

Penn State University
November 14, 2011 – University Park, PA

Forensic DNA Research at the U.S. National Institute of Standards and Technology

Becky Hill

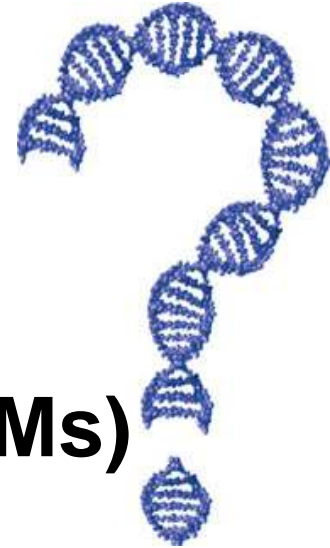
Applied Genetics Group
National Institute of Standards and Technology
Gaithersburg, Maryland



Presentation Outline

Please ask
questions

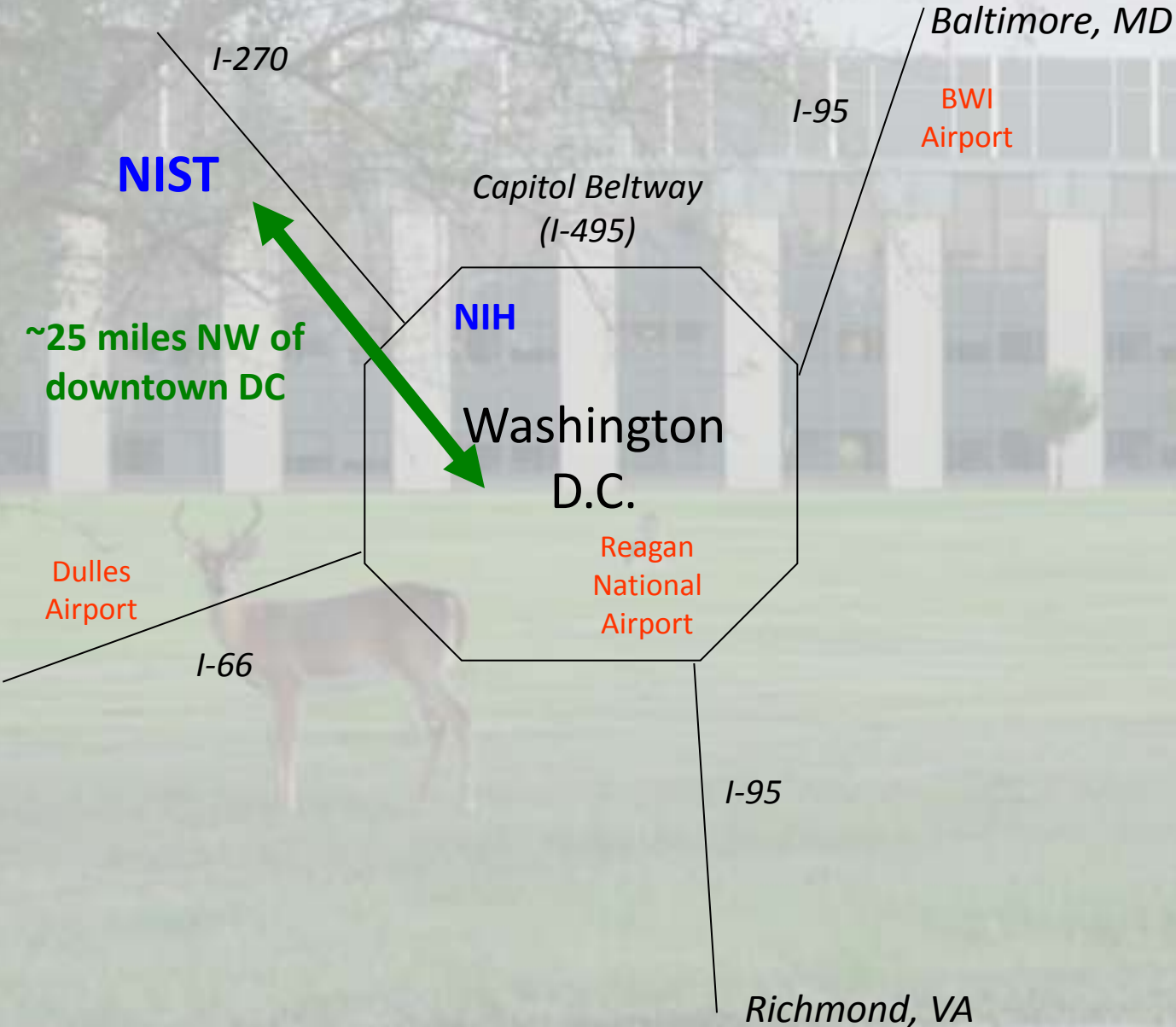
- **NIST**
 - location, role, organizational structure, funding
- **Applied Genetics Group**
 - members, expertise, funding
- **Standard Reference Materials (SRMs)**
 - SRM 2391c: DNA Profiling Standard
- **Forensic DNA Research**
 - Concordance studies, miniSTRs and 26plex
- **Final thoughts and some advice for you...**



NIST Background

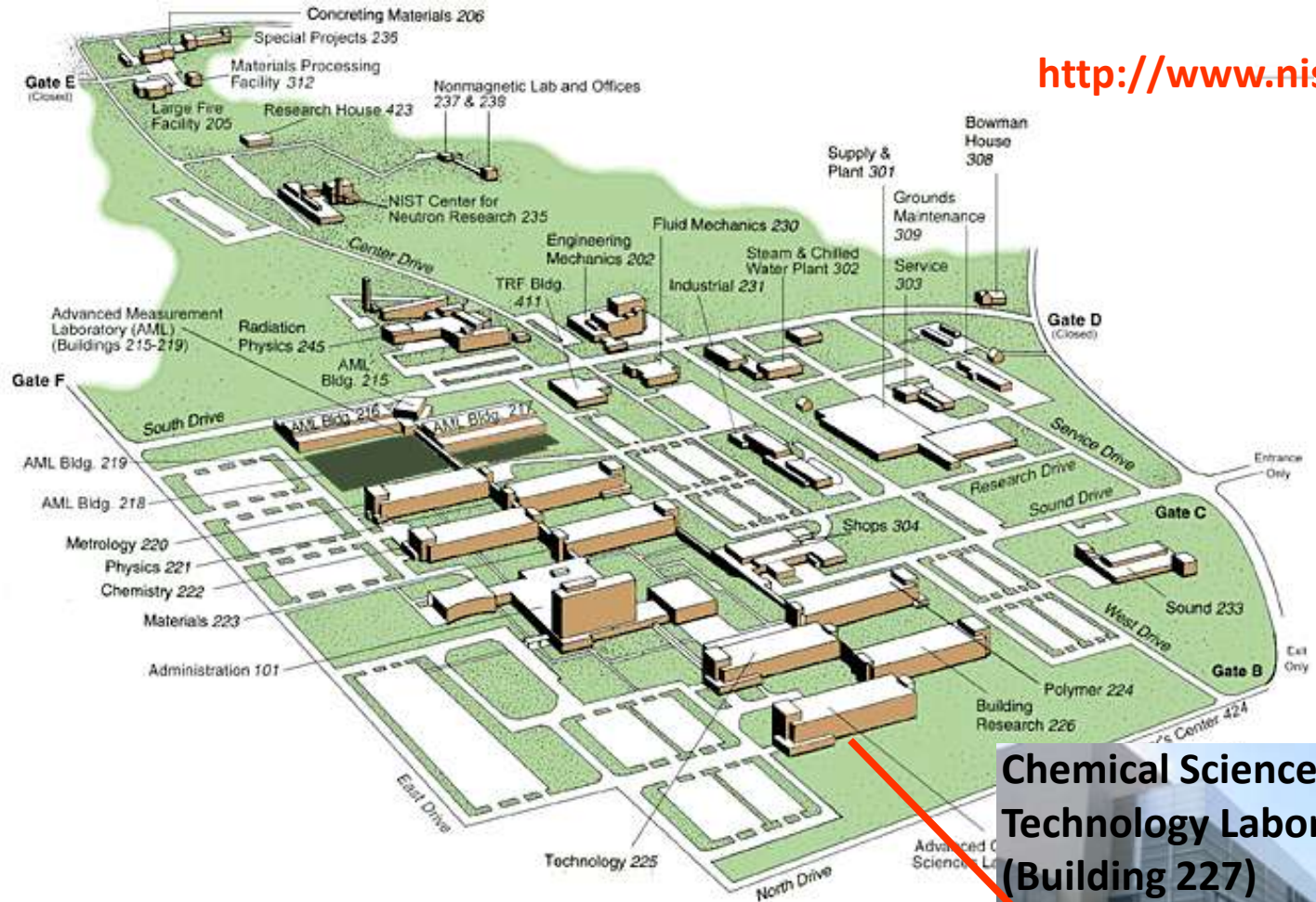
U.S. National Institute of Standards
and Technology
Department of Commerce

Location of NIST



NIST Gaithersburg Campus

<http://www.nist.gov>



**Chemical Sciences and
Technology Laboratory
(Building 227)**



**Human Identity
Project Team**

National Institute of Standards & Technology (NIST)

- **Non-regulatory agency** established in 1901 in the US Department of Commerce.
- Mission to promote US innovation and industrial competitiveness by advancing measurement science, standards & technology.
- NIST is at the top of the US standards pyramid for a wide variety of physical standards, test methods, and calibrations.



Early Driver for U.S. Standards



1904

- Out-of-town fire companies arriving at a Baltimore fire cannot couple their hoses to the hydrants. 1526 buildings razed.

1905

- National Fire Protection Association adopted NBS-developed national hose coupling standard.

NIST Today

Major Assets

- ~ 2,900 employees
- ~ 2600 associates and facilities users
- ~ 400 NIST staff on about 1,000 national and international standards committees
- 3 Nobel Prizes in past 15 years



Major Programs

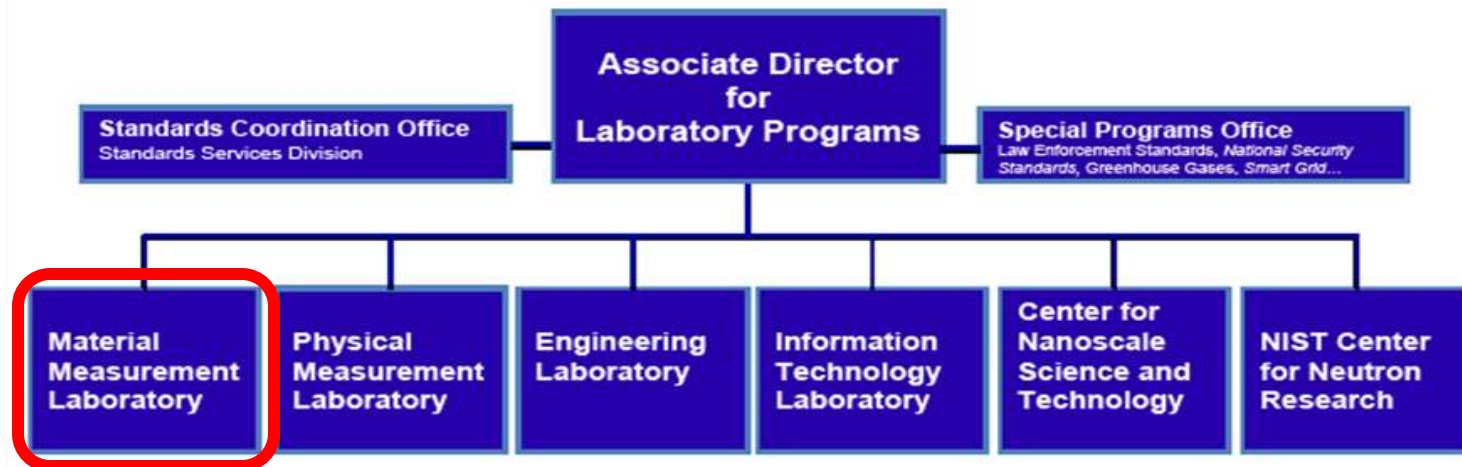
- **NIST Laboratories**
- Baldrige National Quality Program
- Hollings Manufacturing Extension Partnership
- Technology Innovation Program

Joint NIST/University Institutes:

- JILA
- Joint Quantum Institute
- Institute for Bioscience & Biotechnology Research
- Hollings Marine Laboratory

The NIST Laboratories

New Structure for NIST Laboratory Programs



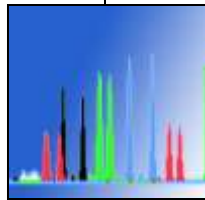
Traditionally focused research and measurement service activities on physical science and engineering disciplines

Bioscience and Health identified as a new area for significant emphasis for NIST labs

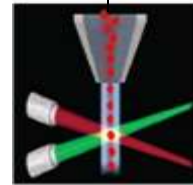
NIST Biochemical Science Division

Laurie Locascio
Division Chief

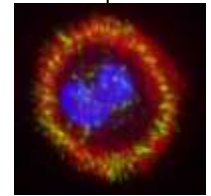
Process Sensing
Dean Ripple
Group Leader



Applied Genetics
John Butler
Group Leader



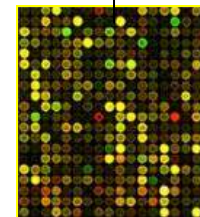
Bioassay Methods
Steve Choquette
Group Leader



Cell Systems Science
Anne Plant
Group Leader



Macromolecular Structure & Function
John Marino
Group Leader



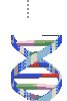
Multiplexed Biomolecular Science
Marc Salit
Group Leader

NIST Applied Genetics Group

Applied Genetics Group Mission Statement

Advancing technology and traceability
through quality genetic measurements to
aid work in

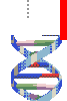
- **forensic DNA testing**
- **clinical diagnostics**
- **cell line authentication**
- **agricultural biotechnology**
- **DNA biometrics**



