AmpFℓSTR[™] Identifiler[™] Plus PCR Amplification Kit USER GUIDE

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Revision	Date	Description
Н	19 September 2018	Updated branding and trademarks, no technical changes.
G	16 May 2015	Added the 3500/3500xL Genetic Analyzer and theGeneScan [™] 600 LIZ [™] Size Standard v2.0.
F	February 2015	Add information for the ProFlex [™] PCR System.
E	August 2014	Add information about 50-, 100-, and 1,000-reaction kit sizes.
D	March 2012	Update legal information.
С	April 2011	Remove patent numbers.
В	September 2010	Remove patent numbers.
A	September 2008	New document.

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Preface

Purpose

The Applied BiosystemsTM AmpF ℓ STRTM IdentifilerTM Plus PCR Amplification Kit User Guide provides information about the Applied BiosystemsTM instruments, chemistries, and software associated with the AmpF ℓ STRTM IdentifilerTM Plus PCR Amplification Kit.

Pull-out chapters

This guide is designed to allow users to pull out chapters 2, 3, and 4. The pull-out chapters have title and back pages, which indicate the chapter number and title.

This chapter covers:

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Product overview

Purpose	The AmpFℓSTR [™] Identifiler [™] Plus PCR Amplification Kit is a short tandem repeat (STR) multiplex assay that amplifies 15 tetranucleotide repeat loci and the Amelogenin gender-determining marker in a single PCR amplification:
	• All thirteen of the required loci for the Combined DNA Index System (CODIS) loci are included in this kit for known-offender databasing in the United States (Budowle <i>et al.</i> , 1998).
	 Two additional loci, D2S1338 and D19S433, are included. These loci are consistent with the AmpFℓSTR SGM Plus[™] PCR Amplification Kit.
	• The combination of the 15 loci are consistent with several worldwide database recommendations.
	The AmpF <i>l</i> STR Identifiler Plus Kit delivers a 16-locus multiplex with the same power of discrimination as, better sensitivity than, and better robustness than the earlier generation of the AmpF <i>l</i> STR Identifiler Kit. The kit uses modified PCR cycling conditions for enhanced sensitivity, a new buffer formulation to improve performance with inhibited samples, and an improved process for DNA synthesis and purification of the amplification primers to deliver a much cleaner electrophoretic background.
Product description	The AmpFℓSTR Identifiler Plus Kit uses the same primer sequences as the earlier generation AmpFℓSTR [™] Identifiler [™] Kit. The AmpFℓSTR Identifiler Plus Kit contains all the necessary reagents for the amplification of human genomic DNA.
	The reagents are designed for use with the following Applied Biosystems [™] instruments:
	• Applied Biosystems [™] 3100/3100-Avant Genetic Analyzer
	• Applied Biosystems [™] 3130/3130 <i>xl</i> Genetic Analyzer
	• Applied Biosystems [™] 310 Genetic Analyzer
	• GeneAmp [™] PCR System 9700 with the Silver 96-Well Block
	 GeneAmp[™] PCR System 9700 with the Gold-plated Silver 96-Well Block Veriti[™] 96-Well Thermal Cycler
	• ProFlex [™] PCR System
About the primers	The AmpF <i>l</i> STR Identifiler Plus Kit employs the latest improvements in primer synthesis and purification techniques to minimize the presence of dye-labeled artifacts. These improvements result in a much cleaner electropherogram background that enhances the assay's signal-to-noise ratio and simplifies the interpretation of results.
	Non-nucleotide linkers are used in primer synthesis for the following loci: CSF1PO, D13S317, D16S539, D2S1338, and TPOX. For these primers, non-nucleotide linkers are placed between the primers and the fluorescent dye during oligonucleotide synthesis (Butler, 2005, Grossman <i>et al.</i> , 1994, and Baron <i>et al.</i> , 1996). Non-nucleotide linkers enable reproducible positioning of the alleles to facilitate

interlocus spacing. The combination of a five-dye fluorescent system and the inclusion of non-nucleotide linkers allows for simultaneous amplification and efficient separation of the 15 STR loci and Amelogenin during automated DNA fragment analysis.

Loci amplified by the kit The following table shows the loci amplified, their chromosomal locations, and the corresponding fluorescent marker dyes. The AmpFℓSTR[™] Identifiler[™] Plus Allelic Ladder is used to genotype the analyzed samples. The alleles contained in the allelic ladder, and the genotype of the AmpFℓSTR[™] Identifiler[™] Plus Control DNA 9947A are also listed in the table.

Locus designation	Chromosome location	Alleles included in Identifiler [™] Plus Allelic Ladder	Dye label	Control DNA 9947A
D8S1179	8	8, 9 10, 11, 12, 13, 14, 15, 16, 17, 18, 19	6-FAM [™]	13 [‡]
D21S11	21q11.2-q21	24, 24.2, 25, 26, 27, 28, 28.2, 29, 29.2, 30, 30.2, 31, 31.2, 32, 32.2, 33, 33.2, 34, 34.2, 35, 35.2, 36, 37, 38		30 [§]
D7S820	7q11.21-22	6, 7, 8, 9, 10, 11, 12, 13, 14, 15	-	10, 11
CSF1PO	5q33.3-34	6, 7, 8, 9, 10, 11, 12, 13, 14, 15	-	10, 12
D3S1358	Зр	12, 13, 14, 15, 16, 17, 18, 19	VIC™	14, 15
TH01	11p15.5	4, 5, 6, 7, 8, 9, 9.3, 10, 11, 13.3	-	8, 9.3
D13S317	13q22-31	8, 9, 10, 11, 12, 13, 14, 15	-	11#
D16S539	16q24-qter	5, 8, 9, 10, 11, 12,13, 14, 15	-	11, 12
D2S1338	2q35-37.1	15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28	-	19, 23
D19S433	19q12-13.1	9, 10, 11, 12, 12.2, 13, 13.2, 14, 14.2, 15, 15.2, 16, 16.2, 17, 17.2	NED™	14, 15
vWA	12p12-pter	11,12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24	-	17, 18
TPOX	2p23-2per	6, 7, 8, 9, 10, 11, 12, 13	-	8 ^{‡‡}
D18S51	18q21.3	7, 9, 10, 10.2, 11, 12, 13, 13.2, 14, 14.2, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27	-	15, 19
Amelogenin	X: p22.1-22.3	Х, Ү	PET™	Х
	Y: p11.2			
D5S818	5q21-31	7, 8, 9, 10, 11, 12, 13, 14, 15, 16		11 ^{§§}
FGA	4q28	17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 26.2, 27, 28, 29, 30, 30.2, 31.2, 32.2, 33.2, 42.2, 43.2, 44.2, 45.2, 46.2, 47.2, 48.2, 50.2, 51.2		23, 24

Table 1	AmpFℓSTR [™]	[®] Identifiler [™]	" Plus Kit	loci and	alleles
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‡ For CODIS purposes, profile reported as 13, 13.

§ For CODIS purposes, profile reported as 30, 30.

For CODIS purposes, profile reported as 11, 11.

‡‡ For CODIS purposes, profile reported as 8, 8.

§§ For CODIS purposes, profile reported as 11, 11.

Allelic ladder
profileFigure 1 shows the allelic ladder for the AmpF/STR Identifiler Plus Kit. See "Allelic
ladder requirements" on page 36 for information on ensuring accurate genotyping.



Figure 1 GeneMapper[™] *ID-X* Software plot of the AmpF/STR[™] Identifiler[™] Plus Kit Allelic Ladder

Control DNAFigure 2 shows amplification of Control DNA 9947A using the AmpF/STR9947A profileIdentifiler Plus Kit.



Figure 2 1 ng of Control DNA 9947A amplified with the AmpFℓSTR[™] Identifiler[™] Plus Kit and analyzed on the Applied Biosystems[™] 3130*x*/ Genetic Analyzer

Workflow overview



Instrument and software overview

This section provides information about the Data Collection Software versions required to run the AmpF ℓ STRTM IdentifilerTM Plus PCR Amplification Kit on specific instruments.

Data Collection and GeneMapper[™] *ID* or *ID-X* Software

Instrument and

software compatibility

The Data Collection Software provides instructions to firmware running on the instrument and displays instrument status and raw data in real time. As the instrument measures sample fluorescence with its detection system, the Data Collection Software collects the data and stores it. The Data Collection Software stores information about each sample in a sample file (.fsa), which is then analyzed by the GeneMapperTM *ID* or *ID-X* Software.

Instrument	Data Collection Software	Analysis software
3500/3500xL	3500 Series Data Collection Software v1.0 or later	GeneMapper [™] <i>ID-X</i> Software v1.2 or later
3130/3130 <i>x</i> /‡	3.0 or later	GeneMapper [™] ID Software v2.2.1
3100/3100-	1.1 (3100)	and
Avant	1.0 (3100-Avant)	 GeneMapper[™] ID-X
	2.0	Software v1.0.1 or later
310	3.1	1
	3.0	

‡ We conducted validation studies for the AmpFℓSTR[™] Identifiler[™] Plus Kit using this configuration.

About multicomponent analysis

Applied Biosystems[™] fluorescent multi-color dye technology allows the analysis of multiple loci, including loci that have alleles with overlapping size ranges. Alleles for overlapping loci are distinguished by labeling locus-specific primers with different colored dyes.

Multicomponent analysis is the process that separates the five different fluorescent dye colors into distinct spectral components. The four dyes used in the AmpFℓSTR[™] Identifiler[™] Plus PCR Amplification Kit to label samples are 6-FAM[™], VIC[™], NED[™], and PET[™] dyes. The fifth dye, LIZ[™] dye, is used to label the GeneScan[™] 500 LIZ[™] Size Standard.

How multicomponent analysis works

Each of these fluorescent dyes emits its maximum fluorescence at a different wavelength. During data collection on the Applied BiosystemsTM and ABI PRISMTM instruments, the fluorescence signals are separated by diffraction grating according to their wavelengths and projected onto a charge-coupled device (CCD) camera in a predictably spaced pattern. The 6-FAMTM dye emits at the shortest wavelength and it is displayed as blue, followed by the VICTM dye (green), NEDTM dye (yellow), PETTM dye (red), and LIZTM dye (orange).

Although each of these dyes emits its maximum fluorescence at a different wavelength, there is some overlap in the emission spectra between the dyes (Figure 3). The goal of multicomponent analysis is to correct for spectral overlap.



Figure 3 Emission spectra of the five dyes used in the AmpFℓSTR[™] Identifiler[™] Plus Kit

Materials and equipment

Kit contents and
storageThe AmpFℓSTR[™] Identifiler[™] Plus PCR Amplification Kit contains materials
sufficient to perform 50, 100, 200, or 1000 amplifications at 25 μL reaction volumes.

IMPORTANT! The fluorescent dyes attached to the primers are light sensitive. Protect the primer set from light when not in use. Amplified DNA, AmpFℓSTR[™] Identifiler[™] Plus Allelic Ladder, and GeneScan[™] 500 LIZ[™] Size Standard should also be protected from light. Keep freeze-thaw cycles to a minimum.

Component	Description	50× Volume (A26182)	100× Volume (A26364)	200X Volume (4427368)	1000X Volume (4486467)	Storage
AmpF ℓ STR [™] Identifiler [™] Plus Master Mix	Contains enzyme, salts, dNTPs, carrier protein, and 0.04% sodium azide.	1 tube, 0.5 mL	2 tubes, 0.5 mL/tube	2 tubes, 1.0 mL/tube	10 tubes, 1.0 mL/tube	-15 to -25°C on receipt, 2 to 8 °C after initial use
AmpF ℓ STR [™] Control DNA 9947A	Contains 0.10 ng/µL human female 9947A DNA in 0.05% sodium azide and buffer [‡] .	1 tube, 0.3 mL	1 tube, 0.3 mL	1 tube, 0.3 mL	5 tubes, 0.3 mL/tube	
	See Table 1 on page 11 for profile.					
AmpF ℓ STR [™] Identifiler [™] Plus Primer Set	Contains forward and reverse primers to amplify human DNA targets.	1 tube, 0.25 mL	2 tubes, 0.25 mL/tub e	1 tube, 1.0 mL	5 tubes, 1.0 mL/tube	-15 to -25°C on receipt, 2 to 8 °C after initial use.
AmpF / STR [™] Identifiler [™] Plus Allelic Ladder	Contains amplified alleles. See Table 1 on page 11 for a list of alleles included in the allelic ladder.	1 tube, 0.025 mL	1 tube, 0.025 mL	1 tube, 0.05 mL	5 tubes, 0.05 mL/tube	Store Protected from light.

‡ The AmpFℓSTR[™] Control DNA 9947A is included at a concentration appropriate to its intended use as an amplification control (i.e., to

provide confirmation of the capability of the kit reagents to generate a profile of expected genotype). The AmpFℓSTR Control DNA 9947A is not designed to be used as a DNA quantitation control, and laboratories may expect to see variation from the labelled concentration when quantitating aliquots of the AmpFℓSTR[™] Control DNA 9947A.

Standards for
samplesFor the AmpFlSTR Identifiler Plus Kit, the panel of standards needed for PCR
amplification, PCR product sizing, and genotyping are:

- Control DNA 9947A A positive control for evaluating the efficiency of the amplification step and STR genotyping using the AmpFℓSTR[™] Identifiler[™] Plus Allelic Ladder.
- GeneScan[™] 500 LIZ[™] Size Standard Standard used for obtaining sizing results. It contains 16 single-stranded labeled fragments of: 35, 50, 75, 100, 139, 150, 160, 200, 250, 300, 340, 350, 400, 450, 490 and 500 nucleotides. This standard, which has been evaluated as an internal lane size standard, yields precise sizing results for AmpFℓSTR Identifiler Plus Kit PCR products. Order the GeneScan 500 LIZ Size Standard (PN 4322682) separately.
- AmpFℓSTR[™] Identifiler[™] Plus Allelic Ladder Allelic ladder for accurate characterization of the alleles amplified by the AmpFℓSTR Identifiler Plus Kit. The AmpFℓSTR[™] Identifiler[™] Plus Allelic Ladder contains most of the alleles reported for the 15 autosomal loci. Refer to Table 1 on page 11 for a list of the alleles included in the AmpFℓSTR[™] Identifiler[™] Plus Allelic Ladder.

Equipment and
materials not
includedTables 2 and 3 list required and optional equipment and materials not supplied with
the AmpFtSTR Identifiler Plus Kit. Unless otherwise noted, many of the items are
available from major laboratory suppliers (MLS).

Table 2 Equipment

Equipment	Source
ABI PRISM [™] 3100/3100-Avant Genetic Analyzer	Contact your local sales representative
Applied Biosystems [™] 3130/3130 <i>xI</i> Genetic Analyzer	
Applied Biosystems [™] 310 Genetic Analyzer	
GeneAmp [™] PCR System 9700 with the Silver 96-Well Block	N8050001
GeneAmp [™] PCR System 9700 with the Gold-plated Silver 96-Well Block	4314878
ProFlex [™] PCR System	4484075
Veriti [™] 96-Well Thermal Cycler	4375786
Silver 96-Well Sample Block	N8050251
Gold-plated Silver 96-Well Sample Block	4314443
Tabletop centrifuge with 96-Well Plate Adapters (optional)	MLS

Table 3 User-supplied materials[‡]

Item	Source
50-reaction AmpFℓSTR [™] Identifiler [™] Plus PCR Amplification Kit	A26182
100-reaction AmpFℓSTR [™] Identifiler [™] Plus PCR Amplification Kit	A26364
200-reaction AmpF / STR [™] Identifiler [™] Plus PCR Amplification Kit	4427368
1000-reaction AmpFℓSTR [™] Identifiler [™] Plus PCR Amplification Kit	4486467
3100/3100-Avant Genetic Analyzer materials	
96-Well Plate Septa	4315933
Reservoir Septa	4315932
3100/3100-Avant Genetic Analyzer Capillary Array, 36-cm	4333464
POP-4 [™] Polymer for 3100/3100-Avant Genetic Analyzers	4316355
3100/3100-Avant Genetic Analyzer Autosampler Plate Kit, 96-well	4316471
GeneScan [™] 500 LIZ [™] Size Standard	4322682
Running Buffer, 10×	402824
DS-33 Matrix Standard Kit (Dye Set G5)	4345833
MicroAmp [™] Optical 96-Well Reaction Plate	N8010560

Table 3	User-supplied materials [‡]	(continued))
Tuble 0	ocor ouppriou materialo	continuou	,

Item	Source	
250-μL Glass Syringe (array-fill syringe)	4304470	
5.0-mL Glass Syringe (polymer-reserve syringe)	628-3731	
For a complete list of parts and accessories for the 3100 instrument, refer to Appendix B of the A 3100 Genetic Analyzer and 3100-Avant Genetic Analyzer User Reference Guide (PN 4335393).	BI PRISM [™]	
3130/3130x/ Analyzer materials		
96-Well Plate Septa	4315933	
Reservoir Septa	4315932	
3100/3130x/ Genetic Analyzer Capillary Array, 36-cm	4315931	
POP-4 [™] Polymer for 3130/3130x/ Genetic Analyzers	4352755	
3130/3130x/ Genetic Analyzer Autosampler Plate Kit, 96-well	4316471	
GeneScan [™] 500 LIZ [™] Size Standard	4322682	
Running Buffer, 10×	402824	
DS-33 Matrix Standard Kit (Dye Set G5)	4345833	
MicroAmp [™] Optical 96-Well Reaction Plate	N8010560	
For a complete list of parts and accessories for the 3130 <i>xl</i> instrument, refer to Appendix A of the <i>Applied</i> Biosystems [™] 3130/3130 <i>xl</i> Genetic Analyzers Maintenance, Troubleshooting, and Reference Guide (PN 4352716).		
310 Analyzer materials		
310 DNA Analyzer Capillary Array, 47-cm	402839	
0.5 mL Sample Tray	5572	
96-Well Tray Adaptor (for 9700 thermal cycler trays)	4305051	
GeneScan [™] 500 LIZ [™] Size Standard	4322682	
Running Buffer, 10×	4335643	
Genetic Analyzer Septa Retainer Clips for 96-Tube Sample Tray	402866	
Genetic Analysis Sample Tubes (0.5-mL)	401957	
Septa for 0.5-mL Sample Tubes	401956	
DS-33 Matrix Standard Set (6-FAM [™] , VIC [™] , NED [™] , PET [™] , and LIZ [™] dyes) for ABI PRISM [™] 310/377 systems	4318159	
MicroAmp [™] 8-Tube Strip, 0.2-mL	N8010580	
MicroAmp [™] 96-Well Base (holds 0.2-mL reaction tubes)	N8010531	
MicroAmp [™] 96-Well Full Plate Cover	N8010550	

Table 3	User-supplied	materials [‡]	(continued)

Item	Source	
MicroAmp [™] 96-Well Tray/Retainer Set	403081	
POP-4 [™] Polymer for the 310 Genetic Analyzer	402838	
For a complete list of parts and accessories for the 310 instrument, refer to Appendix B of the ABI PRISM [™] 310 Genetic Analyzer User Guide (PN 4317588).		
PCR Amplification		
MicroAmp [™] 96-Well Tray	N8010541	
MicroAmp [™] Reaction Tube with Cap, 0.2-mL	N8010540	
MicroAmp [™] 8-Tube Strip, 0.2-mL	N8010580	
MicroAmp [™] 8-Cap Strip	N8010535	
MicroAmp [™] 96-Well Tray/Retainer Set	403081	
MicroAmp [™] 96-Well Base	N8010531	
MicroAmp [™] Clear Adhesive Film	4306311	
MicroAmp [™] Optical Adhesive Film	4311971	
MicroAmp [™] Optical 96-Well Reaction Plate	N8010560	
Other user-supplied materials		
Hi-Di [™] Formamide, 25-mL	4311320	
Aerosol resistant pipette tips ML		
Microcentrifuge tubes	MLS	
Pipettors	MLS	
Tape, labeling	MLS	
Tube, 50-mL Falcon	MLS	
Tube decapper, autoclavable	MLS	
Deionized water, PCR grade	MLS	
Tris-HCl, pH 8.0	MLS	
EDTA, 0.5 M	MLS	
Vortex	MLS	

‡ For the Material Safety Data Sheet (MSDS) of any chemical not distributed by Applied Biosystems[™], contact the chemical manufacturer. Before handling any chemicals, refer to the MSDS provided by the manufacturer, and observe all relevant precautions.

Chapter 2

PCR Amplification

AmpFtSTR[™] Identifiler[™] Plus User Guide

AmpFℓSTR[™] Identifiler[™] Plus User Guide

This chapter covers:

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DNA quantification	25
Prepare the amplification kit reactions	28
Perform PCR	30

PCR work areas

Work area setup	Many resources are available for the appropriate design of a PCR laboratory:		
and lab design	 If you are using the AmpFℓSTR[™] Identifiler[™] Plus PCR Amplification Kit for forensic DNA testing, refer to <i>Forensic Laboratories: Handbook for Facility Planning, Design, Construction and Moving</i>, National Institute of Justice, 1998 (http://nij.ncjrs.gov/publications/pubs_db.asp). 		
	• If you are using the AmpFℓSTR [™] Identifiler [™] Plus Kit for parentage DNA testing, refer to the <i>Guidance for Standards for Parentage Relationship Testing Laboratories, American Association of Blood Banks</i> , 7th edition, 2004.		
	The sensitivity of the AmpFℓSTR [™] Identifiler [™] Plus Kit (and other PCR-based tests) enables amplification of minute quantities of DNA, necessitating precautions to avoid contamination of samples yet to be amplified (Kwok and Higuchi, 1989).		
	To prevent contamination by human DNA, be careful while handling and processing samples. Wear gloves at all times and change them frequently. Close sample tubes when not in use. Limit aerosol dispersal by handling sample tubes and reagents carefully.		
	Note: These laboratory design resources and guidances constitute only a sample of the precautions that need to be observed when using PCR technology. Refer to your laboratory's internal policies and procedures for additional information and references.		
PCR-setup tools	IMPORTANT! These items should never leave the PCR setup work area.		
	Calculator		
	Gloves, disposable		
	• Marker pen, permanent		
	Microcentrifuge		
	• Microcentrifuge tubes, 1.5-mL, or 2.0-mL, or other appropriate clean tube (for Master Mix preparation)		
	Microcentrifuge tube rack		
	Pipette tips, sterile, disposable hydrophobic filter-plugged		
	• Pipettors		
	Tube decapper, autoclavable		
	• Vortex		
Amplified DNA	The following PCR systems should be placed in the amplified DNA work area.		
work area tools	• GeneAmp [™] PCR System 9700 with the Silver 96-Well Block		
	• GeneAmp [™] PCR System 9700 with the Gold-plated Silver 96-Well Block		
	• Veriti [™] 96-Well Thermal Cycler		
	• ProFlex [™] PCR System		

Required user-supplied reagents

In addition to the AmpFℓSTR[™] Identifiler[™] Plus Kit reagents, the use of low TE buffer (10 mM Tris, 0.1 mM EDTA, pH 8.0) is recommended. You can prepare the buffer as described in the procedure below or order it from Teknova (Cat # T0223).

To prepare low TE buffer:

- 1. Mix together:
 - 10 mL of 1 M Tris-HCl, pH 8.0
 - 0.2 mL of 0.5 M EDTA, pH 8.0
 - 990 mL glass-distilled or deionized water

Note: Adjust the volumes accordingly for specific needs.

- 2. Aliquot and autoclave the solutions.
- 3. Store at room temperature.

DNA quantification

Importance of quantification

Quantifying the amount of DNA in a sample before amplification allows you to determine whether or not sufficient DNA is present to permit amplification and to calculate the optimum amount of DNA to add to the reaction. The optimum amount of DNA for the AmpFℓSTR[™] Identifiler[™] Plus Kit is 1.0 ng in a maximum input volume of 10 μ L for 28 PCR cycles and 0.5 ng in a maximum input volume of 10 μ L for 29 PCR cycles.

