



Typing Single Nucleotide Polymorphisms (SNPs) Located on the Y Chromosome and in the Mitochondrial Genome

NIST Division Seminar

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DNA Technologies Group

Overview

SNPs

Assay Platforms and Instrumentation

Multiplexing

U.S. Population Samples

Y Chromosome and Mitochondrial Markers

Results

mtSNP 11 plex

Y-SNP multiplexes



SNP

Single Nucleotide Polymorphism

The diagram illustrates three types of SNP mutations:

- Sequence variation:** A red box highlights the first sequence "AGGCTACGT". An arrow points down to the mutated sequence "AGGCCACGT", where the second base ('G') is replaced by a 'C'.
- Deletion:** An arrow points down to the mutated sequence "AGGC-ACGT", where the third base ('G') is deleted.
- Insertion:** An arrow points down to the mutated sequence "AGGCTCACGT", where a 'C' has been inserted after the second base ('G').

Low mutation rate 10^{-8}
Typically Biallelic

SNP Facts

Most common type of variation in the human genome
(90%)

Estimated to occur every 100-300 bases

For a SNP to be defined it must occur in at least 1 % of the population

2 out of every 3 SNPs is a C-T transition

Occur in coding and non coding regions of the genome

http://www.ornl.gov/TechResources/Human_Genome/faq/snps.html

Biomedical Importance of SNPs

Variations in DNA sequence can have a major impact on how humans respond to **disease, bacteria, viruses, toxins**,

According to a Frost & Sullivan report - "U.S. SNP Detection Technology Market," the market share of the diagnostics segment will grow from 11% in 2001 to 33% by 2009.

The report stated that the total SNP detection market generated \$91.28 million in 2001. This market is estimated to reach \$310.76 million by 2008.

from generation to generation --making them easier to follow in population studies



APBiotech - AstraZeneca - Aventis - Bayer - Bristol-Myers Squib - F.Hoffman-La Roche - Glaxo Wellcome
THE SNP CONSORTIUM LTD

IBM - Motorola - Novartis - Pfizer - Searle - SmithKline Beecham - Wellcome Trust

Discovered and characterized nearly 1.8 million SNPs

Forensic Utility of SNPs

Human identification purposes (criminal, **paternity**, evolutionary, population studies, **predicting ethnicity**)

The short PCR amplicons required for typing SNPs may result in success with **degraded samples** and possibly higher sensitivity

Simplicity in testing – typically bi-allelic markers (versus length polymorphisms)

Improve assay development (both multiplex PCR and SNP detection)

For serious forensic usage parallel high-throughput methods will be required for typing

Forensic Utility of SNPs

Short tandem repeat (STR)

CTGATGCTA(**GATA**)_n GACTACTTA

$n = 5 \text{ to } 15 = 66$ possible allelic combinations

SNP

CTGATGCTA(**G/A**)GACTACTTA

3 possible allelic combinations

For human ID purposes more
SNPs would be needed than STRs
Multiplexing is essential

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Instrumentation

PCR & primer extension



Multi-Color Capillary Electrophoresis
(ABI 310 or 3100)

Luminex Beads
hybridization



Luminex 100 Flow Cytometer

Primer Extension



Time-of-Flight Mass Spectrometer

TaqMan



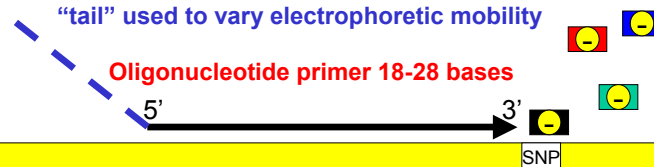
ABI 7000 SDS

Allele-Specific Primer Extension (ASPE)

Primer is extended by one base unit

"tail" used to vary electrophoretic mobility

Oligonucleotide primer 18-28 bases



ABI PRISM® SNaPshot™
Multiplex System

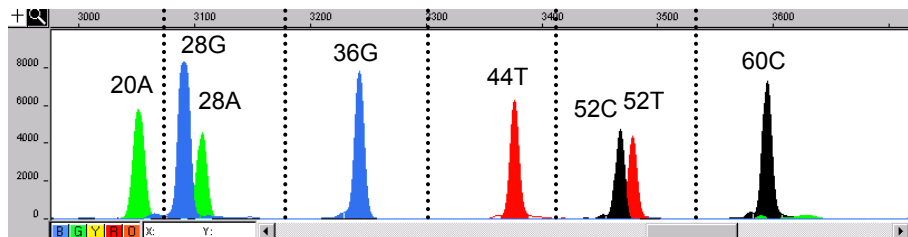
Fluorescently
labeled ddNTPs +
polymerase

PCR Amplified DNA Template

ddNTP	Dye label	Color
A	dR6G	Green
C	dTAMRA	Black
G	dR110	Blue
T	dROX	Red

25 Cycles
96°C 10s
50°C 5s
60°C 30s

Detection of SNPs with ABI 310/3100



SNaPshot™ CEPH Control Reaction

20 nucleotides ddA

36 nucleotides

ddG

Multiplexing possible by use
of different length primers

44 nucleotides

ddT

60 nucleotides

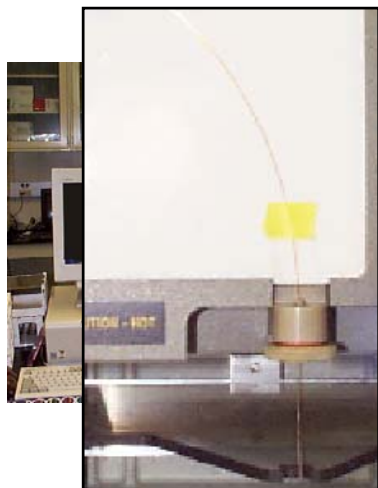
ddC

Poly(T) tail or non-nucleotide linker
to aid separation

Priming site

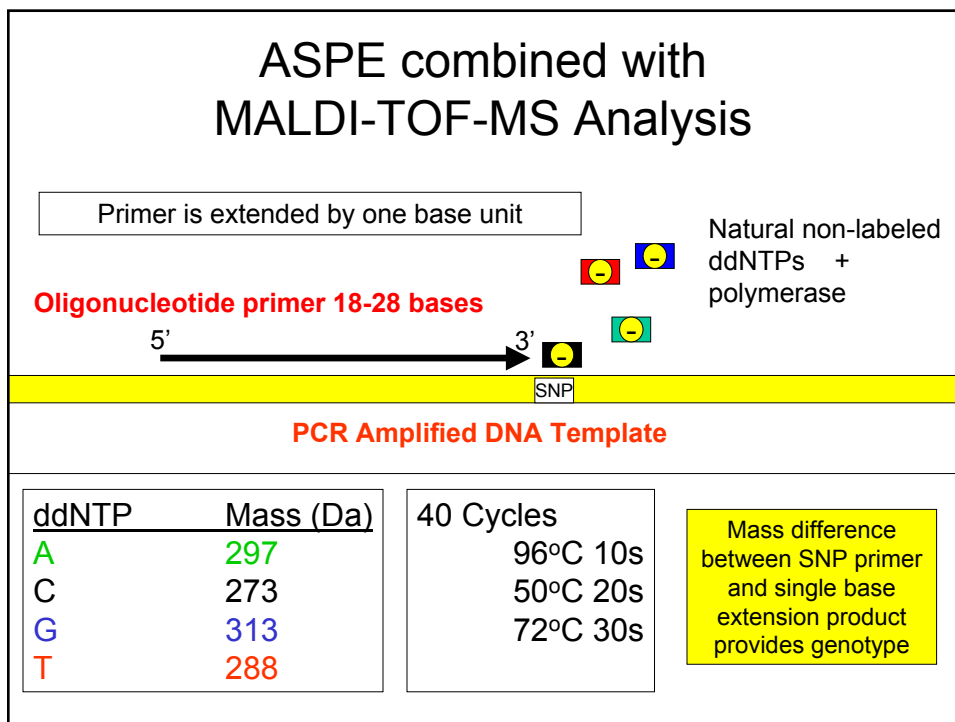
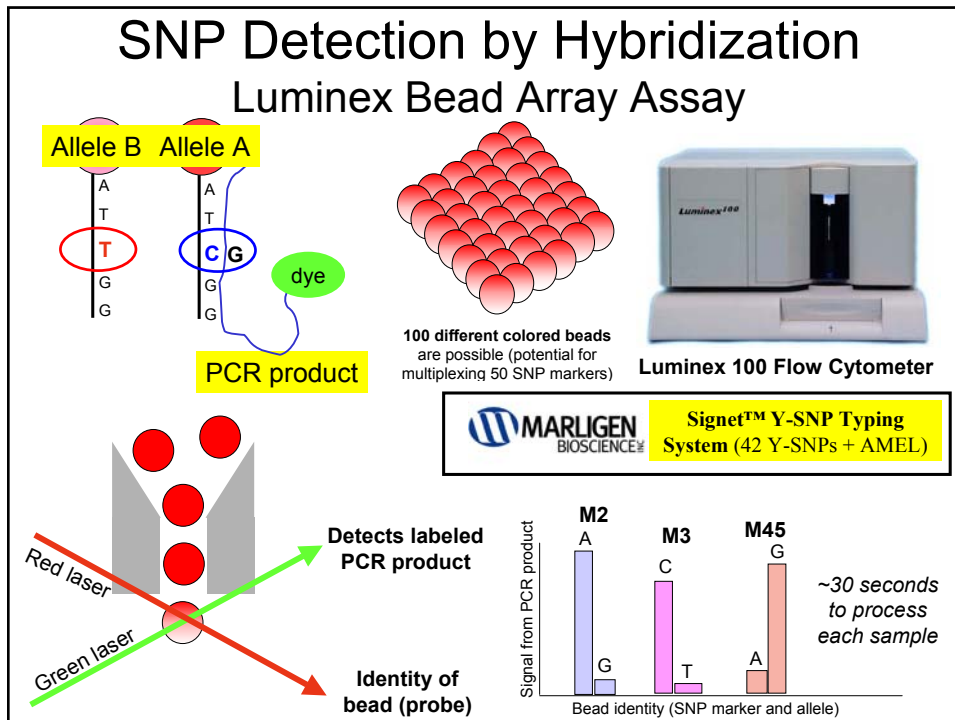
Capillary Electrophoresis Instrumentation

ABI 310
single capillary

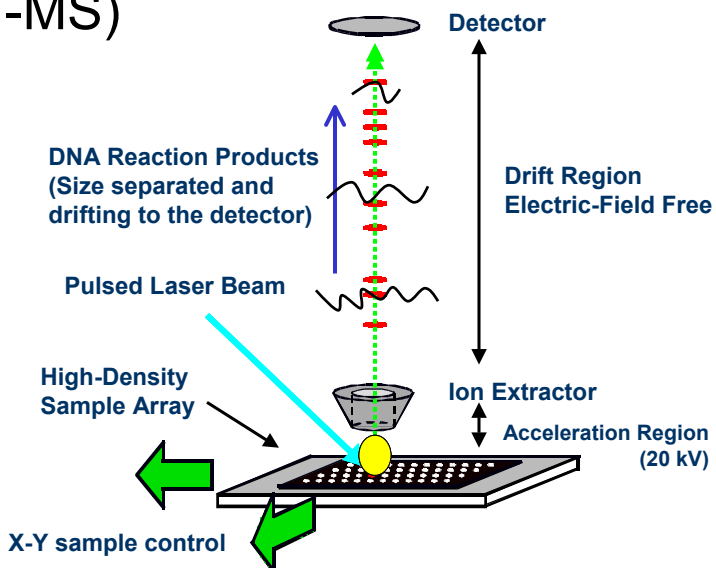


ABI 3100
16-capillary array

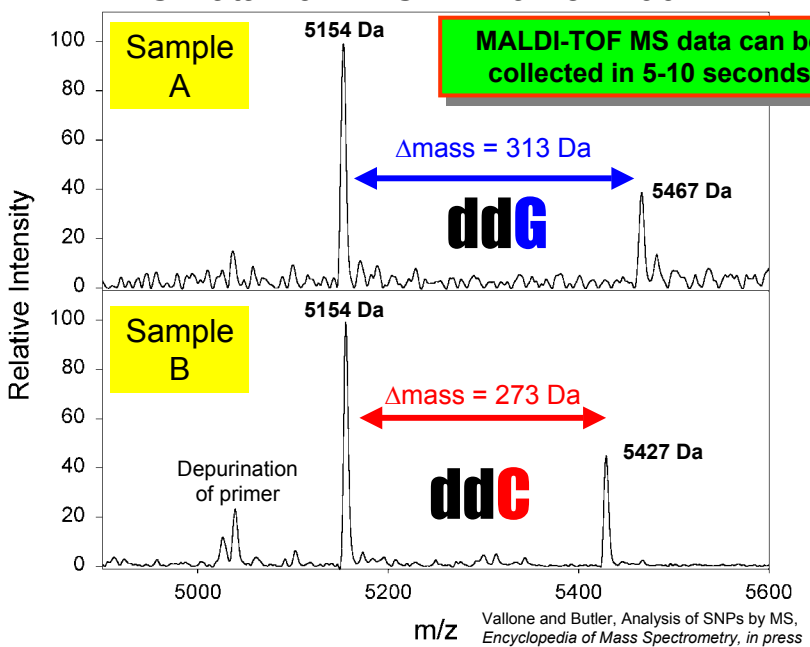




Time-of-Flight Mass Spectrometry (TOF-MS)

