defined marker size ranges. Does not indicate if spikes are detected before the first marker or after the last marker.	
		The software uses a proprietary algorithm that detects spikes based on the peak morphology.	
CGQ (samples)	Composite Genotype Quality	Indicates overall sample genotype quality. Considers the individual marker GQ values.	
CGQ (allelic	Composite Genotype	Indicates overall allelic ladder quality. Considers the allelic ladder quality assessment (see page 10).	
ladders)	Quality	<b>Note:</b> Allelic ladder samples with <b>()</b> CGQ are not used to create bin offsets.	
		<b>IMPORTANT!</b> If the Allelic Ladder GQ Weighting for Spikes is set to 0 (off) in the analysis method, the CGQ may be even if spikes are present in the allelic ladder.	



## GeneMapper<sup>®</sup> *ID-X* Software Security

The GeneMapper<sup>®</sup> *ID-X* Software contains a security system that includes the following features relevant to the procedures outlined in this guide:

Security System Component	Description	
User accounts	A required component of the security system that allows only authorized users to access the software.	
	This guide assumes that you log in using the with the software (see the <i>GeneMapper<sup>®</sup> ID-, Administrator's Guide</i> for a description of use detailed instructions for logging in.	X Software Version 1.0
Security groups	A component of the security system that resitems (project, panels, analysis methods, and	
	When using the Practice User account, data is assigned to the Practice Security Group automatically (you are not prompted to select a security group).	Save Project       Name:     Getting Started AB       Security Group:     Practice Security Group
		OK Cancel Help
Audit trail	A component of the security system that keeps track of changes made and can require users to provide a reason for a change.	
	The audit trail is set up by default to require a reason for change when you edit allele labels. If additional auditing is set up on your system, you may be prompted to specify a reason when you create or edit data items.	Enter the Research() for Charge: Ricroveriant OBR
E-signature	system that requires users to provide a valid user name and password when they make a change.	Electronic Signature Verification
	This guide assumes that E-signature <i>is not</i> set up on your system.	by entering your UserID and Password, you are signing the action
	If E-signature is set up on your system, you are prompted to provide your user name and password when you create or edit data items.	Liser 20 Password CK Concol



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For information on:

- Setting up security, audit trail, and E-signature on your system, see the *GeneMapper*<sup>®</sup> *ID-X Software Version 1.0* Administrator's Guide.
- Using security groups (and the impact that security groups have on the data you can access), using the audit trail, and using E-signature, see the *GeneMapper*<sup>®</sup> *ID-X Software Help*.

## GeneMapper<sup>®</sup> ID-X Software Terms

Term	Definition
allele	Variant form of a marker (locus).
allelic ladder	A sample that contains a set of alleles that are representative of those found in a particular STR marker. Allelic ladders are generated with the same primers as tested samples, providing a reference DNA size for each allele included in the ladder. Unknown samples are compared against them to determine their genotype.
AmpF <b>/</b> STR <sup>®</sup> Chemistry kit	Applied Biosystems Human Identification PCR amplification kits for short tandem repeat (STR) analysis.
analysis method	A collection of user-defined settings that determine the sizing, genotyping and quality value algorithms used by the GeneMapper <sup>®</sup> <i>ID-X</i> Software to analyze sample files in a project.
bin	A fragment size ( $\pm$ 0.5 bp) that defines an allele within a marker.
bin set	A collection of bins (allele definitions), typically specific to a set of panels.
CODIS	The FBI Laboratory Combined DNA Index System. For more information, see: http://www.fbi.gov/hq/lab/codis/index1.htm
custom control	A positive amplification control other than the control DNA supplied with the AmpF\ell STR $^{\mbox{\tiny B}}$ kits.

Some common terms used in this guide are:



Term	Definition
expert system	A computer program that:
	Interprets alleles and DNA profiles
	Uses knowledge acquired from Qualified Analysts
	Is Rule-based
	<ul> <li>Acts as an "assistant" to Qualified Analysts</li> </ul>
	Automated-minimal human intervention
	<ul> <li>Is as good or better than human experts</li> </ul>
	Documents reasoning behind decisions
	<ul> <li>Does not require manual review of "passing" samples</li> </ul>
	Source: National DNA Index System (NDIS) DNA Data Acceptable Standards, Appendix B, "Guidelines for Submitting Requests for Approval of an Expert System for Review of Offender Samples," 15 July 2004.
genotype	Allele designations for a genetic locus.
genotyping	Labeling of alleles based on allelic ladder bin comparisons and filtering of alleles based on analysis method settings.
lab reference profile	A genotyped profile of an analyst or other lab personnel.
marker	A genetic locus. A name, fragment size range in base pairs, dye color, repeat length, and physical allelic ladder alleles are defined for each marker.
negative control	A sample expected to generate no allele calls. Can be an extraction blank, reagent blank or an amplification negative control.
panel	A group of markers and properties (size ranges, dye label color, expected positive control genotypes). Each panel provided with the GeneMapper <sup>®</sup> <i>ID-X</i> Software represents a specific AmpF <i>l</i> STR <sup>®</sup> Chemistry kit.
positive control	A sample of known genotype. Each AmpF <i>l</i> STR <sup>®</sup> Chemistry kit includes a positive control. You can also run custom positive controls.
PQV (Process Quality Value)	Process quality values (PQVs) assess the quality of data at the sample and marker levels.
profile	The genotype (allele designations) of a sample.
project	In the GeneMapper <sup>®</sup> <i>ID-X</i> Software, a collection of sizing and genotyping results for a set of data.
run	The electrophoretic injection of a set of samples from a single plate (48- or 96-well) and the resulting sample files.
run folder	A folder containing a set of sample files from a capillary-electrophoresis run.



Term	Definition	
sample files	Capillary-electrophoresis data files (*.fsa) generated by Data Collection Software.	
size standard definition	A list of fragment sizes in base pairs that a size standard sample contains. Only those sizes required for accurate sizing are contained in the size standard definition.	
stutter ratio	Percent value used to filter stutter peaks (remove labels).	
traditional manual review	Visual inspection of samples regardless of quality.	

For additional definitions, see the *GeneMapper*<sup>®</sup> *ID-X Software Help* (select **Glossary** in the Contents tab).

## How to Use This Guide

Before You Start	<b>IMPORTANT!</b> Before using the procedures in this guide, make sure the GeneMapper <sup>®</sup> <i>ID-X</i> Software has been successfully installed and registered. See the <i>GeneMapper</i> <sup>®</sup> <i>ID-X</i> Software Version 1.0 Installation Guide for more information.
	When performing the procedures described in this guide, keep in mind the following:
	• The steps in each chapter are designed to flow from start to finish, and from one chapter to the next
	• Complete each chapter as a single unit before stopping your work, if possible
	• Make sure you perform each step as it is described
	• Carefully review any previously performed steps if you observe any differences between what is shown in this guide and what is displayed on your own system
User Account Requirements	You must use the Practice User account provided to perform the procedures in this guide.

### Using the Guide with the Example Data Provided

Example data (.fsa) generated using the Applied Biosystems AmpF*l*STR<sup>®</sup> Identifiler<sup>®</sup> PCR amplification kit, and a reference project (.ser) containing analyzed lab reference samples, custom controls, and QC samples are installed with the GeneMapper<sup>®</sup> *ID-X* Software.

To perform the exercises described in this guide, use the example files (.fsa and .ser) located on your computer as shown below.

Install Configuration	File Location		
	Reference project (.ser)	Sample files (.fsa)	
Client install	<pre><drive>:\AppliedBiosystems\ GeneMapperID-X\Client\Example Data\ Projects</drive></pre>	<pre><drive>:\AppliedBiosystems\ GeneMapperID-X\Client\Example Data\ Identifiler Samples</drive></pre>	
Full install	<drive>:\AppliedBiosystems\ GeneMapperID-X\Example Data\ Projects</drive>	<drive>:\AppliedBiosystems\ GeneMapperID-X\Example Data\ Identifiler Samples</drive>	

**Note:** The drive will vary depending on the installation of the GeneMapper<sup>®</sup> *ID-X* Software. The default installation drive is the Local Disk drive. See the *GeneMapper*<sup>®</sup> *ID-X* Software Version 1.0 *Installation Guide* for more information on installation options.

### Using the Guide as a Tutorial

This guide is a tutorial designed to help you follow a typical analysis workflow. Using the example data provided with the GeneMapper *ID-X* Software, follow the procedures in Chapters 2-7:

Chapter	Description
Chapter 2, Setting Up the Software	1. View default panels and bins
	2. Create an analysis method
	3. View default table and plot settings
	4. Create new table and plot settings
	5. View default size standard definitions
	6. Add custom control and lab reference profiles to database
	7. Set project options



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Chapter	Description
Chapter 3, Creating a Project, Analyzing, and Reviewing Analysis Workflow Summaries	1. Create a project and add samples
	2. Analyze the data
	3. Review the Analysis Requirement Summary
	4. Review the Analysis Summary
Chapter 4, Manually Reviewing and Interpreting Data	1. View electropherograms
	2. Interpret anomalies using quality value flags and details
	3. Edit allele labels
	4. Manually accept marker genotypes
Chapter 5, Performing Quality Control of Sample Results	1. Perform Sample Concordance check
	2. Perform Sample Comparison check
	3. Perform Lab Reference Comparison check
	4. Perform Control/QC Sample Comparison check
Chapter 6, Performing Peer/Technical Review of Electronic Data	1. Use filtered tables to find edited samples
	2. Use Label Edit Viewer to view edited labels
	3. Manually accept sample profiles
	4. View edit comments in tables
Chapter 7, Reporting, Exporting and Printing Results	1. Generate a custom report
	2. Export reports and tables

For More<br/>InformationThis guide contains basic procedures. It does not describe all features<br/>and parameters in the GeneMapper<sup>®</sup> ID-X Software. For detailed<br/>information on topics presented in this guide, see the GeneMapper<sup>®</sup><br/>ID-X Software Help.



Chapter 1 Getting Started How to Use This Guide

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## **Overview**

This chapter demonstrates how to prepare the GeneMapper<sup>®</sup> ID-X Software for analysis, using the example data set provided with the software.

#### In This Chapter In this chapter you will learn how to:

- Start the software and log in
- Review the panels, bins, and stutter files provided
- Create analysis methods
- Review the table settings, plot settings and size standards provided
- · Create new table settings and plot settings
- Import lab reference and custom control profiles
- Set project options

## Step 1: Start the Software and Log In

Starting the Software and Logging In

- 1. Double-click (GeneMapper<sup>®</sup> ID-X v1.0) on the desktop to launch the software.
- In the Login to GeneMapper<sup>®</sup> ID-X dialog box, enter Practice User for User Name and password for Password, then click OK.

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🧬 Login to GeneMapper® ID-X	X
GeneMapper* D-2X Software Version 10 Magnetic Ministramenedd All rights meanedd	User Name Practice User  Password: ******* Database Host: FOSKOSMANCND01
New Host Delete Host	Default Host OK Exit Help

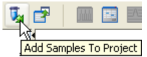
**Note:** If you have logged in before, you can select the user name from the drop-down list.

The Project window opens.

Toolbar									
Menu	Project	name	e Log	iged in use	r Com	puter nai	ne		
🥩 GeneMappe	r® ID-X - Ui	ntitled	Practice Use	er Is Logged In I	Database FOSK	OSMANCNDO	)1)		
File Edit Analys	sis View To	ols Admi	n Help	/					
🐸 🗃 📗	🆫 📄 📗		ш 🛛		🕨 💣 🕴 Table	Setting:		v 🖬 🛛 🔎	ð 🖪   🔤 🕐
🚠 Project	Sample	es Analys	sis Summary Ge	enotypes					
		Status S	5ample File	Sample Name	Sample ID	Comments	Sample Type	Specimen Category	Analysis Method
	1								
	2								
	3						ļ		
	4								
	6								-
	7								
	8	i					1		<u></u>
	9								
	10								
		<							>
						1			
Progress Status									[ Stop ]
			ne that li					owing Sample	s
	folders (	blank	for new p	orojects)		table (b	lank for n	ew projects)	
Status bar									

#### Resizing and Exploring the Project Window

- **1.** Adjust the window to display as many of the table columns as possible:
  - a. Click (Maximize) in the upper right corner of the Project window to expand the window to occupy the full area of the screen.
  - **b.** Resize columns by dragging the separating lines:
    - Position the pointer over the line separating two columns until the pointer changes to sizing arrows.
    - Click-drag the sizing arrows. Dragging to the left narrows the left-hand column. Dragging to the right widens the left-hand column.
- **2.** Place the pointer over toolbar buttons to display tooltips that explain the function of the button.

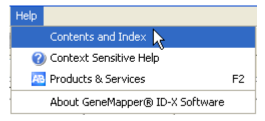


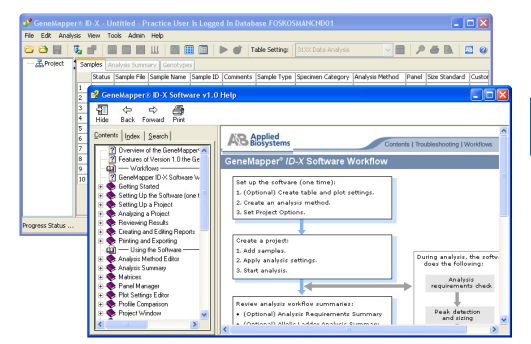
#### Exploring the Context-Sensitive Help System

The GeneMapper<sup>®</sup> ID-X Software context-sensitive Help system provides immediate access to detailed information regarding software views, functions and troubleshooting guidelines.

To access the Help system from the Project window:

1. Click the Help menu, then select Contents and Index.





Help opens in a separate window.

Note: You can also access the Help system anywhere in the software by pressing F1, by clicking @ in the toolbar of the Project window, or by clicking the **Help** button in a window, tab or dialog box to access help topics specific to that particular feature of the user interface. You can then click on the internal links within the specific help topic to navigate to related topics.

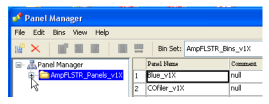
2. Click 🔀 (Close) to close the Help window and return to the Project window.



## Step 2: View Panels, Bins, and Stutter Settings

Overview	Before analyzing data, the software must have access to AmpF <i>l</i> STR <sup>®</sup> kit details such as marker size ranges and dyes, allele sizes, and stutter ratios. The files that contain this information are called Panel, Bin, and Stutter files, respectively.
	<b>IMPORTANT!</b> The panel, bin and stutter values shown in this section are configured specifically for use with AmpFℓSTR <sup>®</sup> kit data. Applied Biosystems recommends you use the provided panels and bins when analyzing AmpFℓSTR <sup>®</sup> data from your laboratory, unless your laboratory has validated alternative values.
When to Import	As part of the GeneMapper <sup>®</sup> $ID$ - $X$ Software installation process, the panel, bin and stutter files are automatically imported into the GeneMapper <sup>®</sup> $ID$ - $X$ Software database.
	<b>Note:</b> If you have installed GeneMapper <sup>®</sup> <i>ID-X</i> Software on the same workstation as Data Collection software (co-installation), you must manually import the panel and bin files. See the <i>GeneMapper</i> <sup>®</sup> <i>ID-X Software Help</i> for information on this procedure.
	<b>Note:</b> You can import new panel, bin and stutter files whenever updated versions are provided.
Viewing Panel, Bin, and Stutter	<ol> <li>If not already started, launch the GeneMapper<sup>®</sup> <i>ID-X</i> Software (see page 22).</li> </ol>
Settings	<b>2.</b> In the Project window toolbar, click 🛄 (Panel Manager).

- 2
- **3.** In the navigation pane of the Panel Manager, select the **AmpFLSTR\_Panels\_v1X** kit folder, then click ⊕ to expand its contents.



**4.** Double-click the **Identifiler\_v1X** panel folder. The markers found in the Identifiler kit are listed in the navigation pane and the marker details are displayed in the content pane.

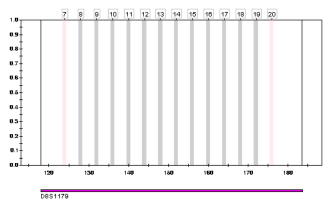
Marker	details
--------	---------

	Edit Bins View Help		Bin Set: An	npFLSTR_E	Bins_v1X		- III V.		<b>A M</b>	
			Marker Name	Dye Color	Min Size	Max Size	Control Alleles	Marker	Comments	Ladder Alleles
Г		1	D851179	Blue	118.0	183.5	13	4	none	8,9,10,11,12,13,14,15,16,:
		2	D21511	Blue	184.5	247.5	30	4	none	24,24.2,25,26,27,28,28.2,2
	⊞ ··· D75820	3	D75820	Blue	251.0	298.5	10,11	4	none	6,7,8,9,10,11,12,13,14,15
		4	CSF1PO	Blue	302.12	348.63	10,12	4	none	6,7,8,9,10,11,12,13,14,15
		5	D351358	Green	98.0	148.0	14,15	4	none	12,13,14,15,16,17,18,19
		6	TH01	Green	159.0	205.0	8,9.3	4	none	4,5,6,7,8,9,9.3,10,11,13.3
	⊞ D165539	7	D135317	Green	205.65	250.16	11	4	none	8,9,10,11,12,13,14,15
		8	D165539	Green	255.3	301.81	11,12	4	none	5,8,9,10,11,12,13,14,15
		9	D251338	Green	304.8	370.31	19,23	4	none	15,16,17,18,19,20,21,22,2
	TPOX	10	D195433	Yellow	101.0	148.0	14,15	4	none	9,10,11,12,12.2,13,13.2,14
	. ± D18551	11	VWA	Yellow	151.0	213.5	17,18	4	none	11,12,13,14,15,16,17,18,1
		12	TPOX	Yellow	216.99	260.99	8	4	none	6,7,8,9,10,11,12,13
	E FGA	13	D18551	Yellow	264.49	350.0	15,19	4	none	7,9,10,10.2,11,12,13,13.2,
	🛓 🔂 SEfiler_v1X	14	AMEL	Red	106.0	114.0	x	9	none	X,Y
<		15	D55818	Red	128.0	180.0	11	4	none	7,8,9,10,11,12,13,14,15,16
	Reference Samples	16	FGA	Red	206.25	360.0	23,24	4	none	17,18,19,20,21,22,23,24,2

#### Markers



**5.** Double-click the **D8S1179** marker in the navigation pane. A plot showing the bins for this marker is displayed in the content pane.



Markers can include two bin types: physical (represent alleles present in the allelic ladder sample) and virtual (represent alleles not present in the allelic ladder sample). The software displays all physical bins in grey, and all virtual bins (except for CODIS bins) in pink. In this example, the D8S1179 marker contains two virtual bins, 7 and 20, displayed in pink in the content pane.

- 6. In the navigation pane, click to expand the Yfiler\_v1X panel folder.
- 7. Click to expand the Y\_DYS392 marker, then select Stutter Ratio & Distance.



The marker-specific stutter ratios defined in the stutter file are displayed in the content pane. In this example, both Minus and Plus stutter ratios are specified for this marker.

	N	/linus Stutter			I	Plus Stutter	
	Ratio	From Distance	To Distance		Ratio	From Distance	To Distance
_	0.1622	2.25	3.75	1	0.0790	2.25	3.75
				2			
)				3			
ł				4			

**Note:** From this window, you can apply up to four minus and four plus stutter ratios per marker, and edit the default stutter percentages provided with the GeneMapper<sup>®</sup> ID-X Software.

8. Click OK to close the Panel Manager window.

## Step 3: Create an Analysis Method

Overview	Analysis methods define the peak detection, sizing, genotyping, and quality assessment parameters applied during analysis of sample data.
In This Section	In this section, you will create a new analysis method, configured specifically for use with the example data provided with the GeneMapper <sup>®</sup> $ID-X$ Software, and with the procedures outlined in this guide.
	<b>IMPORTANT!</b> The values used in this guide may not be suitable for analyzing data generated in your laboratory. You must optimize and validate these values during internal verification.

#### Creating the Analysis Method

- 1. In the Project window toolbar, click 🔲 (GeneMapper ID-X Manager).
- 2. Select the Analysis Methods tab, then click New.
- **3.** Complete the tabs of the Analysis Method Editor for this example analysis method as described in Table 1 on page 30, and use the *GeneMapper*<sup>®</sup> *ID-X Software Help* to learn more about the purpose of specific settings and the effects of changing these settings as needed.

#### Table 1 Analysis method settings for the example data (Identifiler<sup>®</sup> samples)

Tab	Settings							
General	Enter the settings as shown below:							
	Analysis Method Editor							
	General Allele Peak Detector Peak Quality SQ & GQ Settings							
	Analysis Method Description							
	Name: Getting Started _your initials							
	Security Group: Practice Security Group							
	Description: For use with the Getting Started Guide only							
	Instrument:							
	Analysis Type: HID							
	<b>Note:</b> When following the procedures in this guide, Applied Biosystems recommends you add your initials to the names of data objects you create so you can distinguish your data objects from similar objects that may be saved to the same GeneMapper <sup>®</sup> <i>ID-X</i> Software database by other laboratory personnel when performing the same steps in this guide.							
	<b>Note:</b> This analysis method is assigned to the Practice Security Group automatically (you are not prompted to select a security group), and is only available if you log in to the software with the Practice User account. You will verify the Project Options set for the Practice User account in "Step 6: Set the Project Options" on page 52.							