If too much DNA is added to the PCR reaction, then the increased amount of PCR product that is generated can result in:

- Fluorescence intensity that exceeds the linear dynamic range for detection by the instrument ("off-scale" data). Off-scale data are problematic because:
 - Quantitation (peak height and area) for off-scale peaks is not accurate. For example, an allele peak that is off-scale can cause the corresponding stutter peak to appear higher in relative intensity, thus increasing the calculated percent stutter.
 - Multicomponent analysis of off-scale data is not accurate, and it results in poor spectral separation ("pull-up").
- Incomplete A-nucleotide addition.

When the total number of allele copies added to the PCR is extremely low, allelic dropout can occur resulting in a partial profile.

Methods of quantifying DNA

We provide several kits for quantifying DNA in samples. See the reference cited in the following table for details about these kits.

Product	Description	References
Quantifiler [™] Human DNA Quantification Kit (PN 4343895) <i>and</i> Quantifiler [™] Y Human Male DNA Quantification Kit (PN 4343906)	 Properties: The Quantifiler[™] Human and Quantifiler[™] Y Human Male Kits are highly specific for human DNA, and they individually detect total human or male DNA, respectively. The kits detect single-stranded and degraded DNA. How they work: The Quantifiler[™] DNA Quantification Kits consist of target-specific and internal control 5' nuclease assays. The Quantifiler[™] Human and Quantifiler[™] Y Human Male Kits contain different target-specific assays (human DNA or human male DNA, respectively) that each consist of two locus-specific PCR primers and one TaqMan[™] MGB probe labeled with FAM[™] dye for detecting the amplified sequence. The kits each contain a separate internal PCR control (IPC) assay, which consists of an IPC template DNA (a synthetic sequence not found in nature), two primers for amplifying the IPC template DNA, and one TaqMan[™] MGB probe labeled with VIC[™] dye for detecting the amplified IPC DNA. 	Quantifiler [™] Human DNA Quantification Kits User's Manual (PN 4344790)
Quantifiler [™] Duo DNA Quantification Kit (PN 4387746)	Properties: The Quantifiler [™] Duo Kit is highly specific for human DNA. This kit combines the detection of both total human and male DNA in one PCR reaction. The kit detects single-stranded and degraded DNA. How it works: The Quantifiler [™] Duo DNA Quantification Kit consists of target- specific and internal control 5' nuclease assays. The Quantifiler [™] Duo kit combines two human-specific assays in one PCR reaction (for total human DNA and human male DNA). The two human DNA specific assays each consist of two PCR primers and a TaqMan [™] probe. The TaqMan [™] probes for the human DNA and human male DNA assays are labeled with VIC [™] and FAM [™] dyes, respectively. In addition, the kit contains an internal PCR control (IPC) assay similar in principle to that used in the other Quantifiler kits, but labeled with NED [™] dye.	Quantifiler [™] Duo DNA Quantification Kit User's Manual (PN 4391294)

Product	Description	References
Quantifiler [™] HP DNA Quantification Kit (Cat. no. 4482911) and Quantifiler [™] Trio	Properties: The Quantifiler [™] HP Kit is designed to quantify the total amount of amplifiable human DNA in a sample. The Quantifiler [™] Trio Kit is designed to simultaneously quantify the total amount of amplifiable human DNA and human male DNA in a sample.	Quantifiler HP and Trio DNA Quantification Kits User Guide (Pub no. 4485354)
Kit	How it works:	
(Cat. no. 4482910)	The Quantifiler [™] HP and Trio DNA Quantification Kits use multiple- copy target loci for improved detection sensitivity. The human- specific target loci (Small Autosomal, Large Autosomal, and Y- chromosome targets) each consist of multiple copies dispersed on various autosomal chromosomes (Small Autosomal and Large Autosomal), or multiple copies on the Y-chromosome (Quantifiler [™] Trio Kit only).	
	To maximize the consistency of quantification results, genomic targets were selected with conserved primer- and probe-binding sites within individual genomes and also with minimal copy number variability between different individuals and population groups. As a result, the detection sensitivity of the Quantifiler [™] HP and Trio assays is improved over Quantifiler [™] Duo, Human, and Y Human Male DNA Quantification Kit assays. The primary quantification targets (Small Autosomal and Y) consist of relatively short amplicons (75 to 80 bases) to improve the detection of degraded DNA samples. In addition, the Quantifiler [™] HP and Trio Kits each contain a Large Autosomal target with a longer amplicon (>200 bases) to aid in determining if a DNA sample is degraded.	

Prepare the amplification kit reactions

1. Calculate the volume of each component needed to prepare the reactions, using the table below.

DNA sample	Volume per reaction (μ L)
AmpFℓSTR [™] Identifiler [™] Plus Master Mix	10.0
AmpFℓSTR [™] Identifiler [™] Plus Primer Set	5.0

Note: Include additional reactions in your calculations to provide excess volume for the loss that occurs during reagent transfers.

2. Prepare reagents. Thaw the AmpFℓSTR[™] Identifiler[™] Plus Kit Master Mix and the AmpFℓSTR[™] Identifiler[™] Plus Kit Primer Set, then vortex 3 seconds and centrifuge briefly before opening the tubes.

IMPORTANT! Thawing is required only during first use of the kit. After first use, reagents are stored at 2 to 8 °C and, therefore, they do not require subsequent thawing. Do not refreeze the reagents.

- 3. Pipette the required volumes of components into an appropriately sized polypropylene tube.
- 4. Vortex the reaction mix for 3 seconds, then centrifuge briefly.
- 5. Dispense 15 μL of the reaction mix into each reaction well of a MicroAmp[™] Optical 96-Well Reaction Plate or each MicroAmp[™] tube.
- 6. Prepare the DNA samples:

DNA sample	To prepare
Negative control	Add 10 μL of low TE buffer (10mM Tris, 0.1mM EDTA, pH 8.0).
Test sample	Dilute a portion of the test DNA sample with low TE buffer so that 1.0 ng of total DNA is in a final volume of 10 μ L. Add 10 μ L of the diluted sample to the reaction mix.
Positive control	Add 10 µL of 9947A control DNA (0.1 ng/µL).

The final reaction volume (sample or control plus reaction mix) is 25 µL.

- 7. Seal the MicroAmp[™] Optical 96-Well Reaction Plate with MicroAmp[™] Clear Adhesive Film or MicroAmp[™] Optical Adhesive Film, or cap the tubes.
- 8. Vortex the reaction mix for 3 seconds, then centrifuge the tubes at 3000 rpm for about 20 seconds in a tabletop centrifuge (with plate holders if using 96-well plates) to remove bubbles.

Amplify the samples in a GeneAmp[™] PCR System 9700 with the silver 96-well block, a GeneAmp[™] PCR System 9700 with the gold-plated silver 96-well block, a Veriti[™] 96-Well Thermal Cycler, or a ProFlex[™] PCR

Note: The AmpFℓSTR[™] Identifiler[™] Plus Kit is not validated for use with the GeneAmp PCR System 9700 with the aluminium 96-well block. Use of this thermal cycling platform may adversely affect performance of the AmpFℓSTR[™] Identifiler[™] Plus Kit.

Select the appropriate PCR cycle number

All AmpFℓSTR[™] kits are optimized for a specific number of amplification cycles to deliver well-balanced and high quality results. However, increases in the number of low-level DNA samples being submitted for analysis have prompted many laboratories to evaluate increasing the number of amplification cycles to increase the sensitivity of the assay. Before increasing the cycle number, perform a comprehensive validation study to establish new performance criteria for the higher cycle number.

Higher cycle numbers can cause the following to occur:

- Exaggerated stochastic effects resulting from low DNA input amounts
- Greater difference between the presence and absence of an allele
- Greater heterozygote peak imbalance
- Possible differences in expected stutter position and percentage
- Possible increase in artifacts and/or background in the profile to accompany the increase in sample allele signal

The Identifiler[™] Plus Kit offers two PCR-cycle-number options:

- **Standard 28-PCR-cycle protocol** Provides high sensitivity to consistently generate full STR profiles with 125 pg of DNA input. Use with the optimum 1.0 ng DNA input amount in a maximum input volume of 10 µL.
- **29-PCR-cycle protocol** Adds the extra sensitivity when amplifying <125 pg DNA inputs. Recommended for use when the total DNA input amount is <0.5 ng.

The results of the developmental validation at both PCR cycle numbers is presented in Chapter 5 on page 77.

Perform PCR

WARNING! PHYSICAL INJURY HAZARD. Thermal cycler.

- 1. Program the thermal cycling conditions:
 - When using the GeneAmp PCR System 9700 with either 96-well silver or gold-plated silver block, select the **9600 Emulation Mode**.
 - When using the Veriti[™] 96-Well Thermal Cycler, refer to the following document for instructions on how to configure the Veriti instrument to run in the 9600 Emulation Mode: *User Bulletin: Veriti*[™] 96-Well Thermal Cycler AmpFtSTR[™] Kit Validation (PN 4440754).
 - When using the ProFlex[™] PCR System, refer to the *ProFlex[™] PCR System Kit Validation User Bulletin* (Pub. no. 100031595) for more information.

Initial incubation	Cycle (28 or 29 cycles [‡])		Final	Final hold
step	Denature	Anneal/Extend	extension	
HOLD	CY	CLE	HOLD	HOLD
95 °C 11 min	94 °C 20 sec	59 °C 3 min	60 °C 10 min	4 °C Up to 24 hours [§]

‡ Refer to the previous section for selecting the appropriate PCR cycle number

§ The infinity setting (∞) allows an unlimited hold time.

2. Load the plate into the thermal cycler and close the heated cover.

IMPORTANT! If using adhesive clear film instead of caps to seal the plate wells, be sure to place a MicroAmp[™] compression pad (PN 4312639) on top of the plate to prevent evaporation during thermal cycling.

- 3. Start the run.
- 4. On completion of the run, store the amplified DNA and protect from light.

If you are storing the DNA	Then place at
< 2 weeks	2 to 8 °C
> 2 weeks	–15 to –25 °C

IMPORTANT! Store the amplified products so that they are protected from light.

Amplification using bloodstained FTA[™] cards

FTA[™] cards can be useful for collecting, storing, and processing biological samples. A small punch disc of the card containing the sample can be placed directly into an amplification tube, purified, and amplified, without transferring the disc. Our studies indicate that a 1.2-mm bloodstained disc contains approximately 5 to 20 ng DNA. An appropriate cycle number for this high quantity of DNA is 24 cycles as determined by our validation studies. However, it is recommended that each laboratory determine the optimum cycle number based on internal validation studies.

In the example shown in Figure 4, a 1.2-mm disc of a bloodstained FTA card was purified using three washes with FTA Purification Reagent and two washes with $1 \times \text{low TE}$ buffer. The purified punch disc was then amplified in the MicroAmpTM tube for 24 cycles.



Figure 4 AmpF/STR[™] Identifiler[™] Plus PCR Amplification Kit results from a 1.2-mm FTA bloodstain disc (24-cycle amplification), analyzed on the Applied Biosystems[™] 3130x/ Genetic Analyzer

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Chapter 3

Electrophoresis

AmpFℓSTR[™] Identifiler[™] Plus User Guide

This chapter covers:

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Prepare samples for electrophoresis on the 3100/3100-Avant or 3130/3130xl instrument	38
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■ Prepare samples for electrophoresis on the 310 instrument	40

Allelic ladder requirements

To accurately genotype samples, you must run an allelic ladder sample along with the unknown samples. For samples run on the:

- ABI PRISM[™] 310 Genetic Analyzer: Run at least one allelic ladder for every 10 sample injections.
- ABI PRISM[™] 3100 and Applied Biosystems[™] 3130 series Genetic Analyzers: Run at least one allelic ladder per every set of 16 samples.

IMPORTANT! Variation in laboratory temperature can affect fragment migration speed and result in sizing variation. We recommend the following frequency of allelic ladder injections, which should account for normal variation in run speed. However, during internal validation studies, verify the required allelic ladder injection frequency to ensure accurate genotyping of all samples in your laboratory environment.

- Applied Biosystems[™] 3130*xl* or ABI PRISM[™] 3100 systems One ladder per injection; one injection = 16 samples (15 samples + 1 allelic ladder)
- Applied Biosystems[™] 3130 or ABI PRISM[™] 3100-Avant One ladder for every 4 injections; one injection = 4 samples

When genotyping, it is critical to use an allelic ladder run under the same conditions as the samples because:

- Size values obtained for the same sample can differ between instrument platforms because of different polymer matrices and electrophoretic conditions.
- Variation in laboratory temperature can affect migration speed (see IMPORTANT above).
Section 3.1 3100/3100-Avant and 3130/3130xl instruments

Set up the 3100/3100-Avant or 3130/3130xl instrument for electrophoresis

Reagents and
partsTable 3 on page 18 lists the required materials not supplied with the
AmpFℓSTR[™] Identifiler[™] Plus Kit.

IMPORTANT! The fluorescent dyes attached to the primers are light-sensitive. Protect the AmpFlSTR^M Identifiler^M Plus Primer Set from light when not in use. Amplified DNA, AmpFlSTR^M Identifiler^M Plus Allelic Ladder, and GeneScan^M 500 LIZ^M Size Standard should also be protected from light. Keep freeze-thaw cycles to a minimum.

3100/3100-Avant or 3130/3130x/ instrument requirements

The following table lists Data Collection Software and the run modules that can be used to analyze Identifiler[™] Plus PCR products. For details on the procedures, refer to the documents listed in the table.

Operating system	Data Collection Software	Run modules and conditions	References
Windows XP	3.0 [‡] (3130/3130 <i>xl</i> Analyzer)	 HIDFragmentAnalysis36_POP4_1 Injection conditions: 3130 = 3 kV/5 sec 3130xl = 3 kV/10 sec Dye Set G5 	Applied Biosystems [™] 3130/3130xl Genetic Analyzers Using Data Collection Software v3.0, Protocols for Processing AmpF ^ℓ STR PCR Amplification Kit PCR Products User Bulletin (PN 4363787)
Windows 2000	2.0 (3100 Analyzer)	 HIDFragmentAnalysis36_POP4_1 Injection condition: 3 kV/10 sec Dye Set G5 	ABI PRISM [™] 3100/3100-Avant Genetic Analyzers Using Data Collection Software v2.0, Protocols for Processing AmpFlSTR PCR Amplification Kit PCR Products User Bulletin (PN 4350218)
Windows NT [™]	1.1 (3100 Analyzer)	 GeneScan36vb_DyeSetG5Module Injection condition: 3 kV/10 sec GS500Analysis.gsp 	ABI PRISM [™] 3100/3100-Avant Genetic Analyzers Protocols for Processing AmpF l STR PCR Amplification Kit PCR Products User Bulletin (PN 4332345)
	1.0 (3100- <i>Avant</i> Analyzer)	 GeneScan36Avb_DyeSetG5Module Injection condition: 3 kV/5 sec GS500Analysis.gsp 	ABI PRISM [™] 3100/3100-Avant Genetic Analyzers Protocols for Processing AmpF ^ℓ STR PCR Amplification Kit PCR Products User Bulletin (PN 4332345)

‡ We conducted validation studies for the Identifiler[™] Plus Kit using this configuration.

Prepare samples for electrophoresis on the 3100/3100-Avant or 3130/3130x/ instrument

Prepare the
samplesPrepare the samples for electrophoresis on the 3100/3100-Avant or 3130/3130xl
instrument immediately before loading.

1. Calculate the volume of Hi-Di[™] Formamide and GeneScan[™] 500 LIZ[™] Size Standard needed to prepare the samples, using the table below.

Reagent	Volume per reaction (μ L)
GeneScan 500 LIZ Size Standard	0.3
Hi-Di Formamide	8.7

Note: Include additional samples in your calculations to provide excess volume for the loss that occurs during reagent transfers.

IMPORTANT! The volume of size standard indicated in the table is a suggested amount. Determine the appropriate amount of size standard based on your results and experiments.

- 2. Pipette the required volumes of components into an appropriately sized polypropylene tube.
- 3. Vortex the tube, then centrifuge briefly.
- 4. Into each well of a MicroAmp[™] Optical 96-Well Reaction Plate, add:
 - 9 µL of the formamide:size standard mixture
 - 1 µL of PCR product or allelic ladder

Note: For blank wells, add 10 μ L of Hi-DiTM Formamide.

- 5. Seal the reaction plate with appropriate septa, then briefly centrifuge the plate to ensure that the contents of each well are collected at the bottom.
- 6. Heat the reaction plate in a thermal cycler for 3 minutes at 95 °C.
- 7. Immediately place the plate on ice for 3 minutes.
- 8. Prepare the plate assembly on the autosampler.
- 9. Start the electrophoresis run.

Section 3.2 310 instrument

Set up the 310 instrument for electrophoresis

Reagents and
partsTable 3 on page 18 lists the required materials not supplied with the
AmpFℓSTR[™] Identifiler[™] Plus Kit.

IMPORTANT! The fluorescent dyes attached to the primers are light-sensitive. Protect the AmpFℓSTR[™] Identifiler[™] Plus Primer Set from light when not in use. Amplified DNA, AmpFℓSTR[™] Identifiler[™] Plus Allelic Ladder, and GeneScan[™] 500 LIZ[™] Size Standard should also be protected from light. Keep freeze-thaw cycles to a minimum.

310 instrument requirements The following table lists Data Collection Software and the run modules that can be used to analyze Identifiler[™] Plus PCR products. For details on the procedures, refer to the documents listed in the table.

Operating system	Data Collection Software	Run modules and conditions	References
Windows XP or Windows NT [™] and Windows 2000	3.1 [‡] or 3.0	GS STR POP4 (1mL) G5 v2.md5 Injection condition: 15 kV/5 sec	ABI PRISM [™] 310 Genetic Analyzer User's Manual (Windows) (PN 4317588) ABI PRISM [™] 310 Protocols for Processing AmpFlSTR PCR Amplification Kit Products with Microsoft Windows NT Operating System: User Bulletin (PN 4341742)

‡ We conducted concordance studies for the Identifiler[™] Plus kit using this configuration.

Prepare samples for electrophoresis on the 310 instrument

Prepare the
samplesPrepare the samples for capillary electrophoresis on the 310 instrument immediately
before loading.

1. Calculate the volume of Hi-Di[™] Formamide and GeneScan[™] 500 LIZ[™] Internal Size Standard needed to prepare the samples, using the table below.

Reagent	Volume per reaction (μ L)
GeneScan [™] 500 LIZ [™] Size Standard	0.5
Hi-Di [™] Formamide	24.5

Note: Include additional samples in your calculations to provide excess volume for the loss that occurs during reagent transfers.

IMPORTANT! The volume of size standard indicated in the table is a suggested amount. Determine the appropriate amount of size standard based on your results and experiments.

- 2. Pipette the required volumes of components into an appropriately sized polypropylene tube.
- 3. Vortex the tube, then centrifuge briefly.
- 4. Into a 0.2-mL or 0.5-mL sample tube, add:
 - $25 \,\mu L$ of the formamide:size standard mixture
 - 1.5 µL of PCR product or allelic ladder
- 5. Seal the tubes with the appropriate septa, then briefly vortex and centrifuge the tubes to ensure that the contents of each tubes are mixed and collected at the bottom.
- 6. Heat the tubes in a thermal cycler for 3 minutes at 95 °C.
- 7. Immediately place the tubes on ice for 3 minutes.
- 8. Place the sample tray on the autosampler.
- 9. Start the electrophoresis run.

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Chapter 4

Data Analysis

AmpFℓSTR[™] Identifiler[™] Plus User Guide

This chapter covers:

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Section 4.1 GeneMapper[™] ID Software

Overview of GeneMapper[™] ID Software

GeneMapper^m *ID* Software is an automated genotyping software for forensic casework, databasing, and paternity data analysis.

After electrophoresis, the Data Collection Software stores information for each sample in a .fsa file. Using GeneMapperTM *ID* Software v3.2.1 software, you can then analyze and interpret the data from the .fsa files.

Instruments Refer to **"Instrument and software overview" on page 15** for a list of compatible instruments.

Before you start When using GeneMapper^M *ID* Software v3.2.1 to perform human identification (HID) analysis with AmpFlSTR^M kits, be aware that:

- HID analysis requires at least one allelic ladder sample per run folder. Your laboratory can use multiple ladder samples in an analysis, provided individual laboratories conduct the appropriate validation studies.
 For multiple ladder samples, the GeneMapper[™] *ID* Software calculates allelic bin offsets by using an average of all ladders that use the same panel within a run folder.
- Allelic ladder samples in an individual run folder are considered to be from a single run.

When the software imports multiple run folders into a project, only the ladder(s) within their respective run folders are used for calculating allelic bin offsets and subsequent genotyping.

- Allelic ladder samples must be labeled as "Allelic Ladder" in the Sample Type column in a project. Failure to apply this setting for ladder samples results in failed analysis.
- Injections containing the allelic ladder must be analyzed with the same analysis method and parameter values that are used for samples to ensure proper allele calling.
- Alleles that are not in the AmpFℓSTR[™] Allelic Ladders do exist. Off-ladder (OL) alleles may contain full and/or partial repeat units. An off-ladder allele is an allele that occurs outside the ±0.5-nt bin window of any known allelic ladder allele or virtual bin.

Note: If a sample allele peak is called as an off-ladder allele, the sample result needs to be verified according to the laboratory's protocol.

If you are using GeneMapperTM *ID-X* Software to perform Human Identification (HID) analysis with AmpFℓSTR kits, go to "Set up GeneMapperTM ID-X Software for data analysis" on page 61 or refer to the *GeneMapperTM ID-X Software Version* 1.0 Human Identification Analysis Getting Started Guide (PN 4375574).

Set up GeneMapper[™] *ID* Software for data analysis

Workflow Before you can analyze sample (.fsa) files using GeneMapper[™] *ID* Software v3.2.1 for the first time:

- Import panels and bins into the Panel Manager, as explained in "Import panels and bins" on page 48.
- Import an analysis method, as explained in "Import an HID analysis method" on page 51.
- Import a size standard, as explained in "Import an HID size standard" on page 56.
- Define custom views of analysis tables.
- Define custom views of plots.
- **For more info** For details about GeneMapper[™] *ID* Software workflow and features, refer to:
 - GeneMapper[™] ID Software Version 3.1 Human Identification Analysis User Guide (PN 4338775)
 - GeneMapper[™] ID Software Versions 3.1 and 3.2 Human Identification Analysis Tutorial (PN 4335523)
 - Installation Procedures and New Features for GeneMapper[™] ID Software v3.2 User Bulletin (PN 4352543)

Import panels and bins To import the AmpF ℓ STRTM IdentifilerTM Plus Kit panel and bin set from our web site into the GeneMapperTM ID Software v3.2.1 database:

- 1. Download and open the file containing panels and bins:
 - a. From the Support menu of www.appliedbiosystems.com, select Software Downloads, Patches & Updates. Select GeneMapper[™] ID Software v3.2 from the drop-down menu. Select Updaters & Patches, and download the file Identifiler Plus Analysis Files GMID.
 - b. Unzip the file.
- 2. Start the GeneMapper[™] *ID* Software, then log in with the appropriate user name and password.

IMPORTANT! If you need logon instructions, refer to page 2-7 of the *GeneMapper*[™] *ID Software Version 3.1 Human Identification Analysis User Guide* (PN 4338775).

- 3. Select Tools ▶ Panel Manager.
- 4. Find, then open the folder containing the panels and bins:
 - a. Select Panel Manager in the navigation pane.



- b. Select File > Import Panels to open the Import Panels dialog box.
- c. Navigate to, then open the **Identifiler Plus Analysis Files GMID** folder that you unzipped in step 1.
- 5. Select Identifiler_Plus_Panels_v1, then click Import.

Note: Importing this file creates a new folder in the navigation pane of the Panel Manager, AmpFLSTR_Identifiler_Plus_v1. This folder contains the panel and associated markers.



- 6. Import Identifiler_Plus_Bins_v1:
 - a. Select the **AmpFLSTR_Identifiler_Plus_v1** folder in the navigation pane.

💽 Panel Manager		
File Edit Bins View		
	Bin Set:	▼ ■ ■
🖃 🖃 Panel Manager	Panel Name	Comment.
	1 Identifiler_Plus_v1	null
AmpFLSTR_Identifiler_Plus_v1		

- b. Select **File** > **Import Bin Set** to open the Import Bin Set dialog box.
- c. Navigate to, then open the Identifiler Plus Analysis Files GMID folder.
- d. Select Identifiler_Plus_Bins_v1, then click Import.