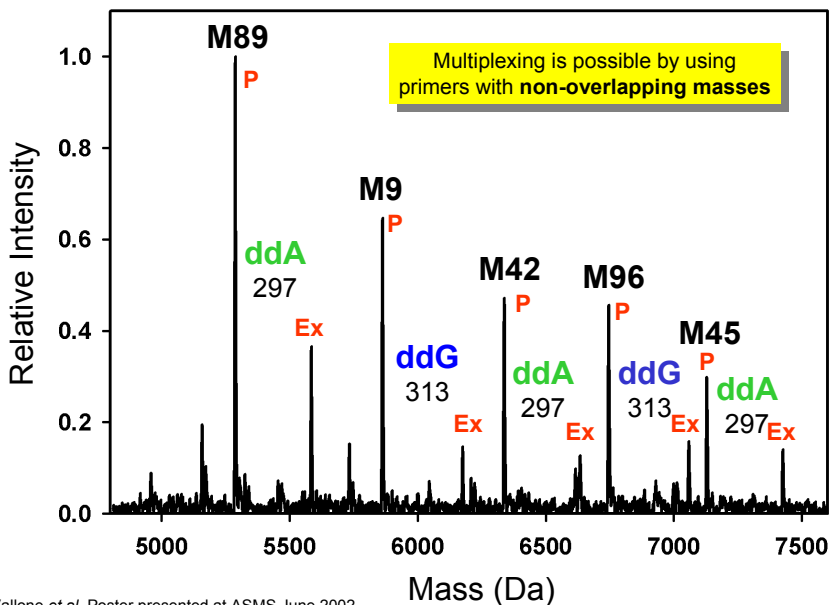


MS Data from Y SNP Marker M96

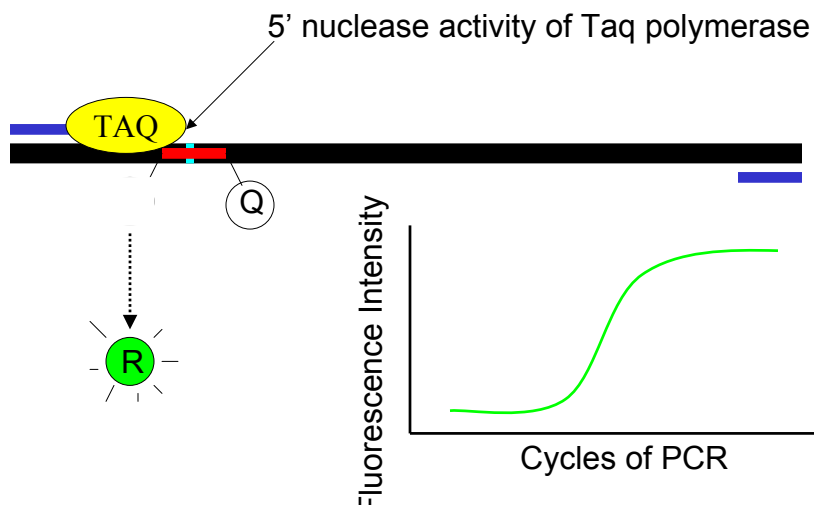


Vallone and Butler, Analysis of SNPs by MS,
Encyclopedia of Mass Spectrometry, in press

SNP (5-plex) Analyzed by TOF-MS



Taq Man – Real Time PCR



Allelic Discrimination Assay using TaqMan

Design **2** Taqman probes for each expected SNP
Each probe will have a different reporter dye (FAM/VIC)

Possible assays outcomes

SNP A = FAM

SNP B = VIC

SNP A/B = FAM/VIC



Typically TaqMan genotyping assays are singleplex!

Summary of SNP Assays

	Advantages	Disadvantages
ASPE-CE	Moderate degree of multiplexing	Development of multiplex
	<u>Other Technologies</u>	
	Pyrosequencing	
ASPE-M	Chip Based – Affymetrix - Agilent	Multiplexing
	Allele specific PCR	more difficult than CE
	Invader-mismatch cleavage	custom
Microbe (Luminescence)	Orchid SNPstream	
	Illumina Bead Arrays	
TaqMan	Rapid - one step Good for one marker on 1000's of sample	No multiplexing Costly for typing many SNPs & few samples

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mtSNP 11 plex

Y-SNP multiplexes



Advantages of Multiplexing

Obtain more information per unit time

Reduce the amount of limited forensic sample used

Save on reagents; enzyme, buffers, DNA oligomers

Reduces labor

Streamlines data analysis

For certain markers it is essential (SNPs, YSTRs)

Coincides with high capacity instrumentation and new SNP typing technologies

Goals for Multiplex Assay Development

Working with collaborators who have markers of forensic interest

Evaluate the forensic utility of newly discovered markers (medium sized multiplexes 5 – 10 loci)

Further the understanding of developing multiplex assays (primer design, QC)

Publish assay details for others to evaluate (commercial and research)

Multiplex PCR Primer Selection

Identify markers of interest (collaborations, literature, research)

Organize sequences with ~250 bases of sequence up- and downstream of the SNP

PCR product size

Short amplicons for degraded samples, SNPs

Longer amplicons for STRs

Use software for selecting singleplex primer pairs

Primer3

www-genome.wi.mit.edu/genome_software/other/primer3.html

Steve Rozen and Helen J. Skaletsky (2000) Primer3 on the WWW for general users and for biologist programmers. In: Krawetz S, Misener S (eds) *Bioinformatics Methods and Protocols: Methods in Molecular Biology*. Humana Press, Totowa, NJ, pp 365-386

Multiplex PCR Design

Select singleplex PCR primers for each amplicon
using Primer 3 software

OLIGO	start	len	tm	gc%	any	3' seq
LEFT PRIMER	27	20	60.06	50.00	4.00	2.00 GGGATAAACAGCGCAATCCTA
RIGHT PRIMER	174	22	60.31	50.00	8.00	3.00 CGGTCTGAACTCAGATCACGTA
SEQUENCE SIZE: 205						
INCLUDED REGION SIZE: 205						
PRODUCT SIZE: 148, PAIR ANY COMPL: 3.00, PAIR 3' COMPL: 2.00						
EXCLUDED REGIONS (start, len)*: 70,65						
1	CTTGACCAACGGAACAAGTTACCCTAGGGATAACAGCGCAATCCTATTCTAGAGTCCATA					
61	TCAACAATAGGGTTTACGACCTCGATGGATCAGGACATCC					
121	TTAAAGGTTCGTTGTTCAACGATTAAAGTCCTACGTGATCTGAGTTCAGACCGGAGTAA					
181	TCCAGGTGGTTCTATCTACCTTC					

Running Primer3 Locally

Sending multiple sequences over the web for primer selection can be tedious

The Primer3 web output is acceptable for the screen viewing or printing but not for organizing in spreadsheets

Primer3 is publicly available and can be run on a Unix, PC (Linux), or Mac (OSX) computer

Developed a program that formats files for Primer3 input

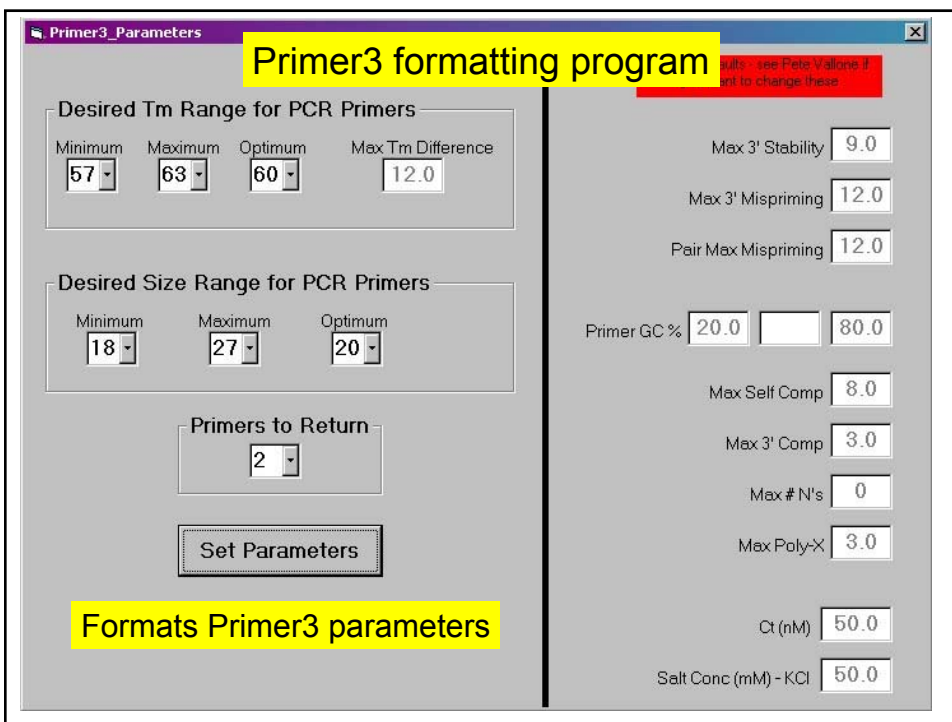
Reference sequences that are stored in Excel can be quickly formatted for Primer3

Format of Template Sequences

Locus	Total Length	Minimum	Maximum	Optimal	Excluded Region	SNP site
M3	255	105	150	125	174,60	204
	TGATTATTTAGAACAAAACAATAACAAACAAATGGTCCCTGTTAAATGTG					
M9	255	105	150	125	237,60	267
	CCTGTGCACGCCAAGCGGAAGCTGAAGTGCAGCGTCTTGATCTCAATCCTGGAG					

Sequences stored in excel

Will be adapted for FASTA format & comma delimited



Example input
format for
Primer3

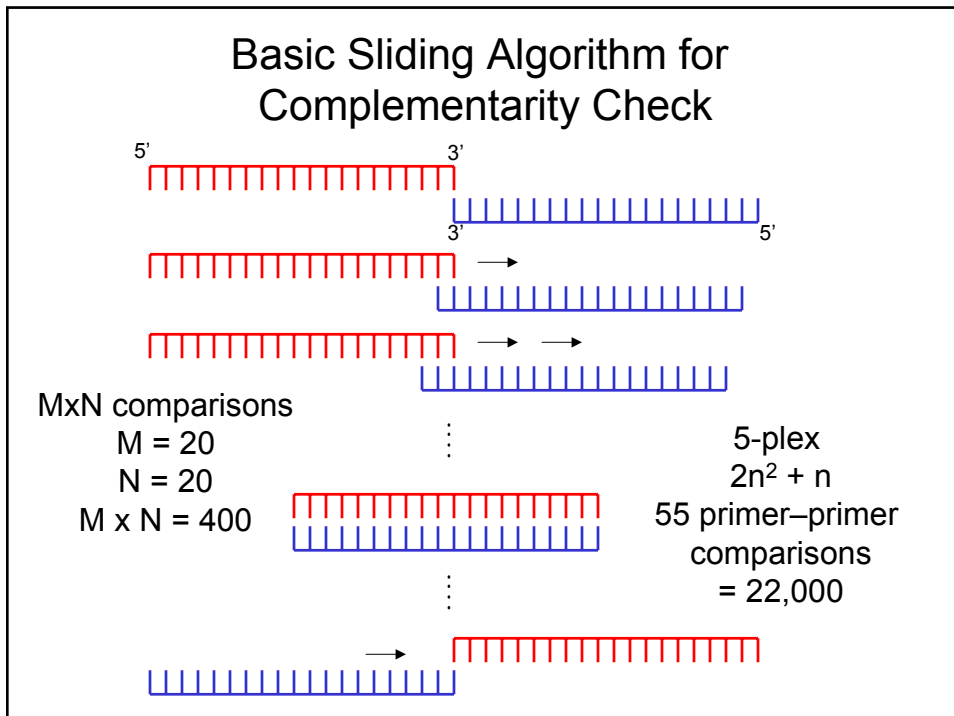
```
PRIMER_SEQUENCE_ID=M9
SEQUENCE=GCAGCATATAAAACTTCAGGACCCGTGAAATACAGAAGT
CAAAGAACGGCTTAAGATGGTTGAATNCTTTATTTCCTTAATTTAG
ACATGTTCAACGTTCAATGCTTACACTTACTAGTTATGTAAGTAG
CGCTTACTTCATTATGCATTTCAACTCAAAAAAATCCTTGTGAAAT
GTTGAAATTTTCTAATCTGTTACGAGCTTCAAAATGAGGAAAAAA
GATTCAAGTTACATTACAGCAGGAAATGCCCTTTTAAATCGGATTATGTTT
ACTTAACATTACAGTACATTACGCTTGAGCAAAGTTAGGTTT
PRIMER_COMMENT=(340 bp); G to C at position 68
PRIMER_MISPRIMING_LIBRARY=/Users/vallone/Desktop/primer3/misprM9
PRIMER_MAX_MISPRIMING=8
PRIMER_PAIR_MAX_MISPRIMING=20
EXCLUDED_REGION=38,60
PRIMER_PRODUCT_SIZE_RANGE=90-150
PRIMER_PRODUCT_OPT_SIZE=105
PRIMER_MIN_SIZE=18
PRIMER_MAX_SIZE=27
PRIMER_OPT_SIZE=20
PRIMER_OPT_TM=60
PRIMER_MIN_TM=57
PRIMER_MAX_TM=63
PRIMER_NUM_RETURN=1
PRIMER_EXPLAIN_FLAG=1
PRIMER_LIBERAL_BASE=1
=
PRIMER_SEQUENCE_ID=M42
SEQUENCE=AAAGCGAGAGATTCAATCCAGGATGACAGAATGCGTTCAC
CTTTAAAGGGATTAAGAAGTATAATACAGTCTGTATTAGATCACCC
AGAGACACACAAAACAAGAACCGTGAATTGAATTAGTGGTATACTAATAG
AGTCTTCTTACCTTAAATTTACACATATCTTACATTTCTTACACACAC
```

Non-Specific Interactions

Primers that interact with non-specific (undesired) regions of a genome OR with each other can degrade PCR performance

Screening for alternate genomic binding regions can be accomplished using **BLAST** <http://www.ncbi.nlm.nih.gov>