APPLIED GENETICS Group

Major Programs Currently Underway

- **Forensic DNA**
 - New loci and assays (26plex)
 - STR kit testing
 - Ancestry SNP assays
 - Low-template DNA studies
 - Mixture interpretation
 - STR nomenclature
 - Variant allele cataloging and sequencing
 - Expert systems review
 - Training workshops to forensic DNA laboratories
 - Validation information and software tools
 - Textbook – 3rd ed. (2 vol.)
- **Clinical Genetics**
 - Huntington’s Disease SRM
 - CMV SRM
 - Exploring future needs
- **Ag Biotech**
 - “universal” GMO detection/quantitation (35S promoter)
- **DNA Biometrics**
 - Rapid PCR methods
 - Efforts to standardize testing of future portable DNA systems
 - Kinship analysis
- **Cell Line Authentication**



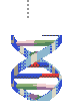
Group Expertise and Funding Sources

Group Expertise

- Reference Material Characterization
- Standard Information Resource Development
- Rapid Multiplex PCR Assay Construction
- Short Tandem Repeat (STR) Genotyping
- Single Nucleotide Polymorphism (SNP) Genotyping
- DNA Sequencing
- Training Materials and Workshops (validation info)

Current Funding Sources

- **National Institute of Justice** (Forensic DNA)
- **FBI Science & Technology Branch** (DNA Biometrics)
- **NIST SRM Program** (SRM development and production)
- **Base funding from Congress (clinical DNA)**



NIST Human Identity Project Teams within the Applied Genetics Group

Forensic DNA Team

Funding from the **National Institute of Justice (NIJ)**
through NIST Office of Law Enforcement Standards



John
Butler



Mike
Coble



Becky
Hill



Margaret
Kline

Workshops &
Textbooks

Concordance &
LT-DNA
Mixtures,
mtDNA & Y

SRM work,
variant alleles &
Cell Line ID



Guest Researcher



Manuel **Fondevila**
Alvarez

*Data
Analysis
Support*



Dave
Duewer

DNA Biometrics Team

Funding from the **FBI S&T Branch**
through NIST Information Access Division



Pete
Vallone

Rapid PCR,
Direct PCR &
Biometrics



Erica
Butts

ABI 3500 &
DNA
Extraction



Kevin
Kiesler

mtDNA &
Mass Spec





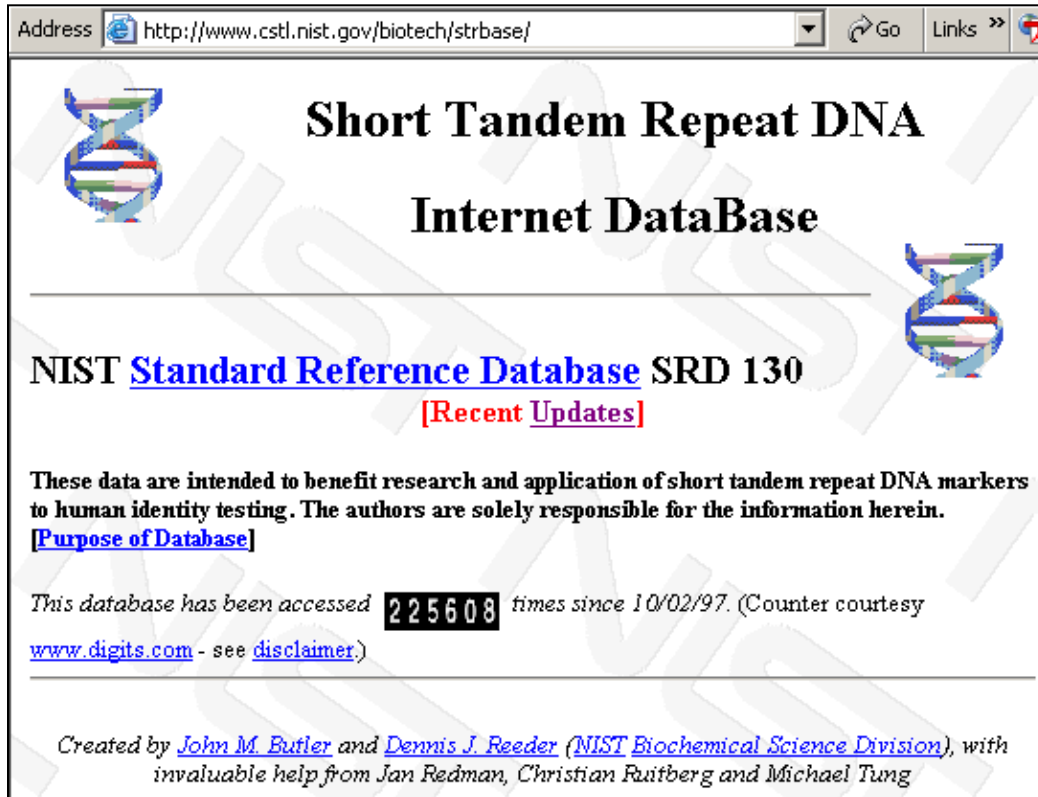
Current Areas of NIST Effort with Forensic DNA

- **Standards**
 - Standard Reference Materials
 - Standard Information Resources (STRBase website)
 - Interlaboratory Studies
- **Technology**
 - Research programs in STRs, SNPs, miniSTRs, Y-STRs, mtDNA, qPCR, LT-DNA, mixtures, rapid PCR
 - Assay and software development, expert system review
- **Training Materials**
 - Textbooks, review articles and workshops on STRs, CE, validation
 - PowerPoint and pdf files available for download
 - Training workshops conducted to scientists, lawyers, and students

<http://www.cstl.nist.gov/biotech/strbase/NIJprojects.htm>

Information Resource

<http://www.cstl.nist.gov/biotech/strbase>



Address <http://www.cstl.nist.gov/biotech/strbase/> Go Links

Short Tandem Repeat DNA Internet DataBase

NIST [Standard Reference Database](#) SRD 130
[[Recent Updates](#)]

These data are intended to benefit research and application of short tandem repeat DNA markers to human identity testing. The authors are solely responsible for the information herein.
[\[Purpose of Database\]](#)

This database has been accessed **225608** times since 10/02/97. (Counter courtesy www.digits.com - see [disclaimer](#).)

Created by [John M. Butler](#) and [Dennis J. Reeder](#) ([NIST Biochemical Science Division](#)), with invaluable help from [Jan Redman](#), [Christian Ruitberg](#) and [Michael Tung](#)

Includes information on:

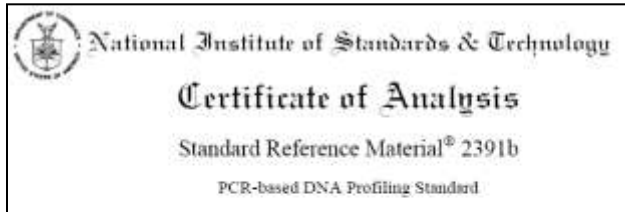
- Core STR loci
- Validation
- STR reference list
- NIST publications
- miniSTRs
- Forensic SNPs
- Variant STR alleles
- Population data resources
- Addresses of scientists

Provides up-to-date information and has been used in court cases to support application of DNA technology

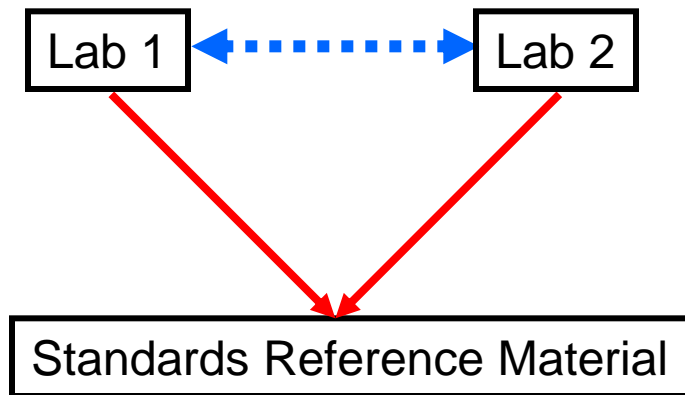
Standard Reference Materials (SRMs)

<http://www.nist.gov/srm>

Traceable standards to ensure accurate and comparable measurements between laboratories



SRM 2391b – autosomal STRs
SRM 2392 & -I – mtDNA sequencing
SRM 2395 – Y-STRs
SRM 2372 – DNA quantitation
SRM 2394 – mtDNA heteroplasmy
SRM 2399 – Fragile X



**Calibration with SRMs
enables confidence in
comparisons of results
between laboratories**

**Helps meet ISO 17025 needs
for traceability to a national metrology institute**

2003: NIST SRM 2391b

Driven primarily by commercial kit loci...



National Institute of Standards & Technology

Certificate of Analysis

Standard Reference Material® 2391b

PCR-based DNA Profiling Standard

This Standard Reference Material (SRM) is intended primarily for use in the standardization of forensic and paternity quality assurance procedures for Polymerase Chain Reaction (PCR)-based genetic testing and for instructional law enforcement or non-clinical research purposes. This SRM can also be used for quality assurance when assigning values to in-house control materials. It is not intended for any human or animal clinical diagnostic use. Note that SRM 2391b is slightly modified from SRM 2391, in that there is more emphasis on Short Tandem Repeats (STRs) and less emphasis on D1S80 [1,2] reflecting the growing interest and utility of STRs [3 to 14]. Additional information on each STR locus can be found at a NIST-sponsored database on the internet: <http://www.csl.nist.gov/biotech/strbase> [14].

This SRM is composed of well-characterized human deoxyribonucleic acid (DNA) in two forms: genomic DNA and DNA to be extracted from cells spotted onto filter paper. A unit of the SRM is composed of 12 frozen components packaged in one box. See the section in this certificate entitled *Descriptive of Components* for a complete listing of the components.

Certified Values: The SRM is certified for genetic loci of forensic interest that were commercially available at the time of production. Genetic types for these loci can be found in Tables 1, 2, and 3. The tables are organized as follows: Table 1 lists the genetic types for the Federal Bureau of Investigation's (FBI's) CODIS (Combined DNA Index System) core STR loci; Table 2 lists additional STR loci of interest, and Table 3 lists the genetic types for D1S80, AmpType® PM + HLADQA1, and Amelogenin.

Expiration of Certification: The certification of this SRM is valid until **31 December 2008**, provided the SRM is handled and stored in accordance with the instructions given in this certificate. However, the certification is invalid if the SRM is contaminated or otherwise modified.

Maintenance of SRM Certification: NIST will monitor this SRM over the period of its certification. If substantive technical changes occur that affect the certification before the expiration of certification, NIST will notify the purchaser. Return of the attached registration card will facilitate notification.

Storage: Store frozen at a temperature of -20 °C. **DO NOT** use a self-defrosting freezer because periodic cycling of temperatures may cause shortened shelf life of this SRM.

The overall direction and coordination of the technical activities leading to certification were under the chairmanship of J.M. Butler of the NIST Biotechnology Division.

Analytical determination and technical measurements leading to the certification of this SRM were performed by M.C. Kline and J.W. Redman of the NIST Biotechnology Division.

The support aspects involved in the preparation, certification, and issuance of this SRM were coordinated through the NIST Standard Reference Materials Group by C.S. Davis.

Vincent Vilker, Acting Chief
Biotechnology Division

John Rumble, Jr., Chief
Measurement Services Division

Gaithersburg, MD 20899
Certificate Issue Date: 06 December 2002

2. Certified Values for Additional STR Loci

F13B	FES/FPS	LPL	Penta D	Penta E	D2S1338	D19S433
10,10	12,12	10,11	10,15	7,12	17,23	13,16,2
8,10	10,11					16
9,10	11,12					4
6,9	10,13					3
8,9	11,13					14
9,10	11,11	10,12	9,12	12,14	25,25	12,14
6,8	11,11*	11,12	3,2,11	12,16	17,22	13,15,2
6,8	10,11	9,11	8,9	5,10	22,22	12,2,15
8,10	10,12	11,12	12,12	12,13	19,23	14,15
8,8	11,11	10,12	8,12	11,11	23,23	13,14
8,10	10,12	11,12	12,12	12,13	19,23	14,15
8,8	11,11	10,12	8,12	11,11	23,23	13,14

**48 autosomal STRs
characterized across
12 DNA samples**

NIST Standard Reference Material (SRM) for Forensic DNA Testing

SRM 2391b (2003-2011)

- **48 autosomal STR loci** with certified values
- **10 liquid genomic DNA components + 2 punches** (cells on 903 paper)
- All single source samples
- 4 males + 6 females
- 9947A & 9948 included

SRM 2391c (2011-future)

- **23 autosomal STR loci and 17 Y-STRs** certified
- **4 liquid genomic DNA components + 2 punches** (cells on **FTA** & 903 paper)
- 5 single source + 1 mixture
- 3 males + 2 females (unique)
- All new samples
 - no 9947A or 9948

SRM 2391c to replace SRM 2391b and SRM 2395 (price reduction)

STR Kits Tested with SRM 2391c

Kit Provider			Primer Mixes
<i>Life Technologies</i>	<i>Promega</i>	<i>Qiagen</i>	<i>NIST</i>
Identifiler	Powerplex 16	ESSplex	26plex [3]
Identifiler Plus	Powerplex 16 HS	IDplex	miniSTRs [4,5]
NGM	Powerplex ESX 17		
NGM SElect	Powerplex ESI 17		
COfiler	Powerplex ES		
Profiler	Powerplex S5		<u>Alleles sequenced:</u> SE33 D12S391 D1S1656 Penta D Penta E D8S1115
Profiler Plus	Powerplex Y		
Profiler Plus ID	FFFL		
SGM Plus			
SEfiler			
MiniFiler			
Yfiler			

22 commercial STR kits examined

NIST developed 26plex and miniplexes

No discordant results observed on SRM 2391c samples

Forensic DNA Research Programs

Concordance Studies
miniSTRs and the 26plex

Steps Involved

Collection

Specimen Storage

Extraction

Quantitation

Multiplex PCR

STR Typing

Interpretation
of Results

Database Storage
& Searching

Calculation of
Match Probability

Steps in DNA Analysis

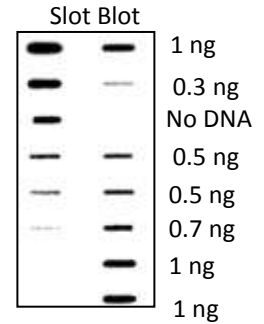
Usually 1-2 day process (a minimum of ~5 hours)