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#### Table 1 Analysis method settings for the example data (Identifiler® samples) (continued)

Tab				Setting	js			
Allele	Select the bin set and revie	w the	defau	It setting	gs as sh	own belov	v:	
	Analysis Method Editor						X	
	General Allele Peak Detector	Peak Qı	uality S	Q & GQ Sel	tings			
	Bin Set: AmpFLSTR_Bins_v1	×				~		
	✓ Use marker-specific stutte	er ratio a	and dista	nce if avail	able			
	Marker Repeat Type:		Tri	Tetra	Penta	Hexa		
	Global Cut-off Value		0.0	0.0	0.0	0.0		
	MinusA Ratio		0.0	0.0	0.0	0.0		
	MinusA Distance	From	0.0	0.0	0.0	0.0		
		То	0.0	0.0	0.0	0.0		
	Global Minus Stutter Ratio		0.0	0.0	0.0	0.0		
	Global Minus Stutter Distance	From	0.0	3.25	0.0	0.0		
		То	0.0	4.75	0.0	0.0		
	Global Plus Stutter Ratio		0.0	0.0	0.0	0.0		
	Global Plus Stutter Distance	From	0.0	0.0	0.0	0.0		
		То	0.0	0.0	0.0	0.0		
	Amelogenin Cutoff	0.0						
	<b>Note:</b> This analysis metho stutter ratios viewed in the Value and a Global Minus S minimize background labeli	Panel Stutter	Mana Ratio	iger will may be	be appli applied	ed. Howe	ver, a Glob -source s	oal Cut-off amples to

#### Table 1 Analysis method settings for the example data (Identifiler<sup>®</sup> samples) (continued)

Tab	Settings
Tab         Peak         Detector	Enter the settings as shown below: Analysis Method Editor          General Allele Peak Detector       Peak Quality       SQ & GQ Settings         Peak Detection Algorithm: Advanced       Peak Detection       Peak Amplitude Thresholds:         Partial Ra       Partial Sizes       Peak Amplitude Thresholds:         Baseline Window:       Start Size:       75         Smoothing       None       None         Olight       Heavy       Min. Peak Half Width:       2         Pack Calling Method       51       pts         Size Calling Method       Size Sigures       Sige Threshold         Order Least Squares       0.0       0.0
	Baseline window:     51     pts       Size Calling Method     Peak Start:     0.0       2nd Order Least Squares     0.0



#### Table 1 Analysis method settings for the example data (Identifiler® samples) (continued)

Tab	Settings
Peak Quality	If a marker does not meet a PQV threshold value set in this tab, the GeneMapper <sup>®</sup> <i>ID-X</i> Software will set the marker PQV flag to yellow <u>(Check)</u> .
	Review the default PQV threshold settings shown below. Click <b>Help</b> at the bottom of the tab and navigate to the <i>Peak Quality</i> help topic to learn more about the parameters presented in this tab.
	Analysis Method Editor
	General Allele Peak Detector Peak Quality SQ & GQ Settings
	Min/Max Peak Height (LPH/MPH)
	Homozygous min peak height 200.0
	Heterozygous min peak height 100.0
	Max Peak Height (MPH) 5000.0
	Peak Height Ratio (PHR)
	Min peak height ratio 0.7
	Broad Peak (BD)
	Max peak width (basepairs) 1.5
	Allele Number (AN)
	Max expected alleles 2
	Allelic Ladder Spike
	Cut-off Value 0.2

#### Table 1 Analysis method settings for the example data (Identifiler® samples) (continued)

Tab	Settings				
SQ & GQ Settings	The values entered in this tab affect the calculation of SQ and GQ. Review the default SQ and GQ settings shown below. Click <b>Help</b> at the bottom of the tab and navigate to the SQ & GQ Settings Tab help topic to learn more about the parameters presented in this tab.				
	Analysis Method Editor       Image: Control GQ Weighting         General Allele Peak Detector Peak Quality SQ & GQ Settings       Image: Control GQ Weighting         Quality weights are between 0 and 1.       Image: Control GQ Weighting				
	Broad Peak (BD)       0.8       Allele Number (AN)       1.0         Out of Bin Allele (BIN)       0.8       Low Peak Height (LPH)       0.3         Overlap (OVL)       0.8       Max Peak Height (MPH)       0.3         Marker Spike (SPK)       0.3       Off-scale (OS)       0.8         Peak Height Ratio (PHR)       0.3       O.3       Off-scale (OS)         Control Concordance (CC) Weight = 1.0 (Only applicable to controls)       Image: Control Concordance (CC) Weight = 1.0 (Only applicable to controls)       Image: Control Concordance (CC) Weight = 1.0 (Only applicable to controls)				
	SQ Weighting   Broad Peak (BD)   0.5     Allelic Ladder GQ Weighting   Spike (SSPK/SPK)   1   Sizing Quality:   From   0.75   to 1.0   From 0.75   to 1.0   From 0.75   to 1.0   From 0.75   to 1.0   From 0.75   to 1.0   From 0.75   to 1.0   From 0.75				

**4.** After completing all tabs, click **Save** to save your changes and close the Analysis Method Editor dialog. The GeneMapper ID-X Manager remains open.



2

# Step 4: Review Default Table Settings, Plot Settings, and Size Standards

**Overview** In this section, you will review the default table, plot, and size standard settings provided with the GeneMapper<sup>®</sup> *ID-X* Software. The default settings are designed to support a logical and efficient data analysis and review workflow using the example data provided with the software and for use when analyzing data generated in your laboratory. Use of the default settings is demonstrated throughout this guide.

**Note:** You can modify the default table, plot, and size standard settings or create new settings to support individual laboratory workflows.

### **Review Default Table Settings**

Overview	Table settings determine the content (columns) displayed in or
	exported from the Samples and Genotypes tables. The default table
	settings shown in this section include specific columns required to
	perform different workflow tasks.

- Reviewing Table Settings1. Open the GeneMapper ID-X Manager if not already open (see page 30).
  - **2.** In the GeneMapper ID-X Manager, select the **Table Settings** tab.
  - **3.** Click a table row to select the table setting to review, then click **Open**.
  - **4.** Review the default table settings provided using Table 2 on page 36.
  - **5.** After you have finished reviewing the default table settings, click **OK** to close the Table Settings Editor. The GeneMapper ID-X Manager remains open.



## **Defaults Provided** The following table settings are installed with the GeneMapper<sup>®</sup> ID-X Software.

#### Table 2 Default table settings

т	able Setting	Column Settings		
Name	Purpose	Samples Table	Genotypes Table	
310 Data Analysis	Used to set up table data to analyze sample files generated on an ABI PRISM <sup>®</sup> 310 Genetic Analyzer.	Displays analysis setting columns and sample-level quality values.	Displays marker and allele columns, and marker-level quality values.	
31XX Data Analysis	Used to set up table data to analyze sample files generated on ABI PRISM <sup>®</sup> 3100 Series and Applied Biosystems 3130 Series Genetic Analyzers.	Same as 310 Data Analysis table setting (see entry above), except the Matrix column is not displayed.	Same as 310 Data Analysis table setting (see entry above).	
CODIS Export	Used to enter data in the appropriate columns of the Samples table for exporting a CODIS- supported CMF file.	Displays the columns that may be used to populate CODIS-compatible fields when the table is exported in CMF file format.	Same as 310 Data Analysis table setting (see entry above). <b>Note:</b> No action is required on the Genotypes tab when exporting a CMF file.	
Import Reference Profiles	Used to import lab reference and custom control profiles into the GeneMapper <sup>®</sup> <i>ID-X</i> Software database.	Displays the Profile ID column, which is required to name and enter profiles into the GeneMapper <sup>®</sup> <i>ID-X</i> Software database.	Same as 310 Data Analysis table setting (see entry above). <b>Note:</b> No action is required on the Genotypes tab when importing reference profiles.	
View CGQ Overrides	Used to quickly identify samples that have been manually accepted (this includes samples with and without allele edits).	Displays only samples with CGQ override flag and sample-level quality values.	Displays marker and allele columns, edit comments and marker- level quality values.	
View Edited Samples	Used to quickly identify samples that have one or more allele edit.	Displays only samples with allele edits and sample-level quality values.	Displays only markers with edits, edit comments and marker- level quality values.	



#### Table 2 Default table settings (continued)

Та	ble Setting	Column Settings			
Name	Purpose	Samples Table	Genotypes Table		
Traditional Allele Table	Used to export data into an allele table using Combined Table Export.	Displays only the sample information required for an allele table export format.	Displays only the marker and allele information required for an allele table export format.		
	<b>Note:</b> The allele table export resembles that created in A Genotyper <sup>®</sup> Software. The compatible with spreadshe such as Microsoft <sup>®</sup> Excel <sup>®</sup> .	BI PRISM <sup>®</sup> 1 Sample Name 2 Sample 01 3 Sample 02	2         3         4           D851179         D21511         D75820           14,16         29,31         9           14,16         29,31         9           14,16         29,31         9           15,16         31.2         11,12           13,14         30         8,10		
View Unedited Samples	Used to quickly identify samples that have not been manually manipulated (edited or overridden).	Displays only samples without label edits, or GQ or CGQ override flags and sample-level quality values.	Displays only markers without allele edits and marker-level quality values.		
Yfiler Haplotype DB Export	Used to export the appropriate columns using Combined Table Export for upload into the Yfiler <sup>®</sup> Haplotype Database.	Displays only the sample information required for export in Yfiler <sup>®</sup> Haplotype Database format.	Displays only the sample and marker information required for export in Yfiler <sup>®</sup> Haplotype Database format.		
VALID_GMIDX_ TableSetting-1.0	Used to export the appropriate columns for importing tabular data into VALID <sup>™</sup> Software.	Displays only the sample information and run information required for a VALID software- compatible format. Displays only the marker information required for a VALID software-compatible format.			

**Note:** The columns in the Genotypes table displayed at the bottom of the Samples plot are determined by the table setting selected in the Project window.

#### Creating a New Table Setting

With custom table settings, you can adjust your view of the Samples and Genotypes tables to show or hide specific columns, and apply filters to display only specific samples.

**1.** Open the GeneMapper ID-X Manager if not already open (see page 30).

- **2.** In the GeneMapper ID-X Manager, select the **Table Settings** tab, then click **New** to open the Table Setting Editor.
- **3.** Enter the following information in the General tab:

💕 Table Setting Editor				
General Samples	Genotypes			
Table Settings De	scription:			
Name:	Custom Table Setting_your initials			
Security Group:	Practice Security Group 🛛 👻			
Description:				

**4.** Select the **Samples** tab, then expand the window (click-drag the window border) to display the Column, Filtering and Content columns in the Column Settings list:

Ceneral Samples Genotypes Samples Table Settings:	
Column Settings:         1       Status       Show All Records         2       Sample File       Show All Records         3       Sample Name       Show All Records         4       Sample ID       Show All Records         5       Comments       Show All Records         6       Sample Type       Show All Records         7       Specimen Category       Show All Records         9       Panel       Show All Records         10       Size Standard       Show All Records         11       Custom Control       Show All Records         12       Matrix       Show All Records         12       Matrix       Show All Hide All    OK Cancel Help	nding ding nding ding ding

- **5.** Click **Hide All** to deselect all of the default table column selections in the Column Settings list.
- 6. Select to Show (☑) the individual Samples table columns you wish to display using this custom table setting.



**7.** Click the **Filtering** field for the Sample Type row, then select **Sample** from the drop-down list.

	Show	Column	Filtering	Content
5		Comments	Show All Records	
6	<b>V</b>	Sample Type	Show All Rec 🗸	Show All Records
7		Specimen Category		N/A
8		Analysis Method	Sample Positive Control	
9		Panel	Allelic Ladder Primer Focus	
10		Size Standard	Primer Focus Negative Control	

**Note:** When you view the Samples table using this table setting, the Samples table displays only sample files of sample type Sample, and not Allelic Ladder or Control sample types. You can then export this filtered view of the Samples table (as described in Chapter 7).

- **8.** Select additional filters to apply to the remaining columns in the Samples table, if desired.
- 9. In the Sort by drop-down list, select Run Date & Time.

**Note:** Each sample file records the run date and time for that individual sample. When you view the Samples table using this table setting, the sample files sort by order of injection rather than the default sort option of by Sample File.

**10.** Select the **Genotypes** tab, then follow the same procedures outlined in steps 5 through 8 above to select the columns to display and filter in the Genotypes table.

**Note:** The settings you specify in the Genotypes tab determine the columns displayed in the Genotypes table in the Project window, and the Genotypes table in the Samples plot.



**11.** Verify the default sort options for the Genotypes table.

Sort by Sample File	<ul> <li>Ascending</li> <li>Descending</li> </ul>
Then by Dye	<ul> <li>Ascending</li> <li>Descending</li> </ul>
Then by Marker	<ul> <li>Ascending</li> <li>Descending</li> </ul>

**Note:** The GeneMapper<sup>®</sup> *ID-X* Software executes the selected sort options in succession. By default, markers are sorted within a sample by dye color (B, G, Y, R), then by size.

**12.** Click **OK** to apply your changes and close the Table Setting Editor. The GeneMapper ID-X Manager remains open.

## **Review Default Plot Settings**

Overview	Plot settings determine the number of panes, headers, labels and tables displayed in the Samples and Genotypes plot windows. The default plot settings shown in this section include a specific set of display elements required to perform different analysis workflow tasks. These elements are designed for efficient data review.
Reviewing Plot Settings	<ol> <li>Open the GeneMapper ID-X Manager if not already open (see page 30).</li> </ol>
	2. In the GeneMapper ID-X Manager, select the Plot Settings tab.
	<b>3.</b> Click on a table row to select the plot setting you wish to review, then click <b>Open</b> .
	<b>4.</b> Review the default plot settings provided using Table 3 on page 41.



**5.** Click **OK** to close the Plot Settings Editor after reviewing all plot settings. The GeneMapper ID-X Manager will remain open.

**Defaults Provided** The following plot settings are installed with the GeneMapper<sup>®</sup> *ID-X* Software.

Plot Setting				
Name	Purpose	Display Settings		
Check LIZ Size Standard	Used to display the GeneScan <sup>™</sup> LIZ <sup>®</sup> size standard in the same format as the Check GS500 Macro in the ABI PRISM <sup>®</sup> Genotyper <sup>®</sup> Software templates.	Displays the GeneScan <sup>™</sup> LIZ <sup>®</sup> size standard fragments with labels per sample in separate electropherogram panes.		
Check ROX Size Standard	Used to display the GeneScan <sup>™</sup> ROX <sup>™</sup> size standard in the same format as the Check GS500 Macro in the ABI PRISM <sup>®</sup> Genotyper <sup>®</sup> Software templates.	Displays the GeneScan <sup>™</sup> ROX <sup>™</sup> size standard fragments with labels per sample in separate electropherogram panes.		
Data Interpretation	Used during manual review of sample data, to enable quick interpretation of anomalies and marker-level quality values.	Displays the electropherogram plots for the selected sample(s), the Genotypes table, and the Quality Value Details (QVD) pane.		
Overlay LIZ Dye	Used to perform sizing precision checks with the GeneScan <sup>™</sup> LIZ <sup>®</sup> size standards.	Overlays all selected size standard fragments in one electropherogram pane, and displays the Sizing table.		
Overlay ROX Dye	Used to perform sizing precision checks with the GeneScan <sup>™</sup> ROX <sup>™</sup> size standards.	Overlays all selected size standard fragments within a project in one electropherogram pane, and displays the Sizing table.		
Sizing Data	Used to display data in a format similar to the ABI PRISM <sup>®</sup> GeneScan <sup>™</sup> Software plots.	Displays all dyes per sample in one electropherogram pane, and the Sizing table.		
Traditional Genotype Plot	Used to display data in a format similar to the ABI PRISM <sup>®</sup> Genotyper <sup>®</sup> Software plots.	Displays each dye for a sample in a separate electropherogram pane.		
View Label Edits	Used to display allele edits for the selected sample(s) in a table below the electropherogram for electronic data review.	Displays the electropherogram plots for the selected sample(s), and the Label Edit Viewer table.		



#### Creating a New Plot Setting

Custom plot settings save frequently used combinations of display elements to minimize the time spent adjusting display settings during manual review.

- 1. Open the GeneMapper ID-X Manager if not already open (see page 30).
- **2.** In the GeneMapper ID-X Manager, select the **Plot Settings** tab, then click **New**.
- **3.** Enter the following information in the General tab of the Plot Setting Editor:

Plot Settings Editor						
eneral	Sample H	leader	Genotype Header	Sizing Table	Labels	Display Settings
When opening the Plot Window:						
Name: Custom Plot Setting_your initials						
Securit	y Group:	Practic	e Security Group	~		
Des	cription:					
e	eneral /hen op Securit	eneral Sample H	eneral Sample Header /hen opening the Plot W Name: Custo Security Group: Practic	eneral Sample Header Genotype Header /hen opening the Plot Window: Name: Custom Plot Setting_your Security Group: Practice Security Group	eneral Sample Header Genotype Header Sizing Table then opening the Plot Window: Name: Custom Plot Setting_your initials Security Group: Practice Security Group	eneral Sample Header Genotype Header Sizing Table Labels then opening the Plot Window: Name: Custom Plot Setting_your initials Security Group: Practice Security Group

- **4.** Select the **Sample Header** tab, then select the columns to display in the Samples plot header.
- **5.** Select the **Genotype Header** tab, then select the columns to display in the Genotypes plot header.

**Note:** The columns in the Genotypes table displayed at the bottom of the Samples plot are determined by the table settings selected in the Project window.

6. Select the Sizing Table tab, then select the columns to display and the font and font size to use for the Sizing table displayed at the bottom of the Samples plot.

**7.** Select the **Labels** tab, then select the label information to display for the detected peaks in the Samples and Genotypes plots:

lot Sett	ings Editor							
General	Sample Header	Genotype Header Sizir	ng Table Labels Disp	lay Settings				
⊢Show I		s and Genotypes Plot:						
Label								
		Assigned Allele	Custom Alle	ele	Allelic Ladder	,	Artifact	$\overline{}$
	Label 1:	Allele Call	✓ Allele Call	~	Allele Call	*	Artifact Label	~
	Label 2:	NONE	NONE	~	NONE	~	NONE	~
	Label 3:	NONE	NONE	*	NONE	~	NONE	~
	Label 4:	NONE	NONE	~	NONE	~	NONE	~
Font								
Font	t: Times New R	oman 🗸						
Size	e: 10 💙							
	opening the Plot \	Nindow:					Specify mult	
□ st	how PQV trigger p	eak (LPH,MPH,BD,OS)				1	abels per ca	ategor
Show data type prefixes								
Display virtual allele label in black								
📃 SI	Show type of edit							
Label	Label Color: Dye Color-Border							

This tab allows you to choose a different set of labels for different types of samples and different types of peak labels. For instance, you may choose to only display the allele call for all allelic ladder samples, but you want to display both the allele call and height on all other sample types and have them both be displayed in the same plot window. Each peak can have up to four labels.



#### 8. Select the **Display Settings** tab, then specify settings:

General	Sample Header Genotype Header Sizing	J Table Labels Display Settings
⊤When oj	oening the Plot Window:	for this plot
	<ul> <li>O Use these display settings:</li> <li>For both Sample and Genotype plots:</li> <li>Panes: 4</li> <li>V</li> <li>Labels</li> <li>V</li> <li>Labels</li> <li>V</li> <li>Labels</li> <li>V</li> <li>V</li> <li>Vertical Labels</li> <li>V</li> <li>Select Dyes</li> <li>For Sample plot only:</li> <li>Select Dyes</li> <li>Far Range</li> <li>Verlow</li> <li>Red</li> <li>Orange</li> <li>Size St</li> <li>Size St</li> </ul>	Select a checkbox or radial button to enable the associated toolbar button
	Marker Margin: 5 bp	Set the x-axis zoom range
	* Will be overridden if Retain X-axis Zo	om Range is enabled on Plots ->Zoom menu

Click **Help** at the bottom of the tab and navigate to the *Display Settings Tab* help topic to learn more about the parameters presented in this tab.

**9.** Click **OK** to apply your changes and close the Plot Setting Editor. The GeneMapper ID-X Manager remains open.

### **Review Default Size Standards**

Overview	base stan Gen fron	ize standard definition file provides a list of fragment sizes in e pairs and the dye color associated with a particular size dard. During peak detection and size-calling, the meMapper <sup>®</sup> <i>ID-X</i> Software matches an observed fragment peak in the size standard run with the sample with a corresponding size he definition file.
Reviewing Size Standard Settings	1.	Open the GeneMapper ID-X Manager if not already open (see page 30).
	2.	In the GeneMapper ID-X Manager, select the <b>Size Standards</b> tab.
	3.	Click a table row to select the size standard setting to review, then click <b>Open</b> .
	4.	Review the default size standard settings provided using Table 4 on page 46.
	5.	When you are finished reviewing all size standard settings, click <b>Done</b> to close the GeneMapper ID-X Manager.
		<b>Note:</b> You can also create a new size standard definition using the GeneMapper ID-X Manager.

**Defaults Provided** The following default size standard definition files are provided with the GeneMapper<sup>®</sup> *ID-X* Software for analysis of AmpF*l*STR<sup>®</sup> kit data:

#### Table 4Default size standards

Size Standard	Description	When to Use
CE_G5_HID_GS500	Includes fragments present in the GeneScan <sup>™</sup> 500 LIZ <sup>®</sup> size standard (75 to 450-bp), excluding the 250-bp fragment.	Use with data generated on ABI PRISM <sup>®</sup> 310 and 3100 Series Genetic Analyzers, and Applied Biosystems 3130 Series Genetic Analyzers, and run with the GS500 LIZ <sup>®</sup> Size Standard.



#### Table 4 Default size standards (continued)

Size Standard	Description	When to Use
CE_F_HID_GS500 (75-400)	Includes fragments present in the GeneScan <sup>™</sup> 500 ROX <sup>™</sup> size standard (75 to 400-bp), excluding the 250-bp fragment.	Use with data generated on ABI PRISM <sup>®</sup> 310 and 3100 Series Genetic Analyzers, and Applied Biosystems 3130 Series Genetic Analyzers, and run with the GS500 ROX <sup>™</sup> Size Standard and all AmpFℓSTR <sup>®</sup> 4-dye kits (except the SGM Plus <sup>®</sup> kit).
CE_F_HID_GS500 (75-450)	Includes fragments present in the GeneScan <sup>™</sup> 500 ROX <sup>™</sup> size standard (75 to 450-bp), excluding the 250-bp fragment.	Use with data generated on ABI PRISM <sup>®</sup> 310 and 3100 Series Genetic Analyzers, and Applied Biosystems 3130 Series Genetic Analyzers, and run with the GS500 ROX <sup>™</sup> Size Standard and the AmpF <i>l</i> STR <sup>®</sup> SGM Plus <sup>®</sup> kit.
GS600_LIZ	Includes fragments present in the GeneScan <sup>™</sup> 600 LIZ <sup>®</sup> size standard (80 to 460-bp).	Use with data generated on ABI PRISM <sup>®</sup> 310 and 3100 Series Genetic Analyzers, and Applied Biosystems 3130 Series Genetic Analyzers, and run with the GS600 LIZ <sup>®</sup> Size Standard.

# Step 5: Import Lab Reference and Custom Control Profiles

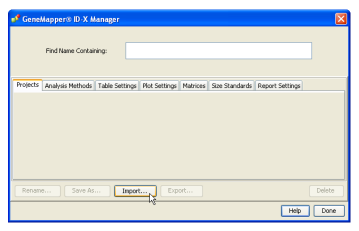
**Overview** In this section, you will import a project that contains analyzed lab reference samples, custom controls, and QC samples that are used to illustrate the quality control features of the GeneMapper<sup>®</sup> *ID-X* Software (see Chapter 5). This project has been edited to remove any off-ladder (OL) labels.

**IMPORTANT!** Before adding your own lab reference and custom control samples using the procedure described below, review the samples manually and edit allele labels as needed to ensure that the profile is accurate. Profiles that include OL labels are not imported into the Profile Manager. Profiles that include numeric allele labels on peaks that are not true DNA peaks will affect concordance results.

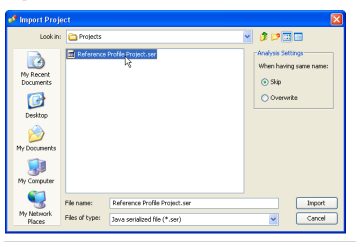


## Importing the Reference Project

- **1.** Open the GeneMapper ID-X Manager if not already open (see page 30).
- **2.** In the GeneMapper ID-X Manager, select the **Projects** tab, then click **Import**.



**3.** Navigate to and select **Reference Profile Project.ser**, then click **Import**.



**Note:** The reference project file path shown above is for Full install configurations only. For the reference project file path for Client install configurations, see "Using the Guide with the Example Data Provided" on page 18.