Screening for potential primer-dimer interactions is accomplished using in house software - **AutoDimer**



Auto Dimer Check

File Help About

Primer Dimer Checker	Cancel
Hairpin Checker	
<hr/>	
Minimum SCORE Requirement	6
<hr/>	
# of Sequences	22
# of Hits	6
Total Number of Primer-Primer Comparisons	253
Na ⁺ (Molar)	0.085
Total Strand Conc (micromolar)	1.0

AutoDimer $\rightarrow 2n^2+n$

C:\Documents and Settings\petev\My Documents\SNP\mSNP\afdl\Paper\nPCR_primers H1.txt

7202-F ACGCCAAAATCCATTCACT versus 16519-F ACCACCATCCTCCGTGAAAT

Matches = 7

Score = 6

ATTCACN

est. tm = 3.6 oC

DeltaG @37 degrees = -3.85 kcal/mole

3'-TAAAGTGCCTCCTACCACCA-5'
||||||| x

5'-ACGCCAAAATCCATTCACT-3'

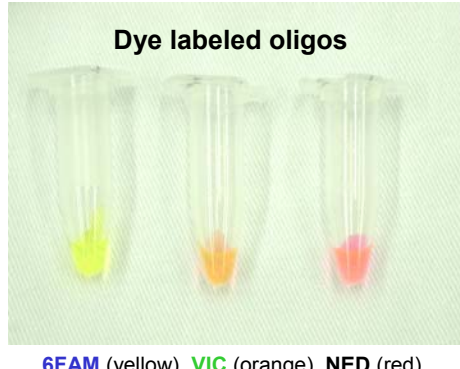
5'-TGTTGGGTCTCATGACTTTGGA-3'

10211-F ACCACAACTAACGGCTACA versus 3010-R TCACGTAGGACTTTAATCGTTGA

Matches = 9

Score = 6

PCR Primer Quality Control



- **UV Spec** to determine concentration
- **HPLC** to evaluate purity
- **TOF-MS** to confirm correct sequence

6FAM (yellow), VIC (orange), NED (red)

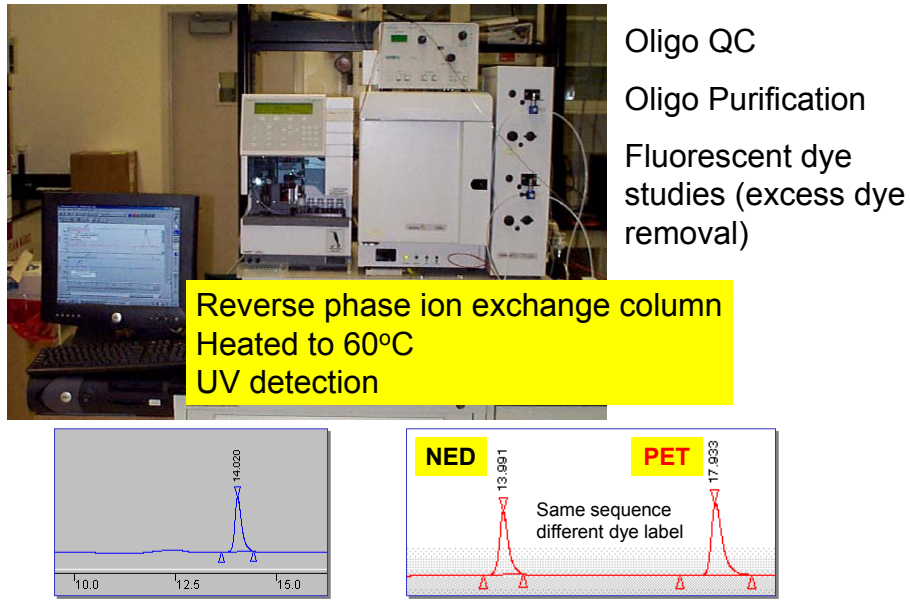
Butler *et al.* (2001) *Forensic Sci. Int.* 119: 87-96

Determination of DNA Oligomer Concentrations

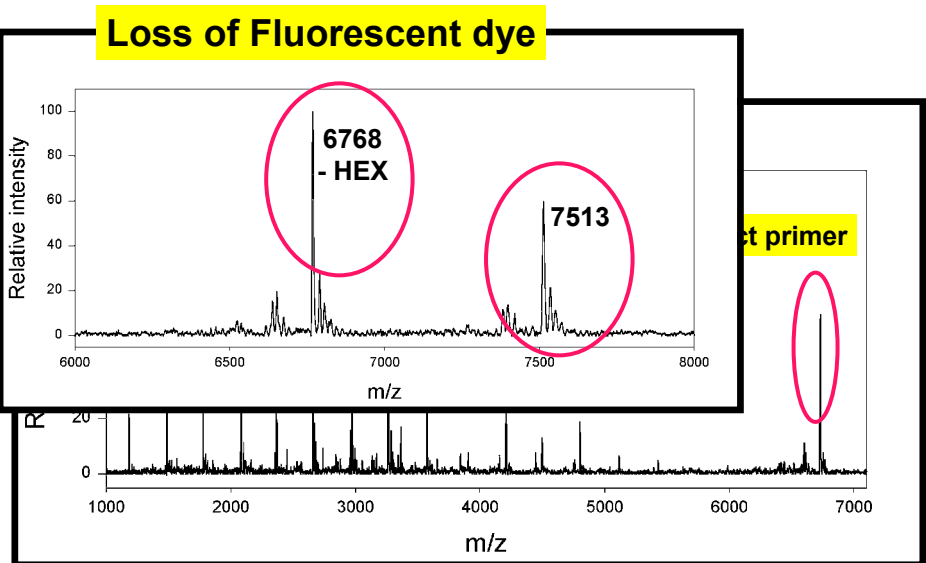
Expected 100 μ M	μ M	% deviation
1	173.3	42.3
2	164.8	39.3
3	155.0	35.5
4	124.1	19.4
5	116.4	14.1
6	98.5	-1.5
7	108.6	7.9
8	103.1	3.0
9	120.8	17.2
10	79.6	-25.7
11	83.0	-20.5

Concentrations were estimated by UV Spec readings @260 using extinction coefficients determined from nearest-neighbor values

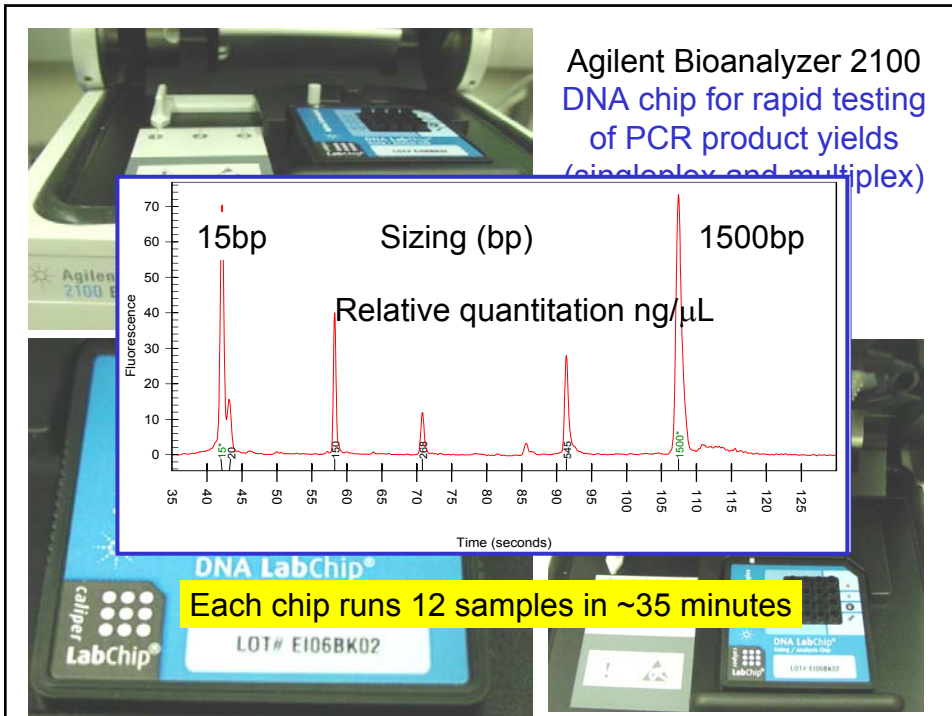
Varian Helix DHPLC System



MALDI QC of Commercial Oligos



Vallone and Butler (Oct 2000) International Symposium on Human Identification (Biloxi, MS)



Publications Describing Multiplex Assay Design

Schoske, R., Vallone, P.M., Ruitberg, C.M., Butler, J.M. (2003) Multiplex PCR design strategy used for the simultaneous amplification of 10 Y chromosome short tandem repeat (STR) loci. *Anal. Bioanal. Chem.*, 375: 333-343.

Butler, J.M., Schoske, R., Vallone, P.M. Highly multiplexed assays for measuring polymorphisms on the Y-chromosome. (2003) *Progress in Forensic Genetics 9* (Brinkmann, B. and Carracedo, A., eds.), Elsevier Science: Amsterdam, The Netherlands, International Congress Series 1239, pp. 301-305.

Schoske, R., Vallone, P.M., Kline, M.C., Redman, J.W., Butler, J.M. (2003) High-throughput Y-STR typing of U.S. populations with 27 regions of the Y chromosome using two multiplex PCR assays, *Forensic Sci. Int., in press*

Butler, J.M. (2003) Constructing STR multiplex assays. *Methods in Molecular Biology: Forensic DNA Typing Protocols* (Carracedo, A., ed.), Humana Press: Totowa, New Jersey, *in press*.

Butler, J.M., Schoske, R., Vallone, P.M., Kline, M.C., Redd, A.J., Hammer, M.F. (2002) A novel multiplex for simultaneous amplification of 20 Y chromosome STR markers. *Forensic Sci. Int.* 129: 10-24.

Butler, J.M., David, V.A., O'Brien, S.J., Menotti-Raymond, M. (2002) The MeowPlex: a new DNA test using tetranucleotide STR markers for the domestic cat. *Profiles in DNA*, Promega Corporation, Volume 5, No. 2, pp. 7-10. http://www.promega.com/profiles/502/ProfilesInDNA_502_07.pdf

Butler, J.M., Devaney, J.M., Marino, M.A., Vallone, P.M. (2001) Quality control of PCR primers used in multiplex STR amplifications. *Forensic Sci. Int.*, 119: 87-96.

Butler, J.M., C.M. Ruitberg, Vallone, P.M. (2001) Capillary electrophoresis as a tool for optimization of multiplex PCR reactions, *Fresenius J. Anal. Chem.* 369: 200-205.

Overview

- SNPs
- Assay Platforms and Instrumentation
- Multiplexing
- U.S. Population Samples
- Y Chromosome and Mitochondrial Markers
- Results
 - mtSNP 11 plex
 - Y-STR multiplexes



NIST U.S. Population Samples

As of 08/2003 **666** males (anonymous; self-identified ethnicities)

260 Caucasians
260 African Americans
143 Hispanic
3 Asian

Whole blood received from
Interstate Blood Bank (Memphis, TN)
**~80 µg total extracted
genomic DNA**
Working plates 1 ng/uL



To date: (35,139 allele calls)
Identifiler (15 autosomal markers + Amelogenin) (10,608)
Roche Linear Arrays (HV1 HV2 10 regions) (6,630)
Y-STRs 27 markers (17,901)

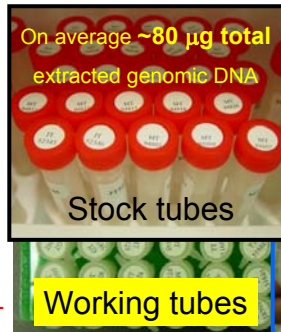
NIST U.S. Population Samples

As of 09/2003 **666 males** (anonymous; self-identified ethnicities)

260 Caucasians
260 African Americans
143 Hispanics
3 Asians

Whole blood received from
Interstate Blood Bank (Memphis, TN)

Working tubes/plates 1 ng/uL



Working tubes



Working plates

To date: (~50,000 allele calls)

Identifier (15 autosomal markers + Amelogenin) (10,608)

Roche Linear Arrays (HV1/HV2 10 regions) (6,630)

Y-STRs 22 loci—27 amplicons (17,388)

Y-SNPs 50 markers on sub-set of samples (11,498)

Samples supplied to
OhioU for miniSTR typing
and **AFDIL** for whole
mtGenome sequencing

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mtSNP 11 plex

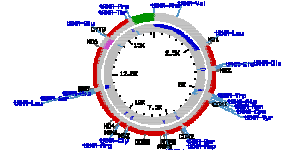
Y-SNP multiplexes



Markers of Interest

- Mitochondrial DNA (mtDNA)

- maternally inherited
 - polymorphic control region (D-loop)
 - ~500-2000 copies per cell
 - coding region



- Y chromosome

- paternally inherited
 - variety of Y-STR and Y-SNP markers
 - ***haplotype rather than genotype***