Blood Stain Buccal swab
Sample Collection &
Storage



DNA
Extraction



DNA
Quantitation



Multiplex PCR Amplification

Genetics

If a match occurs, comparison of DNA profile to population allele frequencies to generate a case report with probability of a random match to an unrelated individual

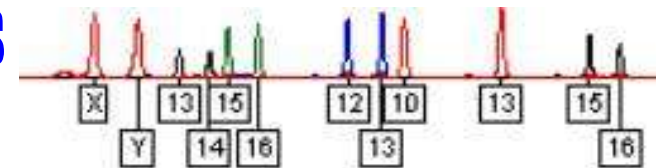


DNA
Database
Search

Biology

Technology

DNA separation and sizing



STR Typing

Male: 13,14-15,16-12,13-10,13-15,16

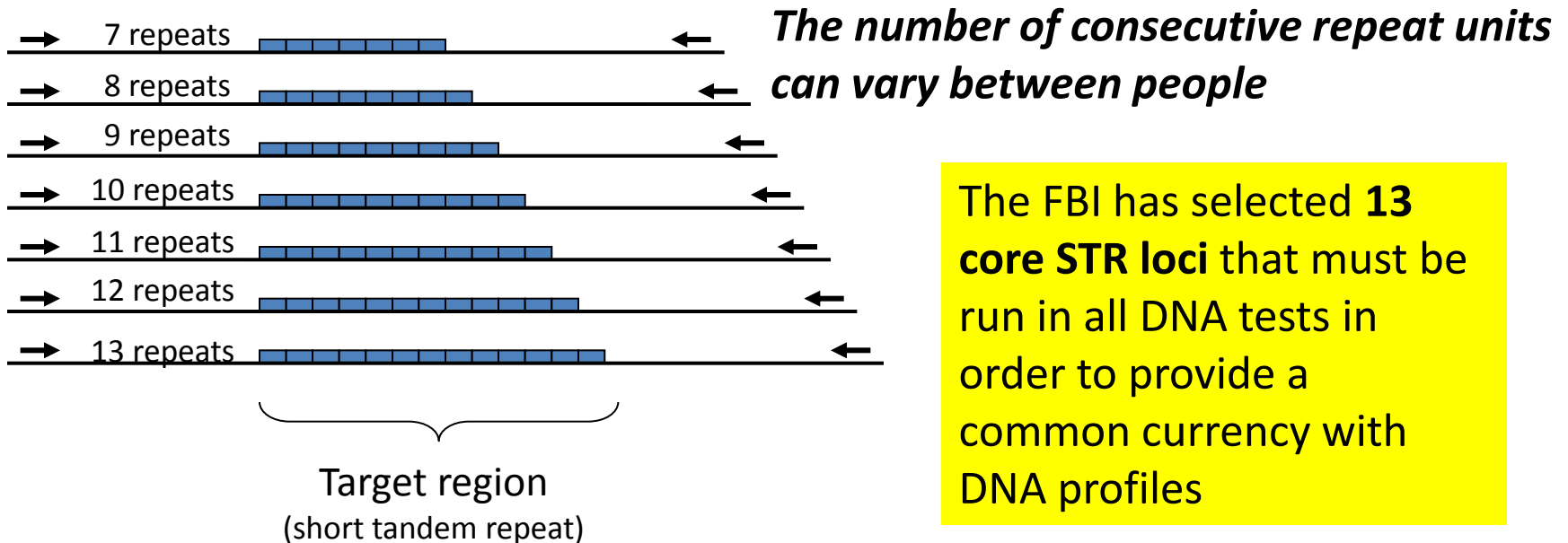
Interpretation of Results

Short Tandem Repeat (STR) Markers

An accordion-like DNA sequence that occurs between genes

TCCAAGCTCTTCCTCTTCCCTAGATCAATACAGACAGAAGACAGGTG**GATAGATA**
GATAGATAGATAGATAGATAGATAGATAGATAGATATCATTGAAAGACAAA
ACAGAGATGGATGATAGATACATGCTTTACAGATGCACAC

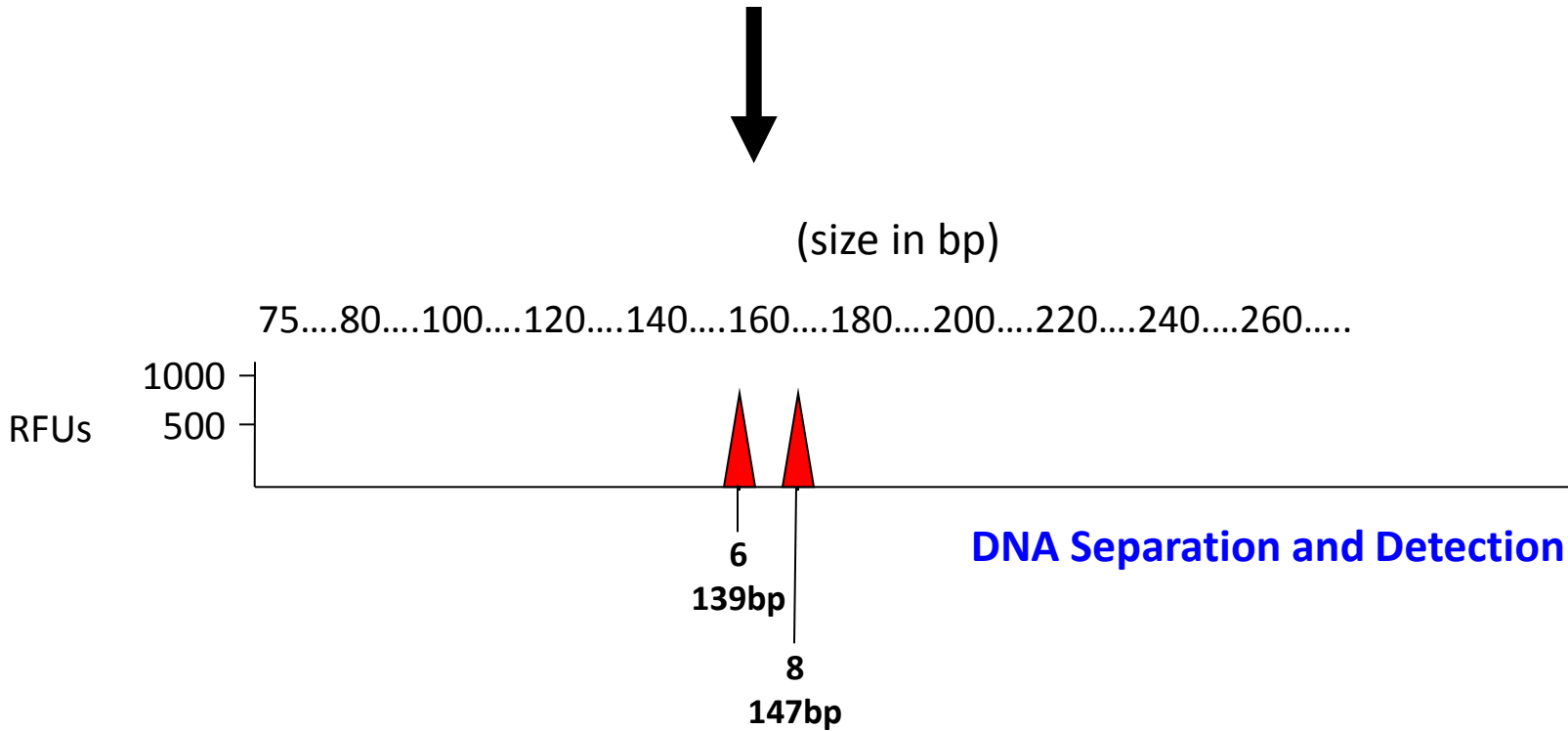
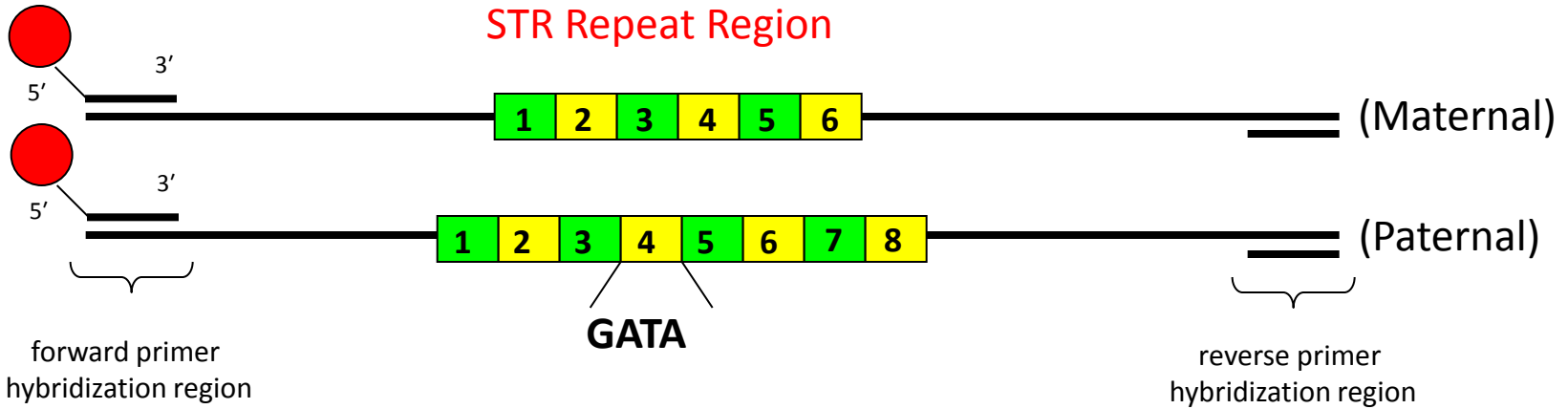
= 12 GATA repeats ("12" is all that is reported)



The FBI has selected **13 core STR loci** that must be run in all DNA tests in order to provide a common currency with DNA profiles

Fluorescent dye-labeled primer

Short Tandem Repeat (STR) Typing



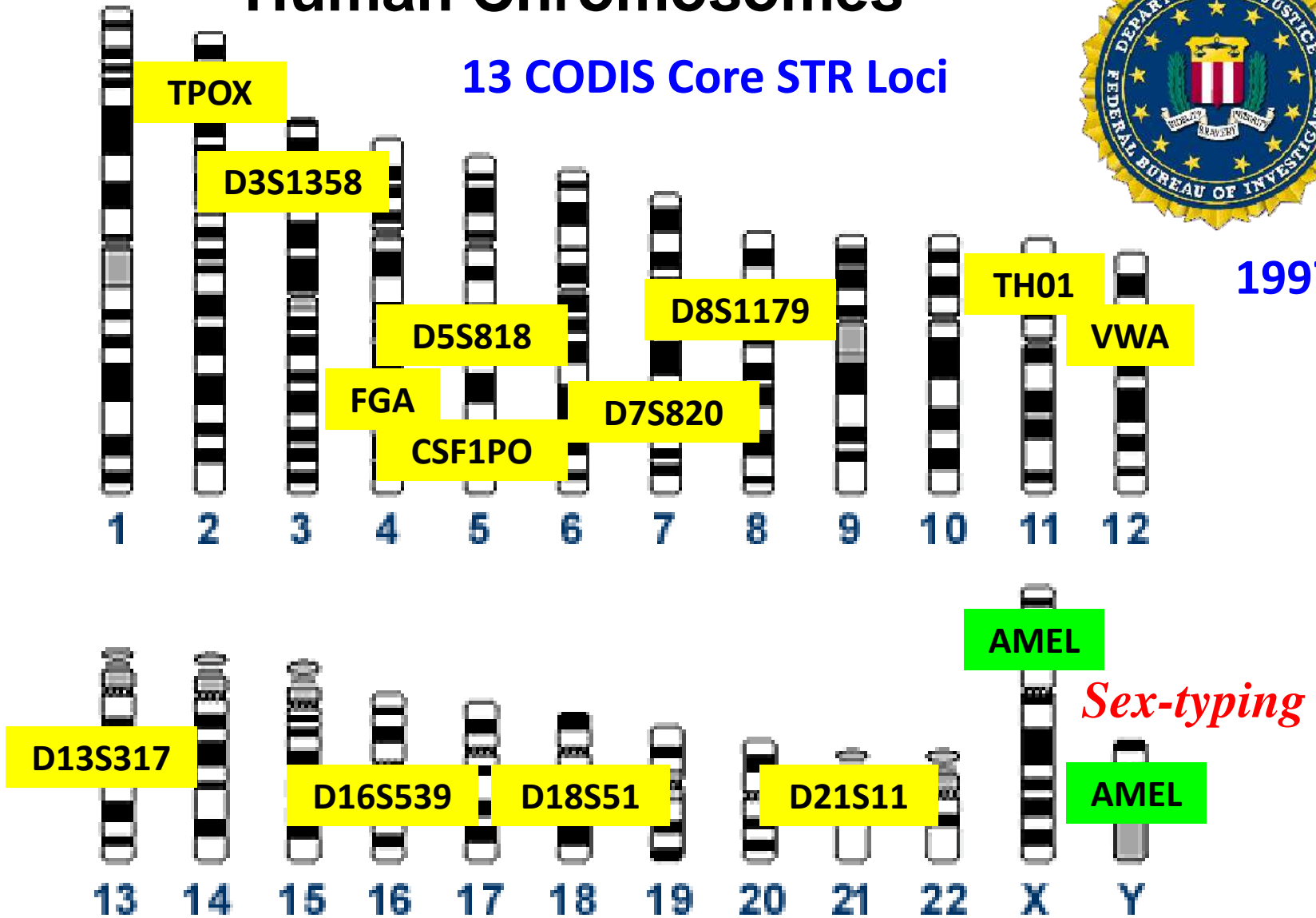
Position of Forensic STR Markers on Human Chromosomes