Note: Importing this file associates the bin set with the panels in the Identifiler_Plus_Panels_v1 folder.

💽 Import Bin Set	×
Look in: 🔁 Identifiler Plus Analysis Files GMID 🗾 🦻	📂 🖽 👘
Image: CE_G5_Identifiler_Plus_GS500.xml My Recent D Image: CE_G5_Identifiler_Plus_GS500.xml Image: CE_G5_Identifiler_Plus_AnalysisMethod_v1.xml Image: CE_G5_Identifiler_Plus_GIS500.xml Image: CE_G5_Identifiler_Plus_AnalysisMethod_v1.xml Image: CE_G5_Identifiler_Plus_Bins_v1.txt Image: CE_G5_Identifiler_Plus_Bins_v1.txt Image: CE_G5_Identifiler_Plus_Bins_v1.txt	
My Documents File name: Identifiler_Plus_Bins_v1.txt	Import
Files of type: All Files	Cancel

- 7. View the imported panels in the navigation pane:
 - a. Double-click the **AmpFLSTR_Identifiler_Plus_v1** folder to view the Identifiler_Plus_Panels_v1 folder.
 - b. Double-click the **Identifiler_Plus_Panels_v1** folder to display the panel information in the right pane.

💽 Panel Manager										
File Edit Bins View										
🎬 🗙 📓 🕷 📗 🛄 Bin S	et:	ldentifiler_Plus_E	Bins_v1	•		; # 🖪 🕷 📗				
🖃 🖃 Panel Manager		Marker Name	Dye Color	Min Size	Max Size	Control Alleles	Marker I	Marker :	Comments	Ladder Alleles
- AmpFLSTR_Panels_v2	1	D8S1179	blue	118.0	183.5	13	4	0.1032	none	8,9,10,11,12,13,1
E-CAmpFLSTR_Identifiler_Plus_v1	2	D21S11	blue	184.5	247.5	30	4	0.1067	none	24,24.2,25,26,27
□ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □	3	D7S820	blue	251.0	298.5	10,11	4	0.0969	none	6,7,8,9,10,11,12;
-D21S11	4	CSF1PO	blue	302.12	348.63	10,12	4	0.092	none	6,7,8,9,10,11,12;
-D7S820	5	D3S1358	green	98.0	148.0	14,15	4	0.1227	none	12,13,14,15,16,1
- CSF1PO	6	TH01	green	159.0	205.0	8,9.3	4	0.0408	none	4,5,6,7,8,9,9.3,10
	7	D13S317	green	205.65	250.16	11	4	0.0993	none	8,9,10,11,12,13,1
D13S317	8	D16S539	green	255.3	301.81	11,12	4	0.1039	none	5,8,9,10,11,12,13
D16S539	9	D2S1338	green	304.8	370.31	19,23	4	0.1244	none	15,16,17,18,19,2
— D2S1338	10	D19S433	yellow	101.0	148.0	14,15	4	0.1121	none	9,10,11,12,12.2,1
	11	WVA	yellow	151.0	213.5	17,18	4	0.1245	none	11,12,13,14,15,1
	12	трох	yellow	216.99	260.99	8	4	0.0638	none	6,7,8,9,10,11,12;
-D18S51	13	D18S51	yellow	264.49	350.0	15,19	4	0.1368	none	7,9,10,10.2,11,12
- AMEL	14	AMEL	red	106.0	114.0	x	9	0.0	none	X,Y
	15	D5S818	red	128.0	180.0	11	4	0.1006	none	7,8,9,10,11,12,13
-rGA	16	FGA	red	206.25	360.0	23,24	4	0.1303	none	17,18,19,20,21,2

8. Select **D8S1179** to display the Bin view for the marker in the right pane.



9. Click **Apply**, then **OK** to add the AmpFℓSTR[™] Identifiler[™] Plus Kit panel and bin set to the GeneMapper[™] *ID* Software database.

IMPORTANT! If you close the Panel Manager without clicking OK, the panels and bins are not imported into the GeneMapper^M *ID* Software database.

Import an HIDThe HID Advanced analysis method for the AmpFℓSTR[™] Identifiler[™] Plus PCRanalysis methodAmplification Kit uses the Identifiler_Plus_Bins_v1 file described in step 6 on
page 49.

Use the following procedure to import the analysis method from the folder that you downloaded from our web site, into the GeneMapperTM *ID* Software database. Refer to **step 1 on page 48** for downloading instructions.

Note: The Identifiler_Plus_AnalysisMethod_v1 has been provided to assist you in getting started with Identifiler Plus kit data analysis. Analysis parameters should be established by each individual laboratory based on the laboratory's internal validation studies.

1. Select **Tools > GeneMapper Manager** to open the GeneMapper Manager.

- 2. Import an analysis method for HID_Advanced:
 - a. Select the Analysis Methods tab, then click Import.

Gen	eMapper Manager						X
Project	ts Analysis Methods Table	Settings Plot Settin	gs Matrices Size :	Standards			
	Name	Last Saved	Owner	Instrument	Analysis Type	Description	
	HID_Advanced	2009-06-18 16:22:2	gmid		HID		
	HID_Classic	2007-08-06 10:03:0	gmid		HID		H
	Microsatellite Default	2004-05-28 11:34:3	gmid		Microsatellite	Factory Provided	-
	4	1	1				
New	v Open Say	/e As Impor	t Export			Dejete	
						Done	

- b. Navigate to, then open the Identifiler Plus Analysis Files GMID folder.
- c. Select Identifiler_Plus_AnalysisMethod_v1, then click Import to import the Identifiler_Plus_AnalysisMethod_v1 into the GeneMapper[™] ID Software database.

👰 Import Analy	sis Method			×
Look in:	🛅 Identifiler Plus Ai	nalysis Files GMID 📃	ø	۲ 🔃
My Recent D Desktop	CE_G5_Identi	filer_Plus_GS500.xml s_AnalysisMethod_v1.xml		
My Documents	File name: Id	lentifiler_Plus_AnalysisMethod_v1.xml ML Files (*.xml)	-	Import Cancel

To view the settings for Identifiler_Plus_AnalysisMethod_v1, select the **Analysis Methods** tab, then select **Identifiler_Plus_AnalysisMethod** in the Name column and click **Open**.

💽 GeneM	lapper Manager							
Projects	Analysis Methods	Table Settings	Plot Settings 🛛 M	atrices Size Sta	ndards			
Na	ame		Last Saved	1	Owner	Instrument	Analysis Ty	/pe [
De	efault		2009-08-3	1 11:27:45.695	gmid		HID	E
Id	lentifiler_Plus_Analy	sisMethod_v1	2009-09-0	2 11:13:05.191	gmid		HID	it
4								F
New	Open	Save As	Import	Export				Delete
								Done

Figure 5 Analysis Method Editor - HID: General tab settings

Figures 6 through 9 below show the settings for each tab of the Analysis Method Editor - HID.

Analysis Method Editor - HID									
General Allele Peak Detector Peak Quality Quality Flags									
Bin Set: Identifiler Plus Bins v1									
Use marker-specific stutter	ratio if ava	ilable							
Marker Repeat Type :	Tri	Tetra	Penta	Hexa					
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					
Minus Stutter Ratio	0.0	0.0	0.0	0.0					
Minus Stutter Distance From	0.0	3.25	0.0	0.0					
То	0.0	4.75	0.0	0.0					
Plus Stutter Ratio	0.0	0.0	0.0	0.0					
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									
Range Filter Factory Defaults									
			OK	Cance	9				

Figure 6 Analysis Method Editor - HID: Allele tab settings

• GeneMapper[™] *ID* Software v3.2.1 allows you to specify four types of marker repeat motifs: tri, tetra, penta, and hexa. You can enter parameter values for each type of repeat in the appropriate column.

• The "Use marker-specific stutter ratio if available" check box is selected by default. Consequently, the software applies the stutter ratio filters supplied in the Identifiler_Plus_Panels_v1 file.

For more information about allele filters, refer to:

- GeneMapper[™] ID Software Version 3.1 Human Identification Analysis User Guide, Chapter 3 (PN 4338775)
- GeneMapper[™] ID Software Versions 3.1 and 3.2 Human Identification Analysis Tutorial (PN 4335523)
- Installation Procedures and New Features for GeneMapper[™] ID Software v3.2 User Bulletin (PN 4352543)

Analysis Method Editor - HID	X
General Allele Peak Detector Peak Quality	Quality Flags
General Allele Peak Detector Peak Quality Peak Detection Algorithm: Advanced Ranges Analysis Sizing Full Range All Sizes Image Start Pt: 0 Start Size: Stop Pt: 10000 Stop Size: Smoothing and Baselining Smoothing None	Quality Flags Peak Detection Peak Amplitude Thresholds: B: 50 G: 50 G: 50 Y: 50 Min. Peak Half Width: 2 Polynomial Degree: 3
C Light C Heavy Baseline Window: 51 pts Size Calling Method 2nd Order Least Squares 3rd Order Least Squares C Cubic Spline Interpolation C Local Southern Method C Global Southern Method	Peak Window Size: 11 pts Slope Threshold Peak Start: 0.0 Peak End: 0.0 Factory Defaults
	OK Cancel

Figure 7 Analysis Method Editor - HID: Peak Detector tab settings

IMPORTANT! Laboratories need to perform the appropriate internal validation studies to determine the peak amplitude thresholds that allow for reliable interpretation of $AmpF\ell STR^{TM}$ IdentifilerTM Plus PCR Amplification Kit data.

The software uses the peak amplitude threshold parameters to specify the minimum peak height to limit the number of detected peaks. Although GeneMapperTM *ID* Software displays peaks that fall below the specified amplitude in electropherograms, the software does not label or determine the genotype of these peaks.

Note: The analysis range is set by the user based on the locations of the primer peaks and size standard peaks.

For more information about peak detection algorithms, refer to:

- GeneMapper[™] ID Software Version 3.1 Human Identification Analysis User Guide (PN 4338775)
- GeneMapper[™] ID Software Versions 3.1 and 3.2 Human Identification Analysis Tutorial, Appendix A (PN 4335523)
- Installation Procedures and New Features for GeneMapper[™] ID Software v3.2 User Bulletin (PN 4352543)

Analysis Method Editor - HID	×
General Allele Peak Detector	Peak Quality 🛛 Quality Flags
Signal level	
Homozygous min peak height	100.0
Heterozygous min peak height	50.0
Heterozygote balance	
Min peak height ratio	0.7
Peak morphology	
Max peak width (basepairs)	1.5
Pull-up peak	
Pull-up ratio	0.05
Allele number	
Max expected alleles	2
·	Easterni Dafaulta
	Factory Defaults
	OK Cancel

Figure 8 Analysis Method Editor - HID: Peak Quality tab settings

IMPORTANT! Laboratories need to perform the appropriate internal validation studies to determine the minimum heterozygous and homozygous minimum peak height thresholds and the minimum peak height ratio threshold that allow for reliable interpretation of AmpFℓSTR[™] Identifiler[™] Plus PCR Amplification Kit data.

Analysis Method Edito	r - HID			×
General Allele Peak D	etector Pea	ak Quality Qualit	y Flags	,
Quality weights are bet Quality Flag Settings—	ween 0 and 1	l.		
Spectral Pull-up Broad Peak Out of Bin Allele Overlap	0.8 0.8 0.8	Cont Low Off-s Peak	rol Concordance Peak Height scale Height Ratio	1.0 0.3 0.8 0.3
PQV Thresholds				
	Pass Ra	nge:	Low Quality	/ Range:
Sizing Quality:	From 0.75	to 1.0	From 0.0 to	0.25
Genotype Quality:	From 0.75	to 1.0	From 0.0 to	0.25
			Eacto	ory Defaults
			<u>o</u> ĸ	Cancel

Figure 9 Analysis Method Editor - HID: Quality Flags tab settings

Import an HID The size standard for the Identifiler[™] Plus Kit uses the following GS500 peaks in its sizing algorithm: 75, 100, 139, 150, 160, 200, 300, 350, 400, and 450.

Use the following procedure to import the size standard for the AmpF ℓ STRTM IdentifilerTM Plus PCR Amplification Kit from the folder that you downloaded from our web site, into the GeneMapperTM *ID* Software database. Refer to **step 1 on page 48** for downloading instructions.

- 1. Select **Tools > GeneMapper Manager** to open the GeneMapper Manager.
- 2. Import a size standard:
 - a. Select the Size Standards tab, then click Import.

Gene	eMapper Manager					X				
Project	Projects Analysis Methods Table Settings Plot Settings Matrices Size Standards									
	Name	Last Saved	Owner	Туре	Description					
	377_F_HID_GS500	2004-05-28 11:34:3	gmid	Basic/Advanced	Factory Provided					
	CE_G5_HID_GS500	2004-05-28 11:34:3	gmid	Basic/Advanced	Factory Provided	H				
	CE_F_HID_GS500	2004-05-28 11:34:3	gmid	Basic/Advanced	Factory Provided					
New	New Open Save As Import Export Delete									
					Done	e				

b. Navigate to, then open the Identifiler Plus Analysis Files GMID folder.

c. Select CE_G5_Identifiler_Plus_GS500, then click Import to import the CE_G5_Identifiler_Plus_GS500 size standard into the GeneMapper[™] *ID* Software database.

Import Size S	itandard Metho	od	×
Look in:	🛅 Identifiler Plus	s Analysis Files GMID 🗾 🦸	i 📂 📰 👘
My Recent D Desktop My Documents	CE_GS_dd	entifiler_Plus_GS500.xml	
	File name:	CE_G5_Identifiler_Plus_GS500.xml	Import
	Files of type:	XML Files (*.xml)	Cancel

Analyze and edit sample files with GeneMapper[™] ID Software

- Analyze a project 1. In the Project window, select File > Add Samples to Project, then navigate to the disk or directory containing the sample files.
 - 2. Apply analysis settings to the samples in the project.

Parameter	Settings
Sample Type	Select the sample type.
Analysis Method	Identifiler_Plus_AnalysisMethod_v1
Panel	Identifiler_Plus_Panels_v1
Size Standard [‡]	CE_G5_Identifiler_Plus_GS500 ^{§#}

‡ For more information about how the Size Caller works, refer to the ABI PRISM[™] GeneScan[™] Analysis Software for the Windows NT[™] Operating System Overview of the Analysis Parameters and Size Caller User Bulletin (PN 4335617).

Size Caller Oser Buildin (FN 4353617).
§ The following fragments are defined for the CE_G5_Identifiler_Plus_GS500 size standard provided with the AmpF4STR[™] Identifiler[™] Plus Kit: 75, 100, 139, 150, 160, 200, 300, 350, 400, and 450. For additional information about size standards, refer to the GeneMapper[™] ID Software Version 3.1 Human Identification Analysis User Guide (PN 4338775), Appendix D.

Neither the 250-nt nor the 340-nt peak are included in the size standard definition. These peaks can be used as an indicator of precision within a run.

- 3. Click ► (Analyze), enter a name for the project (in the Save Project dialog box), then click **OK** to start analysis.
 - The status bar displays the progress of analysis:
 - As a completion bar extending to the right with the percentage completed indicated
 - With text messages on the left

• The Samples table (figure below) displays the row of the sample currently being analyzed in green (or red if analysis failed for the sample).

.2.1 - *	2.1 - *Untitled - gmid Is Logged In											
ew Ta	w Tools Help											
š I	5 🔟 🔟 🕅 🚰 🔁 🕨 🚳 Table Setting: HID_LadderTable_FileName 🔽 🕅 💭 🚭 🗛											
Sampl	Samples Genotypes											
	Status	Sample File	Sample Name	Sample Type	Analysis Method	Panel	Size Standard	Instrument Type				
1	J.	Cerebus_3130XL	IB_0329	Sample	ldentifiler_Plus_AnalysisMethod_v1	ldentifiler_Plus_Panels_v1	CE_G5_Identifiler_Plus_GS500	ABI3130				
2	J.	Cerebus_3130XL	IB_0330	Sample	Identifiler_Plus_AnalysisMethod_v1	Identifiler_Plus_Panels_v1	CE_G5_Identifiler_Plus_GS500	ABI3130				
3	din.	Cerebus_3130XL	IB_0331	Sample	ldentifiler_Plus_AnalysisMethod_v1	Identifiler_Plus_Panels_v1	CE_G5_Identifiler_Plus_GS500	ABI3130				
4	din.	Cerebus_3130XL	IB_0332	Sample	Identifiler_Plus_AnalysisMethod_v1	Identifiler_Plus_Panels_v1	CE_G5_Identifiler_Plus_GS500	ABI3130				
5	din.	Cerebus_3130XL	IB_0333	Sample	Identifiler_Plus_AnalysisMethod_v1	Identifiler_Plus_Panels_v1	CE_G5_Identifiler_Plus_GS500	ABI3130				
6	J.	Cerebus_3130XL	IB_0334	Sample	ldentifiler_Plus_AnalysisMethod_v1	Identifiler_Plus_Panels_v1	CE_G5_Identifiler_Plus_GS500	ABI3130				
7	din.	Cerebus_3130XL	IB_0335	Sample	Identifiler_Plus_AnalysisMethod_v1	Identifiler_Plus_Panels_v1	CE_G5_Identifiler_Plus_GS500	ABI3130				
8	, Im	Cerebus_3130XL	IB_0336	Sample	Identifiler_Plus_AnalysisMethod_v1	Identifiler_Plus_Panels_v1	CE_G5_Identifiler_Plus_GS500	ABI3130				
9	Jh.	Cerebus_3130XL	IB_0337	Sample	ldentifiler_Plus_AnalysisMethod_v1	Identifiler_Plus_Panels_v1	CE_G5_Identifiler_Plus_GS500	ABI3130				
10	J.	Cerebus_3130XL	IB_0338	Sample	ldentifiler_Plus_AnalysisMethod_v1	Identifiler_Plus_Panels_v1	CE_G5_Identifiler_Plus_GS500	ABI3130				
11	J.	Cerebus_3130XL	IB_0339	Sample	ldentifiler_Plus_AnalysisMethod_v1	Identifiler_Plus_Panels_v1	CE_G5_Identifiler_Plus_GS500	ABI3130				
12	J.	Cerebus_3130XL	IB_0340	Sample	ldentifiler_Plus_AnalysisMethod_v1	Identifiler_Plus_Panels_v1	CE_G5_Identifiler_Plus_GS500	ABI3130				
13	J.	Cerebus_3130XL	IB_0341	Sample	Identifiler_Plus_AnalysisMethod_v1	Identifiler_Plus_Panels_v1	CE_G5_Identifiler_Plus_GS500	ABI3130				
14	J.	Cerebus_3130XL	IB_0342	Sample	Identifiler_Plus_AnalysisMethod_v1	Identifiler_Plus_Panels_v1	CE_G5_Identifiler_Plus_GS500	ABI3130				
15	, Im	Cerebus_3130XL	ladder3	Allelic Ladder	Identifiler_Plus_AnalysisMethod_v1	Identifiler_Plus_Panels_v1	CE_G5_Identifiler_Plus_GS500	ABI3130				

• The Genotypes tab becomes available after analysis.

For more information about any of these tasks, refer to the *GeneMapper*[™] *ID* Software Version 3.1 Human Identification Analysis User Guide (PN 4338775).

Examine and edit a project

You can display electropherogram plots from the Samples and Genotypes tabs of the Project window to examine the data. These procedures start with the Samples tab of the Project window (assuming the analysis is complete).

For more information about any of these tasks, refer to:

- GeneMapper[™] ID Software Versions 3.1 and 3.2 Human Identification Analysis Tutorial (PN 4335523)
- GeneMapper[™] ID Software Version 3.1 Human Identification Analysis User Guide (PN 4338775)
- Installation Procedures and New Features for GeneMapper[™] ID Software Version v3.2 User Bulletin (PN 4352543)

Section 4.2 GeneMapper[™] *ID-X* Software

Overview of GeneMapper[™] *ID-X* Software

GeneMapper^m *ID-X* Software is an automated genotyping software for forensic casework, databasing, and paternity data analysis.

After electrophoresis, the Data Collection Software stores information for each sample in a .fsa file. Using GeneMapper^M *ID-X* Software v1.0.1, v1.1, or v1.1.1, you can then analyze and interpret the data from the .fsa files.

Instruments Refer to **"Instrument and software overview" on page 15** for a list of compatible instruments.

Before you start When using GeneMapper^M *ID-X* Software v1.0.1, v1.1, or v1.1.1 to perform human identification (HID) analysis with AmpF*l*STR^M kits, be aware that:

HID analysis requires at least one allelic ladder sample per run folder. Your laboratory can use multiple ladder samples in an analysis, provided individual laboratories conduct the appropriate validation studies. For multiple ladder samples, the GeneMapper[™] *ID-X* Software calculates allelic bin offsets by using an average of all ladders that use the same panel within a run folder.

• Allelic ladder samples in an individual run folder are considered to be from a single run.

When the software imports multiple run folders into a project, only the ladder(s) within their respective run folders are used for calculating allelic bin offsets and subsequent genotyping.

- Allelic ladder samples must be labeled as "Allelic Ladder" in the Sample Type column in a project. Failure to apply this setting for ladder samples results in failed analysis.
- Injections containing the allelic ladder must be analyzed with the same analysis method and parameter values that are used for samples to ensure proper allele calling.
- Alleles that are not in the AmpFℓSTR[™] Allelic Ladders do exist. Off-ladder (OL) alleles may contain full and/or partial repeat units. An off-ladder allele is an allele that occurs outside the ±0.5-nt bin window of any known allelic ladder allele or virtual bin.

Note: If a sample allele peak is called as an off-ladder allele, the sample result needs to be verified according to the laboratory's protocol.

Set up GeneMapper[™] *ID-X* Software for data analysis

Workflow	Before you can analyze sample (.fsa) files using GeneMapper TM <i>ID-X</i> Software v1.0.1, v1.1, or v1.1.1 for the first time:
	• Import panels, bins, and marker stutter into the Panel Manager, as explained in "Import panels, bins, and marker stutter" on page 61.
	• Import an analysis method, as explained in "Import an analysis method" on page 66 .
	• Import a size standard, as explained in "Import a HID size standard" on page 71.
	• Define custom views of analysis tables.
	• Define custom views of plots.
For more info	For quick set up instructions, refer to the GeneMapper [™] <i>ID-X</i> Software <i>Version 1.0 Getting Started Guide</i> (PN 4375574).
	For details about the GeneMapper [™] <i>ID-X</i> Software workflow and features, refer to:
	• GeneMapper [™] ID-X Software Version 1.0 Getting Started Guide (PN 4375574)
	• GeneMapper [™] <i>ID-X</i> Software Version 1.0 Ouick Reference Guide (PN 4375670)
	• GeneMapper ^{\mathbb{M}} <i>ID-X</i> Software Version 1.0 Reference Guide (PN 4375671)
Import panels, bins, and marker stutter	To import the AmpFlSTR TM Identifiler TM Plus Kit panels, bin sets, and marker stutter from our web site into the GeneMapper TM <i>ID-X</i> Software v1.0.1, v1.1, or v1.1.1 database:
	1. Download and open the file containing panels, bins, and marker stutter:
	 a. From the Support menu of www.appliedbiosystems.com, select Software Downloads, Patches & Updates. Select GeneMapper[™] ID-X Software from the drop-down menu. Select Updaters & Patches, and download the file Identifiler Plus Analysis Files GMIDX.
	b. Unzip the file.
	2. Start the GeneMapper [™] <i>ID-X</i> Software, then log in with the appropriate user name and password.
	IMPORTANT! If you need logon instructions, refer to the <i>GeneMapper</i> ^{TM} <i>ID-X Software Version 1.0 Getting Started Guide</i> (PN 4375574).
	3. Select Tools ▶ Panel Manager.

- 4. Find, then open the folder containing the panels, bins, and marker stutter:
 - a. Select **Panel Manager** in the navigation pane.



- b. Select File > Import Panels to open the Import Panels dialog box.
- c. Navigate to, then open the Identifiler Plus Analysis Files GMIDX folder that you unzipped in **step 1 on page 61**.
- 5. Select Identifiler_Plus_Panels_v1X, then click Import.

