





CIB Forensic Science Center
Training Seminar (Taipei, Taiwan)
June 6-7, 2012



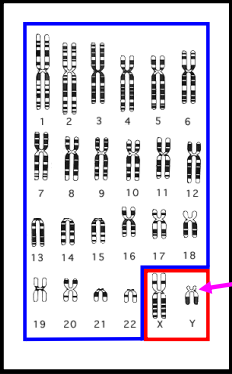
Lineage Markers: Y-STRs, mtDNA, and X-STRs

John M. Butler
NIST Applied Genetics Group
National Institute of Standards and Technology
Gaithersburg, Maryland



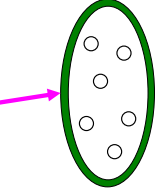
Cell Nucleus – 3.2 billion bp

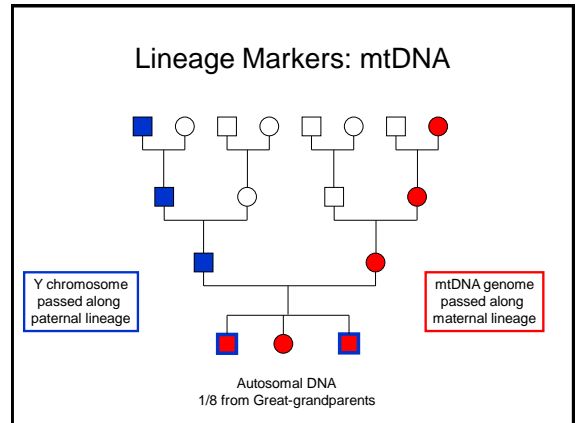
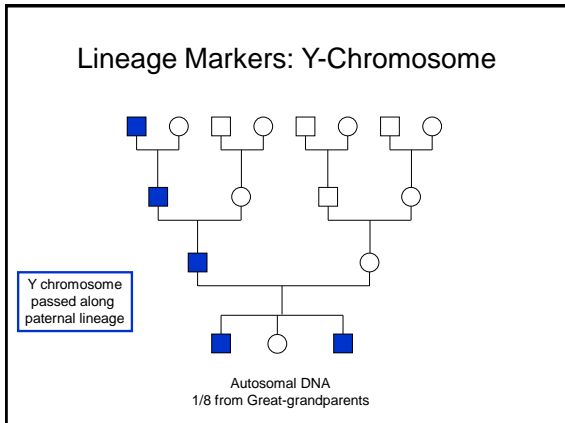


Autosomes – 22 pairs – 2 copies per cell

Sex Chromosomes (XX or XY)

mitochondria – in cell cytoplasm
100s of mtDNA copies per cell





Different Inheritance Patterns

TABLE 15.1 Specific Relationships and the Probability of Transmitting Genetic Information (Barring Mutation). Some of the ChrY Information is Not Applicable (N/A) as Women Do Not Have a Y-Chromosome.

Inheritance	Autosomal Markers	ChrY Markers	mtDNA	ChrX Markers
Mother → Son	50%	N/A	100%	100%
Mother → Daughter	50%	N/A	100%	50%
Father → Son	50%	100%	0%	0%
Father → Daughter	50%	0%	0%	100%
Paternal Grandmother → Granddaughter	25%	N/A	0%	100%
Maternal Grandmother → Granddaughter	25%	N/A	100%	25%
Paternal Grandfather → Grandson	25%	100%	0%	0%

Butler, J.M. (2012) *Advanced Topics in Forensic DNA Typing: Methodology*, p. 457

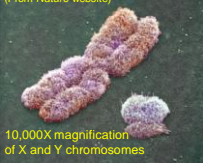
THE HUMAN Y CHROMOSOME: AN EVOLUTIONARY MARKER COMES OF AGE

Mark A. Jobling & Chris Tyler-Smith
Nature Reviews Genetics (2003) 4, 598-612

Abstract

- Until recently, the Y chromosome seemed to fulfill the role of juvenile delinquent among human chromosomes — rich in junk, poor in useful attributes, reluctant to socialize with its neighbors and with an inescapable tendency to degenerate. The availability of the near-complete chromosome sequence, plus many new polymorphisms, a highly resolved phylogeny and insights into its mutation processes, now provide new avenues for investigating human evolution. Y-chromosome research is growing up.

(From Nature website)



10,000X magnification of X and Y chromosomes

What has happened in the past decade...