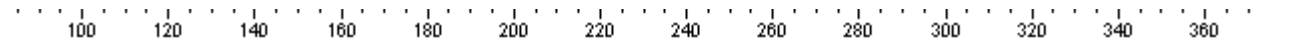
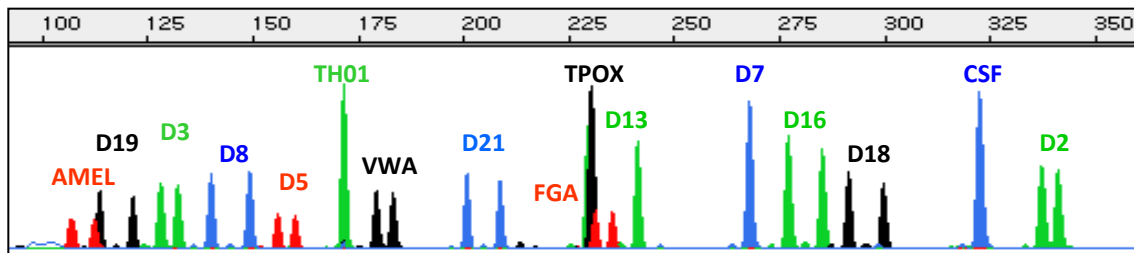
Core STR Loci for the United States



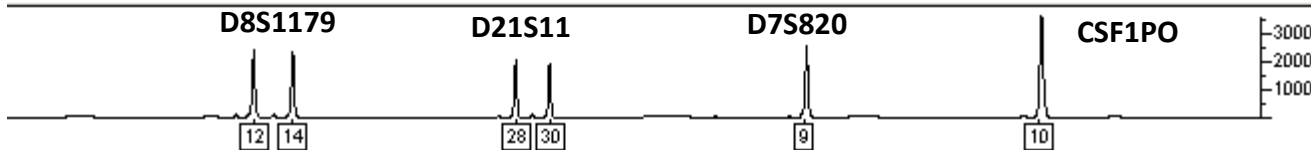
13 CODIS Core STR Loci

1997





6FAM™
(blue)

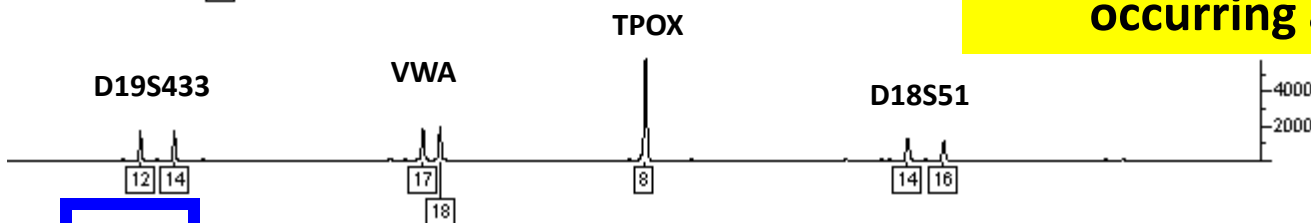


VIC™
(green)



1 in 837 trillion
(probability of this profile occurring at random)

NED™
(yellow)

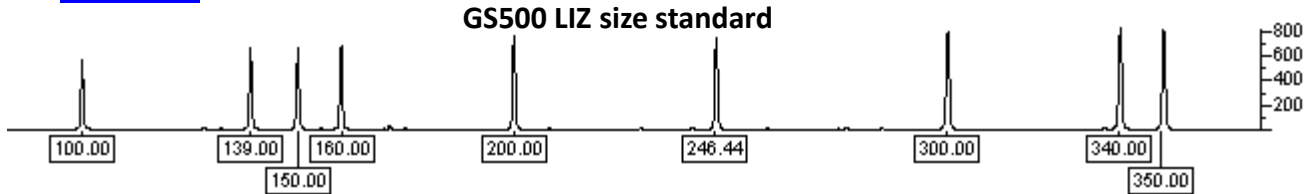


PET™
(red)

AMEL
X
Y



LIZ™
(orange)



Concordance Studies

Commercially Available STR Kits

Applied Biosystems (17)

- ~~AmpFISTR Blue (1996)~~
- ~~AmpFISTR Green I (1997)~~
- Profiler (1997)
- Profiler Plus (1997)
- COfiler (1998)
- SGM Plus (1999)
- **Identifiler** (2001)
- Profiler Plus ID (2001)
- ~~SEfiler (2002)~~
- **Yfiler (2004)**
- MiniFiler (2007)
- SEfiler Plus (2007)
- Sinofiler (2008) – China only
- **Identifiler Direct** (2009)
- NGM (2009)
- **Identifiler Plus** (2010)
- NGM SElect (2010)

Promega Corporation (13)

- PowerPlex 1.1 (1997)
- PowerPlex 1.2 (1998)
- PowerPlex 2.1 (1999)
- **PowerPlex 16** (2000)
- PowerPlex ES (2002)
- **PowerPlex Y (2003)**
- PowerPlex S5 (2007)
- **PowerPlex 16 HS** (2009)
- PowerPlex ESX 16 (2009)
- PowerPlex ESX 17 (2009)
- PowerPlex ESI 16 (2009)
- PowerPlex ESI 17 (2009)
- PowerPlex 18D (2010)

Qiagen (2010)

*Primarily selling kits in Europe
Due to patent restrictions
cannot sell in U.S.*

- ESSplex
- ESSplex SE
- Decaplex SE
- IDplex
- Nonaplex ESS
- Hexaplex ESS
- HD (Chimera)
- Argus X-12
- Argus Y-12
- **DIplex (30 indels)**

**~1/3 of all STR kits were
released in the last year**

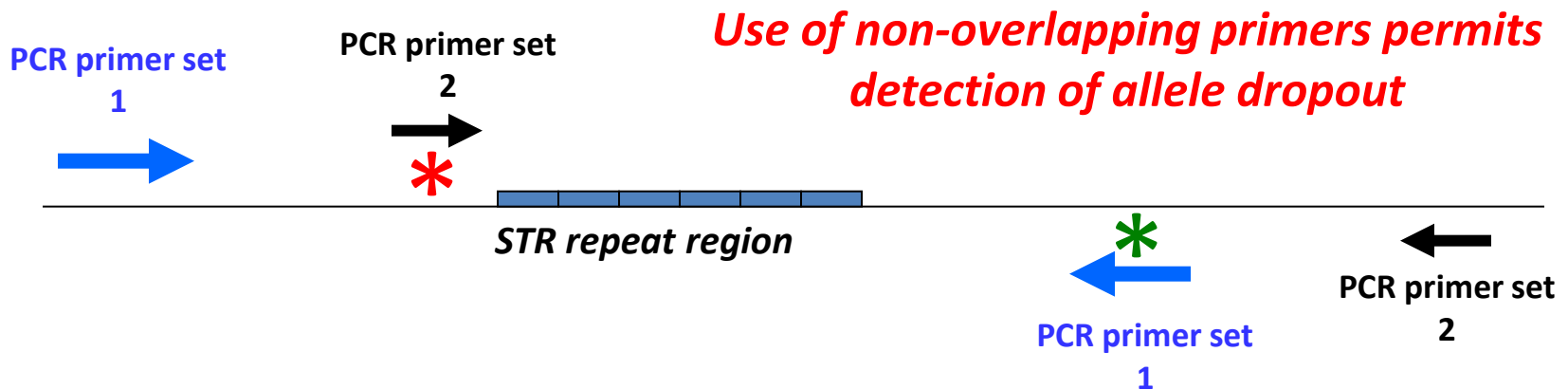
STR Kit Concordance Testing

- Many of these STR kits have different primer sequences for amplifying the same STR locus
- Need to analyze the same DNA samples with different STR typing kits looking for differences
- In some rare cases, allele dropout (null alleles) may occur due to mutations in primer binding regions

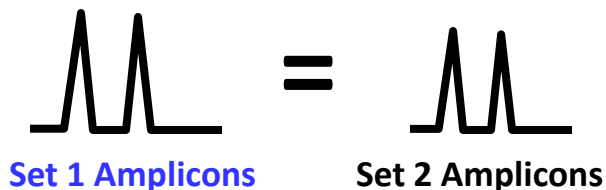
Purpose of Concordance Studies

When different primer sets are utilized, there is a concern that allele dropout may occur due to primer binding site mutations that impact one set of primers but not another

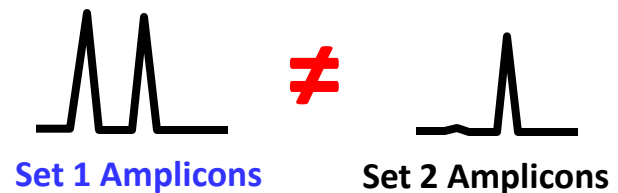
* represents potential mutations impacting primer annealing



If no primer binding site mutations



If a primer binding site mutation exists



STR Kit Concordance Testing

Profiles in DNA Article Published April 2010

Article Type: Feature

Volume 13 No. 1, April 2010

Strategies for Concordance Testing

Carolyn R. Hill, Margaret C. Kline, David L. Duewer and John M. Butler

National Institute of Standards and Technology, Biochemical Science Division, Gaithersburg, Maryland, USA

Concordance evaluations are important to conduct to determine if there are any allelic dropout or "null alleles" present in a data set. These studies are performed because there are a variety of commercial short tandem repeat (STR) multiplex kits with different configurations of STR markers available to the forensic community. The placement of the markers can vary between kits because the primer sequences were designed to amplify different polymerase chain reaction (PCR) product sizes. When multiple primer sets are used, there is concern that allele dropout may occur due to primer-binding-site mutations that affect one set of primers but not another.

http://www.promega.com/profiles/1301/1301_08.html

The 4 “S’s” of Concordance

- NIST Standard **Samples**
 - Run same samples with multiple kits to compare results
- Concordance **Software**
 - Allows comparison of data sets using NIST developed software

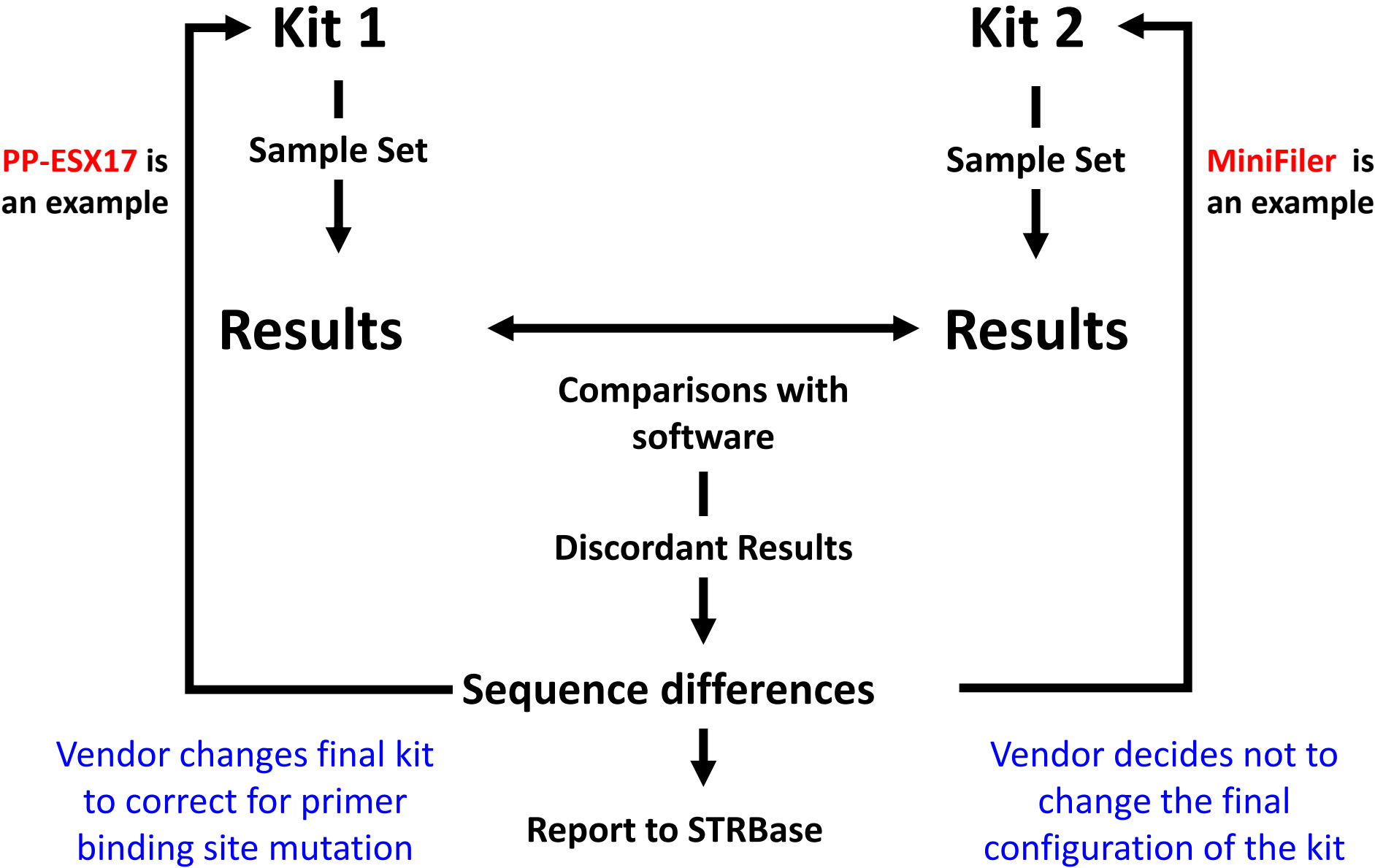
<http://www.cstl.nist.gov/biotech/strbase/software.htm>

- DNA **Sequencing**
 - To validate and determine the exact cause for the null allele

- **STRBase** website
 - To report verified null alleles and discordant results to the forensic community

<http://www.cstl.nist.gov/biotech/strbase/NullAlleles.htm>

NIST Concordance Testing Steps



NIST Sample Set (>1450 Samples)

- **NIST U.S. population samples**
 - 260 African American, 260 Caucasian, 140 Hispanic, 3 Asian
- **U.S. father/son paired samples**
 - ~100 fathers/100 sons for each group: 200 African American, 200 Caucasian, 200 Hispanic, 200 Asian
- **NIST SRM 2391b**, PCR-based DNA Profiling Standard (highly characterized)
 - 10 genomic DNA samples, 2 cell line samples
 - Includes 9947A and 9948
- **NIST SRM 2391c**, PCR-based DNA Profiling Standard
 - 4 genomic DNA (one mixture)
 - 2 cell lines (903 and FTA paper)