Require large databases because recombination does not occur

The Y Chromosome

60,000kb total size

The non-recombining region (NRY) consists of 95% of the Y chromosome

NRY is passed on as a block of information

Variations in the NRY are due to mutation only

Potential for predicting geographical origin

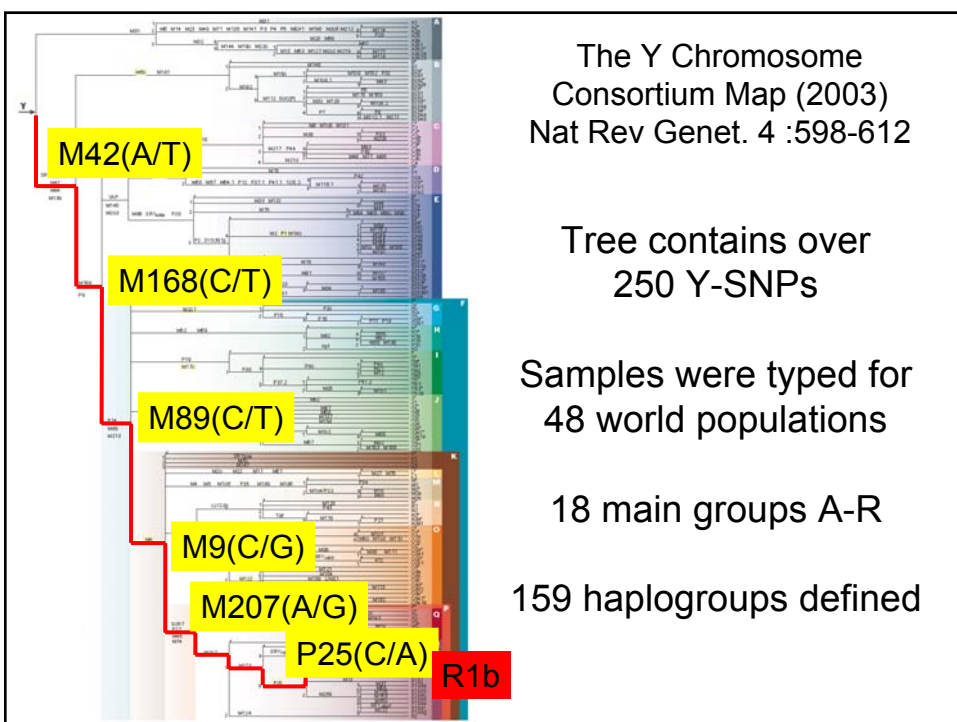
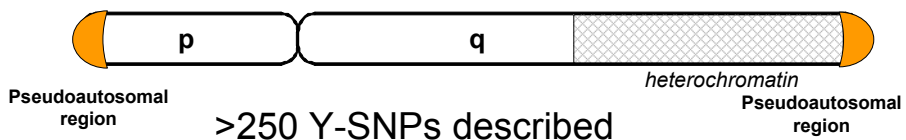


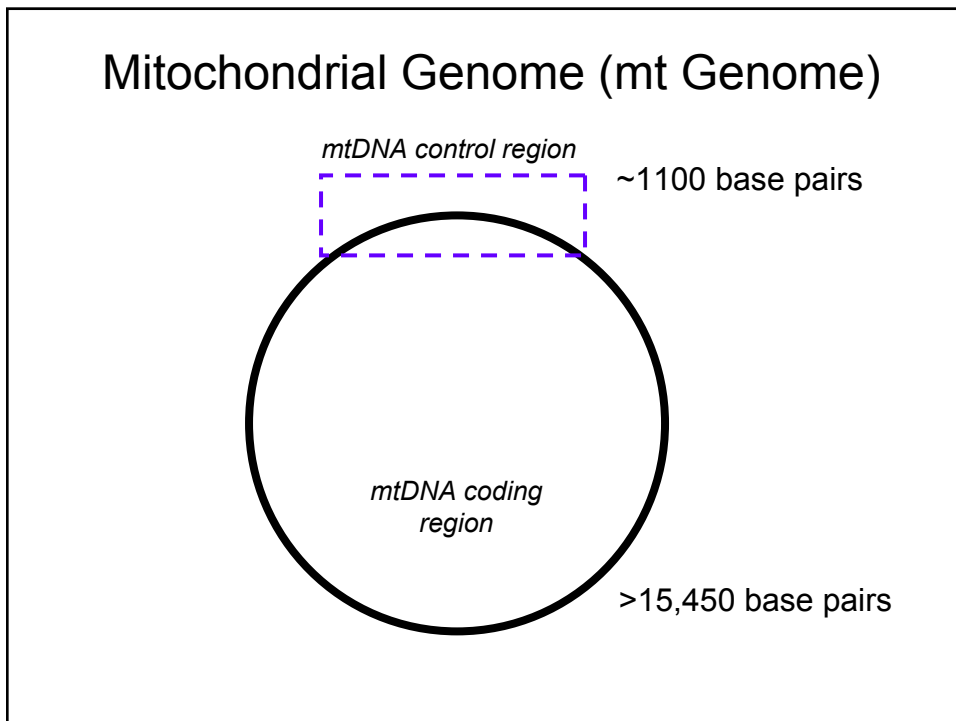
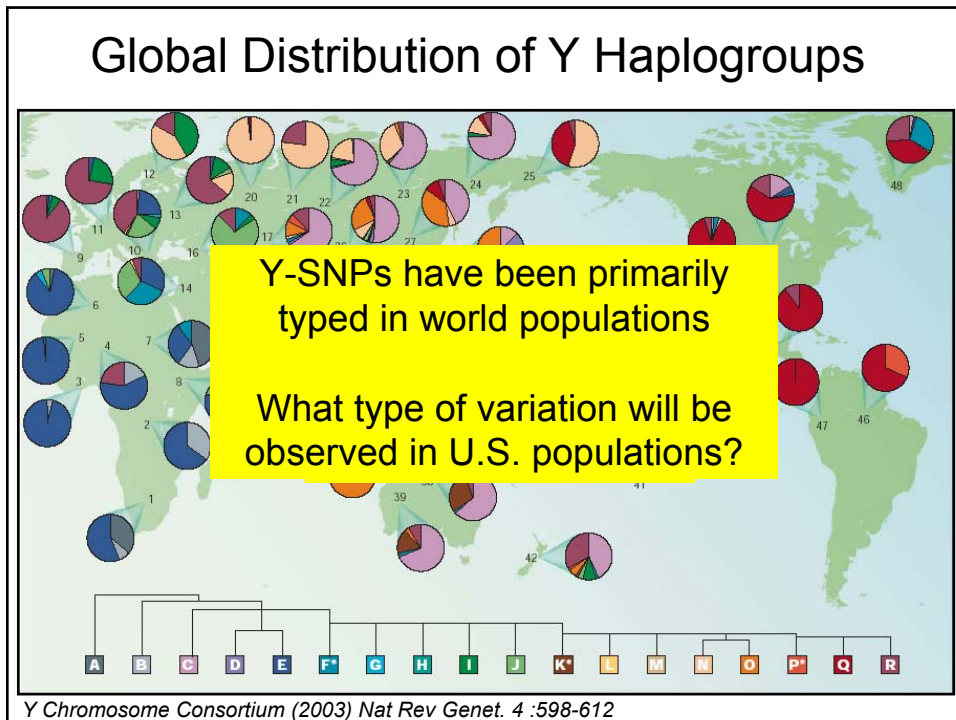
Forensic Utility of Y Chromosome SNPs

Y chromosome markers are useful in mixed male - female samples

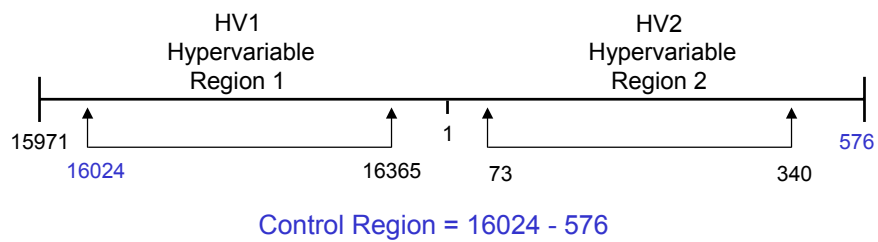
Haplogroups are non-randomly distributed among populations therefore potential exists for predicting population of origin

Low mutation rate of SNPs 2×10^{-8} per base per generation





The Current mtDNA Amplification & Sequencing Strategy Focuses on the Hypervariable Regions of the mitochondrial genome HV1 and HV2



In Caucasians, approximately 7% of HV1 and HV2 sequences are identical

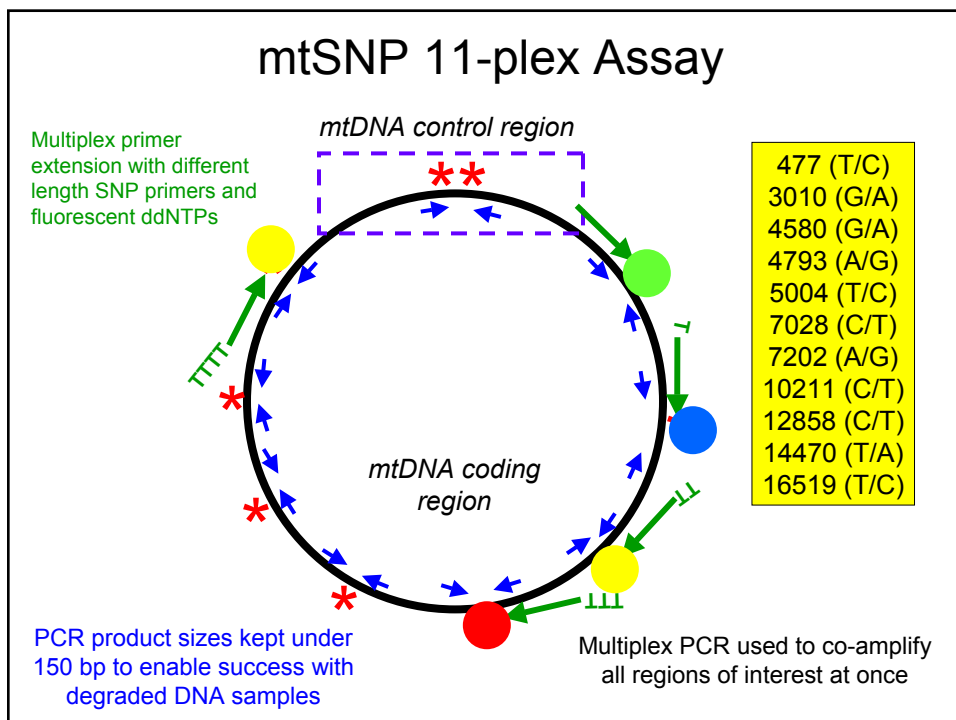
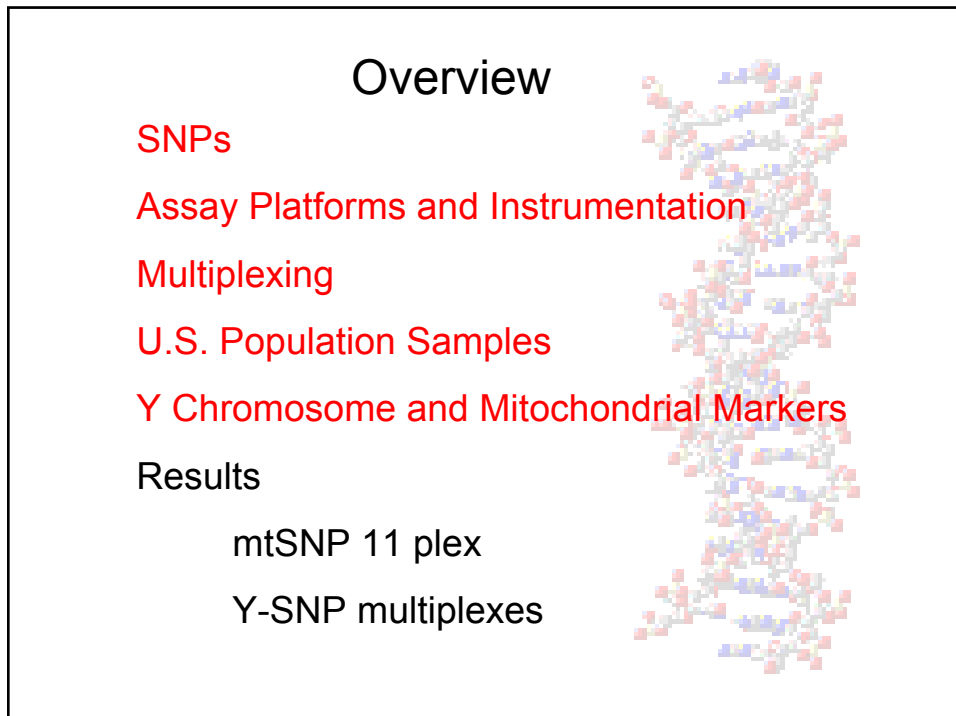
The Use of Full mtGenome Polymorphisms

mtGenome sequencing data reveals numerous SNPs that can help distinguish Caucasians sharing common HV types (Tom Parsons and Mike Coble AFDIL) 241 mt genomes

11 SNP sites were selected to help resolve Caucasian individuals having the most common HV1/HV2 type

mtSNPs: Silent and at third codon positions or fall in the short non-coding regions between genes in the coding region

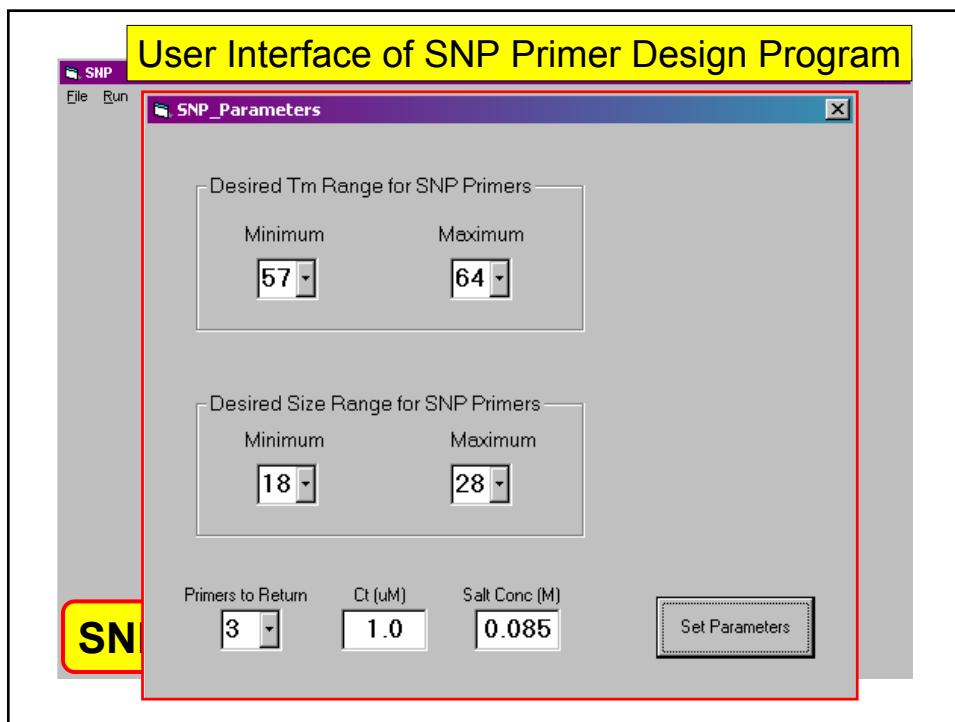
Detect in a multiplex assay run on a common forensic instrumental platform