**4.** Make sure the Practice Security Group is selected, then click **OK**.

Choose Security Group		
Project <referer< th=""><th>nce Profile Project&gt;</th><th></th></referer<>	nce Profile Project>	
Security Group: Practice Security Group		
	ОК Неір	

**5.** Click **Done** to close the GeneMapper ID-X Manager.

Adding Profiles to the Software Database

- 1. In the Project window, click 🙆 (Open Project).
- 2. Select Reference Profile Project, then click OK.
- **3.** From the Table Setting drop-down list, select **Import Reference Profiles**. The Sample table view changes to display only those columns required to add reference profiles to the GeneMapper<sup>®</sup> *ID-X* Software database.

**Note:** All genotypes and edits are saved with the imported project. This project does not require re-analysis.

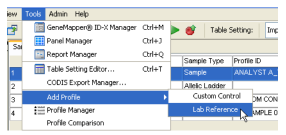
**4.** In the Profile ID column of the Samples tab, click each cell, then enter the Profile ID names as shown below.

Sam	Samples Analysis Summary Genotypes					
	Status	Sample File	Sample Name	Sample Type	Profile ID	
1	<u>N</u>	Analyst A.fsa	Analyst A	Sample	ANALYST A_YOUR INITIALS	
2	<u>N</u>	ID_AllelicLadder.fsa	Ladder	Allelic Ladder		
3	<u>N</u>	ID_CustomControl.fsa	Custom Control	Sample	CUSTOM CONTROL_YOUR INITIALS	
4	<u>N</u>	QC_Sample_01.fsa	QC Sample 01	Sample	QC SAMPLE 01_YOUR INITIALS	

**Note:** Profiles are stored in the GeneMapper<sup>®</sup> *ID-X* Software database under Profile ID, not Sample Name.



 5. Select the Analyst A.fsa row, then select Tools > Add Profile > Lab Reference.



6. Click Close in the Add Profile Results dialog box to save the assigned lab reference profile to the GeneMapper<sup>®</sup> *ID-X* Software database.

1	Add Profile Results	×
The	e following profiles were added successfully.	
	Profile ID	
	ANALYST A_YOUR INITIALS	
	Close Help	

- 7. Shift-click to select the ID\_CustomControl.fsa and QC\_Sample\_01.fsa rows, then select Tools ➤ Add Profile ➤ Custom Control.
- **8.** Click **Close** in the Add Profile Results dialog box to save the assigned custom control profiles to the GeneMapper<sup>®</sup> *ID-X* Software database.



#### Viewing Profiles in the Profile Manager

- 1. In the Project window, select **Tools** > **Profile Manager**.
- View the list of profiles in the Profile Manager window. Click

   to expand at least one Profile ID to view the genotypes
   stored in the GeneMapper<sup>®</sup> ID-X Software database.

Profile Manager			×
· · · · · · · · · · · · · · · · · · ·			
Expand All Collapse All			
Profile ID / Marker	Туре	Alleles	
CUSTOM CONTROL_YOUR INITIALS	Custom Control		
D851179		13, 14	
D21511		28, 29	
• D75820		8, 10	
CSF1PO		10, 11	
🗣 D3S1358		18	
🔶 TH01		6, 9	
- • D135317		12	
- • D165539		12	
🗣 D251338		18	
🏶 D195433		14	
🗢 VWA		15, 16	
🌩 TPOX		8, 11	
- 🗢 D18551		13, 14	
🌩 AMEL		Х, Ү	
🕈 D55818		11, 12	
FGA		20, 24	
😨 🧰 QC SAMPLE 01_YOUR INITIALS	Custom Control		
🗟 🧰 ANALYST A_YOUR INITIALS	Lab Reference		
Delete Export	Close	Help	

**3.** Click **Close** to close the Profile Manager window and return to the Project window.

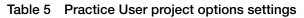


2

## Step 6: Set the Project Options

Overview	Project Options are user-specific project preferences that allow you to specify default security and analysis options for your user account.
In This Section	In this section, you will set the software to:
	• Automatically assign a security group (available selections determined by user account) when you create a project.
	• Automatically assign analysis settings when you add samples to the project.
	• Display the Analysis Requirements Summary (ARS) to identify any conditions that could prevent analysis.
	• Automatically disregard low-quality allelic ladders and proceed with the analysis of run folders containing one or more passing allelic ladders.
	• Display the Analysis Summary, which provides a snapshot of the analysis status of all samples in the project and the quality status of allelic ladders, controls, and samples.
	<b>Note:</b> These options are suggested for an optimized, efficient data review workflow. However, you can modify project options as needed to meet your laboratory workflow requirements.
Setting Project Options	Project options are associated with the user account currently logged in to the GeneMapper <sup>®</sup> $ID-X$ Software (in this example, the Practice User).
	<b>Note:</b> Set project options when you obtain your personal or lab- specific user account. For more information on the GeneMapper <sup>®</sup> <i>ID-X</i> Software security system, see the <i>GeneMapper</i> <sup>®</sup> <i>ID-X</i> Software <i>Version 1.0 Administrator's Guide</i> .

- **1.** Select File > Project Options.
- Complete the Options tabs for the Practice User as described in Table 5 on page 53, using the *GeneMapper<sup>®</sup> ID-X Software Help* as a guide.



Tab	Settings
General	Verify the settings as shown below:
	Options X
	General Add Samples Analysis  Project  Open Blank Project
	Open Previous Project
	Plots Update plots when changing project tab
	Data Access Control         Security group when creating objects         Practice Security Group



2

Tab	Settings	
Add Samples	Enter the settings as shown below: General Add Samples Analysis When adding new samples, automatically Set Analysis Method to: © Read from the Sample. Set Size Standard to: © Et G5_HID_G5500 • or all samples. Read from the Sample. Set Panel to: © Read from the Sample. Set Sample from the Sample. Set Sample from the Sample. • Click in the text field to open the Select a Panel dialog • Click • to view the available Amalysis Read from Data collection 'Info' field. • Click • to view the available Amalysis. • Click • to view the available Amalysis. • Blac_vix • Blac_vix • Profiler_Pus_vix • Coller_vix • Seffer available • Blac_vix • Profiler_Pus_vix • Coller_vix • C	

#### Table 5 Practice User project options settings (continued)

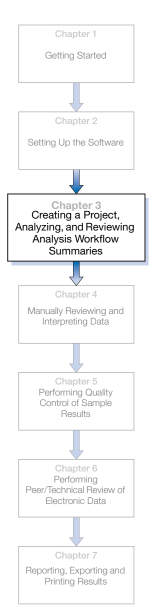


#### Table 5 Practice User project options settings (continued)

Tab	Settings
Analysis	Click <b>Help</b> at the bottom of the tab to learn more about the options presented in the Analysis Summary area of this tab, then enter the settings as shown below:
	Options           Options           General         Add Samples           Analysis
	Analysis Summary If one or more analysis requirements are not met
	Stop analysis and display Analysis Requirements Summary     O Continue analysis
	If one or more allelic ladders do not meet sizing and/or genotyping requirements Stop analysis and display Allelic Ladder Analysis Summary
	Continue analysis of run folders with at least one passing allelic ladder     Continue analysis of all run folders
	After Analysis           O View Analysis Summary
	View Sample Table  Automatic Analysis
	Automatically bring low quality samples to the top     Quality Metrics Display
	Symbols     Numbers
	If only one labelled allele in a genotype, then duplicate the label
	Duplicate homozygous alleles
	OK Cancel Help

**3.** Click **O**K after completing all tabs to close the Options dialog and return to the Project window.

# Creating a Project, Analyzing, and Reviewing Analysis Workflow Summaries



This	chapter	covers:	
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-

Overview	56
Step 1: Create a Project and Add Samples	56
Step 2: View Sample Information and Raw Data	59
Step 3: Select Analysis Settings and Start Analysis	63
Step 4: Review the ARS and Correct the Requirements	64
Step 5: Analyze the Data	66
Step 6: Review the Analysis Summary	68

## **Overview**

During analysis, the GeneMapper<sup>®</sup> *ID-X* Software performs the following tasks, based on the analysis settings and project options set up in Chapter 3:

- Analysis requirements check
- Data analysis (peak detection and sizing, allele-calling)
- Allelic ladder and sample quality assessment
- Analysis workflow summary generation

This chapter will demonstrate these tasks using the example data set provided with the software.

In This Chapter In this chapter, you will learn how to:

- Create a project and add samples
- View sample details
- Select analysis settings
- Analyze the data
- Review the analysis workflow summaries (Analysis Requirements Summary, Analysis Summary)

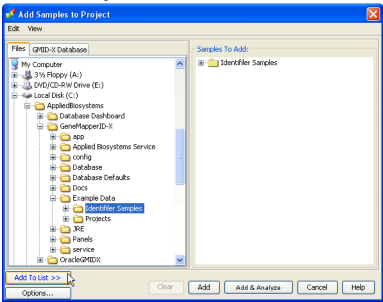
## Step 1: Create a Project and Add Samples

Creating a	If a project is already open in the Project window, click 🖳 (New
New Project	Project).

**Note:** If you have a previous project open you may be prompted to save changes. Click **Yes**.

## Adding Samples from Sample Files

- 1. In the new Project window, click 😼 (Add Samples to Project).
- 2. In the Files tab of the Add Samples to Project dialog box, navigate to the folder containing the example data, select the **Identifiler Samples** folder, then click **Add to List**.



**Note:** The example data file path shown above is for Full install configurations only. For the example data file path for Client install configurations, see "Using the Guide with the Example Data Provided" on page 18.

- **3.** Click **Add**. When you add samples from sample (.fsa) files to a project:
  - You specify the location of the .fsa files on the hard drive or a network drive.
  - The sample files remain in their original location on the drive, and are not stored in the GeneMapper<sup>®</sup> *ID-X* Software project or database.

- The GeneMapper<sup>®</sup> *ID-X* Software reads the information it needs from the .fsa files. No information is written back to the original sample files.
- The added samples are displayed in the Samples table in the content pane of the Project window. The run folder from which you added the samples is displayed in the navigation pane.

Sample table lists all files in selected run folder (only one run folder in this example). Analysis settings are loaded by default as specified in Project Options. Table Setting selected determines the columns displayed in the Samples table

😂 🖨 📙 😼 🖻			E/ E	Ш 🕅 🎞 🗐	🌔 💣 (Table	Setting: Impo	rt Reference Pr	rofiles 🛛 🔽 🛅	P 🖨 🔍	A
🖃 - 🚠 Project		Sam	ples Ar	alysis Summary Genot	ypes					
🛓 🛅 Identifiler Samp	bles		Status	Sample File	Sample Name	Sample Type	Profile ID			
		1	<b>N</b>	ID_AllelicLadder2.fsa	Ladder	Sample				
		2	<b>N</b>	ID_AllelicLadder.fsa	Ladder	Sample				
		3	1	ID_CustomControl.fsa	ID CustomControl	Sample				
		4	<b>1</b>	ID_NegControl.fsa	NegControl	Sample				
		5	1	ID_PosControl.fsa	PosControl	Sample				
		6	<b>N</b>	ID_Sample_01.fsa	Sample 01	Sample				
		7	1	ID_Sample_02.fsa	Sample 02	Sample				
		8	<b>N</b>	ID_Sample_03.fsa	Sample 03	Sample				
		9	1	ID_Sample_04.fsa	Sample 04	Sample				
		10	<b>N</b>	ID_Sample_05.fsa	Sample 05	Sample				
		11	<b>N</b>	ID_Sample_06.fsa	Sample 06	Sample				
		12	6	ID_Sample_07.fsa	Sample 07	Sample				
		13(	1	ID_Sample_08.fsa	Sample 08	Sample				
Progress Status								0%	%	[St

added samples

4. In the Project window, select **31XX Data Analysis** from the Table Setting drop-down list. Only the columns needed for analysis of ABI PRISM<sup>®</sup> 3100 Series and Applied Biosystems 3130 Series Genetic Analyzer data are displayed in the Samples table.

7		🔳 🔤 🛄		] 📄 💕 🛛 Table Sett	ing: 31XX Da	ata Analysis	<b>~</b>	P	) 🖨 🖪		A ()			
Sa	mples 🛛	analysis Summary	Genotypes											
	Status	Sample Name	Sample Type	Analysis Method	Panel	Size Standard	Custom Control	ARNM	SOS	SQ	SSPK	MIX	OMR	CGQ
1	۹Ç	Ladder	Sample	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None							
2	1	Ladder	Sample	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None							
3	<b>1</b>	ID CustomControl	Sample	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None		İ		İ			
4	<b>1</b>	NegControl	Sample	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None		<u> </u>		<u> </u>			
5	<b>1</b>	PosControl	Sample	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None		<u> </u>		<u> </u>			
6	<b>1</b>	Sample 01	Sample	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None		i –		<u> </u>			
7	<b>1</b>	Sample 02	Sample	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None		i –		<u> </u>			
8	<b>1</b>	Sample 03	Sample	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None		i –		<u> </u>			
9	<b>1</b>	Sample 04	Sample	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None		i –		<u> </u>			
10	8	Sample 05	Sample	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None		<u> </u>		<u> </u>			
11	<u> </u>	Sample 06	Sample	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None				İ			
12	<u> </u>	Sample 07	Sample	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None				İ			
13	Ŵ	Sample 08	Sample	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None				<u> </u>			

## Step 2: View Sample Information and Raw Data

You have access to the following sample information from the Project window:

- Sample file, analysis and run parameters
- Raw data
- Electrophoresis, power and temperature (EPT) data

### Viewing Sample Information

Click 
 to expand the Identifiler Samples folder in the navigation pane, then select the ID\_AllelicLadder.fsa sample. The Info tab for the selected sample is displayed in the content pane:



**2.** Click the vertical scroll bar at the right of the Info tab window to review the following *sample-specific* information presented in this tab:

Info Type	Info Listed for Selected Sample
Sample Information	<ul> <li>Sample file name and sample name</li> <li>Sample origin path and file source</li> <li>Status message indicating any changes made to the sample in the Samples table</li> </ul>
Error Message	Errors encountered during analysis (if any)
Last Used Analysis Settings	<ul> <li>Last settings used for analysis</li> <li>Note: This area will be blank if the sample Status is         <ul> <li>(Unanalyzed). After analysis, the settings last used to analyze the sample file are displayed in this tab regardless of the analysis settings selected in the Samples table.</li> </ul> </li> </ul>

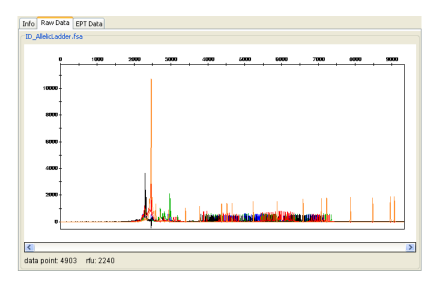


Info Type	Info Listed for Selected Sample
Run Information	<ul> <li>Instrument user name</li> <li>Instrument name and Data Collection version</li> <li>Run date (in yyyy:dd:mm) and time (in hr:min:sec)</li> <li>Run duration</li> <li>Total data points</li> </ul>
	<b>Note:</b> All sample files created during one injection on a multi-capillary instrument (a set of 4 or 16 capillaries) will have the same run date, run time, and injection time.
Data Collection Settings	<ul> <li>Run voltage and injection voltage (in volts)</li> <li>Injection duration (in milliseconds)</li> <li>Laser power (in mW) and temperature (in °C)</li> <li>Run module and run protocol name</li> <li>Dye set name</li> <li>Polymer lot number and expiration date</li> <li>Results Group name</li> <li>Note: Results Groups apply to ABI PRISM<sup>®</sup> 3100 Series Data Collection Software v 2.0 and Applied Biosystems 3130 Series Data Collection Software v 3.0 only.</li> </ul>
Capillary Information	<ul><li>Length and number of capillaries</li><li>Capillary number used for injection</li></ul>

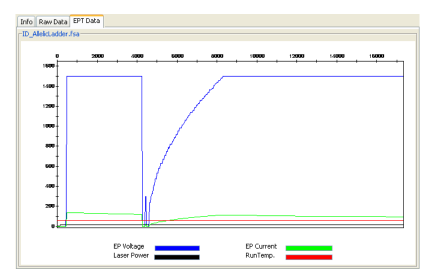
### **Viewing Raw Data**

You can use the raw data view of a sample to help evaluate any anomalies, the causes of poor size-calling, and to determine the start and stop points for analysis.

1. Select the **Raw Data** tab in the content pane. The raw data plot for the ID\_AllelicLadder.fsa sample is displayed.



**2.** Select the **EPT Data** tab in the content pane. The EPT plot is displayed.





**3.** Select the **Project** node in the navigation pane to return to the Samples table view.

	Sample	s Anal	notypes	
Identifiler Samples		Status	Sample Name	Sample Type
ID_AllelicLadder2.fsa	1	Тў	Ladder	Sample
ID_CustomControl.fsa	2	<b>N</b>	Ladder	Sample
ID_NegControl.fsa ID_PosControl.fsa	3	<b>N</b>	ID CustomControl	Sample
ID_FosControl.rsa	4	<b>1</b>	NegControl	Sample

## **Step 3: Select Analysis Settings and Start Analysis**

Analysis settings include the Analysis Method, Size Standard, Panel, and Sample Type selections needed to perform analysis. Based on the Project Options you set in Chapter 2, the samples you added to the Samples table automatically have analysis settings specified.

#### **Selecting Analysis Settings** Note: In this section, you will intentionally alter some of the analysis settings in the Samples tab to trigger the display of the ARS when you start the analysis.

**1.** In the Project window, make sure **31XX Data Analysis** is selected from the Table Setting drop-down list.

Sa	mples 🛛	Analysis Summary	Genotypes				
	Status	Sample Name	Sample Type	Analysis Method	Panel	Size Standard	Custom Control
1	1	Ladder	Allelic Ladder	None	ldentifiler_v1X	CE_G5_HID_GS500	None
2	1	Ladder	Allelic Ladder	Getting Started_your initials	ldentifiler_v1X	None	None
3	1	ID CustomControl	Positive Control	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	CUSTOM CONTROL_YOUR INITIALS
4	1	NegControl	Negative Control	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None
5	1	PosControl	Positive Control	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None
6	1	Sample 01	Sample	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None
7	1	Sample 02	Sample	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None
8	1	Sample 03	Sample	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None
9	1	Sample 04	Sample	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None
10	1	Sample 05	Sample	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None
11	1	Sample 06	Sample	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None
12	1	Sample 07	Sample	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None
13	1	Sample 08	Sample	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None

**2.** Enter the Samples tab analysis settings as shown below:

### Starting Analysis

Click **(**Analyze).

When analysis is started, the software identifies any conditions that may prevent analysis or cause unexpected results, sets a  $\land$  flag for Analysis Requirements Not Met (ARNM) PQV and displays the ARS if chosen in the Project Options. In this example, we have chosen to display the ARS.

The Analysis Requirements Summary dialog box opens because at least one sample in the project does not meet one or more analysis requirements.

🧳 Analysis Requirements Summa	ту 🔀										
2 samples do not meet on	e or more analysis requirements.										
Analysis Requirement Not Met	# Samples That Do Not Meet Requirement										
Analysis Method Not Selected	1										
Size Standard Not Selected	1										
What would you like to do next?	v more analysic requirements										
Continue analysis	<ul> <li>View sample(s) that do not meet one or more analysis requirements</li> <li>Continue analysis</li> </ul>										
ОК	Cancel Help										

# Step 4: Review the ARS and Correct the Requirements

### Reviewing the ARS

From the ARS, you may view the samples that do not meet the analysis requirements or you can continue with analysis. In this section, you will view the flagged samples in the example project.

1. Keep the default selection in the *What would you like to do next*? area of the ARS, then click **OK**.

The Samples table opens. If one or more analysis requirements are not met, the ARNM PQV is set to  $\triangle$ . Based on the analysis settings changes you made in step 2 on page 63, note that only the allelic ladder samples in the example project are listed with a  $\triangle$  ARNM flag in the Samples table.

		s tab lab s table is		<i>bold itali</i> table is fi	<b>cs</b> indicat Itered	es		
7		🛄 🔤 🔛		🗐 📄 💕 Table S	etting: 31X	X Data Analysis		1   ۶
Sar	mples (filt	tered) Analysi:	s Summary Ger	otypes				
	Status	Sample Name	Sample Type	Analysis Method	Panel	Size Standard	Custom Control	ARNM
	6	Ladder	Allelic Ladder	None	Identifiler v1X	CE G5 HID GS500	None	i 🔺 i
1	1	Landon		None		1		
1 2		Ladder	Allelic Ladder	Getting Started_your initials		None	None	

Check ARNM PQV

When you view the Samples table from the ARS, the Samples table is filtered. The current table setting is applied to determine which columns are displayed, but only the samples in the category selected are listed. In this example, samples that do not meet the analysis requirements are listed. The filtered condition of the table is indicated by the status of the Samples tab label (filtered) and the Table Setting label (*bold italics*).

**2.** Place the pointer over a ARNM flag to display a tooltip with analysis requirement information for each sample in the filtered Samples table.

1	5amples (filt	ered) Analysis	s Summary Ger	otypes							
	Status	Sample Name	Sample Type	Analysis Method	Panel	Size Standard	Custom Control	ARNM	SOS	SQ	SSPK
1	<b>N</b>	Ladder	Allelic Ladder	None Identifiler_v1X		CE_G5_HID_GS500 None		4		<b></b>	NA
2	- <b>N</b>	Ladder	Allelic Ladder	Getting Started_your initials	Identifiler_v1X	None	None	Analy:	sis Meth	nod Not S	5elected

Note that the analysis Status for the samples is still **§** (Unanalyzed).

Correcting the Analysis Requirements To correct the unmet requirements listed in the ARS, change the analysis settings in the Samples table back to their original values:

- **1.** Change the Analysis Method for the first Ladder sample to **Getting Started**.
- 2. Change the Size Standard for the second Ladder sample to CE\_G5\_HID\_GS500.

The updated Samples table analysis settings should be:

San	Samples (filtered) Analysis Summary Genotypes												
	Status Sample Name Sample			Analysis Method	Panel	Size Standard	Custom Control						
1	<b>1</b>	Ladder	Allelic Ladder	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None						
2	<b>1</b>	Ladder	Allelic Ladder	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None						

## Step 5: Analyze the Data

Analyzing the<br/>DataIn this section, you will analyze with all analysis requirements<br/>satisfied and display the Analysis Summary.

- 1. Click **>** (Analyze).
- 2. Complete the fields in the Save Project dialog box:
  - a. For Name, type Getting Started <your initials>. For example, Getting Started AB.
  - **b.** Verify that the Practice Security Group is selected from the drop-down list.

Save Project	×
Name: Getting Started AB	
Security Group: Practice Security Group	~
OK Cancel Help	

**c.** Click **OK** to save the project to the GeneMapper<sup>®</sup> *ID-X* Software database and start analysis.



### Viewing Analysis Progress

After saving the project to the database:

• The project name (entered in step 2a on page 66) appears in the title bar of the Project window. For example:

💕 GeneMapper® ID-X - \*Getting Started AB - Practice User Is Logged In Database

• The software then begins analysis.