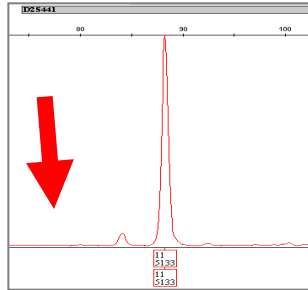
Extra (Degenerate) Primers Added with NGM SElect

NGM SElect and NGM'

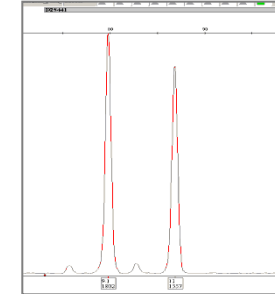
D2S441

9.1 allele missing in 7 Asians

NGM (original)



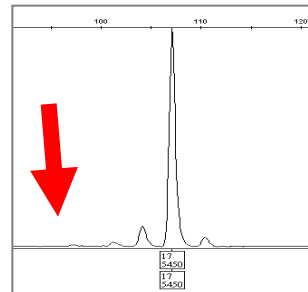
11,11



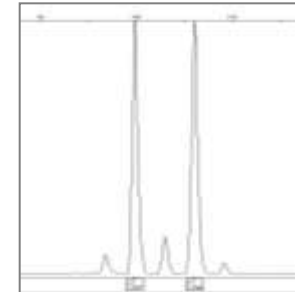
9.1,11

D22S1045

15 allele missing in 4 samples



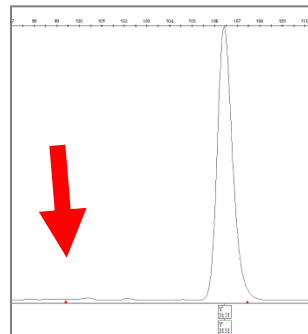
17,17



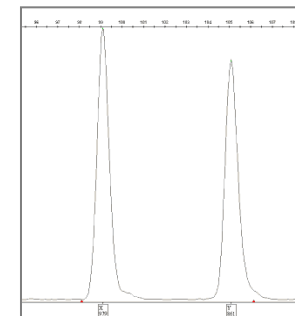
15,17

Amelogenin

X allele missing in 3 samples



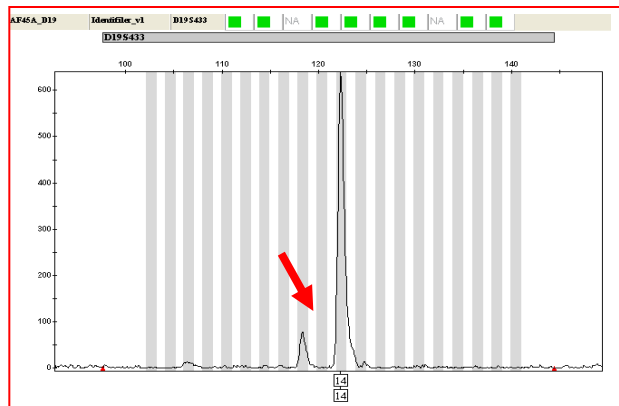
Y,Y



X,Y

D19S433 Discordance

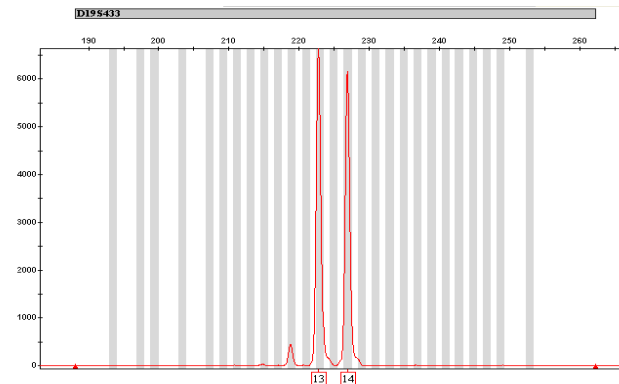
Identifiler & NGM = **14,14**



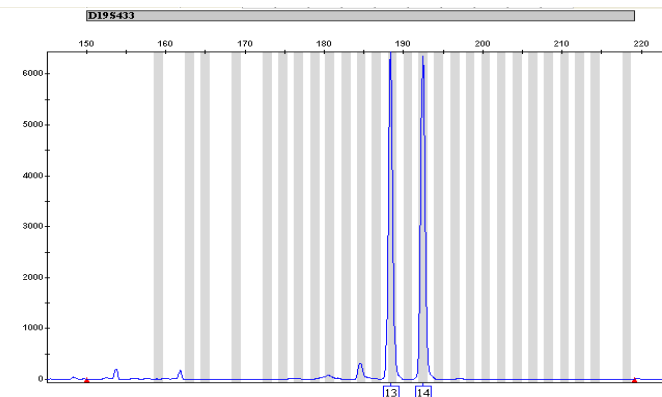
AF45A (Asian)

Allele 13 was missing in two different Asian samples with ABI primers
= $2/2886 = 0.07\%$ discordance

ESX 17 = 13,14



ESI 17 = 13,14



Frequencies [for] the silent allele were determined to be 0.0114 in 176 people from Shizuoka (Honshu) and 0.0128 in 156 people from Okinawa

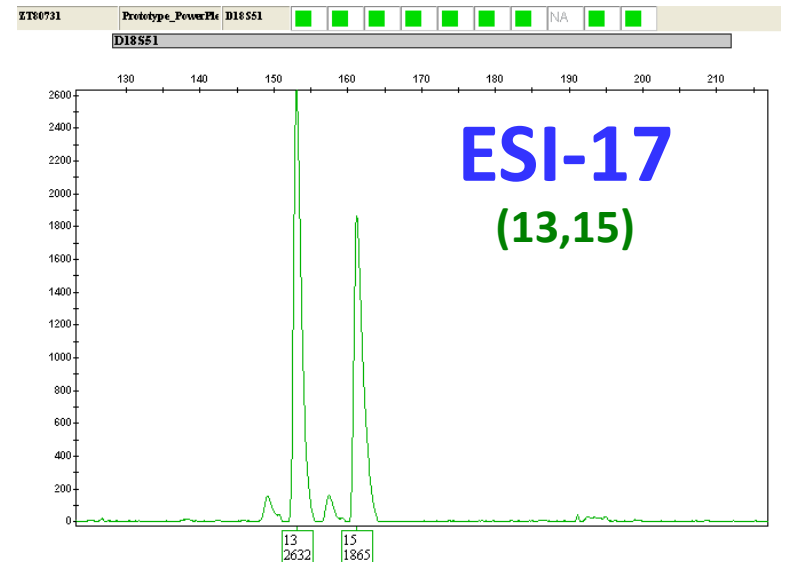
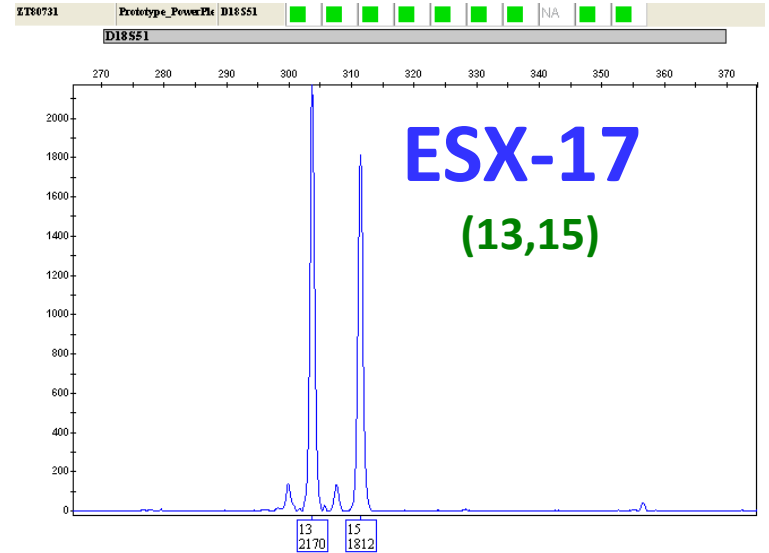
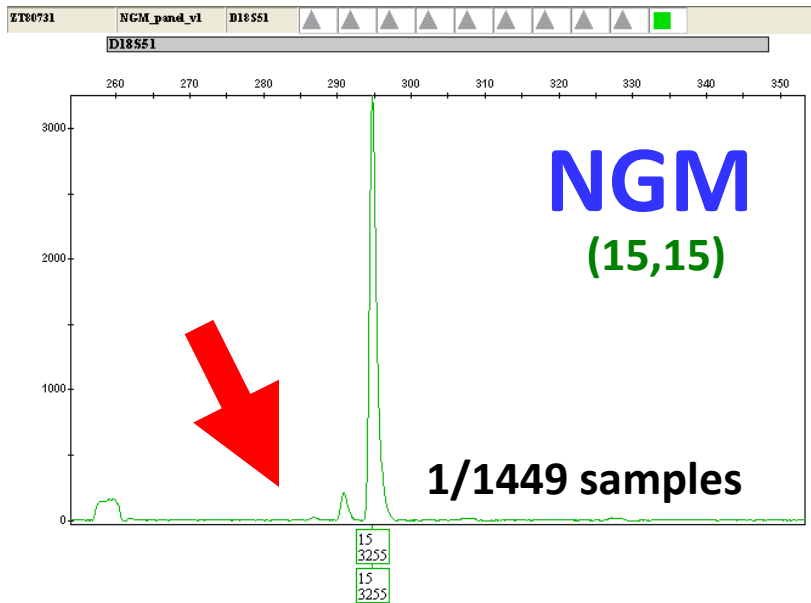
J Forensic Sci, September 2008, Vol. 53, No. 5
doi: 10.1111/j.1556-4029.2008.00806.x
Available online at: www.blackwell-synergy.com

Natsuko Mizuno,¹ D.V.M.; Tetsushi Kitayama,¹ M.Sc.; Koji Fujii,¹ Ph.D.; Hiroaki Nakahara,¹ D.V.M.; Kanako Yoshida,¹ Ph.D.; Kazumasa Sekiguchi,¹ Ph.D.; Naoto Yonezawa,² Ph.D.; Minoru Nakano,² Ph.D.; and Kentaro Kasai,¹ Ph.D.

A D19S433 Primer Binding Site Mutation and the Frequency in Japanese of the Silent Allele It Causes

T→A SNP 8 bp downstream impacting reverse primer binding with Identifiler (and thus SGM Plus)

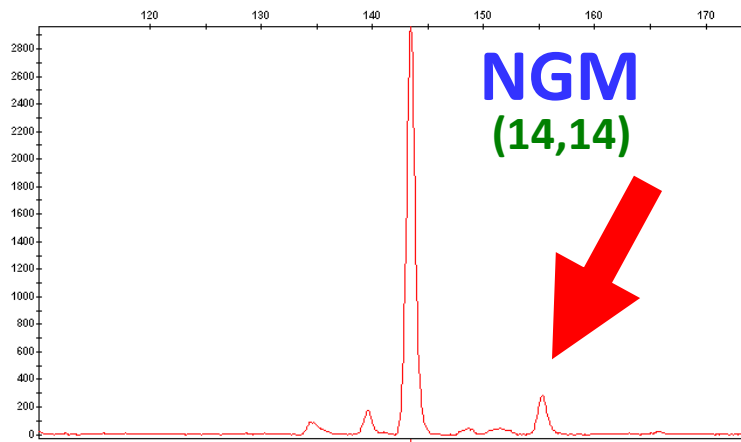
D18S51 Null Allele



Correct type (13,15)

C→T SNP 172 bp downstream from repeat

D3S1358 Null Allele

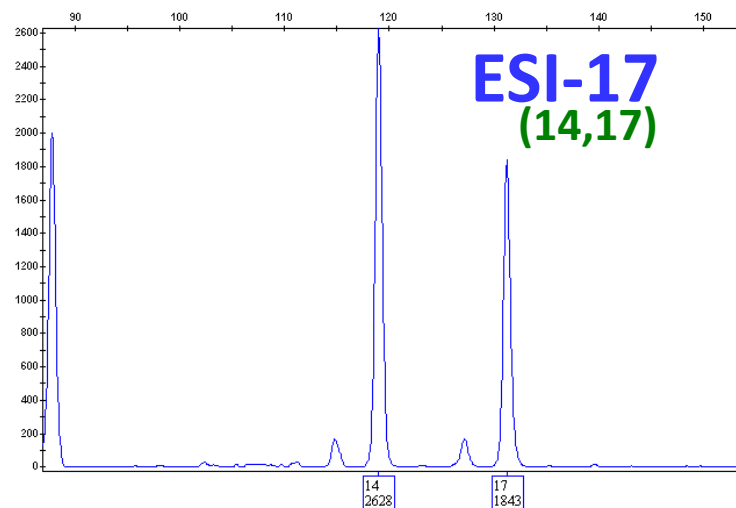
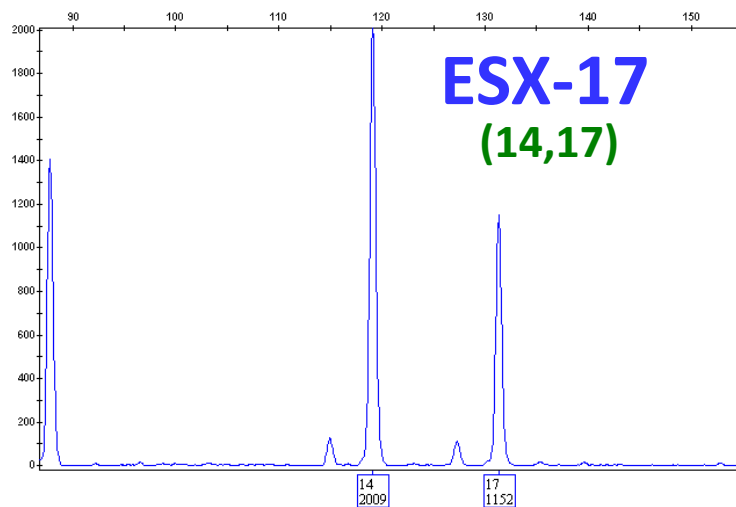