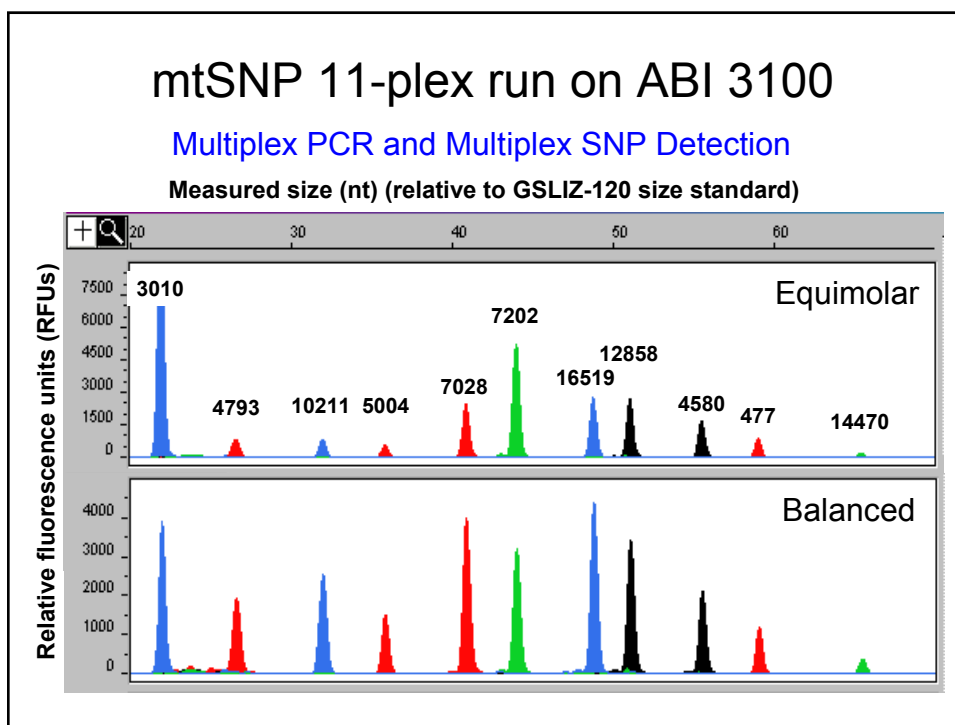
Tailed SNP primers allows for multiplexing in the SNaPshot assay

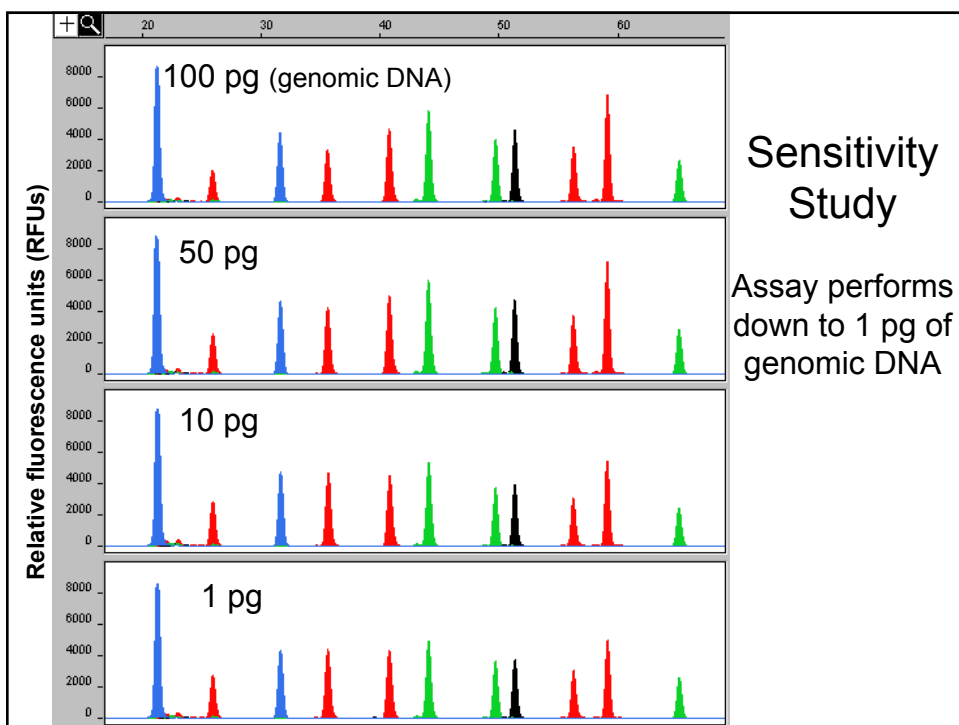
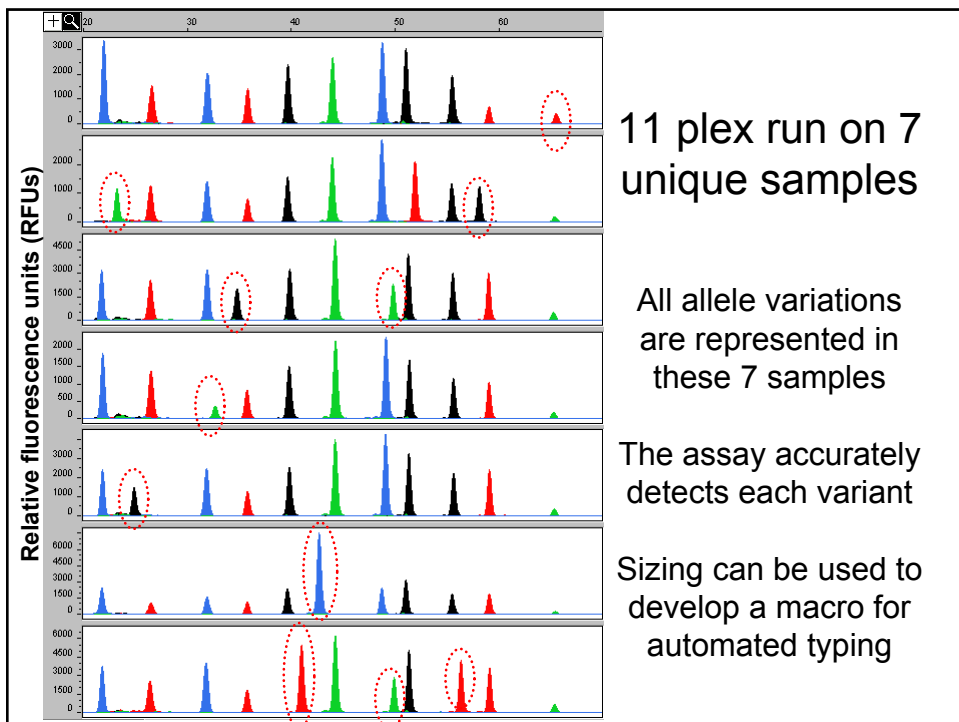
Sequences for 11 extension primers

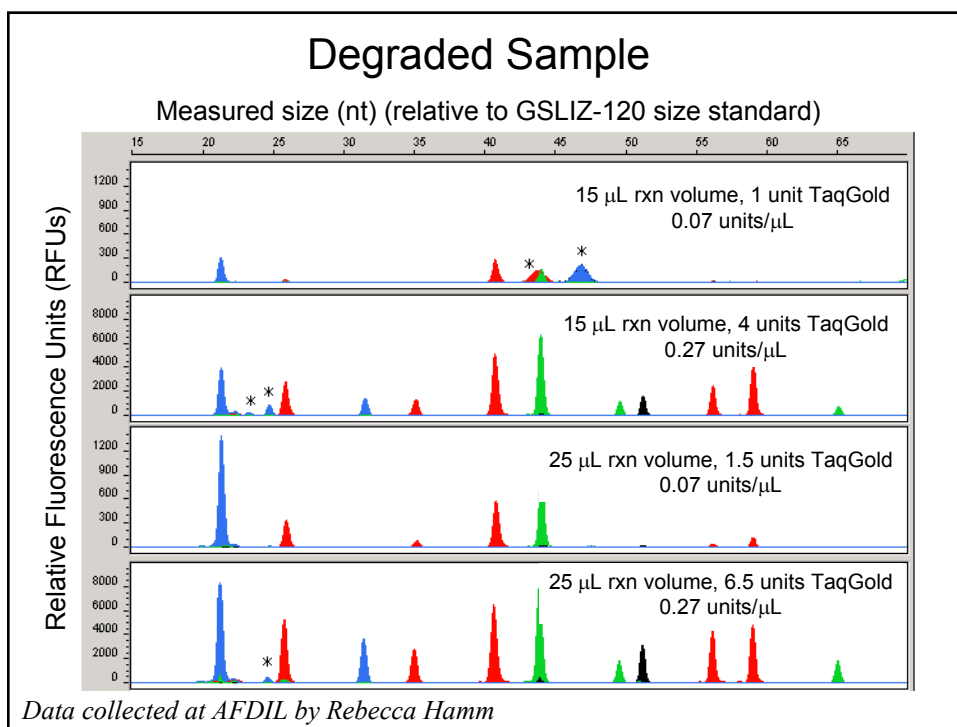
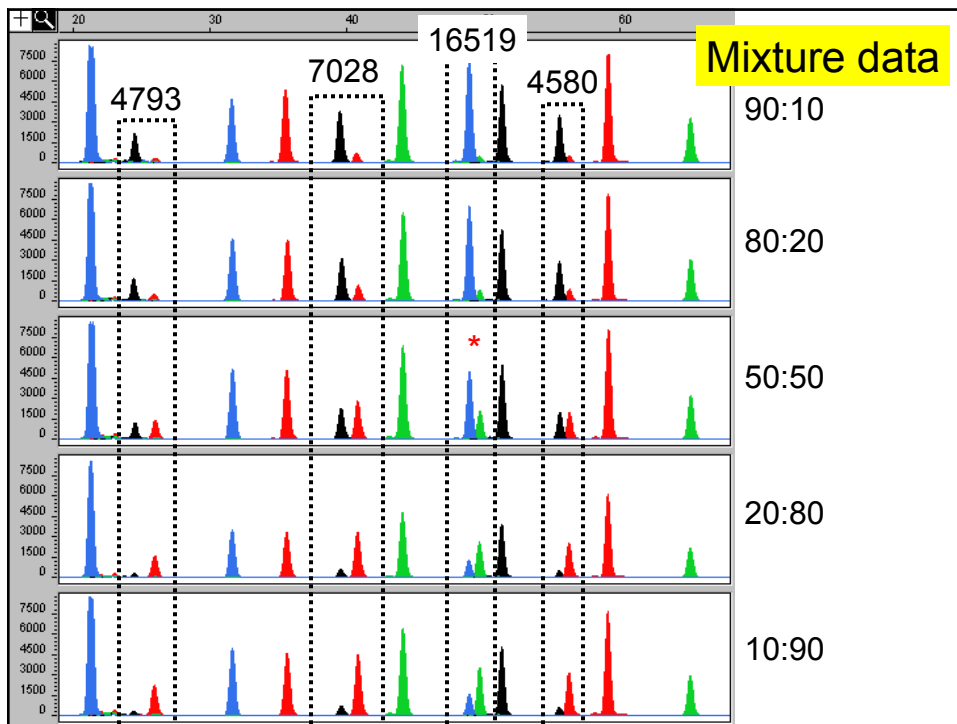
3010-F	TGTTGGATCAGGACATCCC	19 19
4793-R	(T) ₄ – TCAGAAGTGAAAGGGGGC	18 22
10211-R	(T) ₁₀ – ACTAAGAAGAATTATGGA	20 30
5004-F	(T) ₁₄ – AGACCCAGCTACGCAAAATC	20 34
7028-F	(T) ₁₈ – GACACGTACTACGTTGTAGC	20 38
7202-F	(T) ₂₂ – CCACAACACTTCTCGGCCT	20 42
16519-R	(T) ₂₄ – TGTGGGCTATTAGGCTTAGG	22 46
12858-F	(T) ₂₇ – GCAGCCATTCAAGCAATCCTATA	23 50
4580-R	(T) ₂₉ – TGGTTAGAACTGGAATAAGCTAG	25 54
477-F	(T) ₃₈ – CCCTCCCCTCCACTAC	20 58
14470-R	(T) ₄₁ – GGGATGATGGTTGTCTTG	21 62