Note: Importing this file creates a new folder in the navigation pane of the Panel Manager "AmpFLSTR_Identifiler_Plus_v1X." This folder contains the panel and associated markers.



- 6. Import Identifiler_Plus_Bins_v1X:
 - a. Select the **AmpFLSTR_Identifiler_Plus_v1X** folder in the navigation pane.

	📌 P	anel N	1anag	er								
	File	Edit	Bins	View	Help							
	ц ^а	×						Bir	n Set			•
ſ		🚠 Par	iel Mari	ager						Panel Name	Comment.	
)AmpF	LSTR_	Panels_	v1X			1	Identifiler_Plus_Panels_v1X	null	
		÷ 🚞	AmpF	LSTR_J	Identifil	er_Plu	is_v1X					

- b. Select File > Import Bin Set to open the Import Bin Set dialog box.
- c. Navigate to, then open the AmpFLSTR_Identifiler_Plus_v1X folder.

d. Select Identifiler_Plus_Bins_v1X, then click Import.

Note: Importing this file associates the bin set with the panels in the Identifiler_Plus_Panels_v1X folder.

💕 Import Bin Set		
Look in:	🛅 Identifiler Plus Analysis Files GMIDX	💌 🤌 🛤 📰
My Recent Documents Oesktop	CE_GS_Identifiler_Plus_GS500.xml Identifier_Plus_Bins_v1X.txt Identifier_Plus_Panels_v1X.txt Identifier_Plus_Stutter_v1X.txt Identifiler_Plus_AnalysisMethod_v1X.xml	

- 7. View the imported panels in the navigation pane:
 - a. Double-click the **AmpFLSTR_Identifiler_Plus_v1X** folder to view the Identifiler_Plus_Panel_v1X folder.
 - b. Double-click the **Identifiler_Plus_Panel_v1X** folder to display the panel information in the right pane and the markers below it.

💕 Panel Manager									
File Edit Bins View Help									
🚔 🗙 🛛 📰 📓 🖉 🖉 🔡 🛤 Bin Se	et: I	dentifiler_Plus_E	Bins_v1X			•	š	1	
🖃 🖫 🚠 Panel Manager		Marker Name	Dye Color	Min Size	Max Size	Control Alleles	Marker	Comments	Ladder Alleles
🕀 🗁 AmpFLSTR_Panels_v1X	1	D851179	Blue	118.0	183.5	13	4	none	8,9,10,11,12,13,14,15,1
AmpFLSTR_Identifiler_Plus_v1X	2	D21511	Blue	184.5	247.5	30	4	none	24,24.2,25,26,27,28,28.
Desting	3	D75820	Blue	251.0	298.5	10,11	4	none	6,7,8,9,10,11,12,13,14,
⊡ D21511	4	CSF1PO	Blue	302.12	348.63	10,12	4	none	6,7,8,9,10,11,12,13,14,
	5	D351358	Green	98.0	148.0	14,15	4	none	12,13,14,15,16,17,18,19
⊡CSF1PO	6	TH01	Green	159.0	205.0	8,9.3	4	none	4,5,6,7,8,9,9.3,10,11,10
	7	D135317	Green	205.65	250.16	11	4	none	8,9,10,11,12,13,14,15
HI - D135317	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
	10	D195433	Yellow	101.0	148.0	14,15	4	none	9,10,11,12,12.2,13,13.2
	11	VWA	Yellow	151.0	213.5	17,18	4	none	11,12,13,14,15,16,17,18
TPOX	12	TPOX	Yellow	216.99	260.99	8	4	none	6,7,8,9,10,11,12,13
	13	D18551	Yellow	264.49	350.0	15,19	4	none	7,9,10,10.2,11,12,13,13
AMEL	14	AMEL	Red	106.0	114.0	x	9	none	Х,Ү
	15	D55818	Red	128.0	180.0	11	4	none	7,8,9,10,11,12,13,14,15
±™ FGA	16	FGA	Red	206.25	360.0	23,24	4	none	17,18,19,20,21,22,23,24

8. Select **D2S1388** to display the Bin view for the marker in the right pane.

💕 Panel Manager								×
File Edit Bins View Help								
📑 🗙 👔 🖩 📓 📕 Bin	Set: Identifiler_Plus_Bins_	v1X	•	III G B				
Approximate the second s	1.0	7 8	9 10	11 12 13	14 15 16	17 18 19	20	
	0.9-							
	0.8-							
₩ U351358 ₩ TH01 ₩ D135317	0.7							
	0.6 -							
	0.5 -							
H → AMEL H → DSS818 H → FGA	0.4-							
	0.3-							
Reference Samples	0.2 -							
	0.1-							
	0.0 + + + + + + + + + + + + + + + + + +	121 123125 127 12	9131 133135 137 13	9141 143145 14714	9 151 153155 157159 1	61 163165 167 169 171 173	175 177 179 181 183 18	, , , , J 85 187
	D85	\$1179						•
J	Marker: D8S1179 (11	8.00-183.50)						
		OK Cano	cel Apply	Help				

- 9. Import Identifiler_Plus_Stutter_v1X:
 - a. Select the **AmpFLSTR_Identifiler_Plus_v1X** folder in the navigation panel.

💕 P	💕 Panel Manager										
File	Edit	Bins	View	Help							
	\mathbf{X}_{\parallel}			1			Bir	n Set	:: Identifiler_Plus_Bins_v1X		-
🖃 📇 Panel Manager											
	🚠 Pan	el Man	ager						Panel Name	Comment	
	🚠 Pan	el Man (AmpFl	ager LSTR_F	Panels_	v1X			1	Panel Name Identifiler_Plus_Panels_v1X	Comment. null	
	₽an	el Man AmpFl AmpFl	iager LSTR_F LSTR_1	Panels_ Identifik	v1X er_Plu	is_v1X		1	Panel Name Identifiler_Plus_Panels_v1X	Comment null	

- b. Select File → Import Marker Stutter to open the Import Marker Stutter dialog box.
- c. Navigate to, then open the Identifiler Plus Analysis Files GMIDX folder.
- d. Select Identifiler_Plus_Stutter_v1X, then click Import.

Note: Importing this file associates the marker stutter ratio with the bin set in the Identifiler_Plus_Bins_v1X folder.



- 10. View the imported marker stutters in the navigation pane:
 - a. Select the **Identifiler_Plus_Panel_v1X** folder to display its list of markers in the right pane.
 - b. Double-click the **Identifiler_Plus_Panel_v1X** folder to display its list of markers below it.
 - c. Double-click **D5S818** to display the Stutter Ratio & Distance view for the marker in the right pane.



11. Click **Apply**, then **OK** to add the AmpFℓSTR[™] Identifiler[™] Plus Kit panels, bin sets, and marker stutter to the GeneMapper[™] *ID-X* Software database.

IMPORTANT! If you close the Panel Manager without clicking **Apply**, the panels, bin sets, and marker stutter will not be imported into the GeneMapperTM *ID-X* Software database.

Import an analysis method

Use the following procedure to import the analysis method for the AmpF ℓ STRTM IdentifilerTM Plus PCR Amplification Kit from the folder that you downloaded from our web site, into the GeneMapperTM *ID-X* Software database. Refer to **step 1 on page 61** for downloading instructions.

Note: The Identifiler_Plus_AnalysisMethod_v1x has been provided to assist you in getting started with Identifiler Plus kit data analysis. Analysis parameters should be established by each individual laboratory based on the laboratory's internal validation studies.

- 1. Select **Tools** → **GeneMapper**[™] **ID-X Manager** to open the GeneMapper *ID-X* Manager.
- 2. Import an analysis method:
 - a. Select the Analysis Methods tab, then click Import.

💕 GeneM	lapper® ID-X Ma	nager					×
	Find N	ame Containing:					
Projects	Analysis Methods	Table Settings	Plot Settings Matric	es Size Standards	Report Settings		
N	ame		Last Saved	Owner	Instrument	Analysis Type	Description
A	mpFLSTR Analysis M	lethod	2007-12-18 16:18:2	gmidx		HID	
			1	1	1	1	
<u>N</u> ew	Open	Save As	Import	Export			Delete
						Help	Done

- b. Navigate to, then open the Identifiler Plus Analysis Files GMIDX folder.
- c. Select Identifiler_Plus_AnalysisMethod_v1X, then click Import to import Identifiler_Plus_AnalysisMethod_v1X into the GeneMapper[™] *ID-X* Software database.

💕 Import Analy	sis Method				×
Look in	i 🛅 Identifiler	Plus Analysis Files GMIDX	•	ø 🕫 📰	
My Recent Documents Desktop	CE_G5_Id	entifiler_Plus_65500.xml _Plus_AnalysisMethod_v1X.xml			
My Documents	File name: Files of type:	Identifiler_Plus_AnalysisMethod_v1X.xml XML Files (*.xml)		Import Cancel	

3. To view the settings for Identifiler_Plus_AnalysisMethod_v1X, select the Analysis Methods tab, then select Identifiler_Plus_AnalysisMethod_v1X in the Name column and click Open.

💕 GeneMapper® ID-	X Manager					x
	Find Name Containing:					
Projects Analysis Met	hods Table Settings Pla	ot Settings Matrices	Size Standards Repo	rt Settings		
Name		Last Saved	Owner	Instrument	Analysis Type	
Default		2009-08-31 11:27:	45.695 gmidx		HID	
Identifiler_Plus_	AnalysisMethod_v1X	2009-09-02 11:13:	05.191 gmidx		HID	T
•		-	-			
New Open.	Save As	Import	kport		Delete	
					Help Done	

Figure 10 Analysis Method Editor: General tab settings

Figures 11 through 14 below show the settings for each tab of the Analysis Method Editor.

neral Allele Peak Detector Bin Set: Identifiler_Plus_Bin	Peak Q	juality SC) & GQ Sett	ings	
🔽 Use marker-specific stut!	ter ratio	and distar	ice if availal	ble	
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]			
Range Filter				Factor	y Defaults

Figure 11 Analysis Method Editor: Allele tab settings

- GeneMapper[™] *ID-X* Software 1.0.1, 1.1, or 1.1.1 allows you to specify four types of marker repeat motifs: tri, tetra, penta, and hexa. You can enter parameter values for each type of repeat in the appropriate column.
- The "Use marker-specific stutter ratio if available" check box is selected by default. Consequently, the software applies the stutter ratio filters supplied in the Identifiler_Plus_Stutter_v1X file.

For more information about allele filters, refer to:

- GeneMapper[™] *ID-X* Software *Version 1.0 Getting Started Guide* (PN 4375574)
- GeneMapper[™] *ID-X* Software *Version 1.0 Quick Reference Guide* (PN 4375670)
- GeneMapper[™] *ID-X* Software *Version 1.0 Reference Guide* (PN 4375671)

Analysis Method Editor	×
Analysis Method Editor General Allele Peak Detector Peak Quality Peak Detection Algorithm: Advanced Ranges All Sizes Image: Image	SQ & GQ Settings Peak Detection Peak Amplitude Thresholds: B: 50 G: 50 G: 50 Y: 50 Min. Peak Half Width: 2 Polynomial Degree: 3 Peak Window Size: 15 Slope Threshold 0.0 Peak End: 0.0
	Factory Defaults
Save As Save	Cancel Help

Figure 12 Analysis Method Editor: Peak Detector tab settings

The software uses the peak amplitude threshold parameters to specify the minimum peak height to limit the number of detected peaks. Although GeneMapperTM *ID-X* Software displays peaks that fall below the specified amplitude in electropherograms, the software does not label or determine the genotype of these peaks.

IMPORTANT! Laboratories need to perform the appropriate internal validation studies to determine the peak amplitude thresholds that allow for reliable interpretation of AmpFℓSTR[™] Identifiler[™] Plus PCR Amplification Kit data.

For more information about peak detection algorithms, refer to:

- GeneMapper[™] *ID-X* Software *Version 1.0 Getting Started Guide* (PN 4375574)
- GeneMapper[™] *ID-X* Software *Version 1.0 Quick Reference Guide* (PN 4375670)
- GeneMapper[™] *ID-X* Software *Version 1.0 Reference Guide* (PN 4375671)

Analysis Method Editor	×
General Allele Peak Detector Peak Quality	SQ & GQ Settings
Min/Max Peak Height (LPH/MPH)	
Homozygous min peak height	100.0
Heterozygous min peak height	50.0
Max Peak Height (MPH)	8000.0
- 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
	Factory Defaults
Save As Save	Cancel Help

Figure 13 Analysis Method Editor: Peak Quality tab settings

IMPORTANT! Laboratories need to perform the appropriate internal validation studies to determine the minimum heterozygous and homozygous minimum peak height thresholds and the minimum peak height ratio threshold that allow for reliable interpretation of AmpF*l*STR[™] Identifiler[™] Plus PCR Amplification Kit data.

nalysis Method Editor	>
General Allele Peak Detector Peak Quality 5Q & GQ Settings	
Quality weights are between 0 and 1. Sample and Control GQ Weighting	
Broad Peak (BD) 0.8 Allele Number (AN) 1 Out of Bin Allele (BIN) 0.8 Low Peak Height (LPH) 0 Overlap (OVL) 0.8 Max Peak Height (MPH) 0 Marker Spike (SPK) 0.3 Off-scale (OS) 0 Peak Height Ratio (PHR) 0	1.0).3).3).8).3
Control Concordance (CC) Weight = 1.0 (Only applicable to controls) SQ Weighting Broad Peak (BD) Allelic Ladder GQ Weighting	
Spike (SSPK/SPK) 1 Off-scale (OS) SQ & GQ Ranges	1
Pass Range: Low Quality Range: Sizing Quality: From 0.75 to 1.0 From 0.0 to 0.25 Genotype Quality: From 0.75 to 1.0 From 0.0 to 0.25	
Reset De <u>f</u> au	ults
Save As Save Cancel Help	

Figure 14 Analysis Method Editor: SQ and GQ tab settings

Import a HID size standard Uses the fo

The size standard for the AmpFℓSTR[™] Identifiler[™] Plus PCR Amplification Kit uses the following GeneScan 500 LIZ size standard peaks in its sizing algorithm: 75, 100, 139, 150, 160, 200, 300, 350, 400, and 450.

Use the following procedure to import the size standard for the AmpF ℓ STRTM IdentifilerTM Plus PCR Amplification Kit from the folder that you downloaded from our web site, into the GeneMapperTM *ID-X* Software database. Refer to **step 1 on page 61** for downloading instructions.

1. Select **Tools** → **GeneMapper**[™] **ID-X Manager** to open the GeneMapper[™] *ID-X* Manager.

- 2. Import a size standard:
 - a. Select the Size Standards tab, then click Import.

💕 GeneMapper® ID-X Manager					×		
Find Name Containing:							
Projects Analysis Methods Table :	Settings Plot Setting	gs Matrices Size S	tandards Report Se	ettings			
Name	Last Saved	Owner	Туре	Description			
CE_F_HID_GS500 (75-400)	2007-08-09 13:23:5	gmidx	Advanced		-		
CE_F_HID_G5500 (75-450)	2007-08-09 13:24:0	gmidx	Advanced				
CE_G5_HID_GS500	2006-10-11 13:12:2	gmidx	Advanced		•		
New Open Sa	New Open Saye As Import Export Delete						
				Help	Done		

- b. Navigate to, then open the Identifiler Plus Analysis Files GMIDX folder.
- c. Select **CE_G5_Identifiler_Plus_GS500**, then click **Import** to import the CE_G5_Identifiler_Plus_GS500 analysis method into the GeneMapper[™] *ID-X* Software database.

💕 Import Size S	tandard Method 🔀
Look in:	i 🗀 Identifiler Plus Analysis Files GMIDX 💿 🏂 🏂 📰 📰
My Recent Documents Oesktop	CE_G5_Identifiler_Plus_GS500.xml Identifiler_Plus_AnalysisMethod_v1X.xml
My Documents	File name: CE_G5_Identifiler_Plus_GS500.xml Import Files of type: XML Files (*.xml)
Analyze and edit sample files with GeneMapperTM *ID- X* Software

- Analyze a project
- 1. In the Project window, select File ➤ Add Samples to Project, then navigate to the disk or directory containing the sample files.
- 2. Apply analysis settings to the samples in the project.

Parameter	Settings
Sample Type	Select the sample type.
Analysis Method	Identifiler_Plus_AnalysisMethod_v1X
Panel	Identifiler_Plus_Panel_v1X
Size Standard [‡]	CE_G5_Identifiler_Plus_GS500 ^{§#}

[‡] For more information about how the Size Caller works, refer to the ABI PRISM[™] GeneScan[™] Analysis Software for the Windows NT[™] Operating System Overview of the Analysis Parameters and Size Caller User Bulletin (PN 4335617).

§ The following fragments are defined for the CE_G5_Identifiler_Plus_GS500 size standard provided with the AmpFSTR[™] Identifiler[™] Plus Kit: 75, 100, 139, 150, 160, 200, 300, 350, 400, and 450. For additional information about size standards, refer to the *GeneMapper[™] ID Software Version 3.1 Human Identification Analysis User Guide* (PN 4338775), Appendix D.

Neither the 250-nt nor the 340-nt peak are included in the size standard definition. These peaks can be used as an indicator of precision within a run.

- 3. Click ► (Analyze), enter a name for the project (in the Save Project dialog box), then click **OK** to start analysis.
 - The status bar displays the progress of analysis as a completion bar extending to the right with the percentage completed indicated.
 - The table displays the row of the sample currently being analyzed in green (or red if analysis failed for the sample).
 - The Analysis Summary tab (see figure on next page) is displayed upon completion of the analysis.

💕 GeneMapper® ID-X - 09-02-09 Dickens	s Analysis Method Example - gmidx Is Logg	ed In Database FOSWA	NGD1L03	
File Edit Analysis View Tools Admin Help				
😂 😂 🔡 🛛 🍢 🚰 🗍 🛄 🖭 🔤	🛄 📄 🃰 📄 🕨 💕 🛛 Table Set	ting: 31XX Data Analysis) 🖨 🖻 🖉 🥝
⊡	Samples Analysis Summary Genotypes			
⊡ Identifiler Plus Analysis Examples	Analysis Summary			
	Select run folder to display: Identifiler Plus	Analysis Examples	Y	
	Sample Status	To	otal # of Samples	
	🕎 Unanalyzed		0	
	Analyzed		39	
	Malysis Setting Changed		0	
Click a link below to display a filtered Samples Table containing only the samples selected. Allelic Ladder Quality per run folder (based on SQ and CGQ only)				
	Run Folder	Total # of Analyzed La	adders 🔳 🍐	s 🧶
	Identifiler Plus Analysis Examples	<u>3</u>	<u>3</u> 0	0
	Control Quality per project (based on s	ample PQVs: SOS, SSP	K, MIX, OMR, SQ, CGQ)	
	Control Type	Total # of Samples	All thresholds met	One or more thresholds not met
	Positive Control	3	3	0
	Custom Control	0	0	0
	Negative Control	4	4	0
	local		<u> </u>	0
	Sample Quality per project (based on s	ample PQVs: SOS, SSP	K, MIX, OMR, SQ, CGQ)	
		Total # of Samples	All thresholds met	One or more thresholds not met
	Samples	29	17	12
	•			
Analysis Completed.				

For more information about any of these tasks, refer to:

- GeneMapper[™] *ID-X* Software Version 1.0 Getting Started Guide (PN 4375574)
- GeneMapper[™] ID-X Software Version 1.0 Quick Reference Guide (PN 4375670)
- GeneMapper[™] *ID-X* Software Version 1.0 Reference Guide (PN 4375671)

Examine and edit a project

You can display electropherogram plots from the Samples and Genotypes tabs of the Project window to examine the data. These procedures start with the Analysis Summary tab of the Project window (assuming the analysis is complete).

For more information about any of these tasks, refer to:

- GeneMapper[™] ID-X Software Version 1.0 Getting Started Guide (PN 4375574)
- GeneMapper[™] ID-X Software Version 1.0 Quick Reference Guide (PN 4375670)
- GeneMapper[™] ID-X Software Version 1.0 Reference Guide (PN 4375671)

Part Number 4440211 Rev. F 2/2015

This chapter covers:

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Overview

Experiments using the AmpFℓSTR [™] Identifiler [™] Plus Kit	This chapter provides results of the developmental validation experiments performed using the AmpFℓSTR [™] Identifiler [™] Plus PCR Amplification Kit.
Importance of validation	Validation of a DNA typing procedure for human identification applications is an evaluation of the procedure's efficiency, reliability, and performance characteristics. By challenging the procedure with samples commonly encountered in forensic and parentage laboratories, the validation process uncovers attributes and limitations which are critical for sound data interpretation in casework (Sparkes, Kimpton, Watson <i>et al.</i> , 1996; Sparkes, Kimpton, Gilbard <i>et al.</i> , 1996, and Wallin <i>et al.</i> , 1998).
Experiment conditions	Experiments to evaluate the performance of the AmpFℓSTR [™] Identifiler [™] Plus PCR Amplification Kit were performed. The experiments were performed according to the DNA Advisory Board (DAB) Quality Assurance Standards, effective October 1, 1998 (DNA Advisory Board, 1998). The DAB standards describe the quality assurance requirements that a laboratory should follow to ensure the quality and integrity of the data and competency of the laboratory.
	Additional validation was performed according to the revised guidelines from the Scientific Working Group on DNA Analysis Methods (SWGDAM, July 10, 2003). Based on these guidelines, we conducted experiments that comply with guidelines 1.0 and 2.0 and its associated subsections. This DNA methodology is not novel. (Moretti <i>et al.</i> , 2001, Frank <i>et al.</i> , 2001, Wallin <i>et al.</i> , 2002, and Holt <i>et al.</i> , 2000).
	This chapter discusses many of the experiments performed by us and provides examples of results obtained. We chose conditions that produced optimum PCR product yield and that met reproducible performance standards. While these experiments are not exhaustive, they are appropriate for a manufacturer of STR kits intended for forensic and/or parentage testing use.
	IMPORTANT! Each laboratory using the AmpF ℓ STR TM Identifiler TM Plus PCR Amplification Kit must perform internal validation studies.

Developmental validation

SWGDAM guideline 1.2.1	"Developmental validation is the demonstration of the accuracy, precision, and reproducibility of a procedure by the manufacturer, technical organization, academic institution, government laboratory, or other party." (SWGDAM, July 2003)
SWGDAM guideline 2.10.1	"The reaction conditions needed to provide the required degree of specificity and robustness must be determined. These include thermal cycling parameters, the concentration of primers, magnesium chloride, DNA polymerase, and other critical reagents." (SWGDAM, July 2003)
PCR components	We examined the concentration of each component of the AmpF/STR ^{$^{\text{M}}$} Identifiler ^{$^{\text{M}}$} Plus PCR Amplification Kit. The concentration for each individual component was established to be in the window that meets the reproducible performance characteristics of specificity and sensitivity. For example, various magnesium chloride concentrations were tested on the Applied Biosystems ^{$^{\text{M}}$} 3130 <i>xl</i> Genetic Analyzer. The amplification of 1.0 ng of the Control DNA 9947A for 28-cycles is shown in Figure 15. We observed that the performance of the multiplex is robust within a ±30% window of the optimum magnesium chloride concentration.
	+ 20%



Figure 15 1.0 ng of Control DNA 9947A amplified with the AmpFℓSTR[™] Identifiler[™] Plus Kit for 28-cycles in the presence of varying concentrations of magnesium chloride and analyzed on the Applied Biosystems[™] 3130*x*/ Genetic Analyzer

Thermal cycler parameters Thermal cycling parameters were established for amplification of the AmpFℓSTR[™] Identifiler[™] Plus PCR Amplification Kit. Thermal cycling times and temperatures of GeneAmp[™] PCR systems were verified. Varying annealing/extension and denaturation temperature windows were tested to verify that a specific PCR product with the desired sensitivity of at least 1.0 ng of AmpFℓSTR Control DNA 9947A was produced.

For example, annealing/extension temperatures were tested at 55, 57, 59, 61, and 63 °C for 3-minute hold times in the Silver 96-Well GeneAmpTM PCR System 9700 (Figure 16). The PCR products were analyzed using the Applied BiosystemsTM 3130*xl* Genetic Analyzer.