Note the following after analysis begins:

• Allelic ladder sample types are analyzed before all other sample types in the project. The sample currently being analyzed is highlighted in green in the Samples table:

Sample	s Analys	is Summary	Genoty	pes											
	Status	Sample F	Sample N	Sample IE	Comment	Sample T	Specimer	Analysis	Panel	Size Star	Custom C	Matrix	Run Nam	Instrumer	Instrume
1		ID_AllelicI	Ladder		None	Allelic La	No Expor	Getting S	Identifiler	CE_G5_F	None	None	Identifiler	ABI3100	3100
2		ID_AllelicI	Ladder	ca9539cf	None	Allelic La	No Expor	Getting S	Identifiler	CE_G5_F	None	None	Identifiler	ABI3130	1405-04
3		ID_Custor	ID Custon	e1639f28	None	Positive Ç	No Expor	Getting S	Identifiler	CE_G5_F	CUSTOM	None	Identifiler	ABI3130	1405-04
4		ID_NegCc	NegContr		None	Negative	No Expor	Getting S	Identifiler	CE_G5_F	None	None	ldentifiler	ABI3130	1405-04
5	<b>N</b>	ID_PosCc	PosContr	ebddb178	None	Positive Ç	No Expor	Getting S	ldentifiler	CE_G5_F	None	None	Identifiler	ABI3130	3130-16
6	<b>N</b>	ID_Sampl	Sample 0	ebddb19(	None	Sample	No Expor	Getting S	Identifiler	CE_G5_F	None	None	Identifiler	ABI3130	3130-16
7	<b>%</b>	ID_Sampl	Sample 0	ebddb19	None	Sample	No Expor	Getting S	ldentifiler	CE_G5_F	None	None	Identifiler	ABI3130	3130-16

• Analysis progress is displayed at the bottom of the Project window:

Analyzing Samples ...

[ Stop

31%



**Chapter 3** Creating a Project, Analyzing, and Reviewing Analysis Workflow Summaries *Step 6: Review the Analysis Summary* 

## Step 6: Review the Analysis Summary

About the<br/>AnalysisBased on the Project Options you set in Chapter 2, the Analysis<br/>Summary tab is displayed when analysis is complete.Summary Tab

es Analysis Summary Genoty	pes				
lysis Summary					Summary Generation Date
Select run folder to display: Ide	entifiler Samples	<b>~</b>			
Sample	Status	Total # of Samples			
👣 Unanalyzed	ĺ	0			
Analyzed		13			
🧕 Analysis Setting Changed	0				
Run Folder	Total # of Analy				
Run Folder	Total # of Analy	zed Ladders			
Identifiler Samples	2		1 0	1	
ontrol Quality per project (b	ased on sample PQVs: SOS		<b>), CGQ)</b> resholds met	0	or more thresholds not met
	local # or Sampi				
Positive Control	<u> </u>		1		0
Custom Control	1		1		0
Negative Control	1		0		1
Total	3		2		1
ample Quality per project (l	ased on sample PQVs: SOS	, SSPK, MIX, OMR, S	Q, CGQ)		
	Total # of Sampl	es 🛛 🚺 All th	resholds met	🔵 One d	or more thresholds not met
Samples	8		4		4
			-	1	

For efficient data evaluation, the Analysis Summary tab:

- Provides a summary of the analysis status for all or a subset of samples in the project
- Displays an overview of allelic ladder, control, and sample quality
- Visually separates passing samples from samples that do not meet one or more quality thresholds



• Provides interactive links to specific categories of samples (passing/check/low quality, allelic ladder/control/sample)

**IMPORTANT!** Refer to your own laboratory protocol to determine the samples (allelic ladder, positive control, negative control, custom control, and unknown) that require manual review when analyzing data generated in your laboratory.

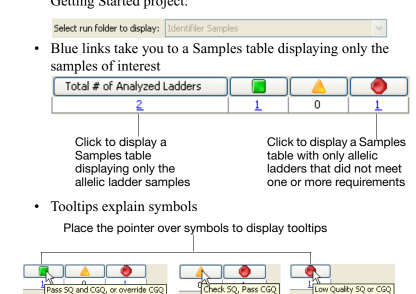
Briefly review the features and areas of the Analysis Summary tab as described below, then continue to Chapter 4, where you will review the functions of this tab in more detail.

Features of the Analysis Summary Tab

- Note the following features of the Analysis Summary tab:
  - You can display an Analysis Summary for the entire project or for an individual run folder in the project by selecting a run folder from the drop-down list at the top of the tab

**Note:** If the project contains only one run folder, this selection is dimmed.

In this example, Identifiler Samples is the only run folder in the Getting Started project:



### Areas of the Analysis Summary Tab

### Analysis Status

The Analysis status area at the top of the Analysis Summary tab indicates that all samples in the Getting Started project are analyzed.

Sample Status	Total # of Samples
Unanalyzed	0
Analyzed	13
🐚 Analysis Setting Changed	0

### Allelic Ladder Quality

The Allelic Ladder Quality area indicates that of the two allelic ladder samples analyzed in the Getting Started project, one met all of the allelic ladder quality requirements (
) and one did not (
).



You can use the allelic ladder quality results to determine the samples that require manual review. For example, if each run folder contains at least one passing allelic ladder and the positive and custom controls meet all quality value thresholds, visual inspection of the allelic ladders may not be required. Depending on validation, you can proceed directly to evaluation of the controls or samples in the project.

### **Control Quality**

The Control Quality area of the Analysis Summary indicates that of the three controls analyzed in the Getting Started project, the positive and custom controls met all quality thresholds ( ) and generated the expected profile, but the negative control did not.

Control Type	Total # of Samples	🔄 All thresholds met	🛛 🥏 One or more thresholds not met 📄
Positive Control	1	1	0
Custom Control	<u>1</u>	<u>1</u>	0
Negative Control	<u>1</u>	0	<u>1</u>
Total	<u>3</u>	<u>2</u>	<u>1</u>

When validated, this area of the Analysis Summary may eliminate the need for you to visually inspect the control samples if they fall under the green All Thresholds Met column. Controls in this column have met all sample-level and marker-level quality values, including



Control Concordance (CC). This means that the sample genotypes for positive and custom controls match the expected known profile without any anomalies. For negative controls, this means that there were no peaks detected above the peak amplitude threshold.

You can also use the control quality results to verify allelic ladder quality. For example, allelic ladders may meet all requirements, but if their migration rate differs from the sample migration rate, samples and controls may contain OL calls. However, controls that met all thresholds (and therefore do not contain OL calls), may indicate consistent migration rates for allelic ladders and samples.

### Sample Quality

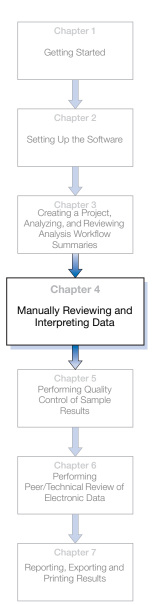
The Sample Quality area of the Analysis Summary indicates that of the eight samples analyzed, four met all quality thresholds ( ) and four did not ( ).

	Total # of Samples	🛛 📘 All thresholds met	One or more thresholds not met
Samples	<u>8</u>	<u>4</u>	<u>4</u>

In Chapter 4, you will use the blue links to visually inspect certain categories of samples.



## Manually Reviewing and Interpreting Data



This	chapter	covers.
1 1115	Chapter	

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4

## **Overview**

	The Analysis Summary discussed in Chapter 3 guides you directly to different types of samples while following a logical data interpretation workflow, regardless of whether the software is being used as an expert system or not. In a traditional manual review workflow, you would most likely visually inspect all unknown sample electropherograms. In a validated expert system workflow, you would only view those samples that did not meet one or more quality value thresholds.					
	<b>Note:</b> Refer to your own laboratory protocol to determine the controls and samples that require manual review.					
In This Chapter	In this chapter, you will review the example data analyzed in Chapter 3 and learn how to:					
	• Use the Analysis Summary tab and filtered Samples table to examine allelic ladder, control and sample quality					
	• Investigate sample-level PQVs, and marker-level PQVs using the QVD pane					
	• Review sample plots and edit peak labels					
	• Adjust plot displays to determine the source of artifacts					

**Note:** Some steps performed in this chapter are included only to demonstrate the use of certain features in the GeneMapper<sup>®</sup> ID-X Software and may not be a part of your routine analysis workflow.

## Step 1: Examine Allelic Ladder Quality

**Overview** The order of information displayed in the Analysis Summary is designed to direct you to evaluate the allelic ladder quality first. The designation is based on a set of allelic ladder requirements. If there is at least one passing allelic ladder per run folder and the positive and custom controls meet all quality value thresholds as displayed in the Analysis Summary, you may not require visual inspection of the

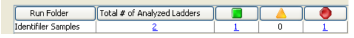


allelic ladders and may proceed directly to evaluation of the controls or samples in the project (depending on validation). However, for the purposes of this guide, you will walk through the process for manually reviewing allelic ladder samples.

### Viewing Allelic Ladder Quality Status

In Chapter 3, the Allelic Ladder Quality area of the Analysis Summary tab indicates that of the two allelic ladder samples analyzed, one has met all allelic ladder requirements () and one has not ().

### Allelic Ladder Quality per run folder (based on SQ and CGQ only)



**Note:** Low-quality () allelic ladders are not used to create bin offsets.

### Examining the Low-Quality Allelic Ladder

1. If the Getting Started project is not already open, click (Open Project) in the Project window.

**Note:** If you have a previous project open you may be prompted to save changes. Click **Yes**.

- In the Open Project window, select the Getting Started <your initials> project, then click OK. The Getting Started project opens in the Project window.
- **3.** In the Samples table, verify that **31XX Data Analysis** is selected from the Table Setting drop-down list.
- **4.** Verify that the Analysis Summary tab is selected in the content pane.

5. In the Allelic Ladder Quality area of the Analysis Summary tab, click the link for the 😑 allelic ladder sample.

Alle	Allelic Ladder Quality per run folder (based on SQ and CGQ only)									
	Run Folder	Total # of Analyzed Ladders								
	Identifiler Samples	2	1	0	<u> </u>					
					40					

The filtered Samples table displays the selected sample. Note that this low-quality allelic ladder sample has a OGQ PQV.

Sa	mples (fil	tered) Analy:	sis Summary 🛛 G	enotypes										
	Status	Sample Name	Sample Type	Analysis Method	Panel	Size Standard	Custom Control	ARNM	SOS	SQ	SSPK	MIX	OMR	CGQ
1		Ladder	Allelic Ladder	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None					NA	NA	۲
		Low Qualit												
										CGQ PQV				

- 6. Select the Ladder sample in the filtered Samples table, then click (Display Plots). The plot for the selected allelic ladder sample opens in the Samples plot.
- 7. Click (Maximize) in the top-right corner of the Samples plot to maximize the display.

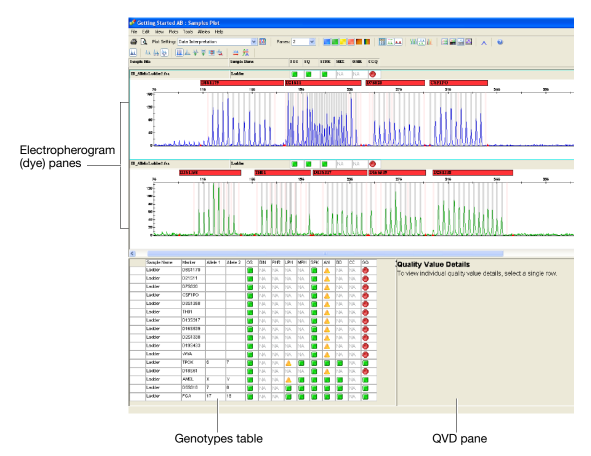


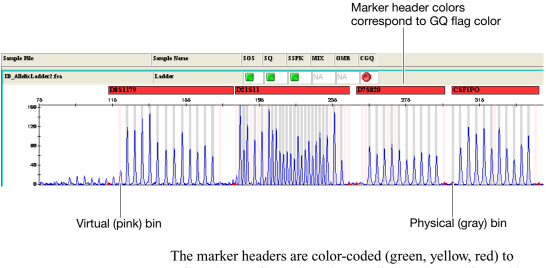
8. Select Data Interpretation from the Plot Setting drop-down list.



Based on the display options of this default plot setting (see Chapter 2), the Samples plot displays:

- Two electropherogram (dye) panes in a single view
- Each pane zoomed to the 75 450 base pair range
- Each of four dyes (blue, green, yellow, red) in a separate pane
- The Genotypes table and QVD pane displayed under the electropherograms



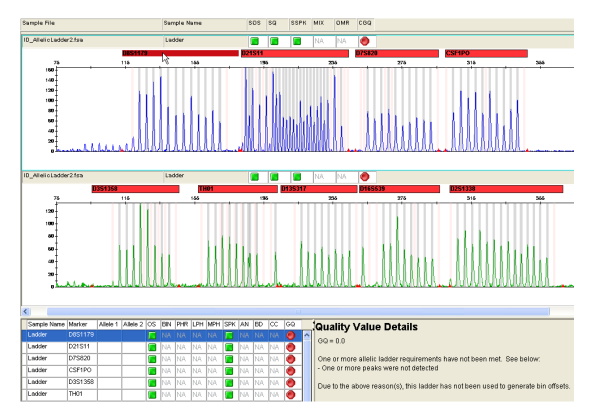


### Note the appearance of the marker headers and bins:

The marker headers are color-coded (green, yellow, red) to reflect the GQ flag color. Bins are displayed according to bin type (see Chapter 2), with virtual bins in pink and physical bins in gray.

**Note:** For more information on virtual and physical bins, see the *GeneMapper*<sup>®</sup> *ID-X Software Version 1.0 Reference Guide*.

**9.** To investigate the reason why a particular marker failed the allelic ladder quality assessment, click the red **D8S1179** marker header in the blue dye pane to display quality assessment information in the QVD pane for this marker.



**Note:** The columns displayed in the Genotypes table are determined by the table setting selected in the Project window.

The QVD pane displays the allelic ladder quality requirement that was not met for the selected marker. In this example, one or more of the expected allelic peaks fell below the peak amplitude threshold set for the D8S1179 marker.

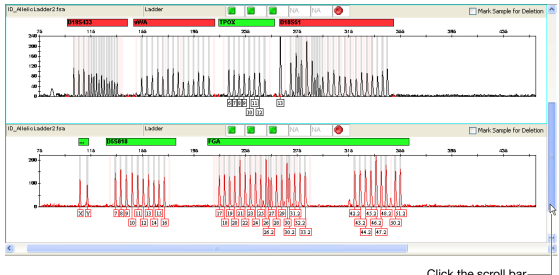
### Quality Value Details

GQ = 0.0

One or more allelic ladder requirements have not been met. See below: - One or more peaks were not detected

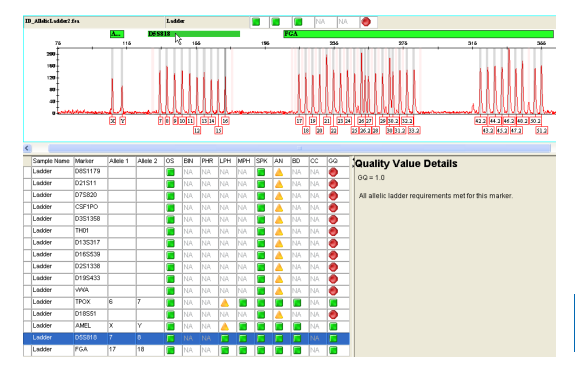
Due to the above reason(s), this ladder has not been used to generate bin offsets.

**10.** Click the vertical scroll bar at the right of Samples plot until you can see the D5S818 marker in the red dye pane.



Click the scroll barto display the next electropherogram



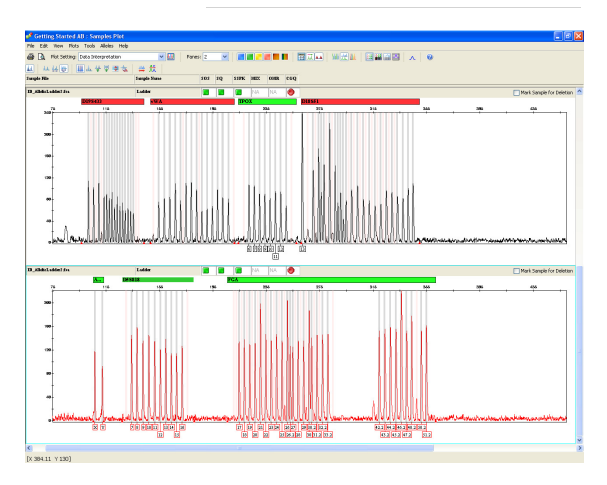


**11.** Click the green **D5S818** marker header to display the quality assessment details for this marker.

The QVD pane indicates that all allelic ladder quality requirements are met for the selected marker. Based on the calculated GQ values shown in the QVD pane, this marker is flagged as Passing quality (). However, since one or more markers within this allelic ladder sample do not meet all quality requirements, the allelic ladder is classified as Low Quality () and it will not be used to create bin offsets.

**12.** Click (No Table) in the Samples plot toolbar to hide the Genotypes table and QVD pane, and view only the dye panes.

**Note:** This toolbar selection does not alter the selected plot settings.





**13.** Select **4** from the Panes drop-down list to display all dyes in the Samples plot.

While this particular ladder would not be usable (since several expected peaks are missing), if another ladder had broad peaks or a spike but was still genotyped accurately, you may override CGQ for the ladder, which will then automatically apply the bin offsets from that ladder. This procedure is recommended only if you do not have another passing ladder.

**14.** Select **File** ► **Close Plot Window** to return to the Project window.

## **Step 2: Examine Control Quality**

Overview	In this section, you will use the features of the Analysis Summary tab and the Samples plot to manually verify the quality of the control samples analyzed in the Getting Started project in Chapter 3.
Viewing Control Quality Status	Select the <b>Analysis Summary</b> tab in the Project window. In Chapter 3, the Control Quality area of the Analysis Summary tab indicates that of the three control samples analyzed, two were of Passing quality ()) and one was of Low Quality ()).
	If a control has met all thresholds, the expected profile was obtained with no other anomalies detected. Therefore, you may not be required to visually inspect the control samples. However, the next two sections demonstrate how to manually review the control samples.



### Examining a Passing Control

1. In the Control Quality area of the Analysis Summary tab, click the link for the 📄 Custom Control sample.

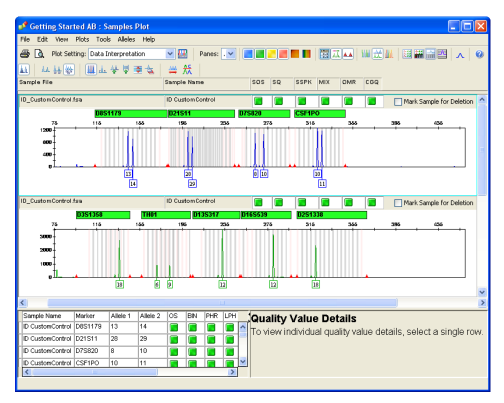
Control Quality per project (based on sample DOVs: SOS\_SSDK\_MIX\_OMP\_SO\_CGO)

Control Type	Total # of Samples	📘 All thresholds met	One or more thresholds not met								
Positive Control	1	1	0								
Custom Control	1	1	0								
Negative Control	1	(h)	1								
Total	<u>3</u>	Σ	<u>1</u>								

The filtered Samples table displays the selected sample. Note that all sample-level PQVs for this custom control sample are  $\boxed{\phantom{a}}$ .

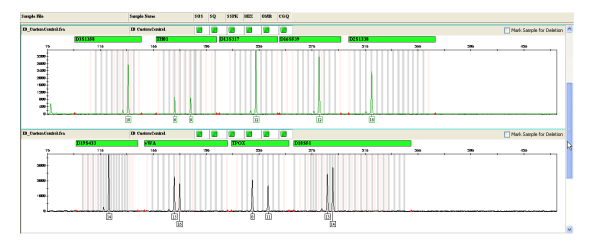
Samples (filtered) Analysis Summary Genotypes														
	Status	Sample Name	Sample Type	Analysis Method	Panel	Size Standard	Custom Control	ARNM	SOS	SQ	SSPK	MIX	OMR	CGQ
3		ID CustomControl	Positive Control	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	CUSTOM CONTROL_YOUR INITIALS							

2. Select the ID CustomControl sample in the filtered Samples table, then click iii (Display Plots). The plot for the selected custom control sample opens in the Samples plot.





**3.** To investigate the quality of this control sample, click the vertical scroll bar at the right of the Samples plot to scroll through all panes. Note that all of the marker headers are colored green, which indicates that this custom control sample has met all marker-level PQV thresholds.



**Note:** For traditional manual review, if you are required to visually inspect the control samples (not rely solely on PQVs), the color-coded marker headers may help reduce the amount of time spent on this task. If all markers are green, you know that this control produced the expected profile and no other anomalies were detected. If a marker header is yellow or red, you know those are the markers where anomalies were detected.

4. Click the green **D8S1179** marker header in the blue dye pane to display the quality assessment details for this marker. Based on the calculated GQ value shown in the QVD pane, this marker is flagged as Passing quality ( ).

Passing CC PQ	V
---------------	---

Sample Name	Marker	Allele 1	Allele 2	os	BIN	PHR	LPH	MPH	SPK	AN	BD	CC	GQ		Quality Value Details
ID CustomControl	D8S1179	13	14											^	GQ = 1.0
ID CustomControl	D21S11	28	29												
ID CustomControl	D7S820	8	10												All quality value thresholds have been met for this marker.

Note the CC PQV, indicating that this marker contains the expected genotype for that marker.

**Note:** The profile for each sample in the project designated as a custom control is compared against the custom positive control profile stored in the Profile Manager. You added the CUSTOM CONTROL profile ID to the Profile Manager in Chapter 2.

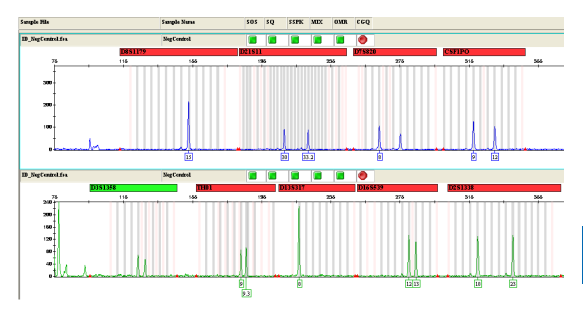
- 5. Select File ➤ Close Plot Window to return to the Project window.
- 1. Select the Analysis Summary tab in the Project window.
- **2.** In the Control Quality area of the Analysis Summary tab, click the link for the **()** Negative Control sample.

Control Quality per p	n oject (based on sa	mple PQVS: 505, 55PK,	MIX, OMR, SQ, CGQ)
Control Type	Total # of Samples	📘 All thresholds met	One or more thresholds not met
Positive Control	1	1	0
Custom Control	1	1	0
Negative Control	1	0	1
Total	3	2	ζm

The filtered Samples table displays the selected sample. Note that this negative control sample shows a 🔴 CGQ PQV.

S	amples (	filtered) Anal-	ysis Summary 🛛 Ge	notypes										
	Status	Sample Name	Sample Type	Analysis Method	Panel	Size Standard	Custom Control	ARNM	SOS	SQ	SSPK	MIX	OMR	CGQ
4		NegControl	Negative Control	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None							۲

Examining a Low-Quality Control 3. To investigate the anomalies detected in this sample, select the NegControl sample in the filtered Samples table, then click
(Display Plots). The plot for the selected negative control sample opens in the Samples plot. Note that several of the marker headers are colored red and there are peaks detected in the sample.