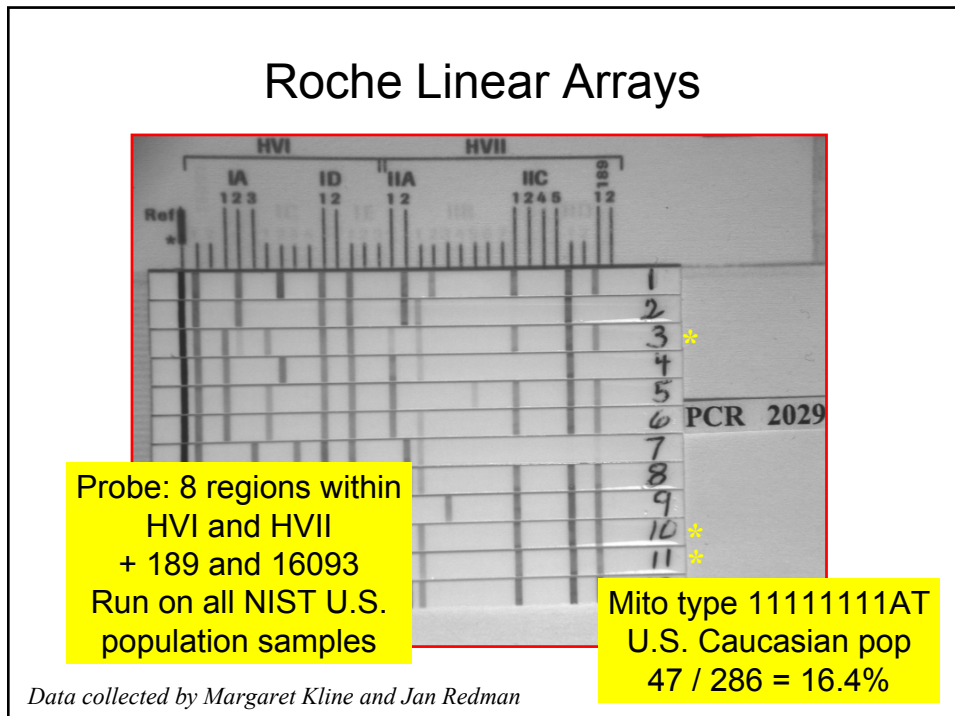


Program Output									
Label	Length	Sequence		Position	Tm				
Forward Primers Salt = 0.3Ct = 10									
M42 340 bp (A/T 297 W) AC010889	18	ATTTAGGACACAAAAGCW		280	60.65398				
M42 340 bp (A/T 297 W) AC010889	19	GATTAGGACACAAAAGCW		279	61.96716				
M42 340 bp (A/T 297 W) AC010889	20	AGATTTAGGACACAAAAGCW		278	63.67808				
Reverse Primers									
M42 340 bp (A/T 297 W) AC010889	23	GCTCTCTTTTCATTATGTAGTW		319	63.5462				
M42 340 bp (A/T 297 W) AC010889	21	TCTCTTTTCATTATGTAGTW		317	59.28964				
M42 340 bp (A/T 297 W) AC010889	20	CTCTTTTCATTATGTAGTW		316	57.50257				
Hairpin	Dimer	Template	Mass	Rank	Mutation	+ddC	+ddT	+ddA	+ddG
4	8	10	5273.48	2.133333	W	N/A	5561.67998	5570.68998	N/A
5	10	10	5602.69	2	W	N/A	5890.889941	5899.899941	N/A
5	10	11	5915.9	2	W	N/A	6204.099902	6213.109902	N/A
4	8	22	6734.42	2.133333	W	N/A	7022.619922	7031.629922	N/A
4	8	20	6116.02	2.133333	W	N/A	6404.22002	6413.23002	N/A
4	8	19	5811.82	2.133333	W	N/A	6100.019824	6109.029824	N/A









Typing 51 samples with mt 11 plex assay

51 (47 cauc/4 hisp) samples were identical by Roche linear array assay (most common Haplogroup observed in NIST U.S. Caucasian population samples)

	G	A	G	G	A	G	G	G	G	G	G	A	
3010	G	A	G	G	A	G	G	G	G	G	G	A	
4793	A	A	A	A	A	A	A	A	G	A	A	A	
10211	C	C	C	C	C	C	C	C	C	C	C	C	
5004	T	T	C	T	T	T	T	T	T	T	T	T	
7028	C	C	C	T	C	T	C	T	C	C	C	C	
7202	A	A	A	A	A	A	A	A	A	A	A	A	
16519	T	C	T	C	T	C	C	T	C	C	C	C	
12858	C	T	C	C	C	C	C	C	C	C	C	C	
4580	G	G	G	G	G	A	G	A	G	G	G	G	
477	T	C	T	T	T	C	T	T	T	T	T	T	
14470	T	T	T	A	T	T	T	T	T	T	T	T	
rCRS		1	1	1	1	2	2	3	4	4	5	12	15

12 haplogroups were observed
4 haplogroups were unique
2 of 11 sites did not vary

11-plex mtSNP assay

Assay is capable of accurately detecting 11 mtSNP in a single assay

The 11-plex assay is currently being validated for case work samples at AFDIL

Manuscript has been submitted

Additional multiplex mtSNP assays are being developed for other common HV1/HV2 types in collaboration with AFDIL

Overview

SNPs

Assay Platforms and Instrumentation

Multiplexing

U.S. Population Samples

Y Chromosome and Mitochondrial Markers

Results

mtSNP 11 plex

Y-SNP multiplexes



Y-SNPs in U.S. populations

What haplogroups will be observed?

How specific will certain Y-SNPs be for a U.S. population group?

Forensic utility in comparison/addition to Y-STRs

Commercial kit (Marligen) 42 Y-SNPs

Medium sized multiplexes developed in-house
(CE or MS)

Y-SNPs Typed at NIST

42 SNPs + Amelogenin present in 5 multiplexes
(commercially available kit from Marligen)

18 SNPs in 3 NIST-designed 6plexes (8 unique)
10 SNPs in 2 NIST-designed 5plexes (1 unique)

19 of the SNP sites overlapped...

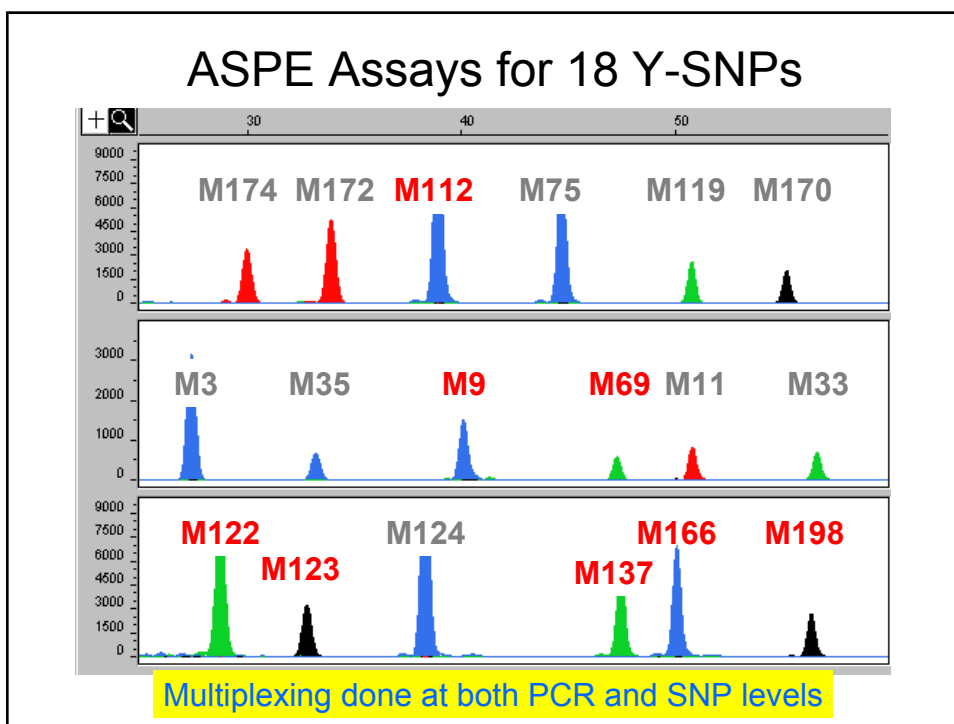
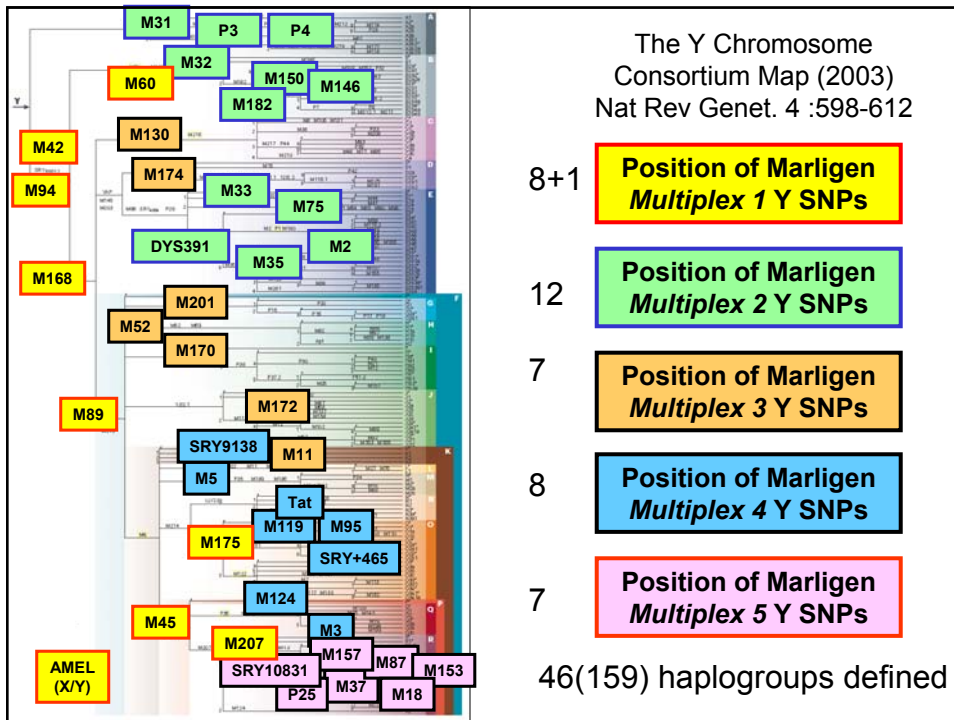


Resulting in a total of 51 Y-SNPs

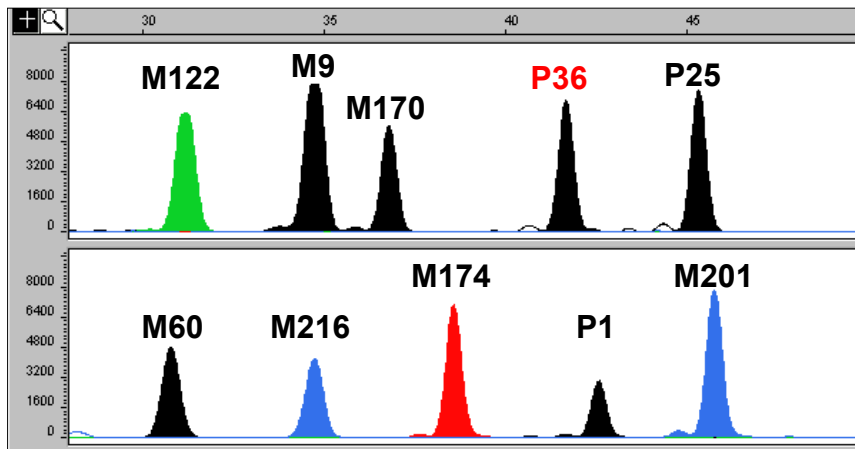
115 African Americans

114 Caucasians

95 Hispanics (presently typed for 10 Y-SNPs)



ASPE Assays for 10 Y-SNPs



Equimolar PCR primer concentration (5plex)
Empirical balancing of extension primers

Summary of Y-SNP Data (115 African Americans and 114 Caucasians)

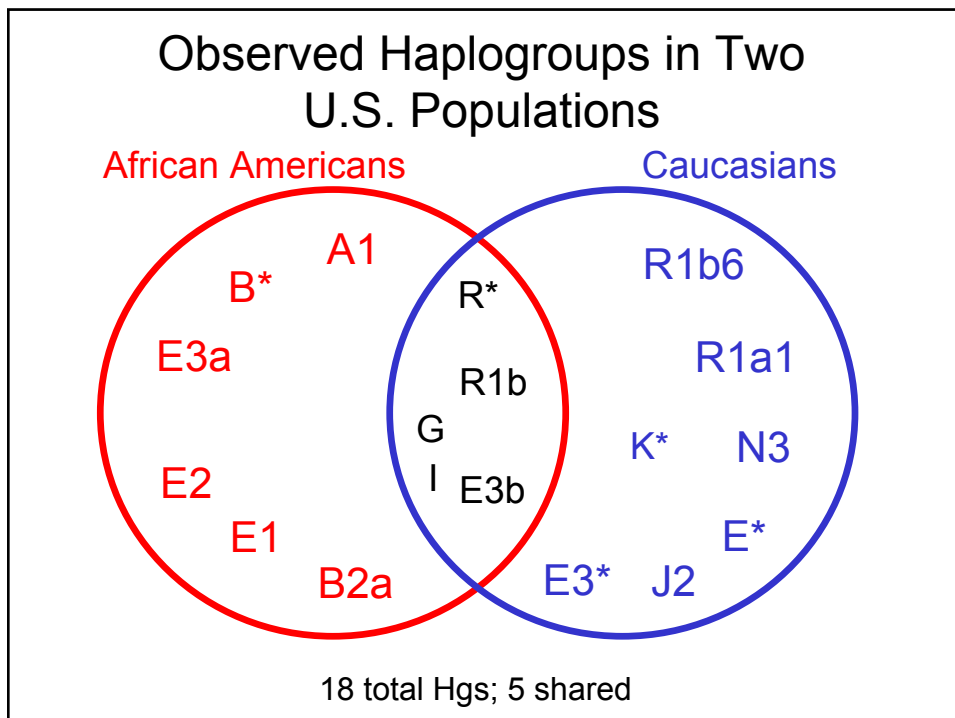
A total of 20 ng of genomic DNA was consumed for the 10 multiplexes