1/1449 samples

14
2962
14
2962

Correct type (14,17)

G → C SNP 11 bp downstream from repeat



Completed Concordance Studies

Kits compared	Samples	Loci Compared	Comparisons	# Differences	Concordance (%)
ID-SGM+	1424	11	15,664	1	99.994
ID-Pro+	1415	10	14,150	1	99.993
ID-IDplex	1426	16	22,816	29	99.873
ID-PP16	662	14	9,268	4	99.957
ID-MiniFiler	1137	9	10,233	26	99.746
ID-NGM	1427	11	15,807	3	99.981
ID-NGMs	663	11	7,293	0	100.000
ID-ESX17	1443	11	15,873	5	99.968
ID-ES17	1443	11	15,873	4	99.975
ID-ES17	1433	11	15,873	28	99.823
ID-ESSplex	662	11	7,262	17	99.767
ID-Hexaplex	653	2	1,306	1	99.923
PP16-SGM+	651	9	5,859	1	99.983
PP16-Pro+	647	10	6,470	3	99.969
PP16-IDplex	657	14	9,198	3	99.967
PP16-MiniFiler	656	8	5,248	14	99.733
PP16-NGM	657	9	5,913	3	99.949
PP16-NGMs	662	9	5,958	1	99.983
PP16-ESX17	662	9	5,958	1	99.983
PP16-ES17	662	9	5,958	0	100.000
PP16-ESSplex	653	9	5,877	16	99.728
PP16-ESSplexSE	662	9	5,958	16	99.731
PP16-Hexaplex	653	2	1,306	1	99.923
SGM+ Pro+	1415	7	9,905	0	100.000
SGM+ IDplex	1424	11	15,664	5	99.968
SGM+ MiniFiler	1137	6	6,822	10	99.853
SGM+ NGM	1424	11	15,664	4	99.974
SGM+ NGMs	651	11	7,161	0	100.000
SGM+ ESX17	1424	11	15,664	6	99.962
SGM+ ES17	1424	11	15,664	5	99.968
SGM+ ESS	1424	11	15,664	5	99.968
SGM+ ESSplexSE	651	11	7,161	5	99.930
SGM+ Hexaplex	651	2	1,302	1	99.923
Pro+ IDplex	1415	10	14,150	5	99.965
Pro+ MiniFiler	1137	6	6,822	16	99.765
Pro+ NGM	1415	7	9,905	4	99.960
Pro+ NGMs	647	7	4,529	0	100.000
Pro+ ES17	1415	7	9,905	4	99.960
Pro+ ES17	1415	7	9,905	3	99.960
Pro+ ESS	1415	7	9,905	4	99.960
Pro+ ESSplexSE	647	7	4,529	4	99.912
Pro+ Hexaplex	647	1	647	1	99.845
IDplex-MiniFiler	1137	9	10,233	48	99.531
IDplex-NGM	1426	11	15,686	30	99.809
IDplex-NGMs	657	11	7,227	17	99.765
IDplex-ESX17	1426	11	15,686	28	99.821
IDplex-ES17	1426	11	15,686	27	99.818
IDplex-ESS	1426	11	15,686	1	99.994
IDplex-ESSplexSE	657	11	7,227	1	99.986
IDplex-Hexaplex	653	2	1,306	1	99.923
MiniFiler-NGM	1137	6	6,822	13	99.809
MiniFiler-NGMs	656	6	3,936	10	99.746
MiniFiler-ESX17	1137	6	6,822	10	99.853
MiniFiler-ES17	1137	6	6,822	9	99.868
MiniFiler-ESS	1137	6	6,822	35	99.487
MiniFiler-ESSplexSE	656	6	3,936	35	99.111
MiniFiler-Hexaplex	653	1	653	1	99.847
NGM-NGMs	657	16	10,512	14	99.867
NGM-ESX17	1417	16	22,992	16	99.930
NGM-ES17	1417	16	22,992	18	99.902
NGM-ESS	1433	16	22,928	42	99.817
NGM-ESSplexSE	657	16	10,512	22	99.791
NGM-Hexaplex	653	7	4,571	9	99.803
NGMs-ES17	662	17	11,254	4	99.964
NGMs-ES17	662	17	11,254	14	99.876
NGMs-ESS	653	16	10,448	17	99.837
NGMs-ESSplexSE	662	17	11,254	34	99.698
NGMs-Hexaplex	653	7	4,571	3	99.934
ESX17-ES17	1443	17	24,531	19	99.923
ESX17-ESS	653	16	10,448	34	99.675
ESX17-ESSplexSE	662	17	11,254	25	99.778
ES17-Hexaplex	657	7	4,599	6	99.870
ES17-ESS	653	16	10,448	28	99.732
ES17-ESSplexSE	662	17	11,254	30	99.733
ES17-Hexaplex	657	7	4,599	3	99.935
ESS-ESSplexSE	653	16	10,448	0	100.000
ESS-Hexaplex	653	7	4,571	3	99.934
ESSplexSE-Hexaplex	653	7	4,571	3	99.934
SE33-ESX17	1443	1	1,443	6	99.584
SE33-ES17	1443	1	1,443	17	99.822
SE33-NGMs	663	1	663	4	99.397
SE33-ESSplexSE	662	1	662	21	96.828
ES17p-ESX17	477	17	8,109	7	99.914
ES17p-NGMs	477	17	8,109	2	99.975
ES17p-ESSplexSE	477	17	8,109	42	99.482
ES17p-SE33	477	1	477	4	99.161
PP180-ID	50	16	800	2	99.750
PP180-PP16	703	16	11,248	1	99.991
ESX17/ESX17	1443	17	24,531	4	99.984
ESX17/ES17p	477	17	8109	3	99.963
ESX17p/NGM	1437	16	22992	22	99.904
ESX17p/NGMs	663	17	11271	4	99.965
ESX17p/ESS	1433	16	22928	30	99.869
ESX17p/ESSplexSE	662	17	11254	44	99.609
ESX17p/Hexaplex	653	7	4571	2	99.956
2plex/ESX17	1443	3	4329	4	99.906
2plex/ES17	1443	3	4329	0	100.000
2plex/NGM	1437	3	4311	11	99.745
2plex/NGMs	663	3	1989	0	100.000
2plex/ESS	1433	3	4299	0	100.000
2plex/ESSplexSE	662	3	1986	0	100.000
2plex/Hexaplex	653	3	1959	2	99.898
2plex/ESX17*	663	3	1989	0	100.000
miniSTR/ESX17	663	3	1989	3	99.849
miniSTR/ES17	663	3	1989	0	100.000
miniSTR/NGM	657	3	1971	3	99.848
miniSTR/NGMs	663	3	1989	0	100.000
miniSTR/ESS	653	3	1959	0	100.000
miniSTR/ESSplexSE	662	3	1986	0	100.000
miniSTR/Hexaplex	653	3	1959	2	99.898
miniSTR/ESX17*	663	3	1989	0	100.000
Totals	102,345	1,021	948,301	1,109	99.883

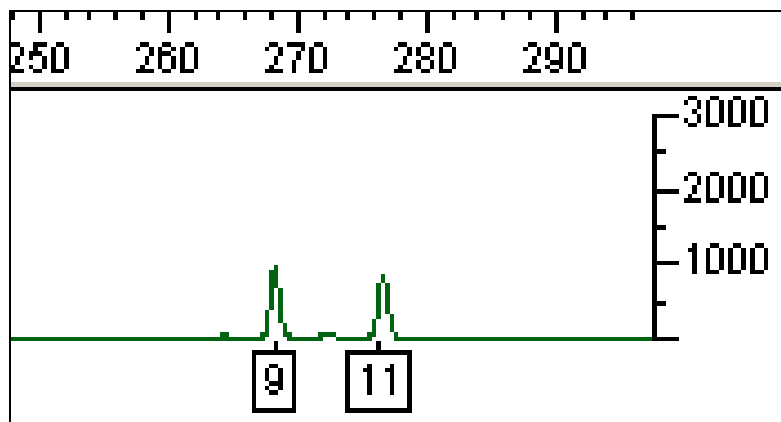
Kits compared	Samples	Loci Compared	Comparisons	# Differences	Concordance (%)
111	102,345	1,021	948,301	1,109	99.883

948,301 allele comparisons
1,109 total differences
99.88% concordance

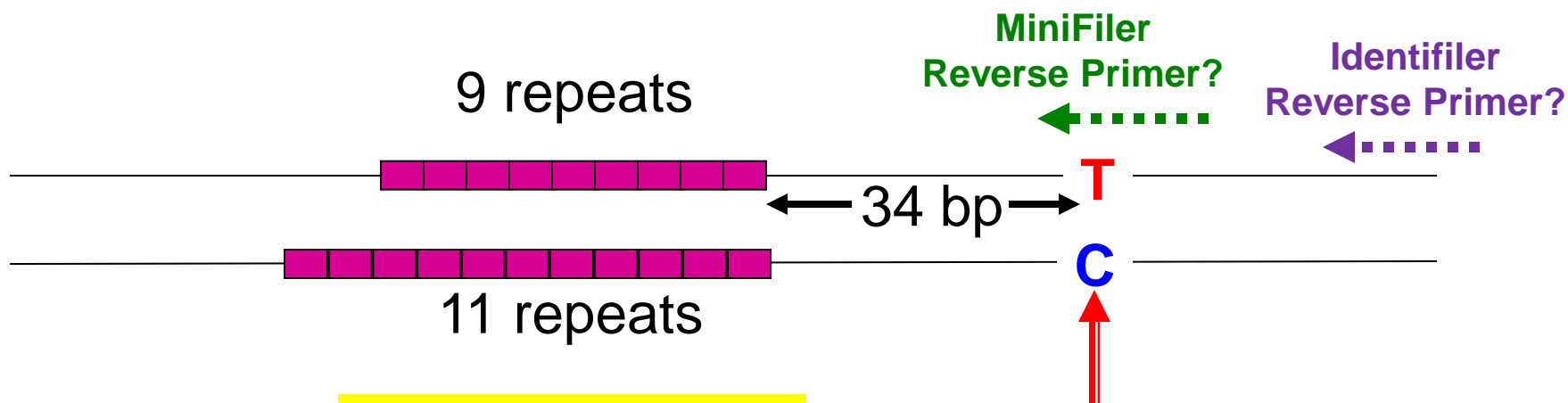
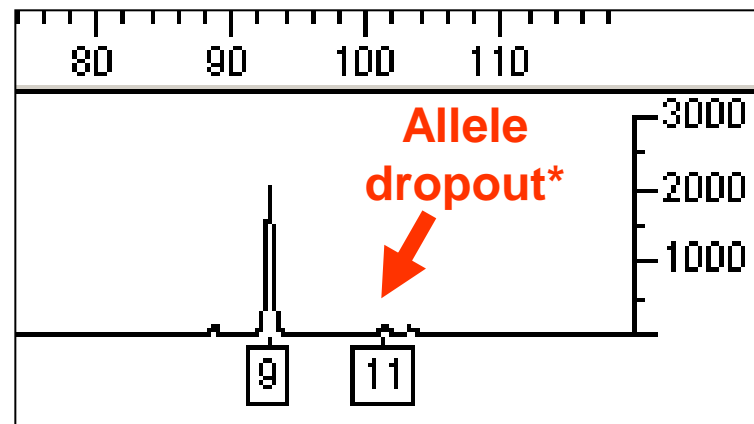
*Kits (except Identifiler) were kindly provided by **Promega, Qiagen and Applied Biosystems** for concordance testing performed at **NIST***

SRM 2391b Genomic 8 with D16S539

Identifiler



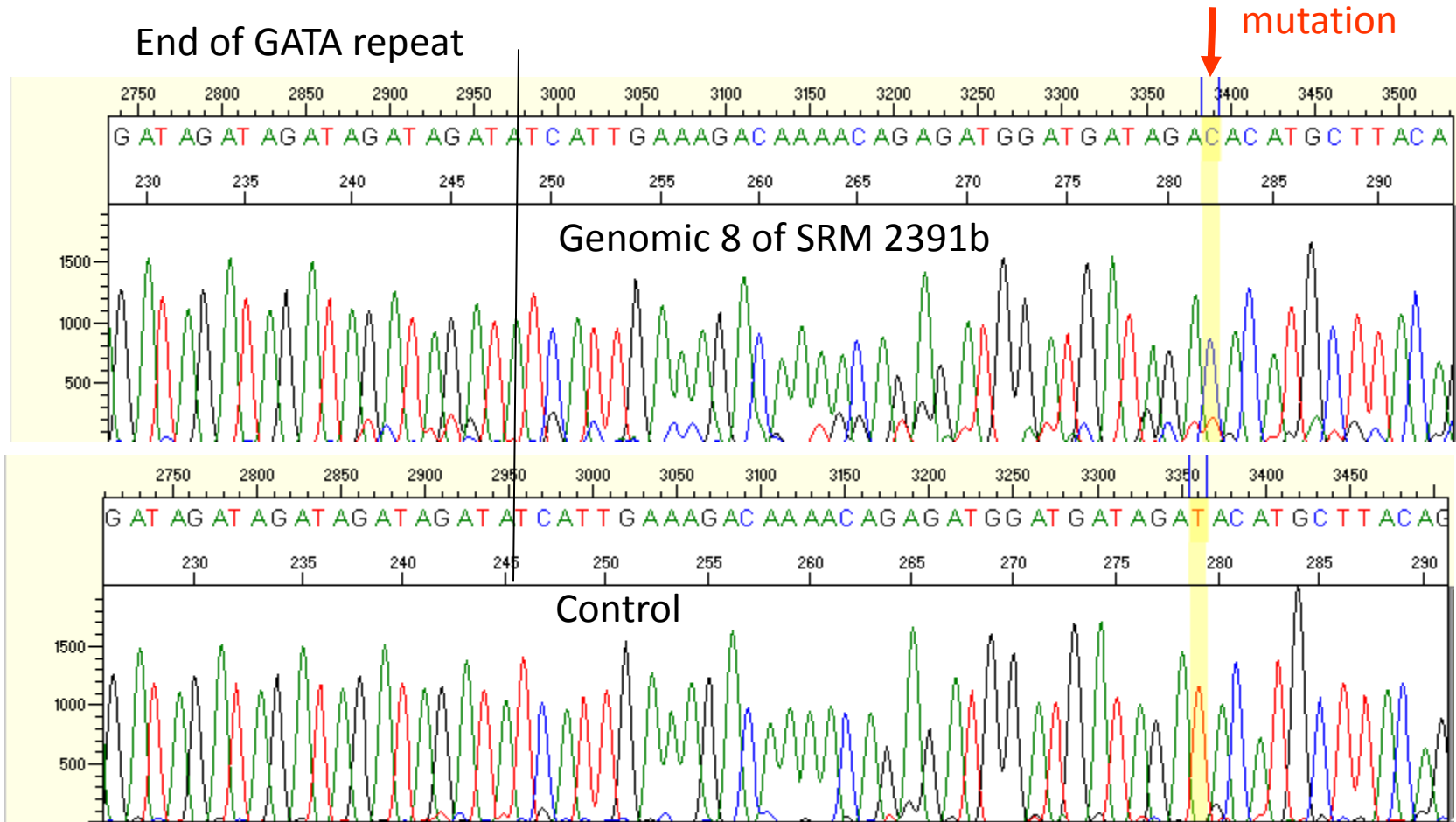
MiniFiler



Type 9**T**, 11**C**

D16S539 SRM 2391b Genomic 8

T→C mutation 34 bp downstream of the repeat



Position of the T→C probably affects the reverse primer of Minifiler and is the 3rd base found the 5' end of the Reverse PP16 primer. This could explain the imbalance of the allele seen when using PP16.