4. Click the red **D8S1179** marker header in the blue dye pane to display the quality assessment details for this marker. Based on the calculated GQ value shown in the QVD pane, this marker is flagged as Low Quality ().

4

Sample Name	Marker	Allele 1	Allele 2	os	BIN	PHR	LPH	MPH	SPK	AN	BD	CC	GQ	Quality Value Details									
NegControl	D8S1179											<u> </u>	۲	GQ =		-							
NegControl	D21S11	30	33.2	NA	NA	NA	NA	NA	NA	NA	NA	<b></b>	۲										
NegControl	D7S820	8		NA	NA	NA	NA	NA	NA	NA	NA			The 1	The following quality value thresholds have not been met:								
NegControl	CSF1PO	9	12	NA	NA	NA	NA	NA	NA	NA	NA		Ă	PQ	V ]	Size	Observed Value	Threshold	GQ Weighting				
	ļ	-											<u> </u>	 C	2	N/A	Non Concordant	N/A	0.5				
NegControl	D3S1358			NA	NA	NA	NA	NA	NA -	NA -	NA												

Check CC PQV

Note the  $\triangle$  CC PQV for this marker, indicating that this marker does not contain the expected result. Since peaks are detected in several markers of this negative control, this negative control is flagged as low quality.

**Note:** You will use the Profile Comparison tool in Chapter 5 of this guide to help determine the possible contributor to this negative control profile.

5. Select File ➤ Close Plot Window to return to the Project window.

## Step 3: Examine Sample Quality

Overview	In this section, you will use the features of the Analysis Summary tab and the Samples plot to manually verify the quality of the unknown samples analyzed with the Getting Started project in Chapter 3.
Viewing Sample Quality Status	Select the <b>Analysis Summary</b> tab in the Project window. In Chapter 3, the Sample Quality area of the Analysis Summary tab indicates that of the eight samples analyzed, four met all quality thresholds ( ) and four did not ( ).
	For the purposes of this guide, you will examine only those samples that did not meet one or more thresholds. Examination of passing

samples would follow the same workflow.



### Examining the Low-Quality Samples

1. In the Sample Quality area of the Analysis Summary tab, click the link for the 🔴 Samples.

Sample Quality per project (based on sample PQVs: SOS, SSPK, MIX, OMR, SQ, CGQ)

	Total # of Samples	📘 All thresholds met	One or more thresholds not met
Samples	<u>8</u>	<u>4</u>	4
			40

San	Samples (filtered) Analysis Summary Genotypes													
	Status	Sample Name	Sample Type	Analysis Method	Panel	Size Standard	Custom Control	ARNM	SOS	SQ	SSPK	MIX	OMR	CGQ
10		Sample 05	Sample	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None				<b></b>			
11		Sample 06	Sample	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None							۲
12		Sample 07	Sample	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None							۲
13		Sample 08	Sample	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None							۲

From the sample-level PQVs shown, you can conclude that:

- Sample 05 contains a spike ( SSPK) within a marker size range or another minor anomaly within the marker size range was detected in addition to spike between two markers ( CGQ)
- **Sample 06** is a potential mixture ( A MIX)
- Sample 07 has a  $\land$  SQ, which indicates there is a resolution issue or a problem with the size standard peak detection
- Sample 08 contains an unexpected peak in between two markers ( OMR) and other anomalies within one or more marker size ranges ( CGQ)

Follow the procedures outlined in "Step 4: Review the Low-Quality Sample Results" on page 90 to individually review the quality of these unknown samples.



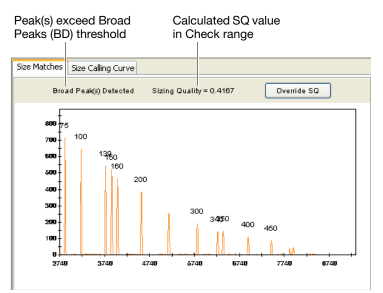
## Step 4: Review the Low-Quality Sample Results

Examining the SQ Sample Results A  $\triangleq$  SQ PQV indicates that the sizing quality is below the passing range specified in the analysis method. This example illustrates:

- How to view SQ using the Size Match Editor
- The SQ PQV flag

### To examine a 📥 SQ PQV:

Select Sample 07 in the Samples table, then click III (Size Match Editor) to view the peak assignments for the size standard peaks in this sample.





Note the following in the Size Match Editor:

- All size standard peaks are present and labeled correctly, as compared to the fragment sizes stored in the CE\_G5\_HID\_GS500 size standard definition file set for this sample
- The calculated SQ value, 0.4167 for this sample, is within the Check ( ) range set in the analysis method (default value = 0.74 - 0.26)
- A "Broad Peak(s) Detected" message indicates there are peaks present in this sample that exceed the Broad Peak (BD) Max Peak Width threshold set in the analysis method

In this example, the SQ is  $\triangle$  due to the broad peaks.

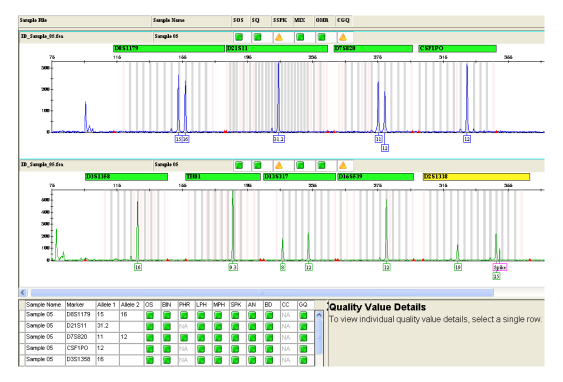
**Note:** For more information on the calculation of SQ, see the GeneMapper<sup>®</sup> ID-X Software Version 1.0 Reference Guide.

- **2.** Click **OK** to close the Size Match Editor.
- 1. Shift-click to select all samples in the filtered Samples table, then click **M** (Display Plots).
- 2. Verify that **Data Interpretation** is selected from the Plot Setting drop-down list.



4

Displaying the Low-Quality Sample Plots **3.** The plots for all of the selected samples are shown in the Samples plot.



The first two panes display the blue dye and green dye for the first sample selected in the Samples table (in this example, Sample 05).

**Note:** The selection order in the Samples table determines the display order in the Samples plot.

### Examining the SSPK Sample Results

A A SSPK (Sample Spike) PQV indicates that one or more spike peaks have been detected within a marker range or between two markers. This example illustrates:

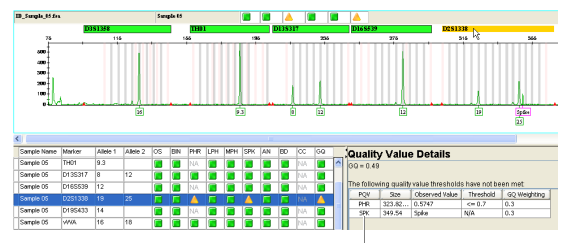
- Automatic labeling of spikes
- The Marker Spike (SPK) and Peak Height Ratio (PHR) PQV flags
- How to confirm a spike in raw data



- How to override GQ
- How to override CGQ

#### To investigate a marker with a yellow or red marker header bar:

1. Click the yellow **D2S1338** marker header in the green dye pane of Sample 05 to display the quality assessment details for this marker.



Check PHR and SPK PQVs

Note the following:

 There is a peak labeled as "Spike" present in this marker and the marker-level SPK PQV is triggered, indicating that one or more spikes are detected within the marker size range (in this case, one spike was detected)

Note: This peak is automatically labeled as a Spike artifact and displayed with a pink label border by the GeneMapper<sup>®</sup> ID-X Software. Peaks labeled as artifacts are not considered true alleles by the software and are not listed in the Genotypes table.

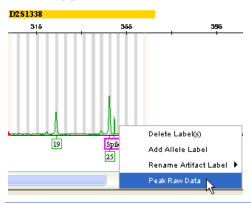
- The marker-level A PHR PQV is also triggered for this marker, indicating that the peak height ratio calculated between the lowest and highest allele peaks within the marker is less than the Min Peak Height Ratio threshold defined in the analysis method (in this example, 0.7)
- Click-drag the Size column margin in the QVD pane to the right to expand its contents and view the peak sizes contributing to the A PHR and SPK PQVs for this marker.

		ag colun to resize	nn							
Quality	/ Value	Details								
GQ = 0.49 The follow	ing quality	value thr	esholds	have n	otbee	en me	et:			
[ PQV ]	Size 🔶	bserve	d Value ]	Threst	nold ]	GQ Weighting				
PHR	323,82,	0.5747	1	<= 0.7	,	0.3	1			
SPK	349.54	Spike		N/A		0.3				
pea	culated ak sitions (bp)	Calcula PHR thresho		PHR thresh define Analy Methe	ed in sis	def Ana	Weighting ined in alysis thod			

**Note:** The SPK and PHR PQV status are used to determine the GQ PQV. Based on the calculated GQ value shown in the QVD pane, this marker is flagged as Check ( $\triangle$ ).

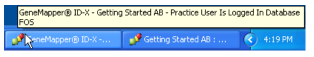


**3.** In the green dye pane, left-click the **Spike** artifact label to select the peak, then right-click the label to open a drop-down menu containing peak edit actions.



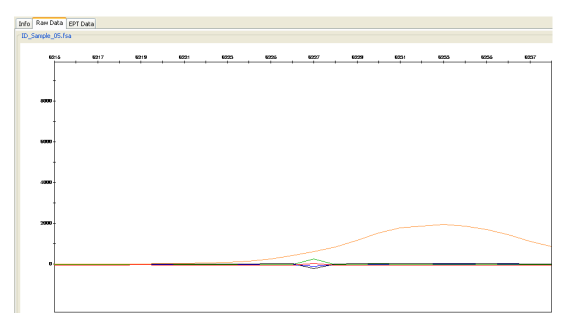
**IMPORTANT!** You must left-click the label before you right-click the label.

- **4.** To confirm the spike peak, select **Peak Raw Data** from the drop-down menu to open the raw data plot for this peak in the Project window.
- 5. To view the raw data plot, click the GeneMapper<sup>®</sup> ID-X Getting Started <your initials> taskbar button.



4

**6.** Review the raw data and peak morphology to confirm that the selected peak is a spike.



**Note:** Once you have validated the spike detection PQVs using your own data, you may not be required to view the raw data to confirm a spike peak.

7. To return to the Samples plot, click the Getting Started <your initials> : Samples Plot taskbar button.

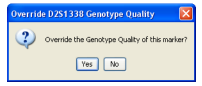


**8.** Right-click the  $\triangle$  GQ PQV in the D2S1338 marker row highlighted in the Genotypes table.

											R	ight-	-clic	k PC	QV
Sample Name	Marker	Allele 1	Allele 2	os	BIN	PHR	LPH	MPH	SPK	AN	BD	CC	GQ		Quality Value Details
Sample 05	D16S539	12				NA						NA		^	To view individual quality value
Sample 05	D2S1338	19	25			<b></b>			<b></b>			NA	Ì <u>≜</u> ¦'	.	
Sample 05	D19S433	14				NA						NA			



**9.** Click **Yes** in the dialog box to override the genotype quality for this marker.



Note that after overriding:

- The GQ PQV changes to
- All other PQVs for the marker turn gray and maintain their original shape (
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- The marker header turns green **D2S1338**

**Note:** Overriding GQ allows you to manually accept the genotype for a particular marker and also provides evidence that the marker was visually inspected by the analyst.

10. To accept the entire sample profile for Sample 05, you can override its ▲ CGQ PQV. To do so, first right-click the ▲ CGQ PQV for Sample 05 in the genotypes header of the Samples plot.

Sample File	sos	SQ	SSPK	MIX	OMR	ceð
ID_Sample_05_fsa			<b></b>			

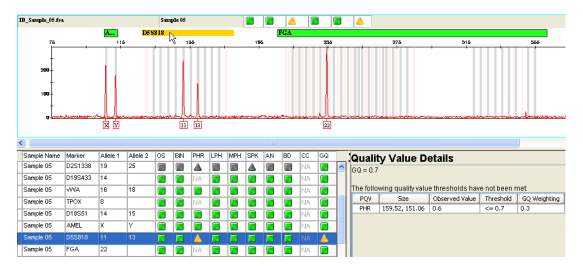
Right-click PQV

The next dialog box displays a message indicating that there are still  $\land$  or o marker-level GQ PQVs present in this sample. Unless you have viewed the rest of the markers and confirmed the genotypes, you should not continue with the CGQ override.



Click **No** to return to the Samples plot and confirm the genotypes for the remaining markers.

- **11.** Click the vertical scroll bar at the right of the Samples plot to scroll through the two remaining dye panes (yellow and red) for Sample 05. Note that there is a yellow D5S818 marker header in the red dye pane. It was this marker that triggered the Override CGQ warning message seen in step 10 on page 97.
- **12.** Click the yellow **D5S818** marker header to display the quality assessment details for this marker.



Note the marker-level  $\triangleq$  PHR PQV in the Genotypes table for this marker. Based on the calculated PHR value shown in the QVD pane, this marker is flagged as Check ( $\triangleq$ ).

**13.** Right-click the  $\triangle$  GQ PQV in the highlighted row of the Genotypes table, then click **Yes** in the dialog box to override the genotype quality for this marker.



**14.** Now that all GQ PQVs are for Sample 05, click **Yes** in the next dialog box to override the CGQ for this sample.



Note that after overriding, the CGQ PQV changes to (Manually Overridden).

**Note:** Overriding CGQ allows you to manually accept the entire profile for a particular sample and also provides evidence that the sample was visually inspected by the analyst.

You have now completed reviewing Sample 05. Another use for CGQ override will be demonstrated in Chapter 6 for review purposes.

A A Mixed Source (MIX) PQV indicates a potential mixed-source sample. This example illustrates:

- · How to select peaks and view their calculated peak height ratios
- The marker-level PQVs triggered in mixed samples
- The PHR and Allele Number (AN) PQV flags

Examining the

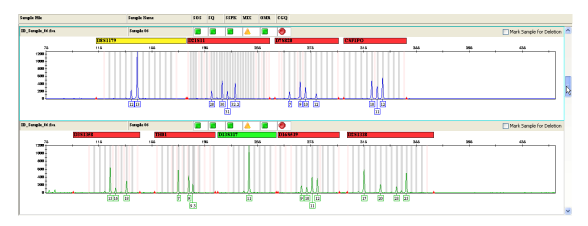
MIX Sample

Results



## To investigate the cause of the yellow MIX PQV flag:

 Click the vertical scroll bar at the right of the Samples plot until you can see the first two dye panes (blue and green) for Sample 06.

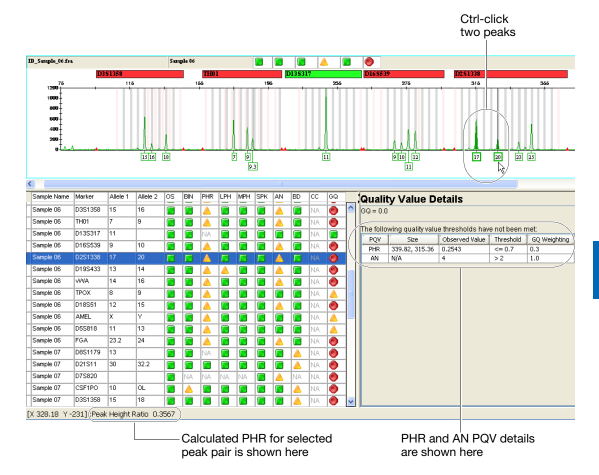


**2.** Click the yellow **D8S1179** marker header in the blue dye pane to display the quality assessment details for this marker.

Sample Name	Marker	Allele 1	Allele 2	OS	BIN	PHR	LPH	MPH	SPK	AN	BD	CC	GQ		Qualit	v Value De	etails		
Sample 05	AMEL	х	Y									NA		^	GQ = 0.7				
Sample 05	D5S818	11	13									NA		1					
Sample 05	FGA	22	1			NA						NA		1	The follow	ving quality valu	e thresholds hav	e not been r	net:
Sample 06	D8S1179	40	42												PQV	Size	Observed Value	Threshold	GQ Weighting
Sample up	Dostina	12													PHR	139.43.143.98	0.1811	<= 0.7	0.3
Sample 06	D21S11	28	30									NA.	۲			,			

Note the marker-level  $\triangle$  PHR PQV in the Genotypes table for this marker. Based on the calculated PHR value shown in the QVD pane, this marker is flagged as Check ( $\triangle$ ).

**3.** Ctrl-click the **17** and **20** allele peaks in the green dye pane for marker D2S1338 to select them both. The software automatically calculates a peak height ratio (PHR) value for the selected peaks and displays the results in the status bar at the bottom of the Samples plot.





Note the following:

- The calculated PHR is less than the minimum PHR threshold of 0.7 specified in the analysis method
- The AN PQV in the Genotypes table for this marker, indicating the number of alleles detected is greater than the Max Expected Alleles defined in the analysis method

**Note:** The AN and PHR PQV status is used to determine the GQ PQV. Based on the observed AN and PHR values shown in the QVD pane, this marker is flagged as Low Quality ().

**4.** Repeat step 3 on page 101 for any of the other peak pairs in this sample.

You have now completed reviewing Sample 06. This sample was expected to be from a single source. You will use the Profile Comparison tool in Chapter 5 to help determine the potential contributor(s) to this mixture.

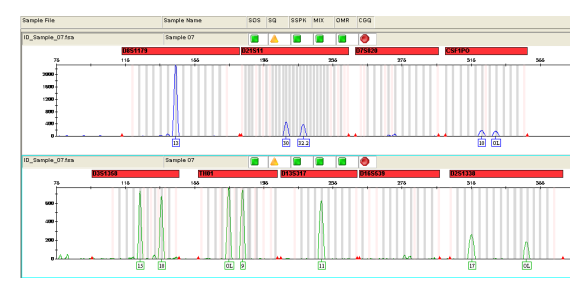
**Note:** For more information on the conditions that must be met for a sample to be considered a potential mixture, see the  $GeneMapper^{\ensuremath{\mathbb{R}}}$  *ID-X Software Version 1.0 Reference Guide.* 

Examining the SQ Sample Results This example illustrates:

- How to view PQV trigger labels
- The Broad Peak (BD) PQV flag
- How to delete allele labels for a sample
- How to add a reason for change to the audit trail
- How to mark a sample for deletion from the project

### To investigate this low-quality sample:

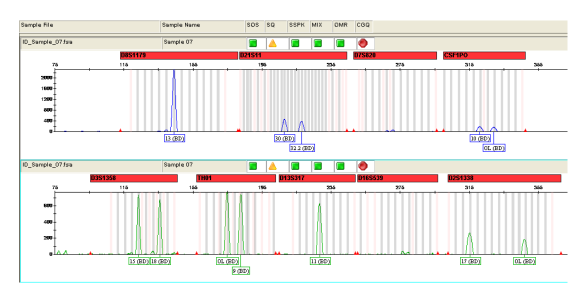
**1.** Click the vertical scroll bar at the right of the Samples plot until you can see the first two dye panes (blue and green) for Sample 07.



**Note:** This is the same sample you viewed using the Size Match Editor earlier in this chapter (see "Examining the SQ Sample Results" on page 90).

In the Samples plot toolbar, click (PQV Trigger Peak) to add PQV suffixes to peak labels for peaks that cause any of the following PQVs: LPH, MPH, BD, OS.

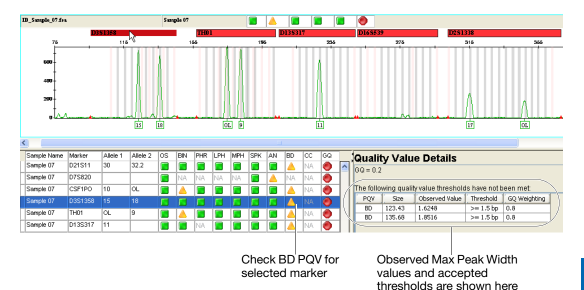
**Note:** Displaying the PQV trigger peak labels can help you quickly find the exact peaks that triggered specific PQV flags.



In this example, (BD) is appended to the label of each peak that exceeds the Max Peak Width threshold.

**3.** Click (PQV Trigger Peak) again to turn off the PQV trigger labels.

4. Click the red D3S1358 marker header to display the quality assessment details for this marker. Note the marker-level ▲ BD PQV in the Genotypes table, and the Max Peak Width threshold and observed values related to this PQV in the QVD pane.



The genotype profile for this sample is not reliable and should not be reported. Therefore, from the Samples plot, you can choose to:

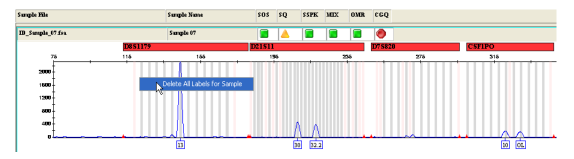
• Delete all labels for a sample

or

• Mark the sample for deletion from the project

The following steps 5 through 8 demonstrate these two options.

**5.** To delete all labels for Sample 07, right-click in any pane for this sample, then select **Delete All Labels for Sample**.



- 6. Click Yes in the Delete All Labels dialog box.
- 7. Enter **Bad injection** in the Reason(s) for Change dialog box (recorded in the audit trail), then click **OK**.

🧬 Reason(s) f	or Change
Reason(s) for Ch	ange
Attribute	RUN.Identifiler Samples.SAMPLE.ID_Sample_07.fsa.ALLELE.13.deleted
Old Value	13 [KitT.AmpFLSTR_Panels_v1X.PANEL.Identifiler_v1X.MARKER.D851179.
New Value	
Enter the Reaso	on(s) for Change:
	Bad injection
	ОК

**Note:** By default, a reason for change audit trail entry is required for all label edits.

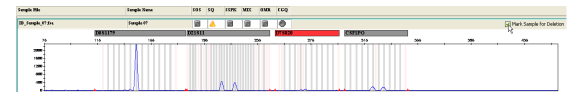


Note that after deleting all allele labels for Sample 07:

- All sample-level and marker-level PQVs for the sample turn gray and maintain their original shapes (
- Affected marker header colors change to gray
- The sample will still be listed in the Samples and Genotypes table, but no allele values are reported



**8.** To delete Sample 07 from the Getting Started project, select **Mark Sample for Deletion** at the top right of any dye pane for this sample.



**Note:** The remaining panes associated with this sample are automatically marked for deletion by the software.

The sample will be deleted from the project when the Samples plot is closed. For now, keep the Samples plot open to continue reviewing the last low-quality sample results in this project.

You have now completed reviewing Sample 07.



A  $\triangleq$  Outside Marker Range (OMR) PQV indicates that one or more peaks were detected between two marker size ranges specified in the panel for the marker. This example illustrates:

- Automatic labeling of OMR peaks
- The Out of Bin Allele (BIN), Low Peak Height (LPH), and Max Peak Height (MPH) PQV flags
- How to change an artifact label to an allele label and assign it to a selected marker
- How to save a custom artifact label for future use
- How to change an allele label to an artifact label



# To investigate the source of the OMR peak and other potential anomalies:

Click the vertical scroll bar at the right of the Samples plot until you can see the last two dye panes (yellow and red) for Sample 08.