Of the tested annealing/extension temperatures, 55 to 61 °C produced robust profiles. At 63 °C the yield of the majority of loci was significantly reduced. Routine thermal cycler calibration is recommended when you follow the amplification protocol. No preferential amplification was observed at the standard annealing/extension temperature of 59 °C.



Figure 16 Electropherograms obtained from amplification of 1.0 ng of Control DNA 9947A at annealing/extension temperatures of 55 °C, 57 °C, 59 °C, 61 °C, and 63 °C, analyzed on the Applied Biosystems[™] 3130x/ Genetic Analyzer, Y-axis scale (0 to 4,000 RFUs)

PCR cycle AmpFℓSTR[™] Identifiler[™] Plus PCR Amplification Kit reactions were amplified for 26, 27, 28, 29, and 30 cycles on the Silver 96-Well GeneAmp[™] PCR System 9700 using 1.0 ng from three DNA samples. As expected, the amount of PCR product increased with the number of cycles. A full profile was generated at 26 cycles and off-scale data were collected for several allele peaks at 30 cycles (Figure 17).



Although none of the cycle numbers tested produced nonspecific peaks, 28 cycles was found to give optimal peak heights with 1 ng of DNA input when the amplified products were examined on Applied BiosystemsTM 3130*xl* Genetic Analyzers.

Figure 17 Representative AmpFℓSTR[™] Identifiler[™] Plus Kit profiles obtained from amplification of 1.0 ng DNA template using 26, 27, 28, 29, and 30 cycles, analyzed on the Applied Biosystems[™] 3130*x*/ Genetic Analyzer, Y-axis scale, (0 to 4,000 RFUs)

Accuracy, precision, and reproducibility

SWGDAM "The extent to which a given set of measurements of the same sample agree with their mean and the extent to which these measurements match the actual values being measured should be determined." (SWGDAM, July 2003)

Accuracy Laser-induced fluorescence detection of length polymorphism at short tandem repeat loci is not a novel methodology (Holt *et al.*, 2000 and Wallin *et al.*, 2002). However, accuracy and reproducibility of AmpFℓSTR Identifiler[™] Plus profiles have been determined from various sample types. Figure 18 shows the size differences that are typically observed between sample alleles and allelic ladder alleles on the Applied Biosystems[™] 3130 𝔄 Genetic Analyzer with POP-4[™] polymer. The x-axis in Figure 18 represents the nominal nucleotide sizes for the AmpFℓSTR Identifiler[™] Plus Allelic Ladder. The dashed lines parallel to the x-axis represent the ±0.25-nt windows. The y-axis represents the deviation of each sample allele size from the corresponding allelic ladder allele size. All sample alleles are within ±0.5 nt from a corresponding allele in the allelic ladder.



Figure 18 Size deviation of 200 samples analyzed on the Applied Biosystems[™] 3130*xI* Genetic Analyzer. For each sample, 1.0 ng of DNA was amplified for 28 PCR cycles.

Precision and
size windowsSizing precision allows for determining accurate and reliable genotypes. Sizing
precision was measured on the Applied BiosystemsTM 3130xl Genetic Analyzer.
The recommended method for genotyping is to employ a ± 0.5 -nt "window"
around the size obtained for each allele in the AmpFlSTR IdentifilerTM Plus Allelic
Ladder. A ± 0.5 -nt window allows for the detection and correct assignment of
alleles. Any sample allele that sizes outside the specified window could be:

- An "off-ladder" allele, that is, an allele of a size that is not represented in the AmpFlSTR Identifiler[™] Plus Allelic Ladder
- or
 - An allele that does correspond to an allelic ladder allele, but whose size is just outside a window because of measurement error

The measurement error inherent in any sizing method can be defined by the degree of precision in sizing an allele multiple times. Precision is measured by calculating the standard deviation in the size values obtained for an allele that is run in several injections on a capillary instrument or in several lanes of one gel.

Table 4 on page 84 shows typical precision results obtained from five runs (16 capillaries/run) of the AmpFtSTR^M Identifiler^M Plus Allelic Ladder on the Applied Biosystems^M 3130*xl* Genetic Analyzer (36-cm capillary and POP-4^M polymer). The internal size standard that was used was GeneScan^M 500 LIZ^M Size Standard. The results were obtained within a set of injections on a single capillary array.

Sample alleles may occasionally size outside of the ± 0.5 -nt window for a respective allelic ladder allele because of measurement error. The frequency of such an occurrence is lowest in detection systems having the smallest standard deviations in sizing. **Figure 18 on page 82** illustrates the tight clustering of allele sizes obtained on the Applied Biosystems^M 3130xl Genetic Analyzer, where the standard deviation in sizing is typically less than 0.15 nt. The instance of a sample allele sizing outside the ± 0.5 -nt window because of measurement error is relatively rare when the standard deviation in sizing is approximately 0.15 nt or less (Smith, 1995).

For sample alleles that do not size within a ± 0.5 -nt window, the PCR product must be rerun to distinguish between a true off-ladder allele versus measurement error of a sample allele that corresponds with an allele in the allelic ladder. Repeat analysis, when necessary, provides an added level of confidence to the final allele assignment.

GeneMapper^{$^{\text{M}}$} *ID* Software and GeneMapper^{$^{\text{M}}$} *ID-X* Software automatically flag sample alleles that do not size within the prescribed window around an allelic ladder allele by labelling the allele as OL (off-ladder).

Maximum precision is obtained with a set of capillary injections on each of the supported platforms however the determined allele sizes will vary between the different platforms. Cross-platform sizing differences occur from a number of factors including type and concentration of polymer, run temperature, and electrophoresis conditions. Variations in sizing can also occur between runs on the same instrument and between runs on different instruments of the same platform type because of these factors.

We strongly recommend that the allele sizes be compared to the sizes obtained for known alleles in the AmpF ℓ STRTM IdentifilerTM Plus Allelic Ladder from the same run and then be converted to genotypes (as described in **"Before you start"** on **pages 46** and **60**). See Table 4 for the results of five runs of the AmpF ℓ STRTM IdentifilerTM Plus Allelic Ladder. For more information on precision and genotyping, see Lazaruk *et al.*, 1998 and Mansfield *et al.*, 1998.

In Table 4, the mean sizes for all the alleles in each run (16 capillaries) were calculated. The mean range shown in the table represents the lowest- and highest-mean size values obtained across all five runs. Similarly, the standard deviation for the allele sizing was calculated for all the alleles in each run. The standard deviation range shown in Table 4 represents the lowest and highest standard deviation values obtained across all five runs.

Applie	Applied Biosystems [™] 3130x/ Genetic Analyzer			
Allele	Mean	Standard Deviation		
Amelogenin				
Х	106.03-106.13	0.033–0.045		
Y	111.69–111.8	0.03-0.042		
	CSF1PO			
6	303.99–304.12	0.041-0.063		
7	308.04–308.17	0.037–0.058		
8	312.1–312.2	0.039–0.065		
9	316.13–316.25	0.035–0.045		
10	320.18-320.3	0.034–0.055		
11	324.24-324.34	0.03–0.046		
12	328.3–328.39	0.025–0.047		
13	332.36-332.44	0.032–0.037		
14	336.39–336.49	0.024–0.039		
15	340.42–340.53	0.038–0.05		
D13S317				
8	216.36-216.48	0.031–0.064		
9	220.34-220.48	0.035–0.051		
10	224.32-224.45	0.034–0.059		
11	228.31-228.45	0.031–0.065		
12	282.42-32.55	0.031–0.063		
13	236.3–236.43	0.038–0.066		
14	240.24–240.37	0.043–0.058		
15	244.23–244.37	0.037–0.066		

Table 4 Precision results of five runs (16 capillaries/run) of the AmpFℓSTR[™] Identifiler[™] Plus Allelic Ladder

Table 4	Precisi	on results o	of five ru	ns (16 c	apillaries	s/run) of
the Amp	FℓSTR™	Identifiler™	Plus All	elic Lad	lder (con	tinued)

Allele	Mean	Standard Deviation	
D16S539			
5	252.01-252.15	0.05–0.06	
8	264–264.15	0.05–0.061	
9	268–268.14	0.05–0.063	
10	272–272.15	0.045-0.059	
11	276.02-276.17	0.04-0.064	
12	280.03-280.18	0.039–0.067	
13	284.05-284.22	0.045–0.06	
14	288.08-288.23	0.044-0.054	
15	292.12-292.26	0.038-0.059	
D18S51			
7	261.8–261.9	0.037-0.049	
9	269.94–270.03	0.037-0.051	
10	274.02–274.12	0.043-0.051	
10.2	276.03-276.13	0.037–0.048	
11	278.11–278.22	0.042-0.059	
12	282.2–282.29	0.037–0.046	
13	286.29–286.39	0.039–0.051	
13.2	288.29–288.4	0.034–0.052	
14	290.38–290.49	0.032-0.051	
14.2	292.39–292.5	0.043-0.057	
15	294.48–294.59	0.029-0.049	
16	298.57–298.69	0.035–0.053	
17	302.69–302.81	0.039–0.056	
18	306.83–306.95	0.039–0.056	
19	310.96–311.07	0.034–0.049	
20	315.08–315.18	0.036-0.045	
21	319.2–319.31	0.036-0.045	
22	323.39–323.5	0.03–0.044	
23	327.46-327.53	0.025-0.052	
24	331.59–331.65	0.026-0.043	
25	335.69–335.76	0.032-0.043	
26	339.8–339.88	0.036-0.047	
27	343.87–343.99	0.031-0.048	

Applied Biosystems[™] 3130x/ Genetic Analyzer

Table 4 Precision results of five runs (16 capillaries/run) of the AmpFℓSTR[™] Identifiler[™] Plus Allelic Ladder *(continued)*

Applied Biosystems 5130X Genetic Analyzer			
Allele	Mean	Standard Deviation	
D19S433			
9	101.25–101.34	0.022–0.034	
10	105.16–105.25	0.028–0.037	
11	109.09–109.17	0.021–0.033	
12	113.04–113.12	0.024–0.036	
12.2	115.06–115.13	0.027–0.035	
13	117.02–117.09	0.026–0.036	
13.2	119.03–119.1	0.027–0.038	
14	121.02–121.07	0.025–0.038	
14.2	123.05-123.1	0.028-0.037	
15	125.03-125.09	0.03–0.041	
15.2	127.08–127.13	0.027–0.04	
16	129.08–129.13	0.031-0.039	
16.2	131.13–131.19	0.023-0.042	
17	133.16–133.21	0.034–0.046	
17.2	135.23–135.28	0.034–0.041	
D21S11			
24	184.41–184.46	0.024-0.042	
24.2	186.39–186.45	0.025-0.04	
25	188.35–188.4	0.024-0.038	
26	192.27-192.34	0.027-0.035	
27	196.21-196.28	0.024–0.039	
28	200.06-200.13	0.026-0.039	
28.2	202.03-202.1	0.026-0.036	
29	204.02-204.09	0.025-0.044	
29.2	206.08-206.14	0.027-0.041	
30	208.06-208.11	0.028-0.04	
30.2	210.03-210.09	0.031-0.037	
31	212.04-212.12	0.031-0.037	
31.2	214.03-214.1	0.023-0.04	
32	216.04-216.11	0.028-0.042	
32.2	218.03-218.09	0.023-0.0351	
33	220.05-220.1	0.031-0.043	
33.2	221.98-222.05	0.033–0.038	
34	224.12-224.18	0.024–0.033	
34.2	226.03-226.09	0.029–0.041	

Applied Biosystems[™] 3130x/ Genetic Analyzer

Table 4	Precisi	on results c	of five runs (16 capilla	aries/run)	of
the Ampl	FℓSTR™	Identifiler™	Plus Allelic	Ladder	(continue	d)

Applied Biosystems [™] 3130 <i>xI</i> Genetic Analyzer			
Allele	Mean	Standard Deviation	
35	228.1-228.17	0.03–0.048	
35.2	230.03-230.09	0.03-0.037	
36	232.02-232.09	0.03-0.047	
37	236.08-236.17	0.026-0.041	
38	240.04–240.1	0.033–0.045	
D2S1338			
15	306.27-306.39	0.033–0.058	
16	310.35–310.47	0.031–0.055	
17	314.39–314.53	0.029-0.042	
18	318.45–318.58	0.029–0.046	
19	322.52-322.63	0.025–0.046	
20	326.58-326.67	0.029–0.039	
21	330.66–330.74	0.034–0.045	
22	334.71–334.8	0.031-0.043	
23	338.74–338.85	0.026-0.045	
24	342.75–342.89	0.026–0.05	
25	346.78-346.92	0.026-0.051	
26	350.77–350.89	0.028-0.049	
27	354.69–354.81	0.026-0.045	
28	358.87–359.01	0.028-0.045	
D3S1358	I I		
12	111.12–111.22	0.024–0.047	
13	115.23–115.32	0.03–0.046	
14	119.2–119.31	0.03-0.044	
15	123.14–123.22	0.031–0.045	
16	127.32-127.41	0.032-0.042	
17	131.54–131.62	0.027-0.039	
18	135.64–135.71	0.021-0.042	
19	139.72–139.81	0.024–0.045	
D5S818	J		
7	133.69–133.75	0.029–0.039	
8	137.8–137.86	0.031–0.037	
9	142.17–142.24	0.022-0.035	
10	146.64–146.71	0.025–0.039	
11	151.05–151.12	0.031-0.043	
12	155.32-155.39	0.028-0.041	

Table 4 Precision results of five runs (16 capillaries/run) of the AmpFℓSTR[™] Identifiler[™] Plus Allelic Ladder *(continued)*

Applied Diosystems 5100/ denetic Analyzei			
Allele	Mean	Standard Deviation	
13	159.55–159.61	0.027-0.051	
14	163.63–163.7	0.032-0.038	
15	167.68–167.74	0.026-0.051	
16	171.7–171.76	0.035–0.04	
D7S820			
6	255.08-255.19	0.029–0.058	
7	259.13-259.22	0.04–0.056	
8	263.16-263.25	0.037–0.053	
9	267.19–267.29	0.046-0.053	
10	271.25–271.34	0.039–0.051	
11	275.28–275.4	0.037-0.06	
12	279.34–279.45	0.034–0.05	
13	283.38–283.49	0.039–0.049	
14	287.44–287.54	0.039–0.051	
15	291.51–291.62	0.043-0.052	
D8S1179			
8	122.49-122.61	0.03–0.044	
9	126.56-126.68	0.037–0.045	
10	130.66–130.76	0.026-0.044	
11	134.8–134.89	0.031-0.041	
12	138.98–139.09	0.019–0.043	
13	143.58–143.68	0.028-0.042	
14	148.03–148.14	0.03–0.046	
15	152.43–152.54	0.025–0.043	
16	156.73–156.83	0.026-0.039	
17	160.93–161.04	0.031-0.042	
18	165.03–165.12	0.024–0.046	
19	169.1–169.2	0.035–0.044	
FGA		-	
17	214.11–214.23	0.041–0.05	
18	218.14–218.26	0.043-0.052	
19	222.17–222.3	0.039–0.054	
20	226.21-226.35	0.044–0.057	
21	230.26–230.38	0.045–0.055	
22	234.29–234.42	0.05–0.058	
23	238.33-238.47	0.038–0.057'	

Applied Biosystems[™] 3130x/ Genetic Analyzer

Table 4	Precisi	on results o	of five runs (16 capilla	aries/run) o	٥f
the Amp	FℓSTR™	Identifiler™	Plus Allelic	Ladder	(continued)

Applied Biosystems 3130x/ Genetic Analyzer					
Allele	Mean	Standard Deviation			
24	242.37–242.52	0.044–0.067			
25	246.42-246.57	0.044–0.056			
26	250.48-250.62	0.038-0.069			
26.2	252.49-252.64	0.046-0.066			
27	254.5-254.65	0.047-0.057			
28	258.55-258.71	0.045-0.064			
29	262.63-262.78	0.049-0.062			
30	266.72-266.88	0.052-0.069			
30.2	268.53-268.7	0.049-0.065			
31.2	272.62-272.78	0.036-0.062			
32.2	276.71-276.86	0.05–0.068			
33.2	280.77–280.94	0.043-0.069			
42.2	317.89–318.06	0.045-0.062			
43.2	322.01-322.16	0.038-0.055			
44.2	326.14-326.27	0.034–0.05			
45.2	330.28-330.39	0.039–0.048			
46.2	334.28-334.4	0.044–0.05			
47.2	338.37–338.49	0.039–0.055			
48.2	342.51–342.66	0.034–0.055			
50.2	350.59–350.76	0.041-0.061			
51.2	354.54–354.7	0.039-0.063			
TH01	1	I			
4	162.72-162.77	0.025–0.04			
5	166.78–166.84	0.027-0.035			
6	170.82–170.87	0.03–0.046			
7	174.83–174.9 0.029				
8	178.84–178.9	0.02-0.046			
9	182.82–182.89	0.027–0.034			
9.3	185.84–185.9	0.022-0.042			
10	186.77–186.83 0.026–0.03				
11	190.71–190.77	0.027-0.034			
12	201.48-201.55	0.026-0.037			
ТРОХ	1	1			
6	221.82-221.91	0.029–0.05			
7	225.8–225.88	0.029-0.053			
8	229.79-229.86	0.034–0.048			

Applied Biosystems[™] 3130*xI* Genetic Analyzer

Applied Biosystems [™] 3130 <i>xI</i> Genetic Analyzer				
Allele	Mean	Standard Deviation		
9	233.77–233.86	0.039–0.048		
10	237.76–237.83	0.025-0.053		
11	241.75–241.83	0.028-0.04		
12	245.78-245.85	0.04–0.049		
13	249.76-249.85	0.036-0.048		
vWA				
11	154.07–154.14	0.028-0.042		
12	158.26–158.34	0.028-0.04		
13	162.42-162.49	0.031-0.043		
14	166.66–166.73	0.031–0.05		
15	170.59–170.66	0.029–0.044		
16	174.62–174.68	0.03–0.04		
17	178.61–178.67	0.028-0.051		
18	182.54–182.61	0.021-0.037		
19	186.5–186.56	0.024–0.043		
20	190.43–190.49 0.028–0			
21	194.29–194.36	0.024-0.044		
22	198.17–198.24	0.023-0.036		
23	202.01-202.09	0.028-0.041		
24	206.36-206.42	0.034–0.041		

Table 4 Precision results of five runs (16 capillaries/run) of the AmpFℓSTR[™] Identifiler[™] Plus Allelic Ladder (continued)

Extra peaks in the electropherogram

Causes of extra peaks

Peaks other than the target alleles may be detected on the electropherogram. Causes for the appearance of extra peaks include stutter products, incomplete 3' A nucleotide addition (at the n-1 position), dye artifacts, and mixed DNA samples (see DAB Standard 8.1.2.2).

Stutter products

A stutter is a well-characterized PCR artifact that refers to the appearance of a minor peak one repeat unit smaller (or less frequently, one repeat larger) than the major STR product (Butler, 2005 and Mulero *et al.*, 2006). Sequence analysis of stutter products at tetranucleotide STR loci has revealed that the stutter product is missing a single tetranucleotide core repeat unit relative to the main allele (Walsh *et al.*, 1996).

The proportion of the stutter product relative to the main allele (percent stutter) is measured by dividing the height of the stutter peak by the height of the main allele peak. Peak heights were measured for samples (n = 500) amplified using the 28-cycle-protocol (DNA input ~1ng) at the loci used in the IdentifilerTM Plus Kit. All data were generated on the Applied BiosystemsTM 3130xl Genetic Analyzer.

Some conclusions from these measurements and observations are:

- For each AmpFℓSTR[™] Identifiler[™] Plus Kit locus, the percent stutter generally increases with allele length, as shown in Figure 19 to Figure 22 on **pages 92** through **94**.
- Smaller alleles display a lower level of stutter relative to the longer alleles within each locus.
- Each allele within a locus displays a percent stutter that is consistent with other alleles in the locus.
- The stutter value for each locus shown in Table 5 on page 94 was determined by taking the mean plus three times the standard deviation. These values are the stutter filter percentages in the Identifiler[™] Plus stutter file; they will be used during the filtering step in the GeneMapper[™] ID Software v3.2.1 or GeneMapper[™] ID-X Software v1.0.1, v1.1, or v1.1.1. Peaks in the stutter position that are above the stutter filter percentage will not be filtered. Peaks in the stutter position that have not been filtered and remain labeled can be further evaluated. For evaluation of mixed samples, see Figure 30 on page 108.
- The percent stutter cannot be accurately measured for allele peaks that are offscale and may appear unusually high relative to the main peak.



Figure 19 Stutter percentages for the D8S1179, D21S11, D7S820, and CSF1PO loci







Figure 21 Stutter percentages for the D19S443, vWA, TPOX, and D18S51 loci



Figure 22 Stutter percentages for the D5S818 and FGA loci

Locus	% Stutter
CSF1PO	9.2041
D13S317	9.9348
D16S539	10.3945
D18S51	13.6799
D19S433	11.2096
D21S11	10.6714
D2S1338	12.4409
D3S1358	12.2719
D5S818	10.0599
D7S820	9.6926
D8S1179	10.3155
FGA	13.028
TH01	4.0813

Table 5 Marker-specific stutter filter percentages for AmpFℓSTR[™] Identifiler[™] Plus PCR Amplification Kit loci[‡]

Locu	IS	% Stutter		
TPO	X	6.3832		
vWA	A	12.446		

Table 5Marker-specific stutter filter percentages forAmpF/STR[™] Identifiler[™] Plus PCR Amplification Kit loci[‡] (continued)

‡ These percentages are used as stutter filters in used in GeneMapper ID v3.2.1 Identifiler_Plus_Panels_v1 and GeneMapper[™] ID-X software v1.0.1, v1.1, or v1.1.1 Identifiler_Plus_Panels_v1x.

Addition of 3' A nucleotide

Many DNA polymerases can catalyze the addition of a single nucleotide (predominately adenosine) to the 3' ends of double-stranded PCR products (Clark, 1988 and Magnuson *et al.*, 1996). This nontemplate addition results in a PCR product that is one nucleotide longer than the actual target sequence. The PCR product with the extra nucleotide is referred to as the "+A" form.

The efficiency of +A addition is related to the particular sequence of the DNA at the 3' end of the PCR product. The Identifiler[™] Plus Kit includes two main design features that promote maximum +A addition:

- The primer sequences have been optimized to encourage +A addition.
- The final extension step is 60 °C for 10 min.

This final extension step gives the DNA polymerase additional time to complete +A addition to all double-stranded PCR products. STR systems (where each allele is represented by two peaks that are one nucleotide apart) that have not been optimized for +A addition may have "split peaks."





Lack of complete +A nucleotide addition may be observed in AmpFℓSTR[™] Identifiler[™] Plus PCR Amplification Kit results when the amount of input DNA is greater than the recommended protocols, because more time is needed for the enzyme to add the +A nucleotide to all molecules as more PCR product is generated. Amplification of too much input DNA may also result in off-scale data.

Artifacts Artifacts and anomalies are seen in all molecular biological systems. Artifacts are typically reproducible while anomalies are non-reproducible, intermittent occurrences that are not observed consistently in a system (for example, spikes and baseline noise). Reproducible artifacts have not been seen in data produced, with either 28 or 29 cycles of amplification, on the genetic analyzers used during developmental validation of the Identifiler[™] Plus Kit.

However, you should consider possible noise and artifacts when interpreting data from the AmpF ℓ STRTM IdentifilerTM Plus PCR Amplification Kit on the Applied BiosystemsTM 3130/3130*xl*, ABI PRISMTM 3100/3100-*Avant*, and ABI PRISMTM 310 Genetic Analyzers.