miniSTRs and the 26plex

More Loci are Useful in Situations Involving Relatives

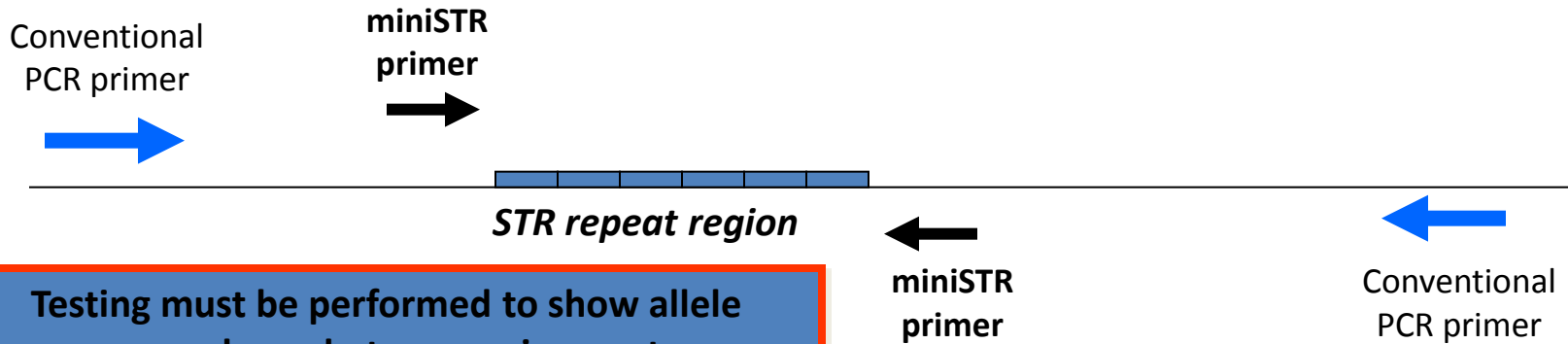
- **Missing Persons** and Disaster Victim Identification (kinship analysis)
- Immigration Testing (often limited references)
 - Recommendations for 25 STR loci
- Deficient Parentage Testing
 - often needed if only one parent and child are tested

Relationship testing labs are being pushed to answer more difficult genetic questions...and **we want to make sure the right tools are in place**

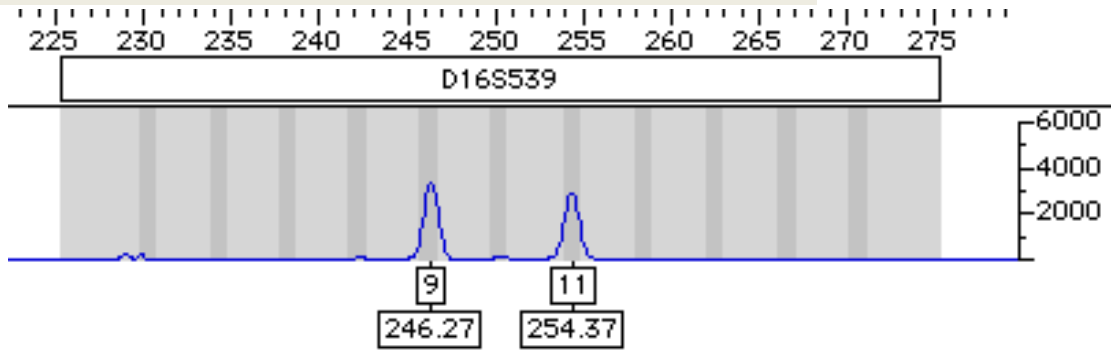
Selection of New Autosomal Loci

- Aim to have candidate sets for optimal **miniSTRs**
 - Obtaining additional information with degraded DNA samples
- Using ~900 STR loci with some literature data as a starting point...
 - Loci with high heterozygosities (>0.7)
 - Loci with small allele ranges (<24 bp) – **low mutation?**
 - Tetra (some tri-)nucleotide repeats without variants
 - Clean flanking regions (PCR products <140 bp)
- **26 loci** met criteria and were fully characterized...

A miniSTR is a reduced size STR amplicon that enables higher recovery of information from degraded DNA samples

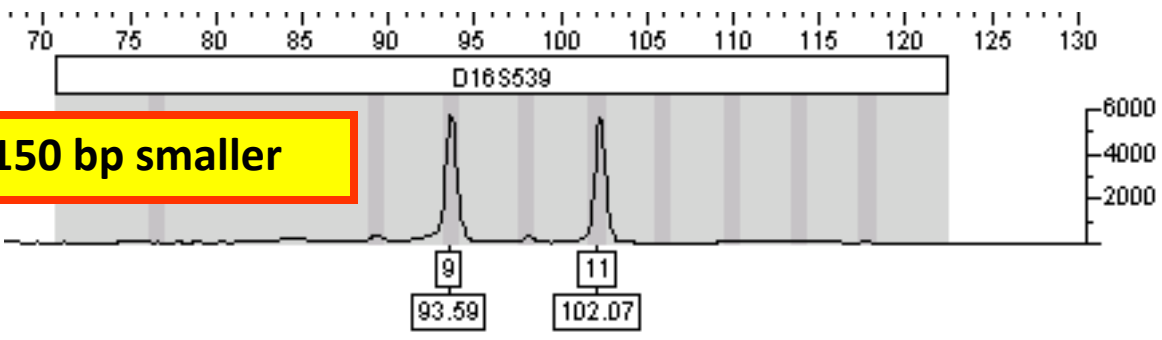


Testing must be performed to show allele concordance between primer sets



Conventional STR test (COfiler™ kit)

~150 bp smaller



MiniSTR assay (using Butler *et al.* 2003 primers)

Characterization of New Loci

“Computer Work”

Candidate STR marker selection

(e.g. literature searches)

Pull down sequence data from the web

(e.g. NCBI)

Identify Chromosome Location

(e.g. Human BLAT Search)

Screen for PCR Primers

(e.g. Primer3)

Test primers for Multiplex-ability

(e.g. AutoDimer - NIST)

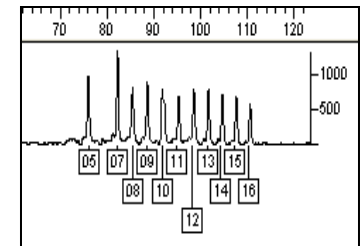
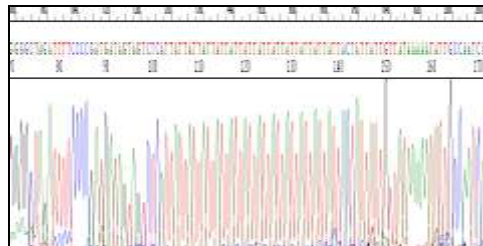
“Laboratory Work”

Test Markers on Population samples

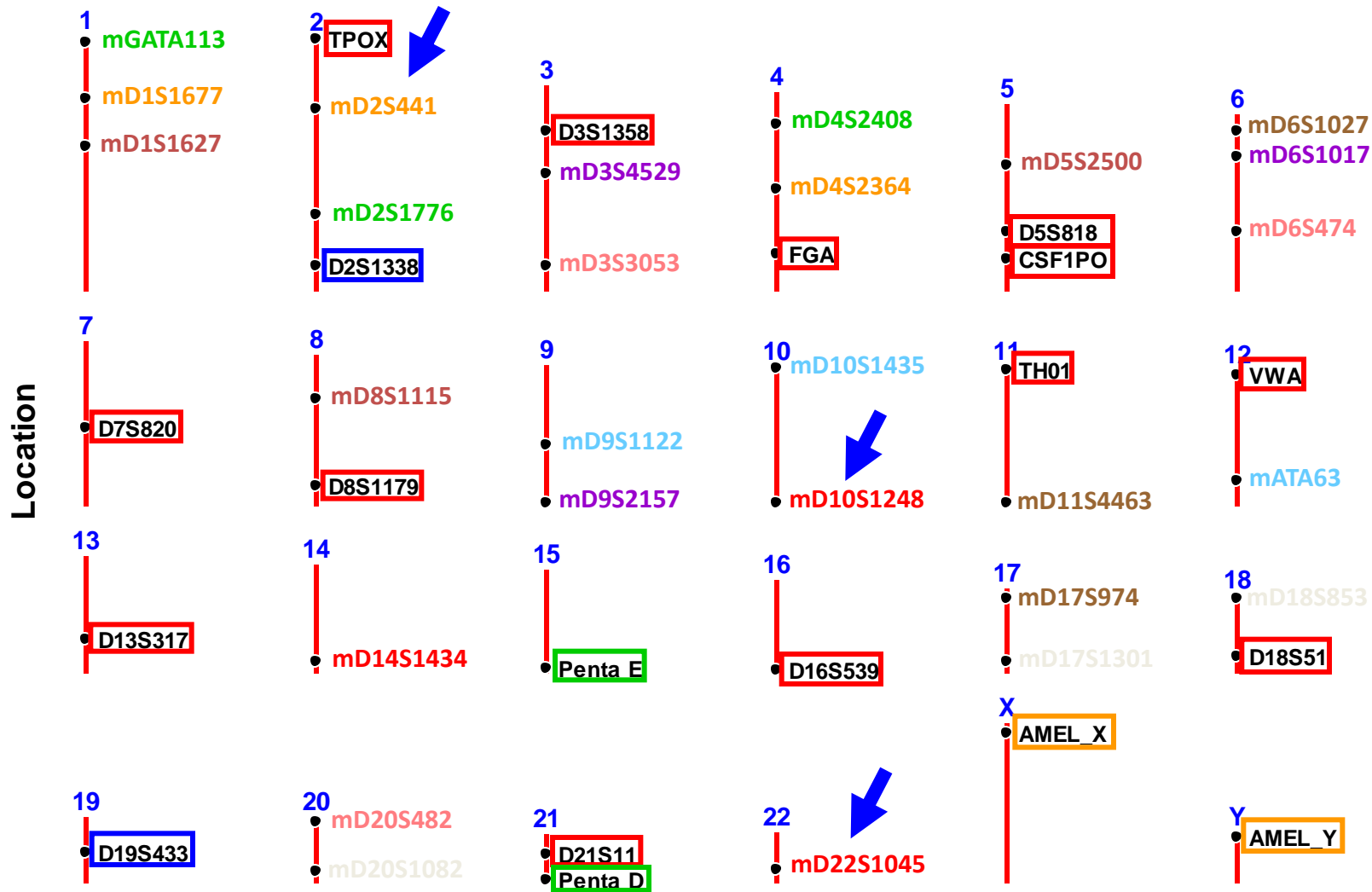
Sequence homozygotes to determine allele sizes

Build Macros for Genotyping

Construct Allelic Ladders



Chromosomal Locations of New miniSTR Loci



NC01	NC04	NC07
NC02	NC05	NC08
NC03	NC06	NC09

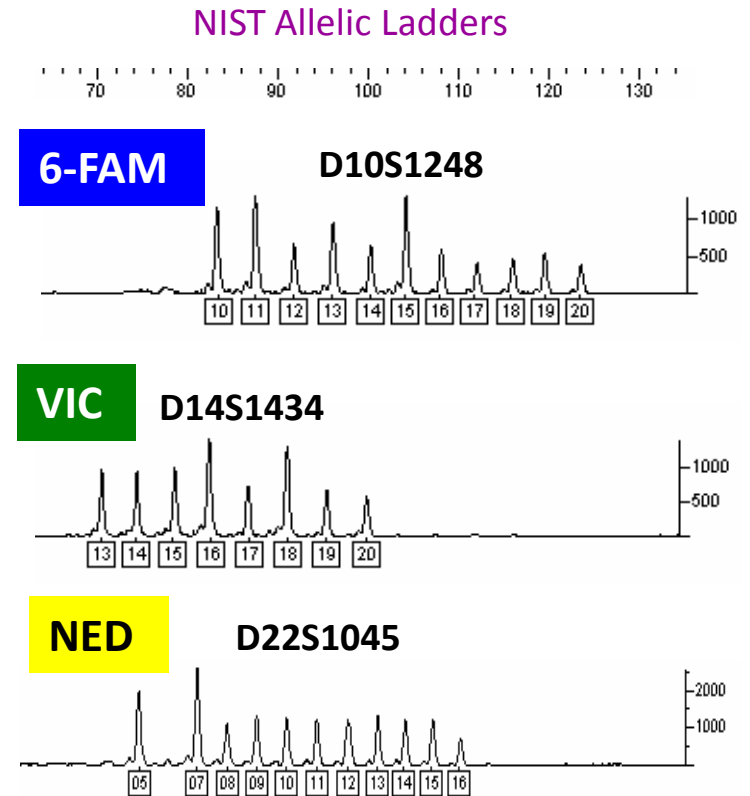
Chromosome

CODIS	PowerPlex 16
Identifier	Sex-Typing

Multiple Miniplexes

- **26 characterized loci** divided into 10 miniplexes
- One locus per dye color
- Allelic ladders created
- **Amplicons <140 bp**
- miniSTRs
- Work with 100 pg DNA
- **For degraded samples**
(bones in missing persons cases)

**NC = Non-CODIS or
non-core**



NC01 Loci

NC Miniplexes

NC01

D10S1248
D14S1434
D22S1045

NC02

D1S1677
D2S441

NC03

D3S3053
D6S474

26 total new loci

NC04

D1GATA113
D2S1776
D4S2408

NC05

D1S1627
D8S1115

D9S324

NC06

D3S4529
D9S2157

D10S1430

NC10

D3S3053
D6S474
D20S482

NC07

D9S1112
D12ATA63
D14S1280

NC08

D17S1301
D18S8534
D20S1082

NC09

D10S2327
D11S4463
D17S974

Removed because they were problematic

In Jan 2008 Issue of *J. Forensic Sci.*

J. Forensic Sci., Jan 2008, 53(1):73-80

J Forensic Sci, January 2008, Vol. 53, No. 1
doi: 10.1111/j.1556-4029.2008.00595.x
Available online at: www.blackwell-synergy.com

Carolyn R. Hill, M.S.; Margaret C. Kline, M.S.; Michael D. Coble,[†] Ph.D.; and John M. Butler, Ph.D.