Note the following:

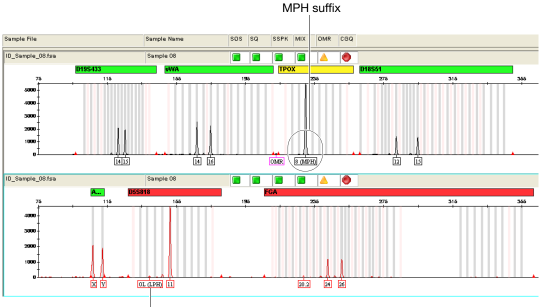
- There is a peak labeled OMR present between the vWA and TPOX markers
- The marker header for TPOX is yellow
- The marker headers for D5S818 and FGA are red
- There is a peak labeled OL present in the D5S818 marker
- There are three labeled peaks present in the FGA marker

Follow the procedures in the following sections (on pages 110 to 128) to individually review the quality of these markers.

You will first investigate the OMR peak between vWA and TPOX and the quality of the TPOX marker.

### To examine TPOX marker and OMR peak:

 In the Samples plot toolbar, click (PQV Trigger Peak) to add PQV suffixes to peak labels for peaks that cause any of the following PQVs: LPH, MPH, BD, OS.

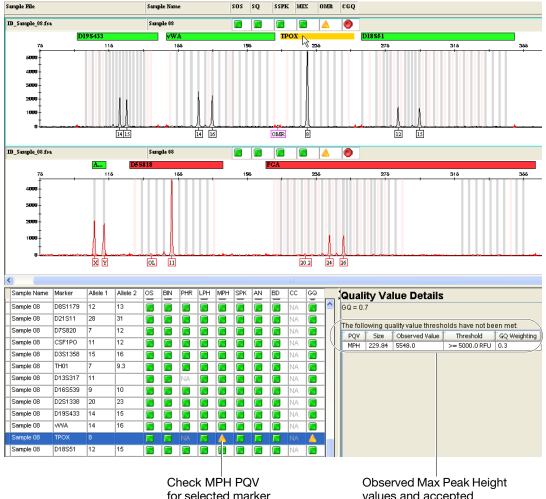


LPH suffix

In this example, (MPH) in TPOX and (LPH) in D5S818 are appended to the label of each peak that does not meet the thresholds related to these PQVs set in the analysis method.

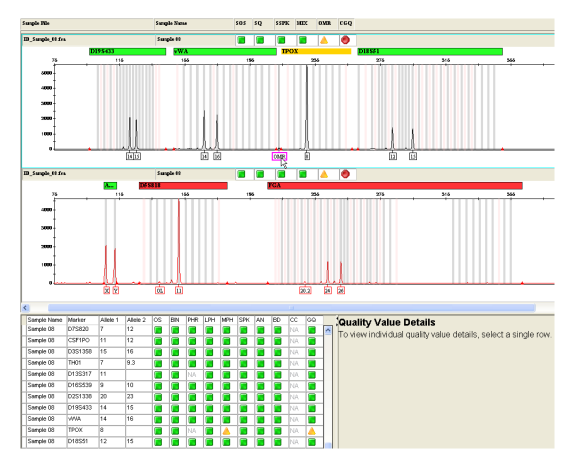
2. Click (PQV Trigger Peak) again to turn off the PQV trigger labels.

**3.** Click the yellow **TPOX** marker header to display the quality assessment details for this marker. Note the marker-level MPH PQV in the Genotypes table for this marker, and the Max Peak Height threshold (in RFU) and observed values related to this PQV in the QVD pane.



values and accepted thresholds are shown here 4

**4.** With the TPOX marker header still selected, left-click the **OMR** label.

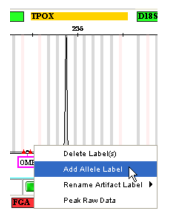


**Note:** This peak is automatically labeled as an OMR artifact by the GeneMapper<sup>®</sup> *ID-X* Software. Peaks labeled as artifacts are not listed in the Genotypes table. In some rare cases, an OMR artifact peak may be a true allele; you may need to verify the classification of an OMR peak if you encounter one in your own data.

4

For this example, you will assume that this OMR artifact peak is indeed a true allele solely to learn how to add an allele label to an artifact peak and assign it to a selected marker (although you are aware this is not a true DNA peak).

5. Right-click the OMR label, then select Add Allele Label.



In the Add Custom Allele Label dialog box, enter < 6, then click OK. Note that the TPOX marker is selected for this allele since this was the marker selected before adding the allele.</li>

Add Custom Allele Label 🛛 🔀								
Custom Allele Label:	< 6							
Marker Name:	IPOX							
ОК	Cancel Help							

**Note:** This custom allele will be added to the TPOX marker and entered as an allele in the Genotypes table.

7. In the Reason(s) For Change dialog box, enter Microvariant OMR, then click OK.

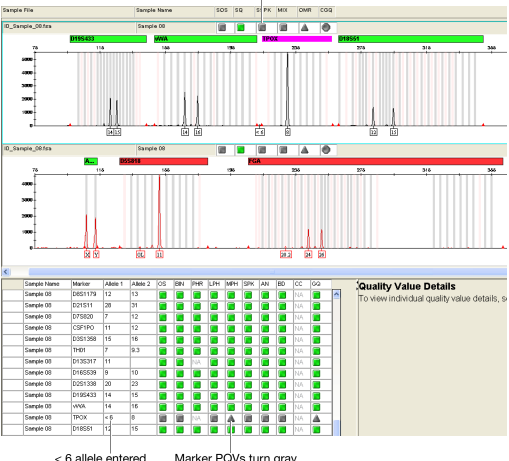
💕 Reason(s) t	for Change 🛛 🛛 🔀						
Reason(s) for Cl	hange						
Attribute	RUN.Identifiler Samples.SAMPLE.ID_Sample_08.fsa.ALLELE.OMR.modified						
Old Value	OMR [KitT.AmpFL5TR_Panels_v1X.PANEL.Identifiler_v1X.MARKER.TPOX.BASEPAIR.212.76.]						
New Value	< 6 [KitT.AmpFLSTR_Panels_v1X.PANEL.Identifiler_v1X.MARKER.TPOX.BASEPAIR.212.76.5]						
Enter the Reas	on(s) for Change:						
	Microvariant OMR						
ОК							

Note that after editing:

- All sample-level PQVs (except SQ) for the sample turn gray (
   () to indicate that the sample has been edited
- All marker-level PQVs for the marker turn gray and maintain their shape (
   (), to indicate that the marker has been edited
- The allele is listed in the Genotypes table
- The allele label changes text and color from pink to black
- The marker header color changes from yellow to gray (pink if selected)



Sample PQVs (except SQ) turn gray, but keep original shape



< 6 allele entered in Genotypes table Marker PQVs turn gray, but keep original shape

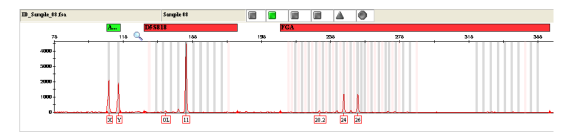
- 8. Select the **TPOX** marker row in the Genotypes table, then rightclick its ▲ GQ PQV.
- **9.** Click **Yes** in the dialog box to override the genotype quality and manually accept the current genotype (<6, 8) for the TPOX marker. The GQ PQV changes to **and the marker header** turns green.

You have completed reviewing the TPOX marker. You will now investigate the OL peak present in the D5S818 marker.

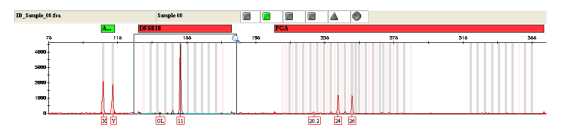
4



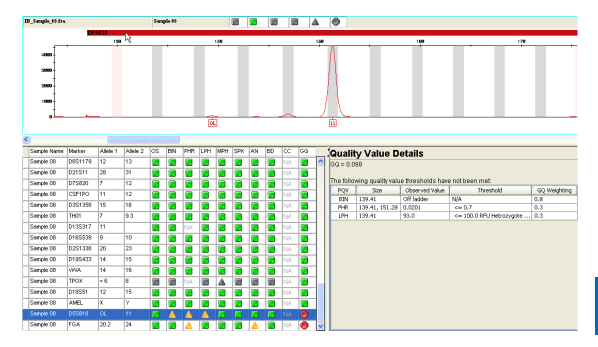
### To examine the cause of an extra peak:



**2.** Click-drag the pointer to the right to create a box that includes the area to the left of the D5S818 marker, then release to zoom in on this marker, as shown below.



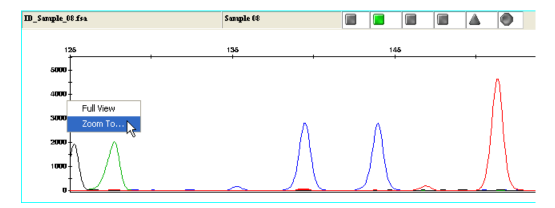
Click the red D5S818 marker header to display the quality assessment details for this marker. Note the marker-level <a>A</a>BIN PQV in the Genotypes table for this marker.



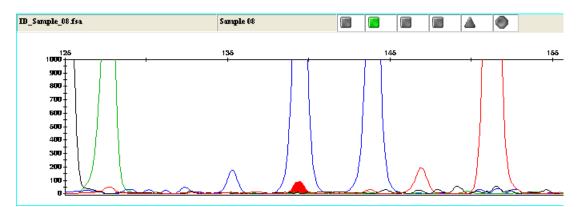
**Note:** A A BIN PQV indicates that one or more labeled peaks are detected outside an offset bin. These peaks are automatically labeled as OL peaks by the software. For information on bin offsetting, see the *GeneMapper*<sup>®</sup> *ID-X Software Version 1.0 Reference Guide.* 

- 4. Click the OL peak label to select the OL peak.
- 5. Click iiii (Combine Dyes) in the Samples plot toolbar to display all dye colors for the sample in one pane.

6. Place the pointer next to the Y-axis until the pointer changes to a , then right-click and select **Zoom to**.



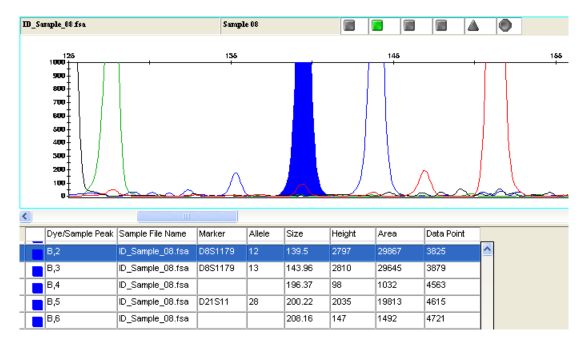
7. Enter 1000 in the Y-Axis Zooming dialog box, then click **OK** to zoom in on the OL peak. Note that the red OL peak is located under a larger blue dye peak.



**8.** Ctrl-click the overlaying blue dye peak to select it (the red OL peak should still be highlighted).

ID_Sample_08.fs	a				Sample	08								4			
1000 900 800 700 600 800 400 200 100 0		1	30 +	~				140					<u> </u>		50		<u> </u>
<	Martin	Allele 4		00	-	DUD	L DU	MOLL	ODK	L a b l	00	00	0.0	_			
Sample Name Sample 08	Marker D8S1179	Allele 1	Allele 2	os	BIN	PHR	LPH	MPH	SPK	AN	BD	CC	GQ		Quality		
Sample 00	D21S11	28	31									NA		<u>^</u>	To view ii	ndividual	quality va
Sample 00	D7S820	7	12		_				_		_			-			
Sample 08	CSF1PO	11	12	_			_			_		NA					
Sample 08	D3S1358	15	16									NA	_				
Sample 08	TH01	7	9.3									NA					
· ·	D13S317	11	9.3									NA					
Sample 08 Sample 08	D135317	9	10			NA						NA					
Sample 08	D165539	20	23									NA					
<u> </u>	D251330	14	15									NA					
Sample 08 Sample 08		14	15									NA					
<u> </u>												NA					
Sample 08	TPOX	< 6	8			NA						NA					
Sample 08	D18S51	12	15									NA		-			
Sample 08	AMEL	x	Y									NA					
Sample 08	D5S818	OL	11										۲	$\mathbf{\mathbf{v}}$			

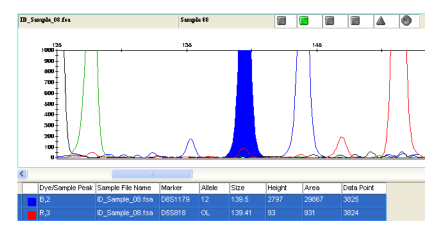
**9.** Click *iii* (Sizing Table) in the Samples plot toolbar to display the Sizing table, which contains the following peak sizing information for all detected peaks in the sample: size (in base pairs), peak height, peak area, and data point.



**Note:** The columns displayed in the Sizing table are determined by the plot setting selected in the Samples plot.



**10.** With the blue dye and red OL peaks still selected, press **Ctrl+G** to display only the selected peaks in the Sizing table.

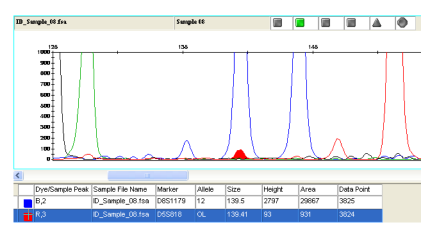


The sizing data indicates that the red OL peak could be caused by spectral pull-up.

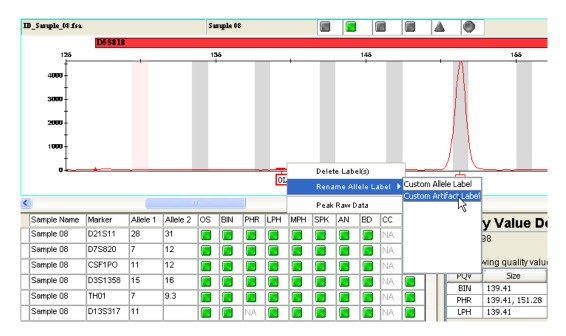
You will change the software-generated OL peak label to a custom artifact label in the next steps.

# To change the OL peak label to a custom artifact label:

1. Select the **D5S818** row in the Sizing table to deselect the overlaying blue dye peak (the red OL peak should still be highlighted).



- 2. Click 🔛 (Separate Dyes) in the Samples plot toolbar to display each dye color for the sample in separate panes.
- **3.** Click (Genotypes Table) in the Samples plot toolbar to replace the Sizing table with the Genotypes table and QVD pane.
- 4. Left-click the OL peak label in the red dye pane, then rightclick the label and select Rename Allele Label → Custom Artifact Label.





- 5. Complete the Custom Artifact Label dialog box:
  - a. Enter SPU <your initials> (for example, SPU AB).
  - b. Select Add to predefined list.

Custom Artifact Label		×
Custom Artifact Label:	SPU AB	
Add to predefined list:		
	Cancel Help	

c. Click OK.

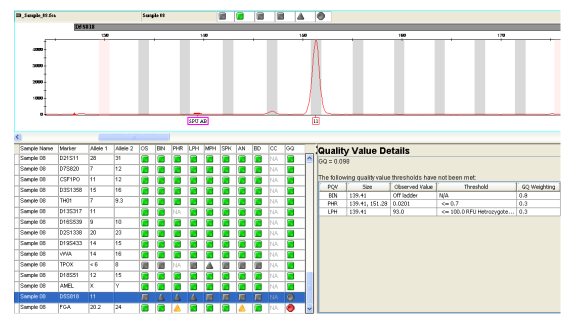
**Note:** This option allows you to select this custom artifact label from a list of labels in the future.

6. In the Reason(s) For Change dialog box, enter **Pull-up**, then click **OK**.

💕 Reason(s) f	for Change 🛛 🔀							
-Reason(s) for Ch	hange							
Attribute	RUN.Identifiler Samples.SAMPLE.ID_Sample_08.fsa.ALLELE.OL.modified							
Old Value	OL [KitT.AmpFLSTR_Panels_v1X.PANEL.Identifiler_v1X.MARKER.DSS818.BASEF							
New Value	SPU AB [KitT.AmpFLSTR_Panels_v1X.PANEL.Identifiler_v1X.MARKER.D55818.B]							
Enter the Reaso	on(s) for Change:							
	Pull-up							
OK Å								

Note that after editing:

- The SPU artifact label is displayed with a pink border
- The SPU artifact peak is not added to the Genotypes table since an artifact is not considered a true allele
- All marker-level and sample-level PQVs (except SQ) turn gray and maintain their original shape (
- The marker header color changes to gray (pink if selected)



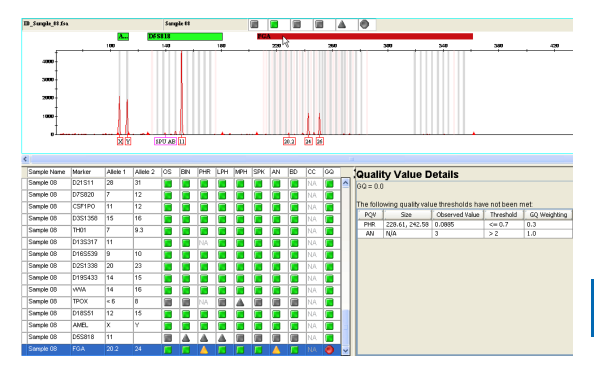
- 7. Verify the **D5S818** marker row is selected in the Genotypes table, then right-click its GQ PQV.
- 8. Click Yes in the dialog box to override the genotype quality and manually accept the current genotype (11) for the D5S818 marker. The GQ PQV changes to and the marker header turns green.

You have completed reviewing the D5S818 marker. You will now investigate the extra allele present in the FGA marker.

1. Press Ctrl+Minus Sign (–) four times to incrementally zoom out on all panes of the Samples plot.



**2.** Click the red **FGA** marker header in the red dye pane of Sample 08 to display the quality assessment details for this marker.



Note the following:

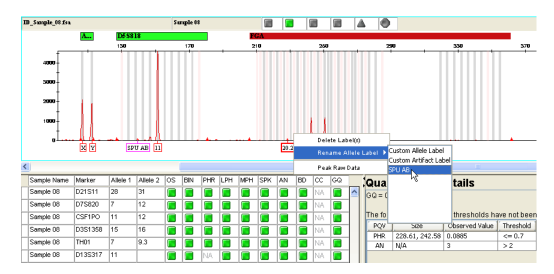
- The software has detected three alleles in the marker range: 20.2, 24 and 26
- The marker-level 📥 AN PQV is triggered

**Note:** A AN PQV indicates that the software detects either no alleles or more alleles than the Max Expected Alleles threshold set in the analysis method (in this example, set to two).

• The marker-level 📥 PHR PQV is triggered

In this example you will assume the extra allele is caused by pull-up as well.

4



**3.** Left-click the **20.2** peak label, then right-click the label and select **Rename Allele Label → SPU <your initials>**.

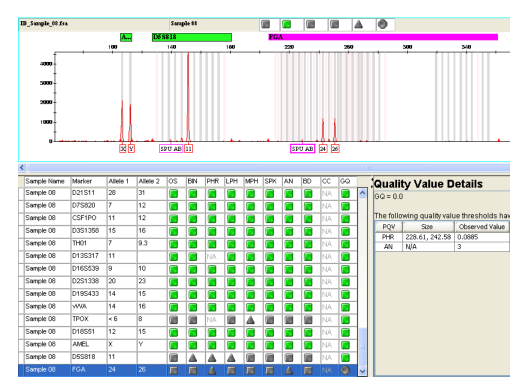
**Note:** Note that the SPU custom artifact label you created in step 5 on page 123 is now displayed in the list of available labels.

**4.** In the Reason(s) For Change dialog box, leave the **Pull-up** text as shown (recorded in the audit trail), then click **OK**.

🥩 Reason(s) t	for Change
Reason(s) for Ch	hange
Attribute	RUN.Identifiler Samples.SAMPLE.ID_Sample_08.fsa.ALLELE.20.2.modified
Old Value	20.2 [KitT.AmpFLSTR_Panels_v1X.PANEL.Identifiler_v1X.MARKER.FGA.BASEPAIR.228.61]
New Value	SPU AB [KitT.AmpFLSTR_Panels_v1X.PANEL.Identifiler_v1X.MARKER.FGA.BASEPAIR.228]
Enter the Reas	on(s) for Change:
	Pull-up
	ОК

Note that after editing:

- The SPU artifact label is displayed with a pink border
- The SPU artifact peak is not added to the Genotypes table since an artifact is not considered a true allele



• The marker header color changes to gray (pink if selected)

- **5.** Verify the row for the FGA marker is selected in the Genotypes table, then right-click its GQ PQV.
- **6.** Click **Yes** in the dialog box to override the genotype quality and manually accept the current genotype (24, 26) for the FGA marker.

The next dialog box displays a message indicating that all marker-level GQ PQVs for sample Sample 08 are now green.



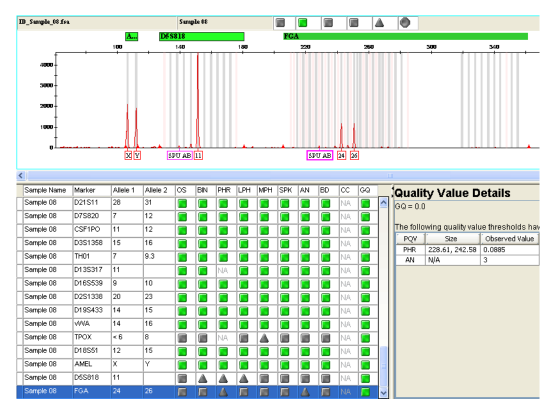
**7.** Click **No** in this Override CGQ dialog box to keep the current CGQ status for this sample.



You will review this sample further in Chapter 6.

Note that after overriding GQ:

- The GQ PQV changes to
- The marker header turns green



You have now completed reviewing Sample 08.

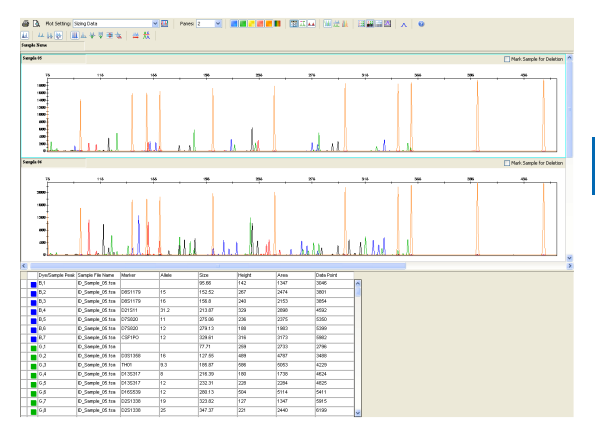


## **Step 5: View Additional Plot Settings**

Additional plot settings are supplied with the GeneMapper<sup>®</sup> ID-XSoftware to assist with performing other functions within the plot window. Follow the procedures on pages 129 through 132 to review the elements of these other plot settings.