Characterization of loci

SWGDAM guideline 2.1	"The basic characteristics of a genetic marker must be determined and documented." (SWGDAM, July 2003)				
	This section describes basic characteristics of the 15 loci and the sex-determining marker, Amelogenin, which are amplified with the AmpFℓSTR [™] Identifiler [™] Plus Kit. These loci have been extensively characterized by other laboratories.				
Nature of the polymorphisms	The primers for the Amelogenin locus flank a 6-nucleotide deletion within intron 1 of the X homologue. Amplification results in 107-nt and 113-nt products from the X and Y chromosomes, respectively. (Sizes are the actual nucleotide size according to sequencing results, including 3' A nucleotide addition.) The remaining AmpFℓSTR [™] Identifiler [™] Plus Kit loci are all tetranucleotide short tandem repeat (STR) loci. The length differences among alleles of a particular locus result from differences in the number of 4-nt repeat units.				
	All the alleles in the AmpFℓSTR [™] Identifiler [™] Plus Allelic Ladder, including microvariants, have been subjected to sequencing. In addition, other groups in the scientific community have sequenced alleles at some of these loci (Nakahori <i>et al.</i> , 1991; Puers <i>et al.</i> , 1993; Möller <i>et al.</i> , 1994; Barber <i>et al.</i> , 1995; Brinkmann and Möller, 1995; Barber <i>et al.</i> , 1996; Barber and Parkin, 1996; Brinkmann <i>et al.</i> , 1998; Momhinweg <i>et al.</i> , 1998; Watson <i>et al.</i> , 1998). Among the various sources of sequence data on the AmpFℓSTR [™] Identifiler [™] Plus Kit loci, there is consensus on the repeat patterns and structure of the STRs.				
Inheritance	The Centre d'Etude du Polymorphisme Humain (CEPH) has collected DNA from families of Utah Mormon, French Venezuelan, and Amish descent. These DNA sets have been extensively studied all over the world and are routinely used to characterize the mode of inheritance of various DNA loci. Each family set contains three generations, generally including four grandparents, two parents, and several offspring. Consequently, the CEPH family DNA sets are ideal for studying inheritance patterns (Begovich <i>et al.</i> , 1992).				
	Three CEPH family DNA sets were examined. One nanogram of DNA from each sample was amplified using the Identifiler [™] Plus Kit, followed by analysis using an Applied Biosystems [™] 3130 <i>xl</i> Genetic Analyzer. The families examined included #1333 (9 offspring), #1340 (7 offspring), and #1345 (7 offspring), representing 23 meiotic divisions.				
	In family #1340, we observed two parent/offspring pairs with mutations at locus D8S1179. The genotypes differed by one repeat unit between the two generations. Calculation of a mutation rate based on these data would be inaccurate due to the small sample size. The other parent-offspring allele transfers were in accordance with Mendelian rules.				

Mapping The Identifiler[™] Plus Kit loci Amelogenin, CSF1PO, D2S1338, D3S1358, D5S818, D7S1179, D13S317, D16S539, D18S51, D19S433, D21S11, FGA, TH01, TPOX, and vWA have been mapped, and the chromosomal locations have been published (Nakahori *et al.*, 1991; Edwards *et al.*, 1992; Kimpton *et al.*, 1992; Kong et al., 2004; Mills *et al.*, 1992; Sharma and Litt, 1992; Li *et al.*, 1993; Straub *et al.*, 1993; Barber and Parkin, 1996).

Species specificity

SWGDAM "For techniques designed to type human DNA, the potential to detect DNA from forensically relevant nonhuman species should be evaluated." (SWGDAM, July 2003)

The AmpF/STR[™] Identifiler[™] Plus PCR Amplification Kit provides the required specificity for detecting human alleles.

Nonhuman studies

Nonhuman DNA may be present in forensic casework samples. The data from AmpFℓSTR[™] Identifiler[™] Plus PCR Amplification Kit experiments on nonhuman DNA sources are shown in Figure 24 on page 99.

Figure 24 shows amplification for: Control DNA 9947A (1.0 ng, panel 1), chimpanzee (1.0 ng, panel 2), dog (10 ng. panel 3), cat (10 ng, panel 4), horse (10 ng, panel 5), microbial DNA pool (equivalent to 105 copies of *Candida albicans*, *Staphylococcus aureus*, *Neisseria gonorrhoeae*, *E. coli* 0157:H7, *Bacillus subtilis*, and *Lactobacillus rhamnosus*, panel 6), and the negative control (panel 7). The extracted DNA samples were amplified with the AmpFtSTRTM IdentifilerTM Plus PCR Amplification Kit and analyzed using the Applied BiosystemsTM 3130xl Genetic Analyzer.

- Primates: gorilla, chimpanzee, orangutan, and macaque (1.0 ng each)
- Non-primates: mouse, dog, sheep, rabbit, cat, horse, hamster, rat, chicken, and cow (10 ng each)
- Microorganisms: Candida albicans, Staphylococcus aureus, Escherichia coli, Neisseria gonorrhoeae, Bacillus subtilis, and Lactobacillus rhamnosus (equivalent to 10⁵ copies)

The chimpanzee and gorilla DNA samples produced partial profiles within the 70 to 350 nucleotide region.

The microorganisms, chicken, hamster, mouse, rabbit, and rat did not yield detectable products. Dog, horse, sheep, and cow produced a 98-bp fragment near the amelogenin locus in the PET^{TM} dye.



Figure 24 Representative electropherograms from a species specificity study including positive and non-template controls (NTC) run for 28 PCR cycles

Sensitivity

"When appropriate, the range of DNA quantities able to produce reliable typing results should be determined." (SWGDAM, July 2003)			
The optimal amount of input DNA added to the AmpF/STR TM Identifiler TM Plus PCR Amplification Kit should be between 0.75 and 1.0 ng for 28-cycle amplification. The DNA sample should be quantitated before amplification using a system such as the Quantifiler TM Human DNA Quantification Kit (PN 4343895) or the Quantifiler TM HP (Human Plus) DNA Quantification Kit (PN 4482911). The final DNA concentration should be 0.075 to 0.1 ng/µL so that 0.75 to 1.0 ng of DNA is added to the PCR reaction in a volume of 10 µL. If the sample contains degraded or inhibited DNA, amplification of additional DNA may be beneficial. In Figures 25 and 26, the control DNA 9947A was serially diluted from 1 ng to 0.031 ng. With the 28-PCR-cycle protocol, full profiles (26 alleles) were consistently obtained at 0.125 ng, but occasional partial profiles that are missing anywhere from 1 to 3 alleles were observed at 0.062 ng. With the 29-PCR-cycle protocol, full profiles (26 alleles) were consistently obtained at 0.062 ng, but occasional partial profiles that are missing anywhere from 1 to 5 alleles were observed at 0.031 ng.			
 If too much DNA is added to the PCR reaction, the increased amount of PCR product that is generated can result in: Fluorescence intensity that exceeds the linear dynamic range for detection by the instrument ("off-scale" data) 			
Off-scale data is a problem because:			
 Quantitation (peak height and area) for off-scale peaks is not accurate. For example, an allele peak that is off-scale can cause the corresponding stutter peak to appear higher in relative intensity, thus increasing the calculated percent stutter. 			
 Multicomponent analysis of off-scale data is not accurate. This inaccuracy results in poor spectral separation ("pull-up"). 			
Incomplete +A nucleotide addition			
The sample can be reamplified using less DNA.			
When the total number of allele copies added to the PCR is extremely low, unbalanced amplification of the alleles may occur because of stochastic fluctuation.			
Individual laboratories may find it useful to determine an appropriate minimum peak height threshold based on their own results and instruments using low amounts of input DNA. For information on selecting the appropriate cycle number for your DNA input amount, see "Select the appropriate PCR cycle number" on page 29.			



Figure 25 Effect of amplifying 1 ng, 0.50 ng, 0.25 ng, 0.125 ng, and 0.062 ng of Control DNA 9947A using the 28-PCR-cycle protocol



Figure 26 Effect of amplifying 0.5 ng, 0.25 ng, 0.125 ng, 0.062 ng, and 0.031 ng of Control DNA 9947A using the 29-PCR-cycle protocol

Note that the y-axis scale is magnified for the lower amounts of DNA, analyzed using the Applied BiosystemsTM 3130xl Genetic Analyzer.

Stability

- **SWGDAM** "The ability to obtain results from DNA recovered from biological samples deposited on various substrates and subjected to various environmental and chemical insults has been extensively documented. In most instances, assessment of the effects of these factors on new forensic DNA procedures is not required. However, if substrates and/or environmental and/or chemical insults could potentially affect the analytical process, then the process should be evaluated using known samples to determine the effects of such factors." (SWGDAM, July 2003)
- **Degraded DNA** As the average size of degraded DNA approaches the size of the target sequence, the amount of PCR product generated is reduced. This reduction is due to the reduced number of intact templates in the size range necessary for amplification.

Degraded DNA was prepared to examine the potential for differential amplification of loci. High-molecular-weight Raji DNA was sonicated and incubated with increasing doses of DNase I (0 to 6 Units) for 20 minutes (Bender *et al.*, 2004). The DNA was examined by agarose gel analysis to determine the average size of the DNA fragments at each time point.

One nanogram of degraded DNA was amplified using the AmpFℓSTR[™] Identifiler[™] Plus Kit. As the DNA became increasingly degraded, the loci became undetectable according to size. Preferential amplification was not observed. The loci failed to robustly amplify in the order of decreasing size as the extent of degradation progressed.





Effect of inhibitors hematin

Heme compounds have been identified as PCR inhibitors in DNA samples extracted from bloodstains (DeFranchis *et al.*, 1988; Akane *et al.*, 1994). It is believed that the inhibitor is co-extracted and co-purified with the DNA, and that it subsequently interferes with PCR by inhibiting polymerase activity.

To examine the effects of hematin on the amplification results obtained by the IdentifilerTM Plus Kit, Control DNA 9947A (1 ng input DNA) was amplified with increasing concentrations of hematin for 28 cycles of amplification (Figure 28 on page 103). The concentrations of hematin used were 0 μ M, 100 μ M, 200 μ M, and 300 μ M (see Table 6).



Figure 28 Amplification with the AmpF ℓ STRTM IdentifilerTM Plus Kit in the presence and absence of hematin. Panel 1 corresponds to control samples; panels 2–4 correspond to samples amplified in the presence of 100, 200, and 300 μ M of hematin.

Table 6 Performance in simulated model of hematin inhibition $(n = 3)^{\ddagger}$

Hematin (µM)	Identifiler [™] Plus Kit Alleles detected/Total alleles		
0	26/26, 26/26, 26/26		
100	26/26, 26/26, 26/26		

Hematin (µM)	Identifiler [™] Plus Kit Alleles detected/Total alleles		
200	26/26, 26/26, 26/26		
300	26/26, 26/26, 26/26		

Table 6 Performance in simulated model of hematin inhibition (n = 3) (continued)[‡]

‡ Only those peaks >50 RFUs were counted. A complete profile with Control 9947A DNA yields 26 peaks using the AmpFℓSTR[™] Identifiler[™] Plus Kit

Effect of inhibitors humic acid

Traces of humic acid may inhibit the PCR amplification of DNA evidence collected from soil. In this study, We tested increasing amounts of humic acid in the PCR amplification of 1 ng of Control DNA 9947A with the IdentifilerTM Plus Kit for 28 cycles of amplification (see Figure 29). The concentrations of humic acid tested were 0, 50, 100, and 150 ng/µL (see Table 7).



Figure 29 Amplification with the AmpF/STR[™] Identifiler[™] Plus PCR Amplification Kit in the presence and absence of humic acid. Panel 1 corresponds to control samples; panels 2–4 correspond to samples amplified in the presence of 50, 100, and 150 ng/µL humic acid.

Humic Acid (ng/µL)	ldentifiler [™] Plus Kit
0	26/26, 26/26, 26/26
50	26/26, 26/26, 26/26
100	26/26, 26/26, 26/26
150	26/26, 26/26, 26/26

Table 7 Performance in simulated model of humic acid inhibition $(n = 3)^{\ddagger}$

‡ Only those peaks >50 RFUs were counted. A complete profile with Control 9947A DNA yields 26 peaks using the AmpFℓSTR[™] Identifiler[™] Plus Kit.

Mixture studies

SWGDAM guideline 2.8	"The ability to obtain reliable results from mixed source samples should be determined." (SWGDAM, July 2003)		
	Evidence samples may contain DNA from more than one individual. The possibility of multiple contributors should be considered when interpreting the results. We recommend that individual laboratories assign a minimum peak height threshold based on validation experiments performed in each laboratory to avoid typing when stochastic effects are likely to interfere with accurate interpretation of mixtures.		
Mixture Studies	Evidence samples that contain body fluids and/or tissues originating from more than one individual are an integral component of forensic casework. Therefore, it is essential to ensure that the DNA typing system is able to detect DNA mixtures. Mixed samples can be distinguished from single-source samples by:		
	• The presence of more than two alleles at a locus		
	 The presence of a peak at a stutter position that is significantly greater in percentage than what is typically observed in a single-source sample Significantly imbalanced alleles for a heterozygous genotype 		
	The peak height ratio is defined as the height of the lower peak (in RFU) divided by the height of the higher peak (in RFU), expressed as a percentage. Mean, median, minimum, and maximum peak height ratios observed for alleles in the AmpFtSTR [™] Identifiler [™] Plus PCR Amplification Kit loci in unmixed population database samples are shown in Table 8.		

Locus	Number of Observations (n)	Mean	Median	Minimum	Maximum
Amel	320	90.23	91.29	46.21	99.89
CSF1PO	378	90.37	91.56	67.19	99.96
D13S317	375	90.61	92.09	68.28	99.94
D16S539	389	90.03	91.15	68.11	99.87
D18S51	432	90.06	91.24	62.26	99.94
D19S433	399	90.2	91.2	57.29	99.96
D21S11	428	90.3	91.12	69.05	100
D2S1388	436	90.36	91.56	63.86	100
D3S1358	356	91.39	92.66	62.82	99.94
D5S818	357	91.15	92.21	66.07	100
D7S820	395	90.49	92.07	46.61	99.94
D8S1179	396	91.22	92.7	67.42	100
FGA	429	89.83	91.07	60.38	99.87
TH01	362	91.62	93.04	70.09	100
TPOX	333	91.17	92.18	70.65	100
vWA	414	91.16	92.33	65.22	100

Table 8 Peak height ratios for ~1.0 ng[‡] of input DNA amplified for 28 PCR cycles

‡ Actual DNA input amounts: 0.7 ng to 1.3 ng

If an unusually low peak height ratio is observed for one locus, and there are no other indications that the sample is a mixture, the sample may be reamplified and reanalyzed to determine if the imbalance is reproducible. Possible causes of imbalance at a locus are:

- Degraded DNA
- Presence of inhibitors
- Extremely low amounts of input DNA
- A mutation in one of the primer binding sites
- Presence of an allele containing a rare sequence that does not amplify as efficiently as the other allele

Resolution of genotypes in mixed samples

A sample containing DNA from two sources can comprise (at a single locus) any of the seven genotype combinations (see below).

- Heterozygote + heterozygote, no overlapping alleles (four peaks)
- Heterozygote + heterozygote, one overlapping allele (three peaks)
- Heterozygote + heterozygote, two overlapping alleles (two peaks)
- Heterozygote + homozygote, no overlapping alleles (three peaks)
- Heterozygote + homozygote, overlapping allele (two peaks)
- Homozygote + homozygote, no overlapping alleles (two peaks)
- Homozygote + homozygote, overlapping allele (one peak)

Specific genotype combinations and input DNA ratios of the samples contained in a mixture determine whether or not it is possible to resolve the genotypes of the major and minor component(s) at a single locus.

The ability to obtain and compare quantitative values for the different allele peak heights on Applied Biosystems[™] instruments provides additional valuable data to aid in resolving mixed genotypes. This quantitative value is much less subjective than comparing relative intensities of bands on a stained gel.

Ultimately, the likelihood that any sample is a mixture must be determined by the analyst in the context of each particular case.

Limit of detection of the minor component

Mixtures of two genomic DNA samples were examined at various ratios (0:1, 1:1, 3:1, 7:1, 10:1, 15:1, 1:0). The total amount of genomic input DNA mixed at each ratio was 1 ng. The samples were amplified in a GeneAmpTM PCR System 9700, then electrophoresed and detected using an Applied BiosystemsTM 3130*xl* Genetic Analyzer.

The results of the mixed DNA samples are shown in Figures 30 (28 PCR cycles) and 31 (29 PCR cycles) on page 108 where samples A and B were mixed according to the ratios provided. Using either the 28- or 29-PCR-cycle protocol, the minor component allele calls at non-overlapping loci are highlighted. The amplification of the minor contributor at 3:1, 7:1 (0.875:0.125 ng), and 10:1 (0.9:0.09 ng) mixture ratios was readily typeable. 15:1 (0.9375:0.0625 ng) mixture ratios resulted in full or partial profiles for the minor component. The profiles of these samples are described in Table 9.



Figure 30 Amplification of DNA mixtures at various ratios using the 28-PCR-cycle protocol. Minor allele peaks that do not overlap with the major contributor peaks are highlighted.


Figure 31 Amplification of DNA mixtures at various ratios using the 29-PCRcycle protocol. Minor allele peaks that do not overlap with the major contributor peaks are highlighted.

Table 9	Genotypes of	of mixed	DNA sam	ples
		•••••••		

Locus	Profile Sample A	Profile Sample B
D8S1179	11, 14	10, 11
D21S11	29, 35	31.2, 32.2

Locus	Profile Sample A	Profile Sample B
D7S820	8, 10	11
CSF1PO	8, 10	12
D3S1358	14, 17	15, 16
TH01	7, 8	6, 9.3
D13S317	12, 13	9, 11
D16S539	10, 11	9, 12
D2S1338	17, 23	17, 20
D19S433	11, 17.2	13
vWA	14, 17	17, 19
ΤΡΟΧ	9, 10	8, 11
D18S51	15, 16	13, 14
AMEL	Х, Ү	Х
D5S818	11, 13	8, 10
FGA	19, 25	22, 23

Table 9 Genotypes of mixed DNA samples (continued)

Population Data

SWGDAM	"The distribution of genetic markers in populations should be determined in relevant					
guideline 2.7	population groups." (SWGDAM, July 2003)					
Overview	To interpret the significance of a match between genetically typed samples, you must know the population distribution of alleles at each locus in question. If the genotype of the relevant evidence sample is different from the genotype of a suspects reference sample, then the suspect is excluded as the donor of the biological evidence that was tested. An exclusion is independent of the frequency of the two genotypes in the population.					
	If the suspect and evidence samples have the same genotype, then the suspect is included as a possible source of the evidence sample. The probability that another, unrelated individual would also match the evidence sample is estimated by the frequency of that genotype in the relevant population(s).					
Population samples used in these studies	The AmpFℓSTR [™] Identifiler [™] PCR Amplification Kit, prior to the addition of the D8S1179 degenerate primer, was used to generate the population data provided in this section. Samples were collected from individuals throughout the United States with no geographical preference.					
	African-American					
	357 samples were provided by the Kentucky State Police and the Federal Bureau of Investigation.					
	U.S. Caucasian					
	349 samples were provided by the Kentucky State Police and the Federal Bureau of Investigation.					
	U.S. Hispanic					
	290 samples were provided by the Minnesota Bureau of Criminal Apprehension/Memorial Blood Center of Minneapolis and the Federal Bureau of Investigation.					
	Native American					
	191 samples were provided by the Minnesota Bureau of Criminal Apprehension/Memorial Blood Center of Minneapolis.					
	In addition to the alleles that were observed and recorded in our databases, other					

In addition to the alleles that were observed and recorded in our databases, other alleles have been published or reported by other laboratories (see the STRBase at www.cstl.nist.gov/div831/strbase).

AmpFℓSTR[™] Identifiler[™] Plus Kit allele frequencies

Table 10 shows the AmpFℓSTR[™] Identifiler[™] Plus Kit allele frequencies in four populations, listed as percentages.

Allele	African- American (n = 357)	U.S. Caucasian (n = 349)	U.S. Hispanic (n = 290)	Native American (n = 191)
CSF1PO				
6	+	‡	‡	‡
7	4.62	0.14 [‡]	0.34 [‡]	‡
8	7.56	0.29 [‡]	0.17 [‡]	0.52 [‡]
9	3.78	1.72	0.86 [‡]	8.38
10	27.87	24.21	23.10	30.89
11	20.59	31.91	28.28	21.99
11.3	0.14 [‡]	‡	ŧ	‡
12	29.13	32.81	39.66	32.72
13	5.32	7.31	6.38	4.71
14	0.98	1.43	0.86 [‡]	0.79 [‡]
15	‡	0.29 [‡]	ŧ.	ŧ
D2S1338				
15	0.14 [‡]	‡	‡	ŧ
16	5.32	4.73	2.41	2.62
17	10.78	17.34	21.21	9.95
18	5.60	6.30	4.14	7.07
19	14.15	13.75	22.76	29.58
20	6.02	14.61	13.79	9.69
21	14.01	2.58	2.59	2.38
22	13.17	4.01	7.41	15.18
23	10.78	11.46	11.36	11.78
24	9.80	11.75	8.45	7.85
25	8.12	10.60	5.17	3.14
26	1.96	2.72	0.69 [‡]	0.79 [‡]
27	0.14 [‡]	0.14 [‡]	‡	ŧ
28	+	‡	+	‡
D3S1358				
<11	0.42 [‡]	0.14 [‡]	‡	ŧ
11	+	+	+	0.26 [‡]
12	0.56 [‡]	‡	0.17 [‡]	‡
13	0.70 [‡]	0.29 [‡]	0.17 [‡]	‡
14	12.04	15.76	7.41	6.81
15	30.53	25.36	39.14	40.84
15.2	0.14 [‡]	ŧ	‡	ŧ

 Table 10
 AmpF/STR Identifiler Plus Kit allele frequencies

	African-	U.S.		, Native
Allele	American (n = 357)	Caucasian (n = 349)	U.S. Hispanic (n = 290)	American (n = 191)
16	28.57	22.78	26.72	32.98
17	19.47	18.19	16.03	9.95
18	6.72	16.48	8.97	8.38
19	0.84	1.00	1.03	0.79 [‡]
20	‡	+	0.34 [‡]	‡
D5S818	1	-		
7	0.14 [‡]	+	6.72	15.71
8	5.46	+	0.69 [‡]	‡
9	1.68	4.15	5.17	6.02
10	6.72	5.44	5.17	4.19
11	25.49	39.26	39.14	41.10
12	36.41	35.24	29.31	23.30
13	21.57	15.47	12.59	9.42
14	2.38	0.14 [‡]	0.69 [‡]	0.26 [‡]
15	ŧ	0.29 [‡]	0.18 [‡]	‡
16	ŧ	‡	0.17 [‡]	‡
17	0.14‡	‡	0.17 [‡]	‡
D7S820	1	1		
6	ŧ	0.14 [‡]	0.17 [‡]	‡
7	0.42 [‡]	1.29	1.72	0.52 [‡]
8	18.77	16.48	11.72	13.09
9	13.73	17.62	6.21	8.12
10	34.45	27.22	27.41	21.99
11	19.89	18.05	28.79	28.80
12	10.78	14.76	20.17	24.08
13	1.54	3.72	3.45	3.40
14	0.42 [‡]	0.72	0.34 [‡]	‡
15	ŧ	‡	‡	‡
D8S1179	•	,	· · ·	
8	0.42 [‡]	2.29	0.34 [‡]	0.52 [‡]
9	0.42 [‡]	1.15	0.34 [‡]	0.26 [‡]
10	2.38	9.74	8.45	4.71
11	3.92	6.02	5.86	3.40
12	13.31	14.04	12.07	11.52
13	23.25	32.52	32.93	37.43
14	30.11	21.35	26.21	30.63
15	20.17	9.89	10.86	9.42

 Table 10
 AmpF/STR Identifiler Plus Kit allele frequencies (continued)