18 out of 46 haplogroups observed

Over 99 % success rate for allele calls (both methods)

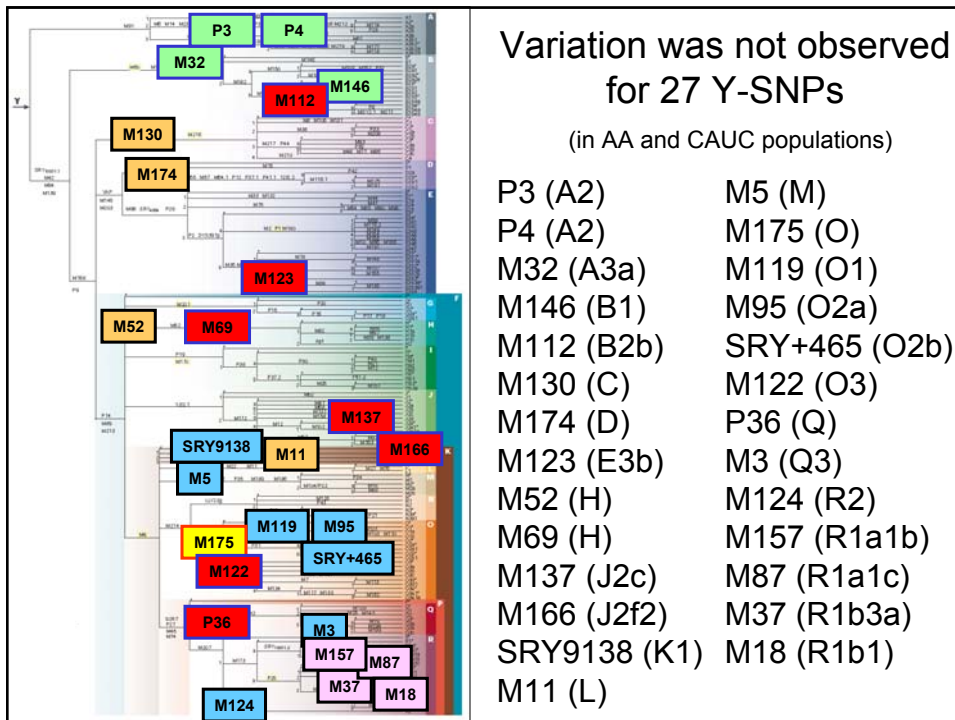
Variation was observed in **24** of the **51** Y-SNPs

100% concordance for the 18 overlapping markers (>3,800 allele calls)



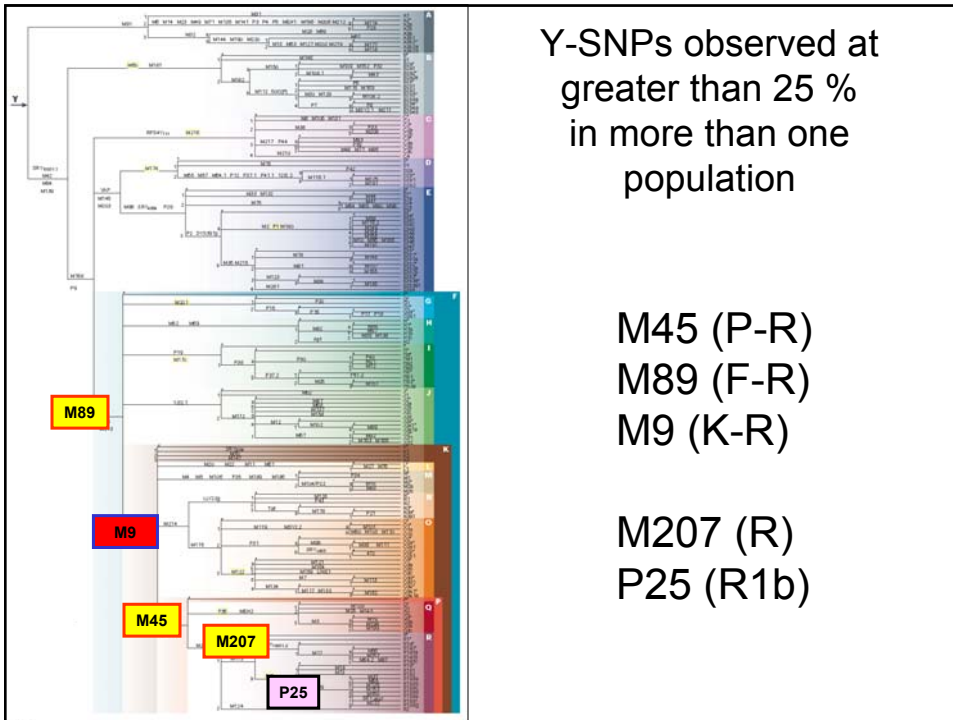
Variation was not observed for 27 Y-SNPs
(in AA and CAUC populations)

<u>M175 +/-</u>	<u>M119 A/C</u>	<u>M37 C/T</u>
<u>M146 A/C</u>	<u>M124 C/T</u>	<u>M87 T/C</u>
<u>M32 T/C</u>	<u>M3 C/T</u>	<u>M69 T/C</u>
<u>P3 (C/T)</u>	<u>M5 C/T</u>	<u>M112 G/A</u>
<u>P4 (G/A)</u>	<u>M95 C/T</u>	<u>M122 T/C</u>
<u>M11 A/G</u>	<u>SRY465 C/T</u>	<u>M123 G/A</u>
<u>M130 C/T</u>	<u>SRY9138 C/T</u>	<u>M137 T/C</u>
<u>M174 T/C</u>	<u>M157 A/C</u>	<u>M166 G/A</u>
<u>M52 A/C</u>	<u>M18 -/+</u>	<u>P36</u>



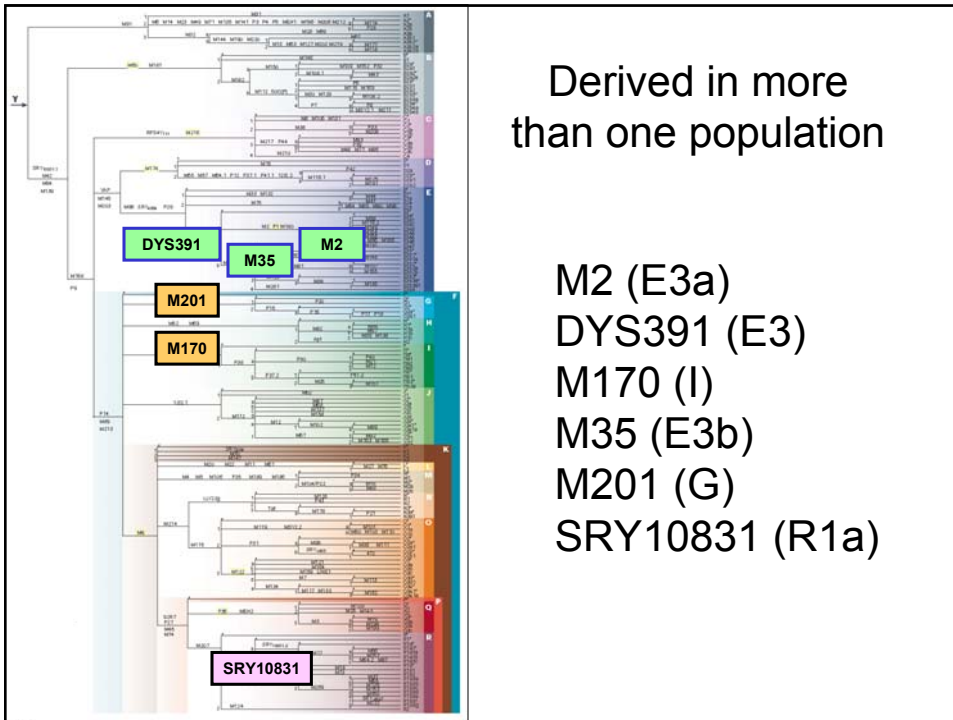
Y-SNPs derived at greater than 25 % in more than one population

Locus	All	AA	Cauc	Hisp	Hap
<u>M207 A/G</u>	0.46	0.27	0.65	na	R
<u>M45 G/A</u>	0.46	0.27	0.64	na	P-R
<u>M89 C/T</u>	0.64	0.32	0.96	na	F-R
<u>P25 C/A</u>	0.47	0.30	0.57	0.53	R1b
<u>M9 C/G</u>	0.53	0.31	0.65	0.64	K-R



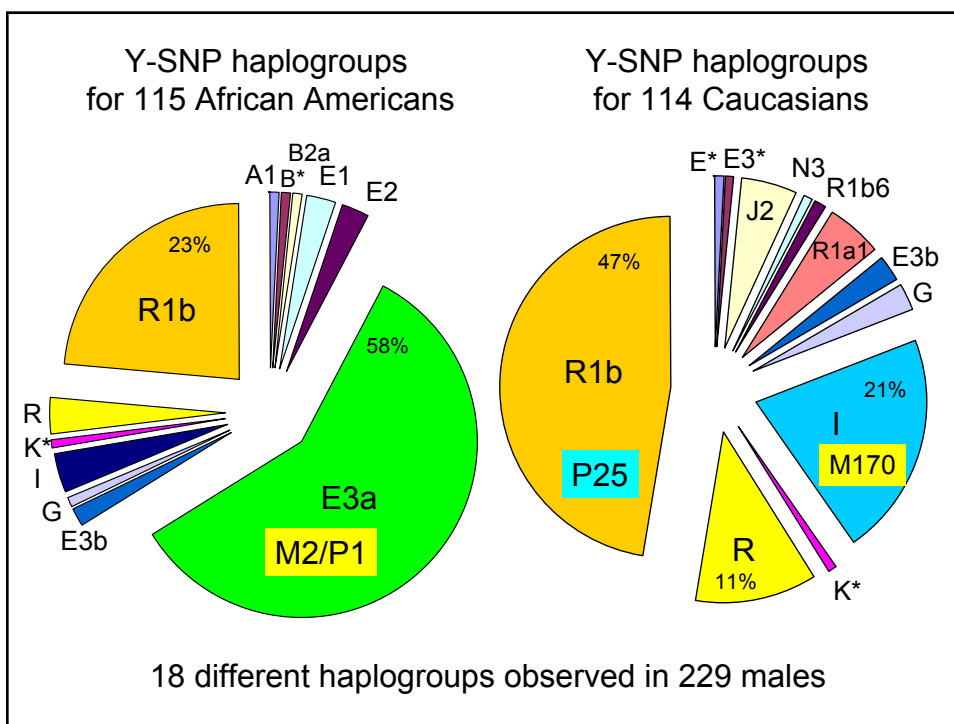
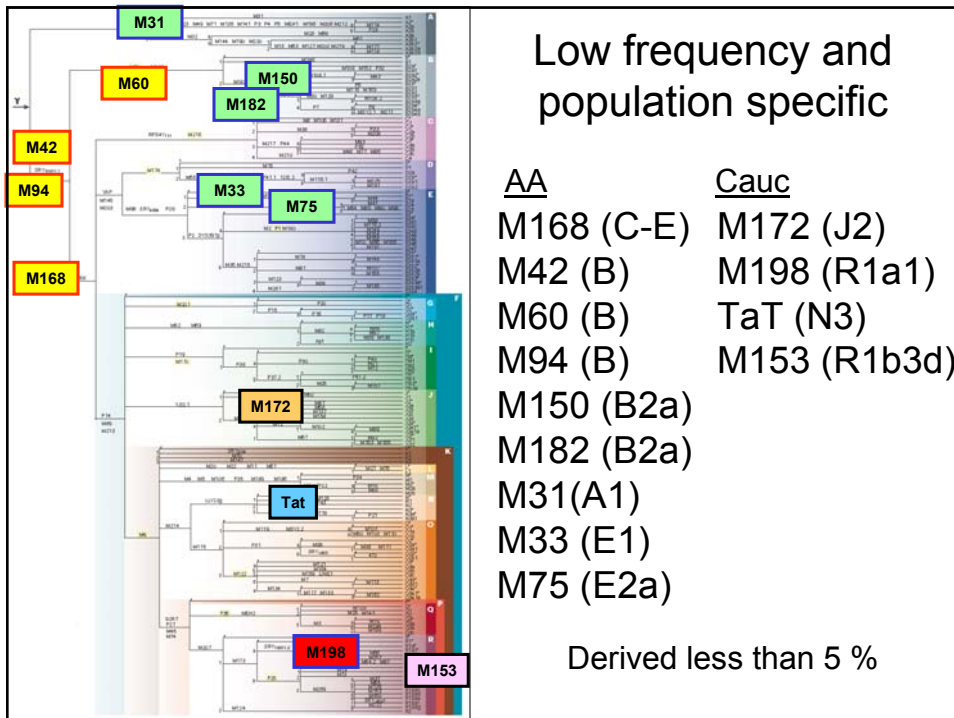
Derived in more than one population