Select each of the following from the Plot Setting drop-down list:

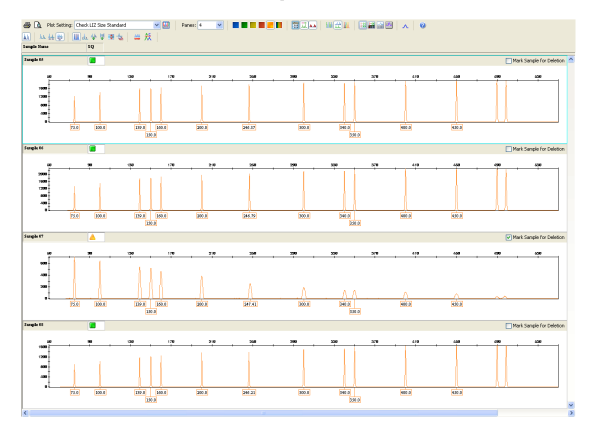
• Sizing Data – Displays peak detection and sizing information in a format similar to the ABI PRISM<sup>®</sup> GeneScan<sup>™</sup> software plots



4

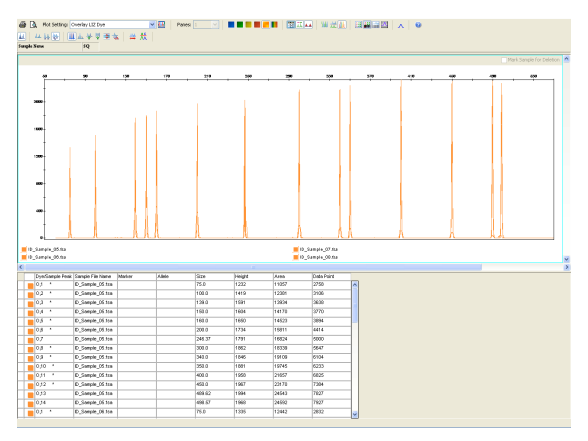


 Check LIZ Size Standard – Displays plots in the same format as the Check GS500 Macro in the ABI PRISM<sup>®</sup> Genotyper<sup>®</sup> Software templates



# 4

• **Overlay LIZ Dye**– Overlays all selected size standard fragments in one electropherogram pane





• **Traditional Genotype Plot** – Displays plots in a format similar to the ABI PRISM<sup>®</sup> Genotyper<sup>®</sup> Software



After you have finished reviewing each plot setting, you are now ready to close the Samples plot.



## Step 6: Delete Samples from the Project

 Click (Bring Marked Samples To Top) in the Samples plot toolbar. The sample (Sample 07) you marked for deletion in step 8 on page 108 is displayed in the top pane of the Samples plot.



You can use this feature to verify which samples are marked for deletion before closing the plot window, since closing the plot window automatically deletes these samples from the project.

2. Select File ➤ Close Plot Window. A dialog box displays a message indicating that one sample is marked for deletion.



**3.** Click **Yes** in the dialog box to close the Samples plot and return to the Project window. The software automatically deletes Sample 07 from the Getting Started project.

**Note:** From step 4 on page 95, the raw data for Sample 05 is still displayed in the Project window.



4. Select the **Project** node in the navigation pane to display the remaining samples of the Getting Started project in the Samples table.

🗅 🔒 🛛 😼 🗗 🛄			u   🖾 🎟 i	3 🕨 🖌	Table Setting: 31XX Da	ta Analysis	-	🔎 🖨 🖪 🖉 🥥							
Project	Sam	ples A	nalysis Summary 0	enotypes											
Identifiler Samples		Status	Sample Name	Sample Type	Analysis Method	Panel	Size Standard	Custom Control	ARNM	SOS	SQ S	SPK	MIX	OMR	CG
ID_AllelicLadder2.fsa	1		Ladder	Allelic Ladder	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None					NA	NA	۲
ID_CustomControl.fsa	2		Ladder	Allelic Ladder	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None					NA	NA	
ID_NegControl.fsa	3		ID CustomControl	Positive Control	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	CUSTOM CONTROL_YOUR INITIALS							
ID_PosControl.fsa ID_Sample_01.fsa	4		NegControl	Negative Control	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None	<u> </u>						۲
ID_Sample_02.fsa	5		PosControl	Positive Control	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None	<u> </u>						Ō
ID_Sample_03.fsa	6	<u> </u>	Sample 01	Sample	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None	<u> </u>						
ID_Sample_04.fsa ID_Sample_05.fsa	7	<u> </u>	Sample 02	Sample	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None	<u> </u>						
ID_Sample_05.fsa	8	<u> </u>	Sample 03	Sample	Getting Started_your initials	ldentifiler_v1X	CE_G5_HID_GS500	None	<u> </u>						
ID_Sample_08.fsa	9		Sample 04	Sample	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None	<u> </u>	<b>i</b>					ī
	10		Sample 05	Sample	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None							ō
	11		Sample 06	Sample	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None		Ē	ā i		$\overline{\Delta}$		۲
	12		Sample 08	Sample	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	None						$\overline{\mathbf{A}}$	ŏ
															-

Note that Sample 07 is removed from this view. You are now ready to save your edits to the Getting Started project.

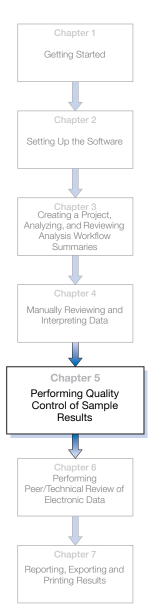
### Step 7: Save the Project

Now that you have finished editing the Getting Started project, click (Save Project) to save all project edits (label edits and comments) to the GeneMapper<sup>®</sup> ID-X Software database so they can be viewed the next time the project is opened.

In Chapter 6, you will review your edits to the Getting Started project.

First, in Chapter 5 you will use the Profile Comparison tool to help determine the potential contributors to the negative control with detected peaks and the mixture sample you just reviewed.

# Performing Quality Control of Sample Results



This chapter covers:

143

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Step 3: Perform Lab Reference Comparison and View Resu	lts.

Step 4: Perform Control/QC Comparison and View Results 144 

## **Overview**

### About Quality Control of Sample Results

Use the Profile Comparison tool to perform quality control of sample results in a project. Only samples and controls with a or SQ are included in comparisons (allelic ladder samples are not included).

The Profile Comparison tool does the following for all samples in a project:

- Groups samples with 100% concordant profiles
- Compares samples in the project against all other samples in the project
- Compares samples in the project against lab reference, custom control, and QC sample profiles stored in the Profile Manager

**IMPORTANT!** Before you use the Profile Comparison tool to evaluate the samples in your project, edit allele labels as needed and ensure that no OL allele labels are present. Samples containing OLlabeled peaks are not considered in comparisons.

**In This Chapter** In this chapter, you will use the Getting Started project, saved in Chapter 4, to determine the potential contributors to the Negative Control and mixture samples observed in Chapter 4 by comparing the unknown sample profiles in the project against each other and against the profiles imported into the GeneMapper<sup>®</sup> *ID-X* Software database in Chapter 2.



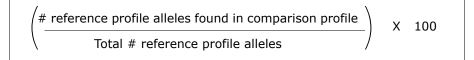
## **Understanding the Profile Comparison Tool**

#### Terms You Need to Know

• **Reference Profile** – The reference profile is the profile against which another profile is compared to determine the % Match. The software performs pairwise comparisons to determine the direction of comparison that yields the higher % Match, then reports only the direction of comparison with the higher % Match.

A mixed-source sample is used as a reference profile only when it is compared to another mixed-source sample.

- **Comparison Profile** The comparison profile is compared to the reference profile to determine the % Match result.
- %-Match Calculated using the following formula:



Each sample or group (determined in Sample Concordance tab) is compared to every other individual sample or single-source or mixed-source group in the project. For a pair of samples or groups, two comparisons are performed. Each sample/group is used as a reference, to which the other sample (comparison sample) is compared. The comparison direction that produces the highest % Match is reported.

For example:

- Sample 1 is a single-source sample
- Sample 2 is a mixed-source sample
- Sample 1 contains 10 alleles (all of which are found in Sample 2)
- Sample 2 contains 20 alleles
- When Sample 1 is used as the reference, the % Match = 100%.
- When Sample 2 is used as the reference, the % Match = 50%.



- In this example, the comparison in which Sample 1 is used as the reference is reported.

**Note:** Results are reported only when the % Match is greater than or equal to the user-defined Percent Match Threshold (range is 50 - 100).

- **Single-source groups** Samples with no more than three alleles in more than one marker in the Genotypes table, and are 100% concordant (all markers and all alleles match).
- Mixed-source groups Samples with two or more markers with three or more alleles in the Genotypes table, and are 100% concordant.
- Individual samples Samples that contain unique profiles.
- Lab Reference and Custom Control Profile Profiles imported and stored in the Profile Manager.

### Sample Concordance

The software performs a sample concordance check to group samples with identical profiles and minimize the number of comparisons performed on the other tabs of the Profile Comparison tool.

To perform sample concordance, the software:

- Considers all analyzed samples in the open project with a 
   (Pass) or 
   (Check) SQ value, except:
  - Allelic Ladder sample types
  - Samples that contain OL labels
- Compares each sample against every other sample to determine 100% concordance (all markers and all alleles match).
- Groups 100% concordant profiles in one or more single-source or mixed-source groups.

**Note:** The Sample Concordance tab only displays grouped samples that are 100% concordant. It does not list individual samples. However, individual samples that meet or exceed the user-defined Percent Match Threshold are considered in the Sample Comparison, Lab Reference Comparison, and Control/QC Comparison tabs. Sample, Lab Reference, Custom Control, and QC Comparison To perform comparisons, the software:

- For sample comparisons: Performs a pairwise comparison of all individual sample, single-source group, and mixed-source group profiles.
- For lab reference and custom control/QC comparisons: Compares all individual sample, single-source group profiles, and mixed-source group profiles to the lab reference or custom control profiles stored in the Profile Manager (QC comparisons use custom control profiles).
- Calculates the % Match for each comparison.
- Determines the reference-to-comparison % Match to report.

**Note:** Only the comparison that yields the highest % Match is reported. The comparison that yields the lower % Match is not used or listed in results.

• Displays the % Match results that are greater than or equal to the user-defined Percent Match Threshold.

5



# Step 1: Examine Sample Concordance

**Overview** In this section, you will use the Profile Comparison tool to check for concordance among all samples in the Getting Started project.

Examining Sample Concordance

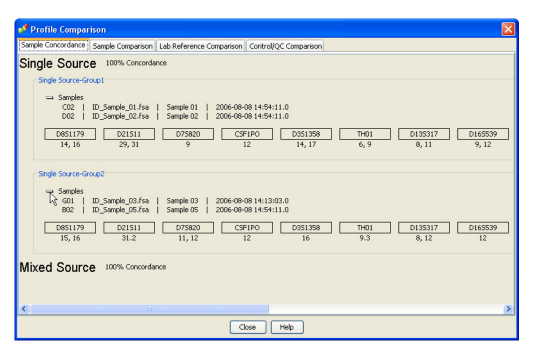
- 1. In the Project window, open the Getting Started project (see step 1 on page 75).
- 2. Select Tools → Profile Comparison. The Profile Comparison tool opens to the Sample Concordance tab.

🧬 Profile Comparison		
Sample Concordance Sample Compari	rison Lab Reference Comparison Control/QC Comparison	
Single Source 100% Conc	cordance	
Single Source-Group1		
슈 Samples		
D851179 D21511 14, 16 29, 31		rs
Single Source-Group2		
D851179 D21511 15, 16 31.2		3
Mixed Source 100% Conc	cordance	
<	<b>N</b>	
	Close Help	

Two single-source groups (Group 1 and Group 2) are listed on this tab for the Getting Started project.



**3.** Click 🔂 to expand the Samples view for both single-source groups.



In the Getting Started project, ID\_Sample\_01 and ID\_Sample\_02 (Single Source-Group 1) are 100% concordant, and ID\_Sample\_03 and ID\_Sample\_05 (Single Source-Group 2) are 100% concordant. All other samples in the project have unique profiles and are not shown on this tab.

# Step 2: Perform Sample Comparison and View Results

**Overview** In this section, you will use the Profile Comparison tool to determine whether any of the sample profiles in the Getting Started project are potential contributors to another sample profile in the project.

Performing	1. Select the Sample Comparison tab.
Sample Comparison	<ol> <li>Keep the Percent Match Threshold at 80, then click Compare Profiles.</li> </ol>
	<b>Note:</b> The default Percent Match Threshold value is set to 80, with an accepted range of 50–100 percent.
Viewing the Results	Review the sample comparison results.
	<b>Note:</b> The matching alleles in the reference sample profile (indicated in <b>bold</b> ) and the comparison sample profile are displayed in blue.
Drafile Companies	
Profile Comparison	
Sample Concordance Sampl	e Comparison Lab Reference Comparison Control/QC Comparison
Percent Match Threshold (Pe	rcent of reference profile alleles detected in the comparison profile)

[	80 Compare Profiles						
	Sample Comparison						
G	Single - Mixed						
			D851179	D21511	D75820	CSF1PO	D3
	B02   ID_Sample_08.fsa   Sample 08 (Reference)	% Match	12, 13	28, 31	7, 12	11, 12	1!
	D02   ID_Sample_06.fsa   Sample 06	96.7%	12, 13	<mark>28</mark> , 30, <mark>31</mark> , 32.2	7, 9, 10, <mark>12</mark>	10, 11, 12	15,
6							
<							>
		Close	Help				
		Close					

From Chapter 4, you know that ID\_Sample\_06 in the Getting Started project is flagged as a potential mixture sample. The sample comparison results indicate that ID\_Sample\_08 may be contributing to the mixed profile of ID\_Sample\_06 because ID\_Sample\_08 (the reference sample) contains 96.7% of the same alleles found in ID\_Sample\_06 (the comparison sample).



# Step 3: Perform Lab Reference Comparison and View Results

Overview	In this section, you will use the Profile Comparison tool to determine
	whether any of the lab reference sample profiles stored in the Profile
	Manager are potential contributors to sample profiles in the project.

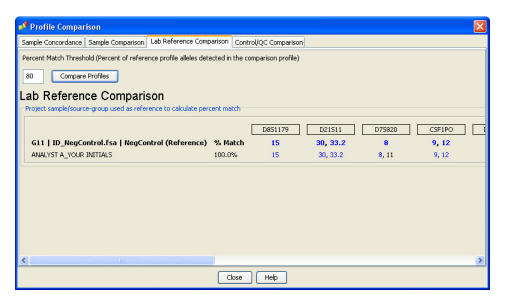
Performing Lab Reference Comparison

- 1. Select the Lab Reference Comparison tab.
- 2. Keep the Percent Match Threshold at 80, then click Compare Profiles.

Viewing the Results

Review the lab reference comparison results.

From Chapter 4, we know that the ID\_Negative Control sample in the Getting Started project contains detected, labeled alleles.



The lab reference comparison results indicate that the Analyst A profile (entered as a lab reference profile in Chapter 2) may be a contributor to the Negative Control sample, because all of the alleles detected in the Negative Control sample profile are found in the Analyst A lab reference profile.



# Step 4: Perform Control/QC Comparison and View Results

**Overview** In this section, you will use the Profile Comparison tool to help perform a blind QC check by comparing the observed profiles for the custom control and QC samples present in the Getting Started project to the custom control profiles stored in the Profile Manager (QC samples use the Custom Control profile type).

Performing Control/QC Comparison

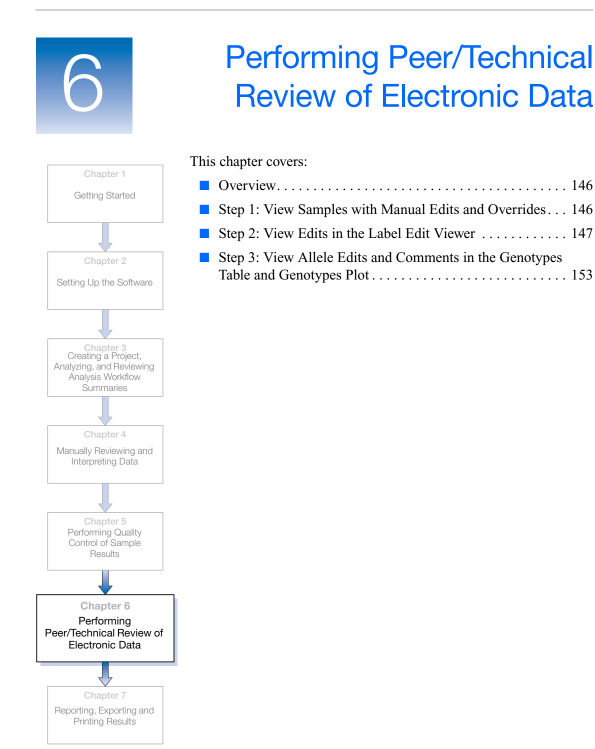
- **1.** Select the **Control/QC Comparison** tab.
- **2.** Keep the Percent Match Threshold at 80, then click **Compare Profiles**.

Viewing the Results Review the control/QC comparison results.

In the Getting Started project, ID\_Sample\_04 is run as a QC sample with profile results expected to match at least one of the custom control profiles stored in the Profile Manager.

🧬 Profile Comparison	×
Sample Concordance Sample Comparison Lab Reference Comparison	Control/QC Comparison
Percent Match Threshold (Percent of reference profile alleles detected in 1 80 Compare Profiles Control/QC Comparison Custom Control profile used as reference to calculate percent match CUSTOM CONTROL_YOUR INITIALS (Reference) % Match	D851179 D21511 D75820 C5F1PO D35
C03   ID_CustomControl.fsa   ID CustomControl 100.0%	13, 14 28, 29 8, 10 10, 11 1
QC SAMPLE 01_YOUR INITIALS (Reference) % Match E01   ID_Sample_04.fsa   Sample 04 100.0%	D851179         D21511         D75820         CSF1PO         D351358           13, 14         30         8, 10         10         14, 15           13, 14         30         8, 10         10         14, 15
	>
	ose Help

The control/QC comparison results indicate that the QC Sample 01 profile (entered as a custom control profile in the Profile Manager in Chapter 2) matches a stored custom control profile as expected (all alleles in ID\_Sample\_04 are also found in the QC Sample 01 profile).





# **Overview**

About Electronic Peer/Technical Review	The GeneMapper <sup>®</sup> <i>ID-X</i> Software provides several data review tools and features designed to improve the efficiency of electronic peer/technical review of projects. These features may minimize the time spent during manual review and eliminate the need to print electropherograms.
In This Chapter	In this chapter, you will use the Getting Started project results edited in Chapter 4 to learn how to:
	• Use table settings to display only those samples containing manual edits or overrides

- Visually confirm manual edits using the Label Edit Viewer
- Export the contents of the Label Edit Viewer
- Use table and plot settings to review manual edits

# Step 1: View Samples with Manual Edits and Overrides

Overview	In this section, you will apply several of the default table settings provided with the software (see Chapter 2) to the Samples table to display only those samples in the Getting Started project that contain manual edits or overrides.
Viewing CGQ Overrides	1. Open the Getting Started project, if not already open (see step 1 on page 75).
	2. In the Project window, select View CGQ Overrides from the

Table Setting drop-down list.



Only those samples in the Getting Started project whose CGQs were manually overridden in Chapter 4 are shown in the Samples table:



If a sample is displayed in this list, it means during the previous interpretation, the analyst manually accepted the sample profile with or without edits.

# Viewing Edited<br/>SamplesIn the Project window, select View Edited Samples from the Table<br/>Setting drop-down list.

Only those samples in the Getting Started project that had at least one allele or artifact label edited in Chapter 4 are shown in the Samples table:



Note that there is a  $\checkmark$  in the Sample Edit (SE) column, and the sample-level PQVs are gray, indicating that peaks within and/or outside of marker ranges have been edited. This table setting allows reviewers to focus on only those samples that have been edited. In an expert system workflow, these may be the only samples that require second review.

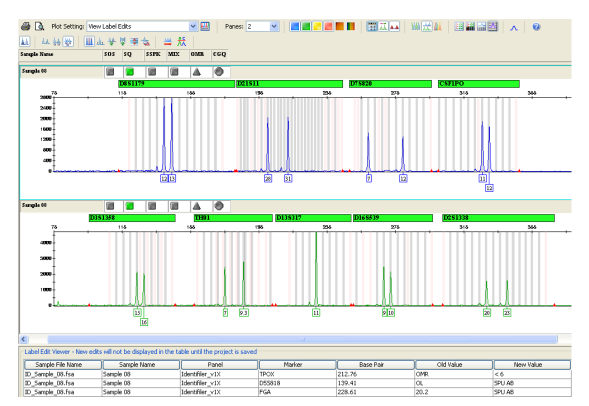
### Step 2: View Edits in the Label Edit Viewer

**Overview** The Label Edit Viewer contains a list of edits made to the allele and artifact labels displayed in the sample electropherogram plots of the Samples plot. In this section, you will use the Label Edit Viewer to assist in the visual confirmation of manual edits you made to the Getting Started project in Chapter 4.

You can view the Label Edit Viewer from the Project window or the Samples plot.

### Viewing Edits from the Samples Plot

- **1.** In the Project window, make sure **View Edited Samples** is selected from the Table Setting drop-down list.
- 2. Select the edited sample in the filtered Samples table, then click (Display Plots).
- **3.** In the Samples plot, select the **View Label Edits** setting from the Plot Setting drop-down list. The Samples plot view changes.



This plot setting displays the sample electropherogram plots above the list of edits shown in the Label Edit Viewer table. In Chapter 4, you made three label edits to Sample 08 in the Getting Started project. The Label Edit Viewer displays detailed information about each of these edits.

**Note:** The Label Edit Viewer is blank if you have not saved the project.

**4.** To visually confirm the displayed peak labels for Sample 08 against the label edit entries in the Label Edit Viewer, select the first row in the table. The corresponding edited peak is highlighted in the electropherogram plot.



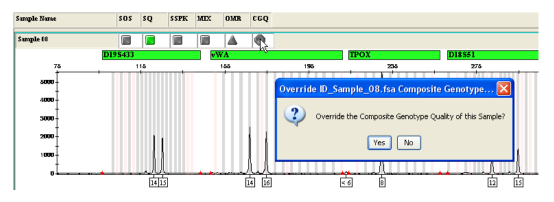
Select row in table

**Note:** Selecting an edited peak in the electropherogram plot will also highlight the corresponding edit in the Label Edit Viewer.

5. Repeat step 4 above for each entry in the Label Edit Viewer.

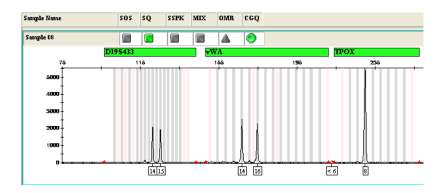
- **6.** After you confirm all label edits, manually accept the sample profile by overriding the CGQ PQV:
  - **a.** Right-click the **()** CGQ in the Samples plot header.

**Note:** A **()** CGQ indicates that one or more peaks within a marker have been edited.



b. Click Yes in the message dialog to override the sample CGQ. The CGQ PQV changes to (Manually Overridden).

**Note:** Overriding CGQ allows the reviewer to manually accept the entire profile for a particular sample and also provides evidence that the sample was visually inspected by the reviewer.





- 7. Select File ➤ Close Plot Window to return to the Project window.
- 8. Click 🕞 (Save Project) to save your changes to the Getting Started project.

#### Viewing Edits from the Project Window

- **1.** In the Project window, make sure **View Edited Samples** is selected from the Table Setting drop-down list.
- Select the edited sample in the filtered Samples table, then click
   (Label Edit Viewer). The Label Edit Viewer opens in a separate window.