Allele	African- American (n = 357)	U.S. Caucasian (n = 349)	U.S. Hispanic (n = 290)	Native American (n = 191)
16	4.62	2.72	2.41	1.57
17	1.12 [‡]	0.29 [‡]	0.52 [‡]	0.52 [‡]
18	0.28 [‡]	‡	‡	‡
19	+	+	‡	+
D13S317				
8	3.08	12.18	9.66	4.97
9	2.52	7.74	21.72	17.80
10	3.78	4.44	9.14	13.61
11	24.51	29.80	23.10	24.35
12	46.22	30.80	20.86	23.04
13	15.41	11.17	10.17	7.85
14	4.34	3.72	5.34	8.12
15	0.14 [‡]	0.14 [‡]	‡	0.26 [‡]
D16S539				
5	‡	‡	‡	‡
8	3.22	1.72	1.72	0.79 [‡]
9	19.05	10.46	9.31	12.30
10	10.92	5.59	15.69	15.45
11	31.51	31.95	30.17	30.89
12	18.77	30.23	29.48	27.75
13	14.85	16.76	11.55	10.73
14	1.54	3.01	2.07	2.09
15	0.14 [‡]	0.29 [‡]	‡	‡
D18S51				
7	‡	‡	‡	‡
9	0.14 [‡]	‡	‡	‡
10	0.28 [‡]	0.86	0.52‡	0.79 [‡]
10.2	0.14 [‡]	‡	‡	‡
11	0.28 [‡]	1.15	1.21 [‡]	‡
12	7.00	13.90	10.34	14.92
13	4.34	12.18	14.48	9.16
13.2	0.42 [‡]	‡	+	‡
14	6.86	16.76	15.52	26.96
14.2	0.28 [‡]	‡	+	‡
15	19.47	13.61	16.55	12.04
16	16.53	13.61	11.72	10.73
17	18.21	12.32	14.14	14.66

 Table 10
 AmpF/STR Identifiler Plus Kit allele frequencies (continued)

Allele	African- American (n = 357)	U.S. Caucasian (n = 349)	U.S. Hispanic (n = 290)	Native American (n = 191)
18	11.90	7.74	6.72	2.62
19	6.02	4.44	4.14	3.93
20	4.90	1.72	2.24	1.83
21	2.10	1.00	1.03	1.31
22	0.70 [‡]	0.43 [‡]	0.52 [‡]	0.79 [‡]
23	0.42 [‡]	0.14 [‡]	0.52 [‡]	0.26 [‡]
24	‡	0.14 [‡]	0.17 [‡]	ŧ
25	‡	‡	0.17 [‡]	‡
26	‡	‡	‡	‡
27	‡	‡	‡	‡
D19S433	I	I	1	
9	‡	0.14 [‡]	0.17 [‡]	‡
10	1.54	‡	‡	‡
11	7.14	0.72	0.52 [‡]	0.52 [‡]
11.2	0.14 [‡]	‡	0.17 [‡]	‡
12	10.78	7.74	6.21	3.14
12.2	6.30	0.57 [‡]	1.90	‡
13	29.83	28.94	16.03	17.80
14	21.01	34.10	31.72	24.87
14.2	4.20	0.86	5.00	3.66
15	4.76	15.76	13.45	13.35
15.2	3.36	2.72	8.79	10.73
16	2.38	4.15	4.31	3.93
16.2	2.38	1.72	2.93	1.83
17	‡	0.29 [‡]	0.17‡	0.79 [‡]
17.2	0.28 [‡]	0.29 [‡]	+	2.88
18.2	0.14 [‡]	0.29 [‡]	‡	1.05 [‡]
D21S11	1	1	1	
24	‡	‡	+	‡
24.2	0.14 [‡]	0.43 [‡]	0.17 [‡]	‡
24.3	‡	‡	‡	‡
25	‡	‡	+	‡
25.2	‡	0.14 [‡]	0.17 [‡]	‡
26	0.14 [‡]	0.14 [‡]	0.17 [‡]	‡
27	5.04	4.58	1.21	0.52 [‡]
28	22.97	16.76	9.14	6.28
28.2	+	‡	+	+

Table 10 AmpF/STR Identifiler Plus Kit allele frequencies (continued)

	African-	11.5		Native
Allele	American (n = 357)	Caucasian (n = 349)	U.S. Hispanic (n = 290)	American (n = 191)
29	19.33	20.49	21.21	16.75
29.2	0.14 [‡]	‡	0.52 [‡]	0.26 [‡]
29.3	0.14 [‡]	‡	‡	‡
30	17.23	25.21	29.31	34.29
30.2	1.40	3.30	2.93	1.83
31	7.98	7.16	6.72	5.76
31.2	7.98	9.46	8.62	18.85
32	1.12	1.43	1.55	0.79 [‡]
32.2	5.88	7.16	12.93	9.69
33	0.56 [‡]	ŧ	‡	0.52 [‡]
33.2	3.78	3.30	4.14	3.66
34	1.26	ŧ	‡	‡
34.1	0.14 [‡]	‡	‡	‡
34.2	0.14 [‡]	0.29 [‡]	0.86‡	0.79 [‡]
35	2.94	ŧ	0.34 [‡]	‡
35.1	0.14 [‡]	‡	‡	‡
35.2	ŧ	0.14 [‡]	ŧ.	‡
36	0.84	ŧ	‡	‡
37	0.28 [‡]	ŧ	‡	‡
38	0.14 [‡]	‡	+	‡
FGA	1	1		I
16	‡	0.14 [‡]	+	+
16.1	0.14 [‡]	‡	+	‡
17	+	0.29 [‡]	0.17 [‡]	‡
17.2	0.14 [‡]	‡	+	+
18	0.70 [‡]	2.72	0.52 [‡]	1.31
18.2	1.40	+	+	‡
19	6.72	6.16	7.07	10.21
19.2	0.28 [‡]	‡	+	+
20	7.00	13.90	7.41	12.30
20.2	‡	0.14 [‡]	+	‡
21	12.89	16.91	14.66	12.83
22	21.57	16.91	17.24	10.47
22.2	0.28 [‡]	1.29	0.34 [‡]	0.26‡
22.3	0.14 [‡]	0.14 [‡]	‡	‡
23	14.99	15.19	11.90	15.97
23.2	0.14 [‡]	0.14 [‡]	0.86 [‡]	0.26 [‡]

 Table 10
 AmpF/STR Identifiler Plus Kit allele frequencies (continued)

·			· ·	•
Allele	African- American (n = 357)	U.S. Caucasian (n = 349)	U.S. Hispanic (n = 290)	Native American (n = 191)
24	17.51	13.75	15.34	15.71
24.2	‡	0.14 [‡]	0.17 [‡]	‡
25	7.98	8.60	14.14	14.14
26	3.50	2.72	6.90	4.45
26.2	ŧ.	‡	ŧ.	0.52
29	0.56 [‡]	‡	ŧ.	‡
30	‡	‡	‡	‡
30.2	0.14 [‡]	‡	‡	‡
31.2	‡	‡	‡	‡
32.2	+	+	+	‡
31.2	‡	‡	‡	‡
33.2	‡	‡	‡	‡
34.2	0.14 [‡]	‡	‡	‡
42.2	+	+	+	‡
43.2	+	+	+	‡
44.2	0.28 [‡]	‡	‡	‡
45.2	‡	‡	‡	0.26 [‡]
46.2	0.14 [‡]	‡	‡	‡
47.2	‡	‡	‡	‡
48.2	0.14 [‡]	‡	‡	‡
50.2	ŧ.	‡	ŧ.	‡
51.2	‡	‡	ŧ.	‡
TH01	I			
4	ŧ	‡	‡	‡
5	0.28 [‡]	0.43 [‡]	0.17 [‡]	‡
6	11.06	20.49	22.76	20.68
7	42.86	21.78	33.62	43.98
8	20.73	11.46	8.45	5.24
8.3	ŧ	0.14 [‡]	ŧ	ŧ
9	12.32	16.19	14.14	6.28
9.3	11.62	29.08	20.34	23.56
10	0.98	0.43 [‡]	0.52 [‡]	0.26 [‡]
11	ŧ	‡	ŧ	ŧ
13.3	0.14 [‡]	+	+	‡
ТРОХ	1	1	1	
6	6.72	0.14 [‡]	0.34 [‡]	‡
7	2.24	‡	0.34 [‡]	0.26 [‡]
	L	1 contract of the second se	1	

Table 10 AmpF/STR Identifiler Plus Kit allele frequencies (continued)

Allele	African- American (n = 357)	U.S. Caucasian (n = 349)	U.S. Hispanic (n = 290)	Native American (n = 191)
8	36.13	53.30	49.66	37.96
9	21.15	11.60	7.24	4.19
10	9.24	4.30	4.66	3.40
11	21.43	25.93	27.24	39.27
12	3.08	4.73	10.52	14.92
13	ŧ	‡	‡	‡
vWA		1		
11	0.28 [‡]	‡	0.17 [‡]	‡
12	ŧ	‡	‡	0.26 [‡]
13	1.26	0.43 [‡]	‡	0.26 [‡]
14	7.14	8.31	6.90	4.45
15	20.03	11.32	10.00	7.07
16	26.75	23.35	34.31	32.98
17	20.59	24.50	21.55	33.51
18	14.71	22.49	18.45	15.45
19	6.72	8.31	7.07	4.71
20	1.96	1.15	1.38	1.05 [‡]
21	0.28 [‡]	+	0.17 [‡]	0.26 [‡]
22	0.28 [‡]	+	+	‡
23	‡	+	‡	‡
24	‡	0.14 [‡]	‡	‡

 Table 10
 AmpF/STR Identifiler Plus Kit allele frequencies (continued)

‡ A minimum allele frequency (0.7% for the African-American database, 0.7% for the U.S. Caucasian database, 0.9% for the U.S. Hispanic database, and 1.3% for the Native American database) is suggested by the National Research Council in forensic calculations.

Low-frequency alleles

Some alleles of the AmpFℓSTR[™] Identifiler[™] Plus Kit loci occur at a low frequency. For these alleles, a minimum frequency (5 divided by 2n, where n equals the number of individuals in the database) was assigned for the AmpFℓSTR Identifiler Plus Kit African-American, Native-American, U.S. Caucasian, and U.S. Hispanic databases, as suggested in the 1996 report of the Committee on DNA Forensic Science

(National Research Council, 1996). These databases are summarized in **Table 10 on page 112**. The minimum reportable genotype frequency at each locus is: 1.19×10^{-4} for the African-American database; 1.19×10^{-4} for the U.S. Caucasian database; 1.70×10^{-4} for the U.S. Hispanic database; and 2.97×10^{-4} for the Native American database [p2 + p(1-p) θ , where $\theta = 0.01$]. Evaluation of Hardy-Weinberg equilibrium Estimates of expected heterozygosity (HExp) were computed as described by Nei, M. (1973) using the program PopGene 1.32. Possible divergence from Hardy-Weinberg expectations (HWE) was tested using various methods:

- By calculating the unbiased estimate of the expected homozygote/heterozygote frequencies (Nei, M., 1978).
- Using chi-square (HW X²p) and likelihood ratio (HW G²p) tests (as implemented in the program PopGene 1.32).
- With an exact test (HW Exact p), which is a Markov chain method, based on 1000 shuffling experiments, to estimate without bias the exact P-value of the Hardy-Weinberg test with multiple alleles (Guo and Thompson, 1992), as implemented in the program GenePop 3.4.
- An inter-class correlation test analysis (Burrows' composite measure of linkage disequilibria between pairs of loci and X² tests for significance [Weir, 1996]) was performed separately in each population to detect any correlations between alleles at any of the pair-wise comparisons of the 15 loci, using the program PopGene 1.32.

Observed heterozygosity (Ho), expected heterozygosity, information content, and tests for detecting departures from Hardy-Weinberg equilibrium are shown for each population in Table 11. While a number of the chi-square tests gave seemingly significant p-values (putatively indicating departures from Hardy-Weinberg equilibrium), chi-squared tests are very sensitive to small expected values (as in the case of multiple rare alleles where the expected number of certain genotypes is 1 or fewer, such as with some of these markers), and can greatly inflate the test statistic in this situation (Weir, 1990). With the exact test, the number of tests with p-value < 0.05 were 0 in the African American and U.S. Caucasian populations, 1 in the U.S. Hispanic population (D8S1179; p=0.0304) and 2 in the Native Americans (D21S11, p=0.0118; D5S818, p=0.0205). These are no more than would be expected by chance. No more alleles were observed to be in linkage disequilibrium than would be expected by chance alone. The average observed heterozygosity across the 15 STR loci was 0.804 in the African American population, 0.792 in the U.S. Caucasian sample population, 0.793 in the Hispanic sample population, and 0.757 in the Native Americans. The most heterozygous locus was FGA (mean observed heterozygosity across all populations of 0.875), and the least heterozygous STR locus was TPOX (mean observed heterozygosity across all populations of 0.677).

Table 11	Hetero	ozygosity and p-	-values for Hardy	/-Weinberg tests	of the
15 Identifil	er [™] Pl	us Kit STR loci i	n four U.S. popu	llations [‡]	
-		African	11.6		Nati

	African- American (n = 357)	U.S. Caucasian (n = 349)	U.S. Hispanic (n = 290)	Native American (n = 191)
CSF1P0		1		1
HW X ² p	0.13649	0.926431	0.951476	0.839278
HW G ² p	0.08902	0.894972	0.918038	0.728023
HW Exact p	0.0762	0.2688	0.5456	0.6148
HExp	0.7829	0.7267	0.7051	0.7398
Но	0.7703	0.7421	0.7138	0.7958

	African- American (n = 357)	U.S. Caucasian (n = 349)	U.S. Hispanic (n = 290)	Native American (n = 191)
D2S1338				<u> </u>
HW X ² p	0.409878	0.537758	0.975972	0.722543
HW G ² p	0.962501	0.407932	0.973054	0.760953
HW Exact p	0.7838	0.3488	0.9794	0.5825
HExp	0.8936	0.8823	0.8529	0.8428
Но	0.8768	0.8653	0.8379	0.801
D3S1358				
HW X ² p	0.947371	0.670787	0.681659	0.087223
HW G ² p	0.907905	0.654776	0.852278	0.175807
HW Exact p	0.2967	0.2814	0.4684	0.0614
HExp	0.7681	0.7986	0.7361	0.7028
Но	0.7955	0.8166	0.7414	0.7382
D5S818				
HW X ² p	0.993751	0.859805	0.944725	0.073002
HW G ² p	0.989776	0.520417	0.979044	0.08025
HW Exact p	0.958	0.462	0.4662	0.0205
HExp	0.7476	0.6931	0.7351	0.7378
Но	0.7479	0.7077	0.7586	0.6806
D7S820				
HW X ² p	0.987668	0.571989	0.336834	0.324754
HW G ² p	0.969887	0.44694	0.687948	0.289733
HW Exact p	0.9818	0.2286	0.4028	0.1276
HExp	0.7758	0.8117	0.7822	0.7858
Но	0.7955	0.7908	0.7862	0.7487
D8S1179				
HW X ² p	0.067164	0.545414	0.047783	0.446248
HW G² p	0.568837	0.275218	0.302937	0.760077
HW Exact p	0.2176	0.3264	0.0304	0.1656
HExp	0.7925	0.8047	0.7853	0.7403
Но	0.7899	0.8424	0.8	0.6806
D13S317				
HW X ² p	0.014379	0.711127	0.353995	0.813948
HW G ² p	0.609389	0.871173	0.190736	0.814681
HW Exact p	0.3818	0.667	0.2415	0.6851
HExp	0.6977	0.7797	0.8251	0.8222
Но	0.6695	0.7364	0.8207	0.8168

Table 11 Heterozygosity and p-values for Hardy-Weinberg tests of the 15 Identifiler^M Plus Kit STR loci in four U.S. populations *(continued)*[‡]

	African- American (n = 357)	U.S. Caucasian (n = 349)	U.S. Hispanic (n = 290)	Native American (n = 191)
D16S539				
HW X ² p	0.433216	0.67702	0.058631	0.996396
HW G ² p	0.482435	0.594871	0.37601	0.981384
HW Exact p	0.3753	0.4328	0.3068	0.9986
HExp	0.7939	0.7632	0.7747	0.7766
Но	0.8263	0.7822	0.7828	0.7853
D18S51				
HW X ² p	0.999844	0.628334	0.999203	0.343027
HW G ² p	1	0.872113	0.999492	0.798859
HW Exact p	0.978	0.0982	0.9152	0.2265
HExp	0.8694	0.8769	0.8761	0.8463
Но	0.8824	0.8682	0.8862	0.8377
D19S433				
HW X ² p	0.91703	0.806717	0.731222	0.810711
HW G² p	0.83419	0.999765	0.975476	0.898389
HW Exact p	0.4517	0.69	0.3475	0.4301
HExp	0.8364	0.7659	0.8310	0.8430
Но	0.8011	0.7622	0.8414	0.822
D21S11				
HW X ² p	0.985687	0.936146	0	0
HW G ² p	1	0.999757	0.999794	0.712937
HW Exact p	0.7627	0.7861	0.6476	0.0118
HExp	0.8585	0.8427	0.8290	0.8003
Но	0.8711	0.8567	0.7931	0.801
FGA				
HW X ² p	0	0.904953	0.263223	0.999686
HW G² p	1	0.999812	0.960137	0.999946
HW Exact p	0.9761	0.4459	0.0891	0.9161
HExp	0.8659	0.8686	0.8751	0.8746
Но	0.8824	0.8854	0.8724	0.8482
TH01				
HW X ² p	0.961911	0.997905	0.649467	0.329461
HW G ² p	0.940414	0.99169	0.617212	0.318591
HW Exact p	0.8286	0.9716	0.4495	0.1377
HExp	0.7323	0.7866	0.7666	0.7016
Но	0.7395	0.7822	0.8103	0.6492

Table 11 Heterozygosity and p-values for Hardy-Weinberg tests of the 15 Identifiler[™] Plus Kit STR loci in four U.S. populations *(continued)*[‡]

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	African- American (n = 357)	U.S. Caucasian (n = 349)	U.S. Hispanic (n = 290)	Native American (n = 191)
ТРОХ				-
HW X ² p	0.765163	0.801518	0.875348	0.333914
HW G ² p	0.611014	0.757735	0.913091	0.229017
HW Exact p	0.7247	0.5775	0.8356	0.0647
HExp	0.7643	0.6311	0.6607	0.6765
Но	0.7563	0.6304	0.6759	0.6178
vWA				
HW X ² p	0.925176	0.005048	0.641684	0.994248
HW G ² p	0.964308	0.218817	0.934427	0.997184
HW Exact p	0.7033	0.0564	0.7066	0.8845
HExp	0.8141	0.8081	0.7818	0.7457
Но	0.8571	0.8138	0.7759	0.7277

Table 11 Heterozygosity and p-values for Hardy-Weinberg tests of the 15 Identifiler[™] Plus Kit STR loci in four U.S. populations *(continued)*[‡]

⁺ HW X² p, probability value of X² test for Hardy-Weinberg equilibrium; HW G² p, probability value of the G-statistic of the Likelihood Ratio test for multinomial proportions; HW Exact p; A Markov chain unbiased exact test to estimate the P-value of the Hardy-Weinberg test with multiple alleles; Hexp, Expected heterozygosity; Ho, observed heterozygosity

Concordance studies We analyzed >500 samples by comparing allele calls between the AmpFℓSTR Identifiler and Identifiler[™] Plus Kits. The genotype data from all the analyzed samples showed 100% concordance between the Identifiler and Identifiler[™] Plus Kits.

Mutation Rate

Estimation of spontaneous or induced germline mutation at genetic loci can be achieved by comparing the genotypes of offspring to those of their parents. From such comparisons the number of observed mutations are counted directly.

In previous studies, genotypes of ten STR loci that were amplified by the AmpF/STR SGM Plus[™] PCR Amplification Kit were determined for a total of 146 parent-offspring allelic transfers (meioses) at the Forensic Science Service, Birmingham, England. One length-based STR mutation was observed at the D18S11 locus; mutations were not detected at any of the other nine STR loci. The D18S11 mutation was represented by an increase of one 4-nt repeat unit, allele 17 was inherited as allele 18 (single-step mutation). The maternal/paternal source of this mutation could not be distinguished.

Additional mutation studies

Additional studies (Edwards *et al.*, 1991; Edwards *et al.*, 1992; Weber and Wong, 1993; Hammond *et al.*, 1994; Brinkmann *et al.*, 1995; Chakraborty *et al.*, 1996; Chakraborty *et al.*, 1997; Brinkmann *et al.*, 1998; Momhinweg *et al.*, 1998; Szibor *et al.*, 1998) of direct mutation rate counts produced:

- Larger sample sizes for some of the AmpFℓSTR[™] Identifiler[™] Plus Kit loci.
- Methods for modifications of these mutation rates (to infer mutation rates indirectly for those loci where the rates are not large enough to be measured directly and/or to account for those events undetectable as Mendelian errors).

Probability of Identity

Table 12 shows the Probability of Identity (PI) values of the AmpFℓSTR[™] Identifiler[™] Plus Kit loci individually and combined.

Locus	African- American	U.S. Caucasian	U.S. Hispanic	Native American
CSF1PO	0.079	0.132	0.141	0.123
D2S1338	0.023	0.027	0.038	0.043
D3S1358	0.097	0.076	0.112	0.158
D5S818	0.104	0.147	0.115	0.110
D7S820	0.085	0.063	0.083	0.081
D8S1179	0.074	0.064	0.089	0.104
D13S317	0.132	0.079	0.056	0.056
D16S539	0.077	0.097	0.090	0.082
D18S51	0.033	0.031	0.031	0.046
D19S433	0.042	0.087	0.049	0.044
D21S11	0.037	0.044	0.047	0.074
FGA	0.034	0.035	0.032	0.031
TH01	0.109	0.079	0.097	0.134
TPOX	0.089	0.188	0.168	0.159
vWA	0.066	0.066	0.080	0.103
Combined	1.31 × 10 ^{−18}	5.01 × 10 ⁻¹⁸	7.65 × 10 ^{−18}	3.62×10^{-17}

Table 12Probability of Identity values for the AmpF/STR Identifiler Plus KitSTR loci

The P_I value is the probability that two individuals selected at random will have an identical AmpF4STR Identifiler Plus Kit genotype (Sensabaugh, 1982). The P_I values for the populations described in this section are then approximately $1/7.64 \times 10^{17}$ (African-American), $1/2.00 \times 10^{17}$ (U.S. Caucasian), $1/1.31 \times 10^{17}$ (U.S. Hispanic), and $1/2.76 \times 10^{16}$ (Native American).

Probability of Paternity Exclusion

Table 13 shows the Probability of Paternity Exclusion (P_E) values of the AmpF ℓ STRTM IdentifilerTM Plus Kit STR loci individually and combined.

Locus	African- American	U.S. Caucasian	U.S. Hispanic	Native American
CSF1PO	0.545	0.496	0.450	0.409
D2S1338	0.748	0.725	0.671	0.399
D3S1358	0.591	0.630	0.495	0.510
D5S818	0.506	0.440	0.525	0.601
D7S820	0.591	0.582	0.574	0.492
D8S1179	0.580	0.680	0.599	0.601
D13S317	0.383	0.487	0.638	0.370
D16S539	0.649	0.566	0.567	0.428
D18S51	0.760	0.731	0.767	0.329
D19S433	0.601	0.531	0.678	0.360
D21S11	0.737	0.708	0.586	0.399
FGA	0.760	0.766	0.739	0.309
TH01	0.492	0.566	0.618	0.646
TPOX	0.521	0.329	0.392	0.687
vWA	0.709	0.625	0.555	0.528
Combined	0.9999996	0.9999992	0.9999990	0.9999527

Table 13Probability of Paternity Exclusion values for the AmpFlSTRIdentifiler Plus Kit loci

The P_E value is the probability, averaged over all possible mother-child pairs, that a random alleged father will be excluded from paternity after DNA typing of the AmpF/STR Identifiler Plus Kit STR loci (Chakraborty and Stivers, 1996).

Follow the actions recommended in Table 14 to troubleshoot problems that occur during analysis.