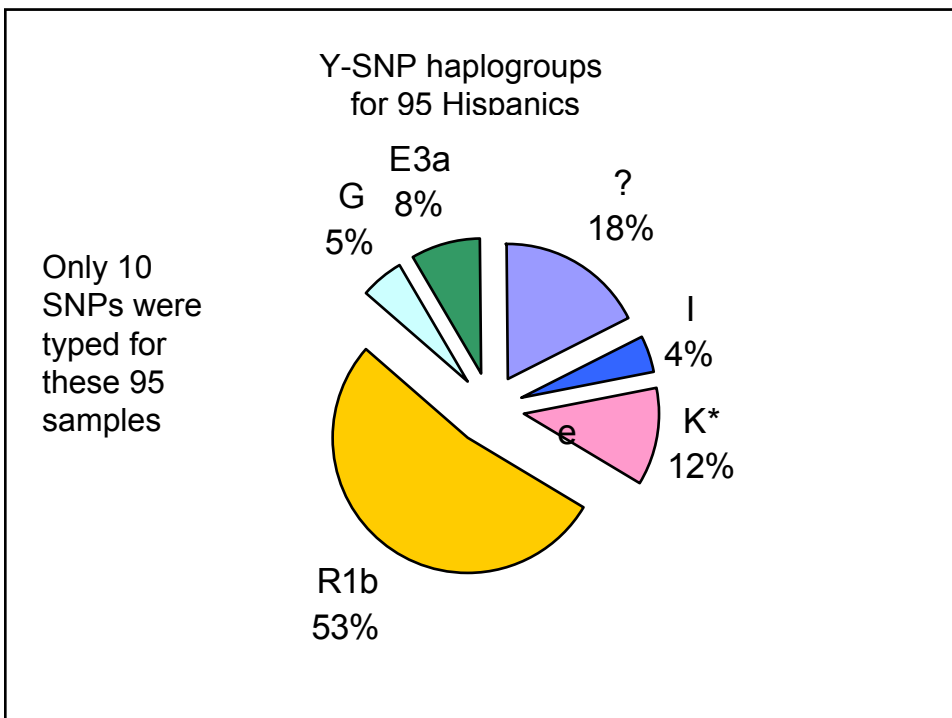
Locus	All	AA	Cauc	Hisp	Hap
<u>M2 A/G</u>	0.23	0.58	not obs	0.08	E3a
<u>DYS391 C/G</u>	0.31	0.60	0.04	na	E3
<u>M170 A/C</u>	0.10	0.04	0.21	0.04	I
<u>M35 G/C</u>	0.02	0.02	0.03	na	E3b
<u>M201 G/T</u>	0.03	0.01	0.03	0.05	G
<u>SRY10831 A/G</u>	0.03	0.01	0.05	na	R1a



Low frequency and population specific

Locus	All	AA	Cauc	Hisp
<u>M168 C/T</u>	0.01	0.03	not obs	na
<u>M42 A/T</u>	0.04	0.01	not obs	na
<u>M60 -/+</u>	0.01	0.02	not obs	not obs
<u>M94 C/A</u>	0.01	0.01	not obs	na
<u>M150 C/T</u>	0.01	0.01	not obs	na
<u>M182 C/T</u>	0.01	0.01	not obs	na
<u>M31 G/C</u>	0.01	0.01	not obs	na
<u>M33 A/C</u>	0.01	0.03	not obs	na
<u>M75 G/A</u>	0.01	0.03	not obs	na
<u>M172 T/G</u>	0.03	not obs	0.05	na
<u>M198 C/T</u>	0.03	not obs	0.05	na
Tat T/C	0.01	not obs	0.01	na
<u>M153 T/A</u>	0.01	not obs	0.01	na





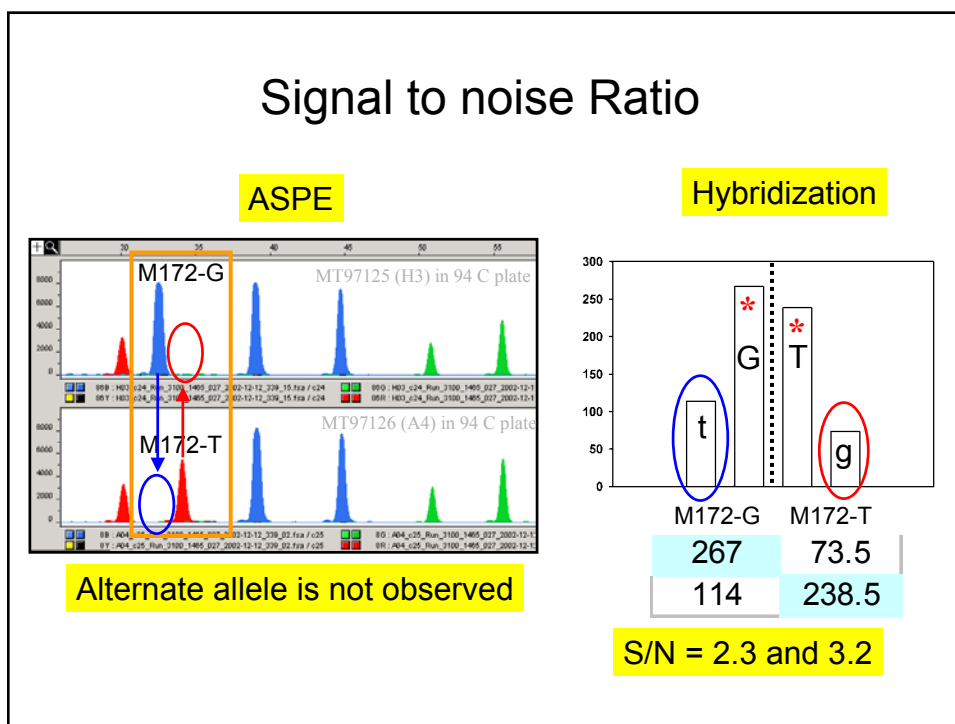
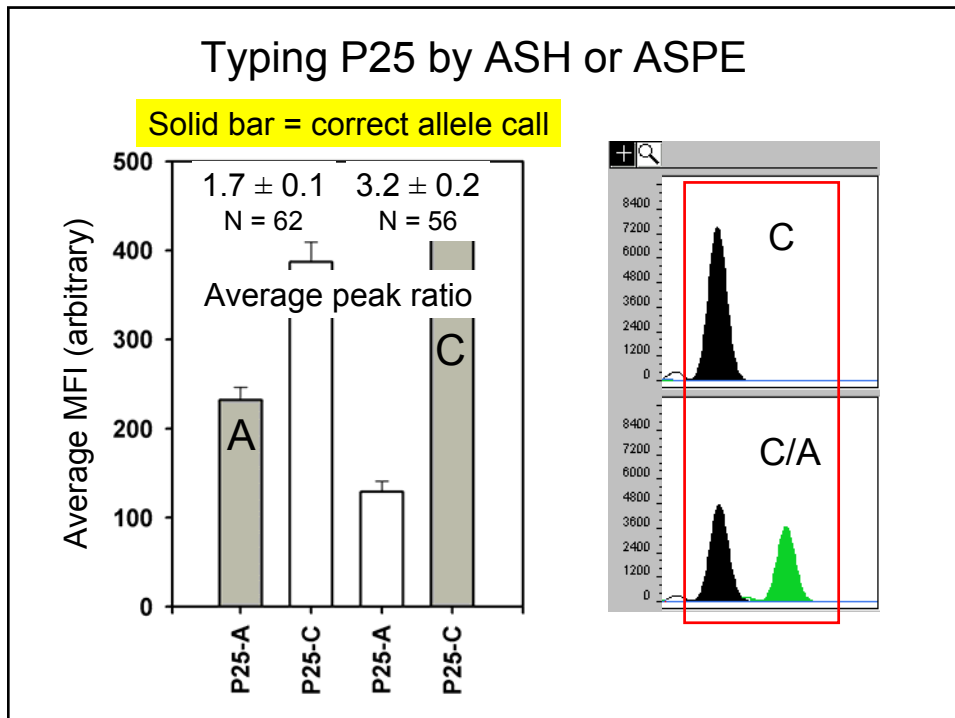
Issues with Y-SNP P25

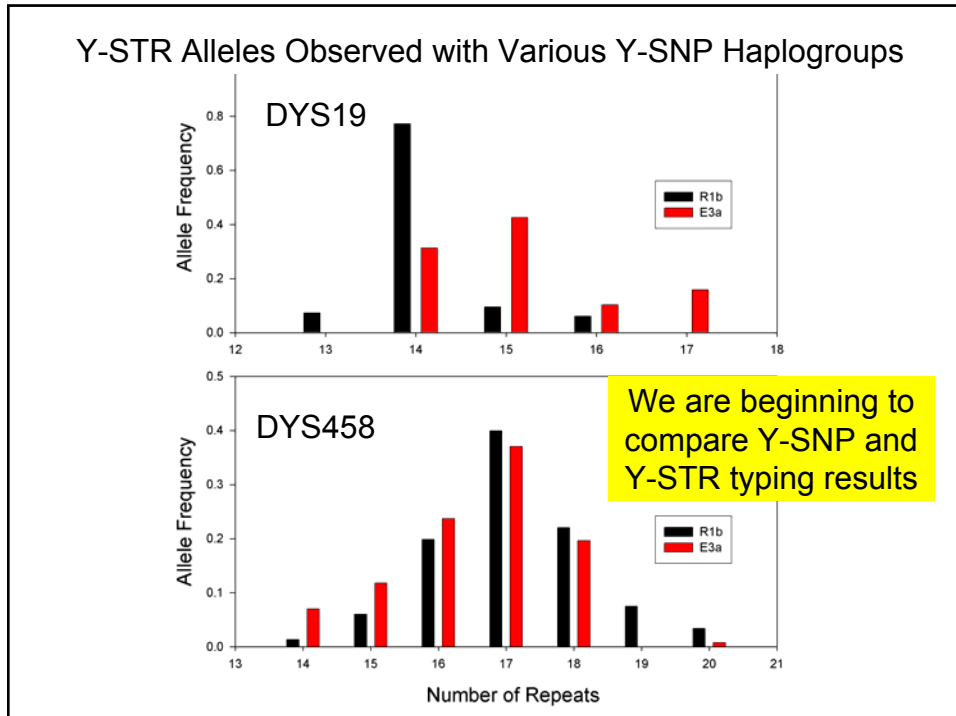
Initially when typing P25 with the Marilgen kit the derived allele (A) was not observed

Alan Redd (Univ of AZ) informed us that P25 is a multi copy locus

After further review of our data we were able to make correct allele call for the P25 marker based on **signal intensity ratio**

BLAST results indicate that the region surrounding P25 is present 3 times on the Y chromosome





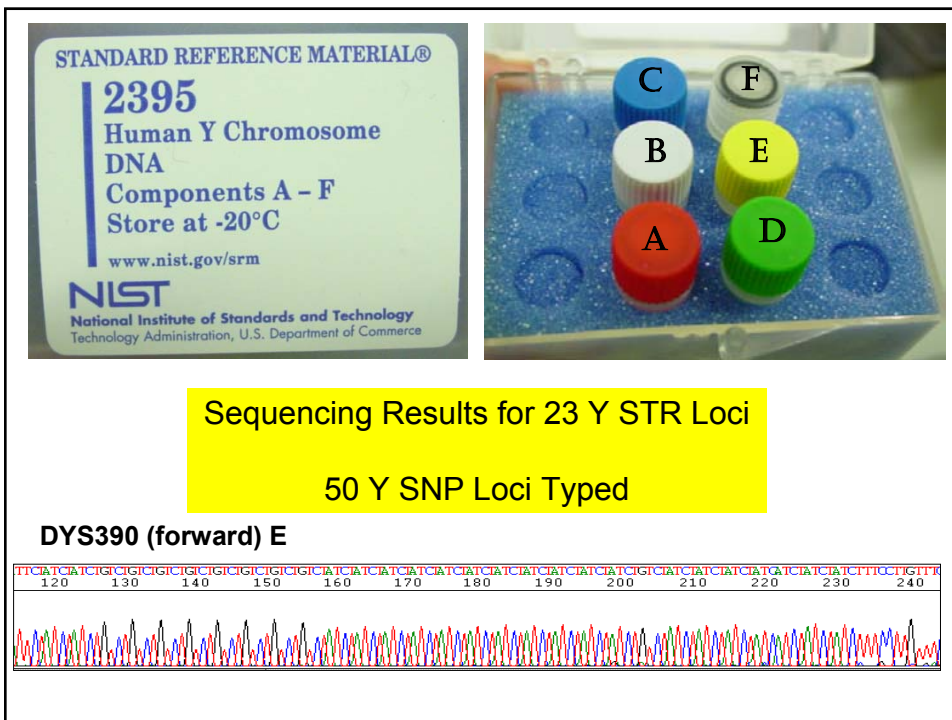
Forensic Utility
51 Y-SNPs versus 1 Y-STR

For N = 211 male samples

	<u>51Y-SNPs</u>	<u>Y-STR DYS464</u>
Amount of sample consumed	10ng	1ng
Number for types observed	18	62
Analysis	Multiple	1 reaction
Degraded samples	+	?

Conclusions

- Full concordance was observed between hybridization and primer extension technologies on 18 different Y-SNPs (>3,800 allele calls)
- Caucasian admixture was observed with our African American population (Hg R and R1b in ~30%)—agrees with Kayser *et al.* (2003) *Genome Res.* 13:624-634 done with 9 Y-STRs
- Y-SNPs may have limited value for ethnic differentiation in U.S. populations
 - One exception: M2 not in Caucasians
- Y-SNPs are not a useful stand-alone assay for forensic purposes, but may be helpful in combination with Y-STRs



Y SNP Results on SRM 2395

SRM 2395	AMEL	M207	M45	M89	DYS391	M2	M170	M172	M201
	(A/G)	(A/G)	(C/T)	(C/G)	(A/G)	(A/C)	(G/T)	(G/T)	
Component A	XY	G	A	T	C	A	A	T	G
Component B	XY	A	G	T	C	A	A	G	G
Component C	XY	A	G	C	G	G	A	T	G
Component D	XY	A	G	T	C	A	A	T	T
Component E	XY	A	G	T	C	A	C	T	G
Component F	XX								

SRM components are all distinguishable from one another with these Y SNPs

50 Y SNPs measured across all samples

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John Butler

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 David Carlson (Marligen)
 Mike Hammer and Alan Redd (U of AZ)



Jan Redman



Margaret Kline