Characterization of 26 MiniSTR Loci for Improved Analysis of Degraded DNA Samples

- Characterization of **26** new autosomal loci
- Primer sequences, GeneMapper bins and panels, genotypes on common samples, and allele frequency information **already available on STRBase**

<http://www.cstl.nist.gov/div831/strbase/miniSTR.htm>

<http://www.cstl.nist.gov/div831/strbase/newSTRs.htm>

European Labs Have Adopted the NIST-Developed NC (non-CODIS) miniSTRs

FSI (2006) **156(2)**: 242-244

Short communication

The evolution of DNA databases—Recommendations
for new European STR loci

**These 3 loci are included in the new European multiplex kits:
Applied Biosystems NGM kits and the Promega PowerPlex ESX 16/17
and ESI 16/17 systems**

^c*Department of Forensic Genetics, Institute of Forensic Medicine, University of Copenhagen, Denmark*

^d*Institute of Legal Medicine, University of Cologne, Germany*

Received 25 May 2005; accepted 26 May 2005

...recommended that existing multiplexes are re-engineered to enable small amplicon detection, and that **three new mini-STR loci with alleles <130 bp (D10S1248, D14S1434 and D22S1045) are adopted as universal**. This will increase the number of European standard Interpol loci from 7 to 10.

(D14 has been replaced with D2S441 from NC02)

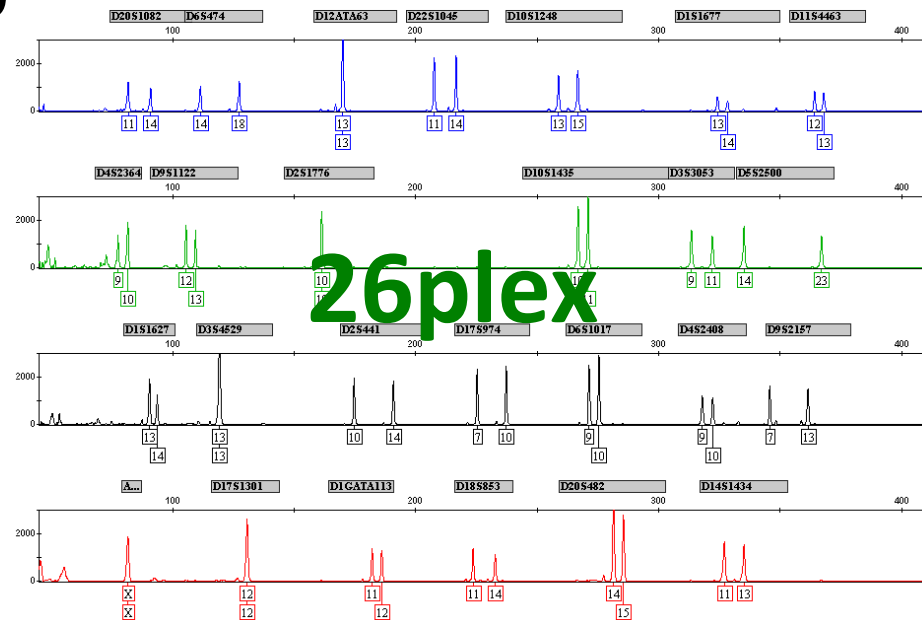
The Design of the Multiplex

- **Goal:** A single amplification 5-dye multiplex to combine the 26 new autosomal loci + Amelogenin in one reaction (27plex)
- How can this be achieved?
 - Initial placement of all loci within 6FAM, VIC, NED, and PET dye channels (the size standard is in the LIZ channel)
 - Primer redesign for all but 7 of the original miniSTR loci
 - Trial and error of primer compatibility, as well as balancing for all working primers

26plex STR Multiplex

- So far **25 STRs and amelogenin** in single multiplex (Eventual goal to have all 26 loci)
- Multiple loci in four dye channels
- **Amplicons 70 to 400 bp**
(No longer ‘miniSTRs’)
- Typically use 1 ng DNA, 30 cycles
- **For reference samples**
(a missing person’s relatives)

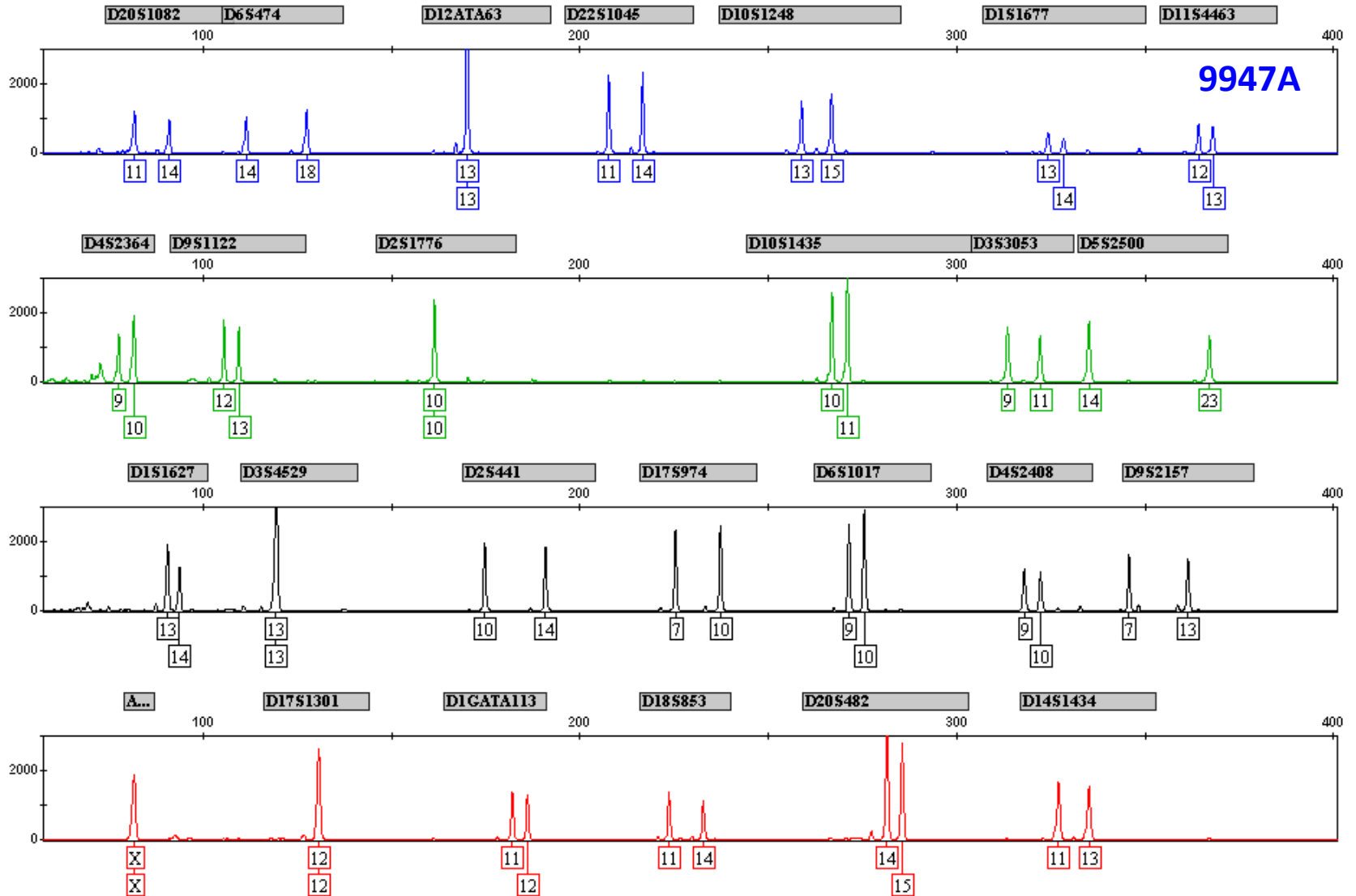
“Autoplex” or
“miniMegaplex”



All loci unlinked from core (CODIS) STRs

NIST STR 26plex

Hill et al. (2009) *Journal of Forensic Sciences*



Gender identification + 25 autosomal STR loci in a single amplification

In Sept 2009 Issue of *J. Forensic Sci.*

J. Forensic Sci., Sept 2009, 54(5):1008-1015

J Forensic Sci, September 2009, Vol. 54, No. 5
doi: 10.1111/j.1556-4029.2009.01110.x
Available online at: www.blackwell-synergy.com

Carolyn R. Hill,¹ M.S.; John M. Butler,¹ Ph.D.; and Peter M. Vallone,¹ Ph.D.

A 26plex Autosomal STR Assay to Aid Human Identity Testing*[†]

- Strategies for building multiplexes
- Primer sequences and PCR conditions listed
- GeneMapper bins and panels, genotypes on common samples, and allele frequency information **available on STRBase**

<http://www.cstl.nist.gov/biotech/strbase/str26plex.htm>

<http://www.cstl.nist.gov/div831/strbase/newSTRs.htm>

Final Thoughts and Advice

Support to the Community

...Bringing traceability and technology to the scales of justice...

- Conduct interlaboratory studies
- Perform beta-testing of new human identity testing products
- **We collaborate with other NIJ grantees**
- We provide input to (or have aided):
 - Scientific Working Group on DNA Analysis Methods (**SWGDM**)
 - Department of Defense Quality Assurance Oversight Committee for DNA Analysis
 - Virginia DFS Science Advisory Committee
 - American Prosecutor's Research Institute (**APRI**) DNA Forensics Program "Course-in-a-Box" for training lawyers
 - WTC Kinship and Data Analysis Panel (**KADAP**) and Hurricane Katrina efforts
 - NIJ Expert System Testbed (**NEST**) Project



A Few Words of Advice

- Hard work and studying are of the utmost importance in any field and really determining where your interests lie
- Penn State is an excellent school that opens many professional doors, especially in the forensic community
- Making contacts and having professional relationships in the field is crucial to getting your foot in the door, but the rest is up to you!
- Having skills in speaking and writing is essential in this field
 - DNA analysts: writing reports and going to court
 - Research scientists: writing journal articles and giving presentations about your research

A Few Useful Websites:

- U.S. Federal Government jobs
 - www.usajobs.gov



- American Academy of Forensic Sciences
 - www.aafs.org



- Mid-Atlantic Association of Forensic Scientists
 - www.maafs.org



- National Institute of Justice
 - www.dna.gov



- **STRBase**
 - www.cstl.nist.gov/strbase



Thank you for your attention

Acknowledgments: Applied Biosystems, Promega, and Qiagen for STR kits used in concordance studies

Contact Information

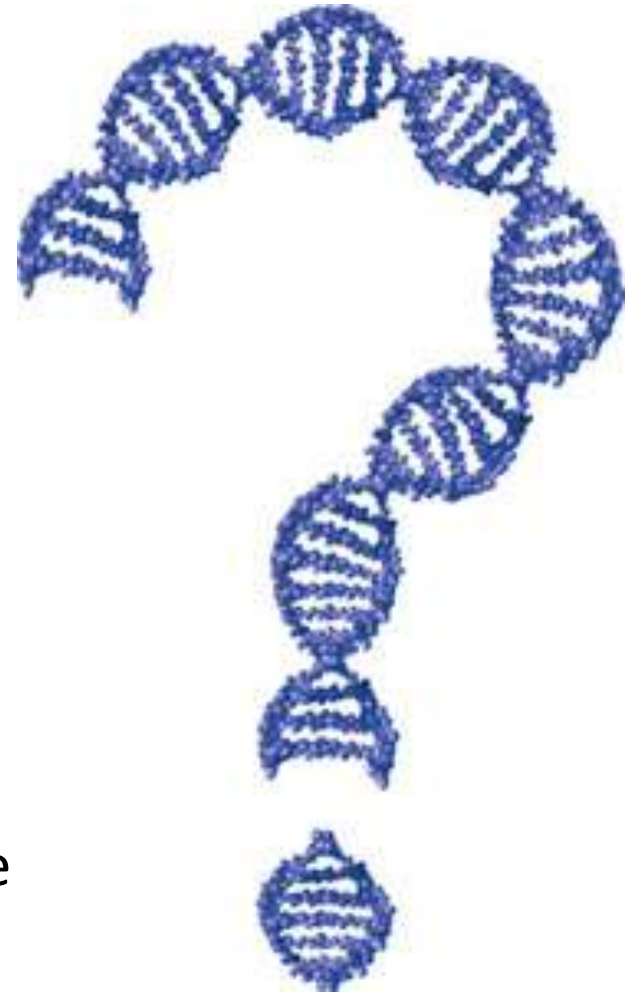
Becky Hill

Research Biologist

becky.hill@nist.gov

301-975-4275

<http://www.cstl.nist.gov/biotech/strbase>



Our team publications and presentations are available at:
<http://www.cstl.nist.gov/biotech/strbase/NISTpub.htm>