Table 14 Troubleshooting

Observation	Possible causes	Recommended actions
Faint or no signal from both the 9947A and the DNA test samples at all loci	Incorrect volume or absence of either AmpFℓSTR Identifiler [™] Plus Master Mix or AmpFℓSTR Identifiler [™] Plus Primer Set	Repeat amplification.
	No activation of enzyme	Repeat amplification, making sure to hold reactions initially at 95 °C for 11 min.
	Master Mix not vortexed thoroughly before aliquoting	Vortex Master Mix thoroughly.
	AmpF ℓ STR Identifiler [™] Plus Primer Set exposed to too much light	Store Primer Set protected from light.
	GeneAmp [™] PCR System malfunction	Refer to the thermal cycler user's manual and check instrument calibration.
	Incorrect thermal cycler parameters	Check the protocol for correct thermal cycler parameters.
	Tubes not seated tightly in the thermal cycler during amplification	Push reaction tubes firmly into contact with block after first cycle. Repeat test.
	Wrong PCR reaction tube	Use Applied Biosystems [™] MicroAmp Reaction Tubes with Caps for the GeneAmp [™] PCR System 9700.
	MicroAmp [™] Base used with tray/ retainer set and tubes in GeneAmp [™] PCR System 9700	Remove MicroAmp Base from tray/retainer set and repeat test.
	Insufficient PCR product electrokinetically injected	For ABI PRISM [™] 3100-Avant or Applied Biosystems [™] 3100/3130x/ runs: Mix 1.0 μL of PCR product and 9 μL of Hi-Di [™] Formamide/ GeneScan [™] 500 LIZ [™] solution.
		For ABI PRISM [™] 310 instrument runs: Mix 1.5 μL of PCR product and 25 μL of Hi-Di [™] Formamide/GeneScan [™] 500 LIZ [™] solution.
	Degraded formamide	Check the storage of formamide; do not thaw and refreeze multiple times. Try Hi-Di [™] Formamide.

Table 14	Troubleshooting	(continued)
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Observation	Possible causes	Recommended actions	
Positive signal from AmpFlSTR Control DNA 9947A but partial or no signal from DNA test samples	Quantity of test DNA sample is below assay sensitivity	Quantitate DNA and add 1.0 ng of DNA. Repeat test.	
	Test sample contains high concentration of PCR inhibitor (for example, heme compounds, certain	Quantitate DNA and add minimum necessary vol- ume. Repeat test.	
	dyes	Wash the sample in a Centricon [™] -100 centrifugal filter unit. Repeat test.	
	Test sample DNA is severely degrad- ed	If possible, evaluate the quality of DNA sample by running an agarose gel. If DNA is degraded, reamplify with an increased amount of DNA or use the AmpFℓSTR [™] MiniFiler [™] Kit.	
	Dilution of test sample DNA in water or wrong buffer (for example, TE formula with incorrect EDTA concentration)	Redilute DNA using low TE Buffer (with 0.1 mM EDTA).	
More than one allele present at a locus	Presence of exogenous DNA	Use appropriate techniques to avoid introducing foreign DNA during laboratory handling.	
	Amplification of stutter product	See "Stutter products" on page 90.	
	Mixed sample		
	Incomplete 3' A base addition (n-1 nt position)	See "Addition of 3´ A nucleotide" on page 95. Be sure to include the final extension step of 60 °C for 10 min in the PCR.	
	Signal exceeds dynamic range of instrument (off-scale data)	Ensure cycle number is optimized according to instructions on page 30. Repeat PCR amplification using fewer PCR cycles or use your laboratory's SOP to analyze off-scale data.	
	Poor spectral separation (bad matrix)	Follow the steps for creating a spectral file.	
		Confirm that Filter Set G5 modules are installed and used for analysis.	
	Too much DNA in reaction	Use recommended amount of template DNA (1.0 ng).	
Some but not all loci visible on electropherogram	Test-sample DNA is severely degraded	If possible, evaluate the quality of DNA sample by running an agarose gel. If DNA is degraded, reamplify with an increased amount of DNA or use the AmpFℓSTR [™] MiniFiler [™] Kit.	
	Test sample contains high concentrations of a PCR inhibitor (for example, heme compounds, certain	Quantitate DNA and add minimum necessary volume. Repeat test.	
	dyes)	Wash the sample in a Centricon [™] -100 centrifugal filter unit. Repeat test.	

Table 14Troubleshooting (continued)

Observation	Possible causes	Recommended actions
Poor peak height bal- ance	Incorrect thermal cycler parameters	Check the protocol for correct thermal cycler parameters.
	GeneAmp [™] PCR System 9700 with Aluminum 96-Well block or third-party thermal cyclers	Use Applied Biosystems [™] GeneAmp [™] PCR System 9700 with silver or gold-plated silver blocks only.

B

This appendix covers:

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Chemical safety

Chemical hazard warning



WARNING! CHEMICAL HAZARD. Before handling any chemicals, refer to the Material Safety Data Sheet (MSDS) provided by the manufacturer, and observe all relevant precautions.



WARNING! CHEMICAL HAZARD. All chemicals in the instrument, including liquid in the lines, are potentially hazardous. Always determine what chemicals have been used in the instrument before changing reagents or instrument components. Wear appropriate eyewear, protective clothing, and gloves when working on the instrument.



WARNING! CHEMICAL HAZARD. Four-liter reagent and waste bottles can crack and leak. Each 4-liter bottle should be secured in a low-density polyethylene safety container with the cover fastened and the handles locked in the upright position. Wear appropriate eyewear, clothing, and gloves when handling reagent and waste bottles.



WARNING! CHEMICAL STORAGE HAZARD. Never collect or store waste in a glass container because of the risk of breaking or shattering. Reagent and waste bottles can crack and leak. Each waste bottle should be secured in a low-density polyethylene safety container with the cover fastened and the handles locked in the upright position. Wear appropriate eyewear, clothing, and gloves when handling reagent and waste bottles.

Chemical safety guidelines

To minimize the hazards of chemicals:

- Read and understand the Material Safety Data Sheets (MSDSs) provided by the chemical manufacturer before you store, handle, or work with any chemicals or hazardous materials. (See "About MSDSs" on page 133.)
- Minimize contact with chemicals. Wear appropriate personal protective equipment when handling chemicals (for example, safety glasses, gloves, or protective clothing). For additional safety guidelines, consult the MSDS.
- Minimize the inhalation of chemicals. Do not leave chemical containers open. Use only with adequate ventilation (for example, fume hood). For additional safety guidelines, consult the MSDS.
- Check regularly for chemical leaks or spills. If a leak or spill occurs, follow the manufacturer's cleanup procedures as recommended in the MSDS.
- Comply with all local, state/provincial, or national laws and regulations related to chemical storage, handling, and disposal.

About MSDSs Chemical manufacturers supply current Material Safety Data Sheets (MSDSs) with shipments of hazardous chemicals to new customers. They also provide MSDSs with the first shipment of a hazardous chemical to a customer after an MSDS has been updated. MSDSs provide the safety information you need to store, handle, transport, and dispose of the chemicals safely.

Each time you receive a new MSDS packaged with a hazardous chemical, be sure to replace the appropriate MSDS in your files.

Obtaining
MSDSsThe MSDS for any chemical supplied by Applied Biosystems is available to you free
24 hours a day. To obtain MSDSs:

- 1. Go to www.appliedbiosystems.com, click Support, then select MSDS.
- 2. In the Keyword Search field, enter the chemical name, product name, MSDS part number, or other information that appears in the MSDS of interest. Select the language of your choice, then click **Search**.
- 3. Find the document of interest, right-click the document title, then select any of the following:
 - **Open** To view the document
 - **Print Target** To print the document
 - Save Target As To download a PDF version of the document to a destination that you choose

Note: For the MSDSs of chemicals not distributed by us, contact the chemical manufacturer.

Chemical waste safety

Chemical waste hazards



WARNING! HAZARDOUS WASTE. Refer to Material Safety Data Sheets and local regulations for handling and disposal.



WARNING! CHEMICAL WASTE HAZARD. Wastes produced by Applied Biosystems instruments are potentially hazardous and can cause injury, illness, or death.



WARNING! CHEMICAL STORAGE HAZARD. Never collect or store waste in a glass container because of the risk of breaking or shattering. Reagent and waste bottles can crack and leak. Each waste bottle should be secured in a low-density polyethylene safety container with the cover fastened and the handles locked in the upright position. Wear appropriate eyewear, clothing, and gloves when handling reagent and waste bottles.

Chem	iical	waste
safety	guic	lelines

To minimize the hazards of chemical waste:

- Read and understand the Material Safety Data Sheets (MSDSs) provided by the manufacturers of the chemicals in the waste container before you store, handle, or dispose of chemical waste.
- Provide primary and secondary waste containers. (A primary waste container holds the immediate waste. A secondary container contains spills or leaks from the primary container. Both containers must be compatible with the waste material and meet federal, state, and local requirements for container storage.)
- Minimize contact with chemicals. Wear appropriate personal protective equipment when handling chemicals (for example, safety glasses, gloves, or protective clothing). For additional safety guidelines, consult the MSDS.
- Minimize the inhalation of chemicals. Do not leave chemical containers open. Use only with adequate ventilation (for example, fume hood). For additional safety guidelines, consult the MSDS.
- Handle chemical wastes in a fume hood.
- After emptying a waste container, seal it with the cap provided.
- Dispose of the contents of the waste tray and waste bottle in accordance with good laboratory practices and local, state/provincial, or national environmental and health regulations.
- **Waste disposal** If potentially hazardous waste is generated when you operate the instrument, you must:
 - Characterize (by analysis if necessary) the waste generated by the particular applications, reagents, and substrates used in your laboratory.
 - Ensure the health and safety of all personnel in your laboratory.

• Ensure that the instrument waste is stored, transferred, transported, and disposed of according to all local, state/provincial, and/or national regulations.

IMPORTANT! Radioactive or biohazardous materials may require special handling, and disposal limitations may apply.

Biological hazard safety

General biohazard



WARNING! BIOHAZARD. Biological samples such as tissues, body fluids, infectious agents, and blood of humans and other animals have the potential to transmit infectious diseases. Follow all applicable local, state/provincial, and/or national regulations. Wear appropriate protective equipment, which includes but is not limited to: protective eyewear, face shield, clothing/lab coat, and gloves. All work should be conducted in properly equipped facilities using the appropriate safety equipment (for example, physical containment devices). Individuals should be trained according to applicable regulatory and company/institution requirements before working with potentially infectious materials. Read and follow the applicable guidelines and/or regulatory requirements in the following:

- U.S. Department of Health and Human Services guidelines published in *Biosafety in Microbiological and Biomedical Laboratories* (stock no. 017-040-00547-4; **bmbl.od.nih.gov**)
- Occupational Safety and Health Standards, Bloodborne Pathogens (29 CFR§1910.1030; www.access.gpo.gov/ nara/cfr/waisidx_01/29cfr1910a_01.html)
- Your company's/institution's Biosafety Program protocols for working with/handling potentially infectious materials.

Additional information about biohazard guidelines is available at:

www.cdc.gov

Chemical alerts

General alerts for all chemicals



Avoid contact with skin, eyes, and/or clothing. Read the MSDS and follow the handling instructions. Wear appropriate protective eyewear, clothing, and gloves.

CAUTION! CHEMICAL HAZARD. AmpFℓSTR[™] Identifiler[™] Plus PCR Amplification Kit may cause eye, skin, and respiratory tract irritation. Sodium azide may react with lead and copper plumbing to form highly explosive metal azides.



WARNING! CHEMICAL HAZARD. POP-4[™] Polymer for 3130/3130*xl* Genetic Analyzers causes skin, eye, and respiratory tract irritation.



WARNING! CHEMICAL HAZARD. Running Buffer, 10× causes skin, eye, and respiratory tract irritation.



WARNING! CHEMICAL HAZARD. Hi-Di[™] Formamide is harmful if swallowed, inhaled or absorbed through skin, and causes irritation to skin, eyes, and respiratory tract. It affects the central nervous system and may affect the reproductive system.



WARNING! CHEMICAL HAZARD. POP-4[™] Polymer for 3100/3100-*Avant* Genetic Analyzers is irritating to eyes, respiratory system, and skin. It causes adverse cardiovascular effects. It contains a known or suspected reproductive toxin and a known or suspected mutagen.

Bibliography

Akane, A., Matsubara, K., Nakamura, H., Takahashi, S., and Kimura, K. 1994. Identification of the heme compound copurified with deoxyribonucleic acid (DNA) from bloodstains, a major inhibitor of polymerase chain reaction (PCR) amplification. *J. Forensic Sci.* 39:362–372.

Barber, M.D., Piercy, R.C., Andersen, J.F., and Parkin, B.H. 1995. Structural variation of novel alleles at the Hum vWA and Hum FES/FPS short tandem repeat loci. *Intl. J. Legal Med.* 108:31–35.

Barber, M.D., McKeown, B.J., and Parkin, B.H. 1996. Structural variation in the alleles of a short tandem repeat system at the human alpha fibrinogen locus. *Intl. J. Legal Med.* 108:180–185.

Barber, M.D. and Parkin, B.H. 1996. Sequence analysis and allelic designation of the two short tandem repeat loci D18S51 and D8S1179. *Intl. J. Legal Med.* 109:62–65.

Baron, H., Fung, S., Aydin, A., Bahrig, S., Luft, F.C., and Schuster, H. 1996. Oligonucleotide ligation assay (OLA) for the diagnosis of familial hypercholesterolemia. *Nat. Biotechnol.* 14:1279–1282.

Begovich A.B., McClure G.R., Suraj V.C., Helmuth R.C., Fildes N., Bugawan T.L., Erlich H.A., and Klitz W. 1992. Polymorphism, recombination, and linkage disequilibrium within the HLA class II region. *J. Immunol.* 148:249–58.

Brinkmann, B., Moller, A. and Wiegand, P. 1995. Structure of new mutations in 2 STR systems. *Intl. J. Legal Med.* 107:201–203.

Brinkmann, B., Klintschar, M., Neuhuber, F., Huhne, J. and Rolf, B. 1998. Mutation rate in human microsatellites: Influence of the structure and length of the tandem repeat. *Am. J. Hum. Genet.* 62:1408–1415.

Budowle, B. *et al.* 1998. CODIS and PCR Based Short Tandem Repeat Loci: Law Enforcement Tools. Second European Symposium on Human Identification. 73–88.

Butler, J.M. 2005. Forensic DNA Typing. Burlington, MA:Elsevier Academic Press.

Chakraborty, R., Stivers, D., and Zhong, Y. 1996. Estimation of mutation rates from parentage exclusion data: applications to STR and VNTR loci. *Mutat. Res.* 354:41–48.

Chakraborty, R. and Stivers, D.N. 1996. Paternity exclusion by DNA markers: effects of paternal mutations. *J. Forensic Sci.* 41:671–677.

Chakraborty, R. Kimmel, M., Stivers, D., Davison, L., and Deka, R. 1997. Relative mutation rates at di-, tri-, and tetranucleotide microsatellite loci. *Proc. Natl. Acad. Sci. USA* 94:1041–1046.

Clark J.M. 1988. Novel non-templated nucleotide addition reactions catalyzed by procaryotic and eucaryotic DNA polymerases. *Nucleic Acids Res.* 16:9677–9686.

DeFranchis, R., Cross, N.C.P., Foulkes, N.S., and Cox, T.M. 1988. A potent inhibitor of Taq DNA polymerase copurifies with human genomic DNA. *Nucleic Acids Res.* 16:10355.

DNA Advisory Board, Federal Bureau of Investigation, U.S. Department of Justice. 1998. Quality assurance standards for forensic DNA testing laboratories.

Edwards, A., Civitello, A., Hammond, H., and Caskey, C. 1991. DNA typing and genetic mapping with trimeric and tetrameric tandem repeats. *Am. J. Hum. Genet.* 49:746–756.

Edwards, A., Hammond, H.A., Lin, J., Caskey, C.T., and Chakraborty, R. 1992. Genetic variation at five trimeric and tetrameric tandem repeat loci in four human population groups. *Genomics* 12:241–253.

Frank, W., Llewellyn, B., Fish, P., *et al.* 2001. Validation of the AmpFℓSTR[™] Profiler Plus[™] PCR Amplification Kit for use in forensic casework. *J. Forensic Sci.* 46:642–646.

Grossman, P.D., Bloch, W., Brinson, E., Chang, C.C., Eggerding, F.A., Fung, S., Iovannisci, D.M., Woo, S., and Winn-Deen, E.S. 1994. High-density multiplex detection of nucleic acid sequences: oligonucleotide ligation assay and sequence-coded separation. *Nucleic Acids Res.* 22:4527–4534.

Guo S.W., and Thompson, E.A. 1992. Performing the exact test of Hardy-Weinberg proportion for multiple alleles. *Biometrics* 48:361–372.

Hammond, H., Jin, L., Zhong, Y., Caskey, C., and Chakraborty, R. 1994. Evaluation of 13 short tandem repeat loci for use in personal identification applications. *Am J. Hum. Genet.* 55:175–189.

Holt, C., Stauffer, C., Wallin, J., Lazaruk, L., Nguyen, T., Budowle, B., and Walsh, P. 2000. Practical applications of genotypic Surveys for forensic STR testing. *Forensic Sci. Int.* 112:91–109.

Kimpton, C., Walton, A., and Gill, P. 1992. A further tetranucleotide repeat polymorphism in the vWF gene. *Hum. Mol. Genet.* 1:287.

Kong, X., Murphy, K., Raj, T., He, C., White, P.S., and Matise, T.C. 2004. A combined linkage-physical map of the human genome. *Am. J. Hum. Genet.* 75:1143–1148.

Kwok, S., and Higuchi, R. 1989. Avoiding false positives with PCR. *Nature* 339:237–238.

Lazaruk, K., Walsh, P.S., Oaks, F., Gilbert, D., Rosenblum, B.B., Menchen, S., Scheibler, D., Wenz, H.M., Holt, C., Wallin, J. 1998. Genotyping of forensic short tandem repeat (STR) systems based on sizing precision in a capillary electrophoresis instrument. *Electrophoresis* 19:86–93.

Li, H. Schmidt, L., Wei, M-H., Hustad, T. Leman, M.I., Zbar, B., and Tory, K. 1993. Three tetranucleotide polymorphisms for loci:D3S1352; D3S1358; D3S1359. *Hum. Mol. Genet.* 2:1327.

Magnuson, V.L., Ally, D.S., Nylund, S.J., Karanjawala, Z.E., Rayman, J.B., Knapp, J.I., Lowe, A.L., Ghosh, S., and Collins, F.S. 1996. Substrate nucleotide-determined non-templated addition of adenine by Taq DNA polymerase: implications for PCR-based genotyping and cloning. *Biotechniques* 21:700–709.

Mansfield, E.S., Robertson, J.M., Vainer, M., Isenberg, A.R., Frazier, R.R., Ferguson, K., Chow, S., Harris, D.W., Barker, D.L., Gill, P.D., Budowle, B., and McCord, B.R. 1998. Analysis of multiplexed short tandem repeat (STR) systems using capillary array electrophoresis. *Electrophoresis* 19:101–107.

Mills, K.A., Even, D., and Murrau, J.C. 1992. Tetranucleotide repeat polymorphism at the human alpha fibrinogen locus (FGA). *Hum. Mol. Genet.* 1:779.

Möller, A., Meyer, E., and Brinkmann, B. 1994. Different types of structural variation in STRs: HumFES/FPS, HumVWA, and HumD21S11. *Intl. J. Legal Med.* 106:319–323.

Momhinweg, E., Luckenbach, C., Fimmers, R., and Ritter, H. 1998. D3S1358: sequence analysis and gene frequency in a German population. *Forensic Sci. Int.* 95:173–178.

Moretti, T., Baumstark, A., Defenbaugh, D., Keys, K., Smerick, J., and Budowle, B. 2001. Validation of short tandem repeats (STRs) for forensic usage: Performance testing of fluorescent multiplex STR systems and analysis of authentic and simulated forensic samples. *J. Forensic Sci.* 46(3):647–660.

Mulero, J.J., Chang, C.W., and Hennessy, L.K. 2006. Characterization of N+3 stutter product in the trinucleotide repeat locus DYS392. *J. Forensic Sci.* 51:826–830.

Nakahori, Y., Takenaka, O., and Nakagome, Y. 1991. A human X-Y homologous region encodes amelogenin. *Genomics* 9:264–269.

National Research Council. 1996. *The evaluation of forensic DNA evidence*. National Academy Press, Washington, D.C.

Nei, M. 1978. Estimation of average heterozygosity and genetic distance from a small number of individuals. *Genetics* 89:583–590.

Nei, M. 1973. Analysis of gene diversity in subdivided populations. *Proc. Natl. Acad. Sci.* USA 70:3321–3323.

Revised Validation Guidelines-Scientific Working Group on DNA Analysis Methods (SWGDAM). *Forensic Science Communications* (July 2004) Volume 6 (3). Available at www.fbi.gov/hq/lab/fsc/current/standards/2004_03_standards02.htm

Puers, C., Hammond, H., Jin, L., Caskey, C., and Schumm, J. 1993. Identification of repeat sequence heterogeneity at the polymorphic short tandem repeat locus HUMTH01 [AATG]n and reassignment of alleles in population analysis using a locus-specific allelic ladder. *Am. J. Hum. Genet.* 53:953–958.

Sensabaugh, G.F. 1982. Biochemical markers of individuality. In: Saferstein, R., ed. *Forensic Science Handbook*. Prentice-Hall, Inc., New York, pp. 338–415.

Sharma, V., and Litt, M. 1992. Tetranucleotide repeat polymorphism at the D21S11 locus. *Hum Mol. Genet.* 1:67.

Smith, R.N. 1995. Accurate size comparison of short tandem repeat alleles amplified by PCR. *Biotechniques* 18:122–128.

Sparkes, R., Kimpton, C., Watson, S., Oldroyd, N., Clayton, T., Barnett, L., Arnold, J., Thompson, C., Hale, R., Chapman, J., Urquhart, A., and Gill, P. 1996a. The validation of a 7-locus multiplex STR test for use in forensic casework. (I). Mixtures, ageing, degradation and species studies. *Int. J. Legal Med.* 109:186–194.

Sparkes, R., Kimpton, C., Gilbard, S., Carne, P., Andersen, J., Oldroyd, N., Thomas, D., Urquhart, A., and Gill, P. 1996b. The validation of a 7-locus multiplex STR test for use in forensic casework. (II), Artifacts, casework studies and success rates. *Int. J. Legal Med.* 109:195–204.

Straub, R.E., Speer, M.C., Luo, Y., Rojas, K., Overhauser, J., Ott, J., and Gilliam, T.C. 1993. A microsatellite genetic linkage map of human chromosome 18. *Genomics* 15:48–56.

Szibor, R., Lautsch, S., Plate, I., Bender, K., and Krause, D. 1998. Population genetic data of the STR HumD3S1358 in two regions of Germany. *Int. J. Legal Med.* 111:160–161.

Wallin, J.M., Buoncristiani, M.R., Lazaruk, K.D., Fildes, N., Holt, C.L., Walsh, P.S. 1998. SWGDAM validation of the AmpFlSTR blue PCR amplification kit for forensic casework analysis. *J. Forensic Sci.* 43:854–870.

Wallin, J.M., Holt, C.L., Lazaruk, K.D., Nguyen, T.H., and Walsh, P.S. 2002. Constructing universal multiplex PCR systems for comparative genotyping. *J. Forensic Sci.* 47:52–65.

Walsh, P.S., Fildes, N.J., and Reynolds, R. 1996. Sequence analysis and characterization of stutter products at the tetranucleotide repeat locus vWA. *Nucleic Acids Res.* 24:2807–2812.

Watson, S., Kelsey, Z., Webb, R., Evans, J., and Gill, P. 1998. The development of a third generation STR multiplex system (TGM). In: Olaisen, B., Brinkmann, B., and Lincoln, P.J., eds. *Progress in Forensic Genetics 7: Proceedings of the 17th International ISFH Congress, Oslo 2-6 September 1997*. Elsevier, Amsterdam, pp. 192–194.

Weber, J. and Wong, C. 1993. Mutation of human short tandem repeats. *Hum. Mol. Genet.* 2:1123–1128.

Weir, B.S. 1996. Genetic data analysis II. Sunderland, MA: Sinauer Associates, Inc.

Documentation

Related documentation

Document title	Part number
Applied Biosystems [™] 3100/3100-Avant Data Collection v2.0 User Guide	4347102
Applied Biosystems [™] 3100/3100-Avant Genetic Analyzers Using Data Collection Software v2.0 User Bulletin	4350218
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