💕 Label Edit Viewer 🛛 🔀													
Label Edit Viewer - New edits will not be displayed in the table until the project is saved													
Sample File Name	Sample Name	Panel	Marker	Base Pair	Old Value	New Value	Action	Reason	User	Date			
ID_Sample_08.fsa	Sample 08	Identifiler_v1X	TPOX	212.76	OMR	< 6	modified	Microvariant OMR	Practice User	2007/08/15 12:28:16			
ID_Sample_08.fsa	Sample 08	Identifiler_v1X	D55818	139.41	OL	SPU AB	modified	Pull-up	Practice User	2007/08/15 12:28:16			
ID_Sample_08.fsa	Sample 08	Identifiler_v1X	FGA	228.61	20.2	SPU AB	modified	Pull-up	Practice User	2007/08/15 12:28:16			
L													
				Expo	rt Close	Help							

**3.** Proceed to the next section if you wish to export the contents of the Label Edit Viewer. Otherwise, click **Close** in the Label Edit Viewer to return to the filtered Samples table.

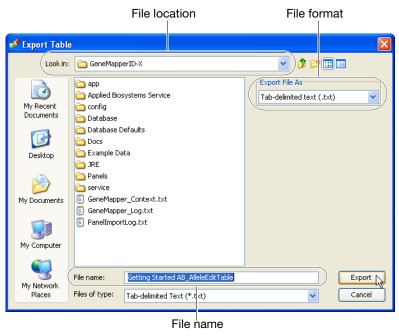
### Exporting the Label Edit Viewer from the Project Window

**Note:** You can also export the contents of the Label Edit Viewer from the Samples plot. See "Exporting from the Samples Plot" on page 168.

- 1. Display the Label Edit Viewer from the Project window (see step 2 above).
- 2. In the Label Edit Viewer, click Export.

6

**3.** In the Export Table dialog, specify a location, name and format (.txt or .csv) for the exported file, then click **Export**.



**Note:** The Files of type selection filters the list of file names displayed in the navigation pane, it does not determine the format of the exported file.

- **4.** Click **Close** in the Label Edit Viewer to return to the filtered Samples table.
- Open the exported file with a spreadsheet software that supports tab-delimited or comma-separated text, such as Microsoft<sup>®</sup> Excel<sup>®</sup>.
- **6.** Print the exported file as needed.



# Step 3: View Allele Edits and Comments in the Genotypes Table and Genotypes Plot

**Overview** In this section, you will use the Genotypes table and Genotypes plot to view the allele edits and comments for the Getting Started project.

Viewing Allele Calls in the Genotypes Table

- **1.** In the Project window, select the **Project** node in the navigation pane, then select the **Genotypes** tab.
  - **2.** Verify that the **View Edited Samples** setting is selected from the Table Setting drop-down list. Only the markers that were edited in the selected sample (in this example, Sample 08) are displayed in the Genotypes table.

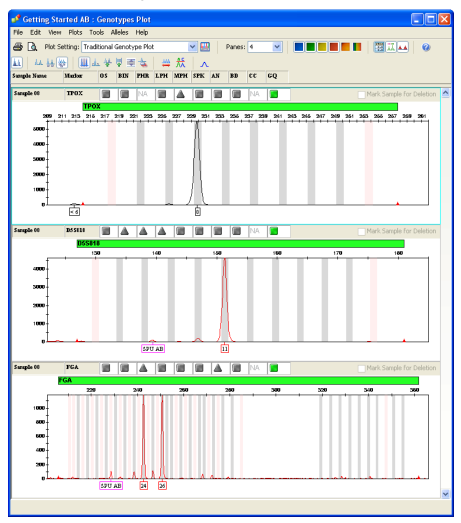
I I I I I I I I I I I I I I I I I I I									<mark>-</mark> 🗖   🔎 🖨 🖪   🔤 🕖									
Sa	Samples Analysis Summary Genotypes																	
	Sample Name	Marker	Allele 1	AE Reason For Change 1	Allele 2	AE Reason For Change 2	MEC	ME	OS	BIN	PHR	LPH	MPH	SPK	AN	BD	CC	GQ
18	3 Sample 08	TPOX		Microvariant OMR	8								▲					
19	Sample 08	D5S818					Pull-up;					▲						
19	2 Sample 08	FGA	24		26		Pull-up;								Δ.			

- **3.** Review the entries in the following table columns:
  - Allele Edit (AE) Reason For Change: Displays the last reason for change entered for an edit that yields an allele label
  - Marker Edit Comment (MEC): Displays the reason for change entered for an edit that yields an artifact label or when alleles are deleted
  - Marker Edit (ME) flag: Displays ✓ (true) if allele or artifact labels are edited within a marker size range

**Note:** You can export the contents of the Genotypes table from the Project window. This procedure is outlined in "Step 4: Export Table Data" on page 165.

Viewing Allele Calls in the Genotypes Plot

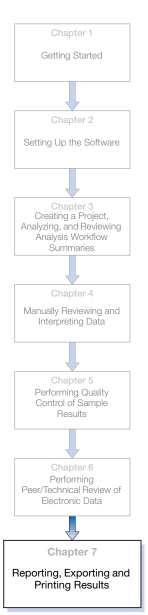
 In the Project window, Shift-click to select all rows in the Genotypes table (if not already selected), then click (Display Plots). The Genotypes plot window opens. **2.** In the Genotypes plot, select the **Traditional Genotype Plot** setting from the Plot Setting drop-down list. The Genotypes plot view changes.



This plot setting displays one marker per pane for each of the markers selected in the Genotypes table. For the Getting Started project, only the markers for Sample 08 are displayed in the Genotypes plot.

3. Select File ➤ Close Plot Window to return to the Project window.

# Reporting, Exporting and Printing Results



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Step 2: Generate the Report 1	60
Step 3: Export the Report	62
Step 4: Export Table Data	65
Step 5: Print Results 1	67
Other Export Options 1	68

# **Overview**

In this chapter, you will learn how to generate custom reports, and export reports and data tables. You will practice these tasks using the Getting Started project you created using the procedures in Chapter 3 of this guide.

**Note:** For information on exporting data in a format suitable for CODIS, see the *GeneMapper*<sup>®</sup> *ID-X Software Help*.

### Step 1: Create a Custom Report Setting

#### Overview

GeneMapper<sup>®</sup> *ID-X* Software can generate table-formatted reports from any combination of the columns in the Samples and Genotypes tables. You can configure the report with custom columns, save the report to the project, print it, or export it as tab-delimited or commaseparated text. You can also save the report settings to generate the same report for other projects.

#### Creating a Custom Report Setting

In this section, you will use the Report Settings Editor to create a new report setting specifically designed for the Getting Started project.

- 1. Launch the software and log in as the Practice User, if not already logged in (see "Starting the Software and Logging In" on page 22).
- **2.** Open the Getting Started project, if not already open (see step 1 on page 75).
- **3.** In the Project window, click 🔚 (GeneMapper ID-X Manager).
- **4.** In the GeneMapper ID-X Manager, select the Report Settings tab, then click **New**. The Report Settings Editor window opens.



- 5. Complete the top portion of the Report Settings Editor:
  - a. Enter Getting Started Report <your initials> as the report Name.
  - **b.** Enter a description for the new report setting.
  - **c.** Verify that the Practice Security Group is selected from the drop-down list.



- 6. Select the Samples table columns to display.
  - **a.** In the Sample tab, Ctrl-click to select the following columns from the Available Columns table.

Sample	Genotype	Custom	
Column(:	5)		
Status			~
Sample Fil	e		
Sample Na	ame		
Sample ID			
Comments	;		
Sample Ty	pe		
Specimen	Category		
Analysis M	lethod		
Panel			
Size Stand	lard		
Custom C	ontrol		
Matrix			
Run Name			
Instrumen			
Instrumen			
Run Date	& Time		
Plate ID			×

b. Click → to add these columns to the Selected Columns table.

**Note:** Columns you add to the Selected Columns table are shown in the Preview Table (at the bottom of the Report Settings Editor).

- 7. Select the Genotypes table columns to display.
  - **a.** In the Genotype tab, Ctrl-click to select the following columns from the Available Columns table.

Sample Genotype Custom	
Column(s)	
Sample File	~
Sample Name	
Sample ID	
Run Name	
Panel	
Marker	
Dye	
Allele 1	
Allele 2	
Size 1	
Size 2	
Height 1	
Height 2	
Peak Area 1	
Peak Area 2	
Data Point 1	
Data Point 2	~

- **b.** Click to add these columns to the Selected Columns table.
- **8.** Add a custom column to the report. The custom column will appear as a column of blank, editable cells in the report table.
  - a. Select the Custom tab, then click Create.
  - b. Enter the following information, then click OK.



c. In the Custom tab, select the **Custom** <**your initials**> column from the Available Columns table, then click → to add this new column to the Selected Columns table.



- 9. Adjust the order that the columns appear in the report.
  - a. In the Selected Columns table, select the **Run Date & Time** column.
  - **b.** Click (Move Up) repeatedly until this column is positioned underneath the Sample Name column.

iample Name Run Date & Time	Sample Name	Image: A start of the start	1 1	
Run Date & Time		<b></b>	Sample	
	Run Date & Time	Image: A start and a start	Sample	
Analysis Method	Analysis Method		Sample	
Panel	Panel	Image: A start of the start	Sample	
5ize Standard	Size Standard	Image: A start of the start	Sample	
Marker	Marker	<ul> <li>Image: A set of the</li></ul>	Genotype	
Allele 1	Allele 1	Image: A start of the start	Genotype	
Allele 2	Allele 2	<ul> <li>Image: A set of the</li></ul>	Genotype	
Custom_your intitials	Custom_your intitials		Custom	
Custom_your intitials	Custom_your intitials		Custom	

- **10.** Adjust the column header names to be displayed in the report.
  - a. In the Display Name column, click the Allele 1 field, then delete this text and enter **Peak 1**.

Column	Display Name
Sample Name	Sample Name
Run Date & Time	Run Date & Time
Analysis Method	Analysis Method
Panel	Panel
Size Standard	Size Standard
Marker	Marker
Allele 1	Peak 1
Allele 2	Allele 2
Custom_your intitials	Custom_your intitials

- b. Replace Allele 2 with Peak 2.
- **11.** In the Number of Alleles field, verify the number of alleles to report is set to **2**.
- **12.** In the Preview Table, review the column display changes you just made to the Getting Started Report setting.

Preview Table									
Sample N	Run Date	Analysis	Panel	Size Stan	Marker	Peak 1	Peak 2	Custom	

7



- **13.** Click **OK** to save your settings and close the Report Settings Editor.
- 14. Click Done to close the GeneMapper ID-X Manager.

## Step 2: Generate the Report

- 1. In the Project window, make sure the Samples tab is selected, then select **View Unedited Samples** from the Table Setting drop-down list.
- 1. Shift-click to select the rows **Sample 01** through **Sample 04** in the Samples table to report.
- 2. Click (Report Manager). A report is displayed in the Report Manager using the first report setting in the drop-down list.

🥩 Керо	rt Manager - *U	ntitled					×
File Edit	: View Tools He	lp					
Report S	etting: Allele Table	~			0		
	Sample File	Sample Name	Marker	Allele 1	Allele 2	Custom_AB	
1	ID_Sample_01.fsa	Sample 01	D851179	14	16		~
2	ID_Sample_01.fsa	Sample 01	D21511	29	31		
3	ID_Sample_01.fsa	Sample 01	D75820	9			
4	ID_Sample_01.fsa	Sample 01	CSF1PO	12			
5	ID_Sample_01.fsa	Sample 01	D351358	14	17		
6	ID_Sample_01.fsa	Sample 01	TH01	6	9		1
7	ID_Sample_01.fsa	Sample 01	D135317	8	11		
8	ID_Sample_01.fsa	Sample 01	D165539	9	12		
9	ID_Sample_01.fsa	Sample 01	D251338	20	25		
10	ID_Sample_01.fsa	Sample 01	D195433	14			
11	ID_Sample_01.fsa	Sample 01	v₩A	14	15		
12	ID_Sample_01.fsa	Sample 01	TPOX	8	10		
13	ID_Sample_01.fsa	Sample 01	D18551	17			
14	ID_Sample_01.fsa	Sample 01	AMEL	Х	Y		
15	ID_Sample_01.fsa	Sample 01	D55818	11			
16	ID_Sample_01.fsa	Sample 01	FGA	23	24		
17	ID_Sample_02.fsa	Sample 02	D851179	14	16		
18	ID_Sample_02.fsa	Sample 02	D21511	29	31		×

**3.** Select the **Getting Started Report <your initials>** setting you created from the Report Setting drop-down list.

7

- 4. Review the report. Note the following:
  - The report contains the columns specified in the report setting, including the Custom column
  - The Allele 1 and Allele 2 columns are reported as Peak 1 and Peak 2

	dit View Tools	Help							
Repor	t Setting: Getting S	Started Report 👻 🧾							
	Sample Name	Run Date & Time	Analysis Method	Panel	Size Standard	Marker	Peak 1	Peak 2	Custom_your intitials
1	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	D851179	14	16	
2	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	D21511	29	31	
3	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	D75820	9		
4	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	CSF1PO	12		
5	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	D351358	14	17	
6	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_G5500	TH01	6	9	
7	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	D135317	8	11	
8	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	D165539	9	12	
9	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	D251338	20	25	
10	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	D195433	14		
11	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	v₩A	14	15	
12	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	TPOX	8	10	
13	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	D18551	17		
14	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_G5500	AMEL	X	Y	
15	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_G5500	D55818	11		
16	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	FGA	23	24	
17	Sample 02	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	D851179	14	16	
18	Sample 02	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	D21511	29	31	
19	Sample 02	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	D75820	9		
20	Sample 02	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	CSF1PO	12		
21	Sample 02	2006-08-08 14:54:11.0	Getting Started your initials	Identifiler v1X	CE G5 HID G5500	D351358	14	17	

**5.** To edit a cell in the Custom column, double-click the cell, type the desired text, then press **Enter**.

	Sample Name	Run Date & Time	Analysis Method	Panel	Size Standard	Marker	Peak 1	Peak 2	Custom_your initials
1	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	D851179	14	16	enter text here
2	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	D21511	29	31	
3	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	D75820	9		
4	Sample 01	2006-08-08 14:54:11.0	Getting Started_your initials	Identifiler_v1X	CE_G5_HID_GS500	CSF1PO	12		

6. Leave the Report Manager window open for the next steps (see "Step 3: Export the Report" on page 162).

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# Step 3: Export the Report

- **Overview** You can export the report generated in the Report Manager as a formatted table. The exported table format depends on the report setting selected when generating the report. By default, the export table format reflects the display in the Report Manager window. However, you can also export in allele table format to list markers in columns or rows, and to group the called alleles for a marker into one or more cells, depending on the number of alleles selected to be displayed.
- Exporting the Report
- **1.** In the Report Manager, select **Allele Table** from the Report Setting drop-down list.

**Note:** You can use this report setting to export genotype data in a traditional allele table format (see step 6 on page 163).

2. Select File > Export to open the Export Report dialog box:

🧬 Export Repo	ort	×
Look in:	💼 GeneMapperID-X	
My Recent Documents	app     Applied Biosystems Service     config     Otabase     Database	Export File As Tab-delimited text (.txt)
Desktop	Docs     Example Data     JRE     Panels     service	Convert to Allele Table By Sample Format     Group alleles in one cell     Separate alleles into individual cells
My Documents	GeneMapper_Context.txt GeneMapper_Log.txt FaneIImportLog.txt	
My Network Places	File name:     Getting Started AB_A       Files of type:     Tab- and Comma-deline	Ilele Table Export nited Text (*.txt,*.csv) Cancel

**3.** Navigate to the location in which to save the file.



**4.** Enter **Export <your initials>** as a suffix to the default Getting Started <your initials>\_Allele Table export File name.



**Note:** The Files of type selection filters the list of file names displayed in the navigation pane, it does not determine the format of the exported file.

5. Verify that **Tab-delimited text (\*.txt)** is selected as the export file format.

**Note:** This selection separates the alleles with tab-stops, and allows you to open the exported file using a spreadsheet application (see step 11 on page 164).

6. Select a format option:



• Apply Selected Report Setting Format (default) – Exports the report as it is displayed in the Report Manager (one marker per row)

	1	2	3	4	5
1	Sample File	Sample Name	Marker	Allele 1	Allele 2
2	ID_Sample_01.fsa	Sample 01	D8S1179	14	16
3	ID_Sample_01.fsa	Sample 01	D21S11	29	31
4	ID_Sample_01.fsa	Sample 01	D7S820	9	
5	ID_Sample_01.fsa	Sample 01	CSF1PO	12	

• Convert to Allele Table by Sample Format – Generates a list of samples (one sample per row) with column(s) for each marker that lists the called alleles for the marker

Note: Markers without called alleles are also listed.



For heterozygote alleles, you can also choose to:

• **Group alleles in one cell** – Generates a single column for each marker and places all alleles for the marker into one cell (that is, a traditional allele table)

	1	2	3	4	5
1	Sample File	Sample Name	D8S1179	D21S11	D7S820
2	ID_Sample_01.fsa	Sample 01	14, 16	29,31	9,
3	ID_Sample_02.fsa	Sample 02	14, 16	29,31	9,
4	ID_Sample_03.fsa	Sample 03	15, 16	31.2,	11, 12
5	ID_Sample_04.fsa	Sample 04	13, 14	30,	8,10

• Separate alleles into individual cells – Generates multiple columns per marker and places only one allele per cell

	1	2	3	4	5	6
1	Sample File	Sample Name	D8S1179 1	D8S1179 2	D21S111	D21S11 2
2	ID_Sample_01.fsa	Sample 01	14	16	29	31
3	ID_Sample_02.fsa	Sample 02	14	16	29	31
4	ID_Sample_03.fsa	Sample 03	15	16	31.2	
5	ID_Sample_04.fsa	Sample 04	13	14	30	

### 7. Click Export.

**Note:** If you are exporting an allele table, a dialog box displays the names of the files created. Click **OK**.

- 8. Select File > Exit to close the Report Manager.
- Click Yes in the Save Report dialog, then enter Getting Started\_Allele Table Report <your initials> in the Save dialog.
- **10.** Click **OK** to save the Allele Table Report to the Getting Started project.
- Navigate to and open the exported Getting Started\_Allele Table Export file in a spreadsheet software (such as Microsoft<sup>®</sup> Excel<sup>®</sup>) or Microsoft<sup>®</sup> Notepad.



## Step 4: Export Table Data

**IMPORTANT!** When exporting or copying and pasting PQV flags, include the ME column (from the Genotypes table) and the SE column (from the Samples table) to indicate whether allele labels were edited, which may affect individual PQV values.

Exporting Individual Tables When you export data from the Samples or Genotypes table individually, the following are exported in a single table:

- All samples in the project (not selected samples)
- Only the displayed columns from the individual table that you are viewing

#### To export individual tables:

- In the Project window, select View > Samples, then select the Samples tab to export the Samples table or the Genotypes tab to export the Genotypes table.
- **2.** Select **View Unedited Samples** from the Table Setting dropdown list.
- **3.** Shift-click a column header in the Samples tab (or Genotypes tab) to sort the table and determine the sample order.
- 4. Click 🖻 (Export Table).
- **5.** In the dialog box, navigate to the location to save the exported table file.
- 6. Select the export file format:
  - Tab-delimited text (\*.txt)
  - Comma-delimited values (\*.csv)

**7.** Enter a name for the exported file.

**Note:** The Files of type selection filters the list of file names displayed in the navigation pane, it does not determine the format of the exported file.

8. Click Export Table.

#### Exporting Combined Tables

When you export data from the Samples table and Genotypes table together, the following are exported in a combined table:

- All samples in the project (not selected samples)
- Only the displayed columns from the Samples table and Genotypes table

#### To export combined tables:

- **1.** In the Project window, select **View Unedited Samples** from the Table Setting drop-down list.
- 2. Select File > Export Combined Table.
- **3.** In the dialog box, navigate to the location to save the exported table file.
- 4. Select the export file format (\*.txt or \*.csv).
- 5. Select a Merge option:
  - Allele table by sample

**IMPORTANT!** This option does not allow you to control the order of columns in the exported file. Columns are exported in the order in which they appear in the Samples and Genotypes tables. To control the order of columns in the exported file, export from the Report Manager. See "Step 3: Export the Report" on page 162.

- One line per sample
- One line per marker (default)



- **6.** Enter a name for the exported file.
- 7. Click Export.

**Note:** If you are exporting an allele table, a dialog box displays the names of the files created. Click **OK**.

**Copying and** You can copy and paste the content of most tables into a spreadsheet or text file.

- 1. In the desired table of the GeneMapper<sup>®</sup> *ID-X* Software, select the cells to copy.
- **2.** To copy the data:
  - a. Without column headers Press Ctrl+C.
  - b. With column headers Press Ctrl+Shift+C.
- In the desired location (spreadsheet or text file), select
   Edit ▶ Paste.

### Step 5: Print Results

Data

Use the print preview function in the GeneMapper<sup>®</sup> *ID-X* Software to examine data items (reports, tables, plots, sample information, raw data, and EPT data) on screen before they print.

When you are satisfied with the results, select **File** > **Print**:

Window/Tab	Access From Project Window By Selecting
Project window – Samples tab	View ► Samples
Project window – Genotypes tab	View ► Genotypes
Project window – Info tab	View ► Sample Info
Project window – Raw Data tab	View ► Raw Data
Project window – EPT Data tab	View ► EPT Data

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Window/Tab	Access From Project Window By Selecting
Samples plot	The Samples tab, then View ▶ Display Plots
Genotypes plot	The Genotypes tab, then View ▸ Display Plots
Report Manager	Tools ▶ Report Manager
Size Match Editor	Tools ► Size Match Editor

## **Other Export Options**

# Exporting from the Samples Plot

You can export results from the following tables by selecting **File > Export Table** in the Samples plot:

Table	Access From Samples Plot By Selecting
Genotypes table	Plots > Tables > Genotypes Table
Label Edit Viewer	Plots > Tables > Label Edit Viewer
Sizing table	Plots > Tables > Sizing Table

#### Exporting from the GeneMapper ID-X Manager

You can export the following data objects stored in the GeneMapper *ID-X* Software database by clicking **Export...** in the GeneMapper ID-X Manager dialog:

- Projects
- Analysis Methods
- Table Settings
- Plot Settings
- Matrices

**Note:** You can export a matrix file for use in the Data Collection Software on the ABI PRISM<sup>®</sup> 310 Genetic Analyzer.

- Size Standards
- Report Settings



Exporting from<br/>the Panel<br/>ManagerYou can export the following data objects stored in the GeneMapper<br/>ID-X Software database by selecting File ▶ Export... in the Panel<br/>Manager window:

- Panels
- Bin Set
- All Kits
- Marker Stutter

#### Exporting from the Profile Manager

You can export the following data objects stored in the GeneMapper *ID-X* Software database by clicking **Export** in the Profile Manager dialog:

- Lab Reference profiles
- Custom Control profiles

**Note:** All of the profiles stored in the Profile Manager will be exported together in a single file.

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**Chapter 7** Reporting, Exporting and Printing Results *Other Export Options* 

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