- **Selection of core Y-STR loci** (SWGAM Jan 2003)
- "Full" Y-chromosome sequence became available in June 2003; over 400 Y-STR loci identified (only ~20 in 2000)
- **Commercial Y-STR kits released**
 - Y-**PLEX-6,5,12** (2001-03), **PowerPlex Y** (9/03), **Yfiler** (12/04), **PPY23** (6/12)
- Many population studies performed and online databases generated with thousands of Y-STR haplotypes
- Forensic casework demonstrations showing value of Y-STR testing along with court acceptance
- Some renewed interest in Y-STRs to aid familial searching

Y-STR Information

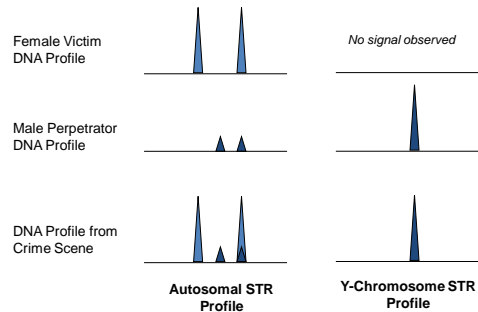
- Why the Y?
- Y-STR Loci & Kits
- Y-STR Databases
- Y-STR Stats & Interpretation Issues
- Genetic Genealogy & Familial Searching

Value of Y-Chromosome Markers

J.M. Butler (2005) *Forensic DNA Typing, 2nd Edition*; Table 9.1

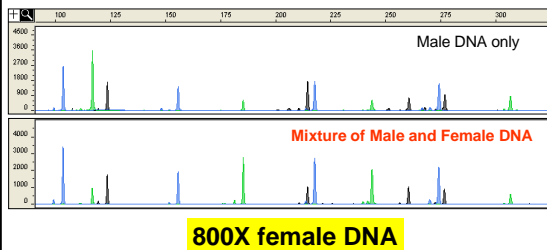
Application	Advantage
Forensic casework on sexual assault evidence	Male-specific amplification (can avoid differential extraction to separate sperm and epithelial cells)
Paternity testing	Male children can be tied to fathers in motherless paternity cases
Missing persons investigations	Patrilineal male relatives may be used for reference samples
Human migration and evolutionary studies	Lack of recombination enables comparison of male individuals separated by large periods of time
Historical and genealogical research	Surnames usually retained by males; can make links where paper trail is limited

Y-STRs can permit simplification of male DNA identification in sexual assault cases



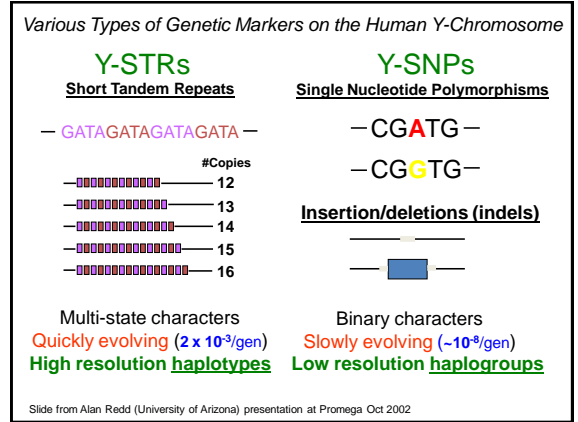
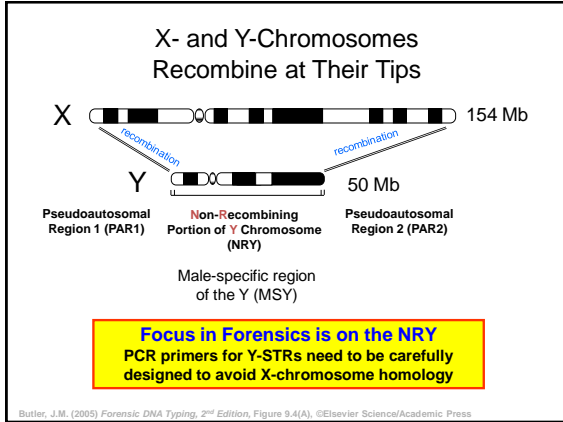
Butler, J.M. (2005) *Forensic DNA Typing, 2nd Edition*, Figure 9.2, ©Elsevier Science/Academic Press

Y-STRs Identify the Male Component even with Excess Female DNA

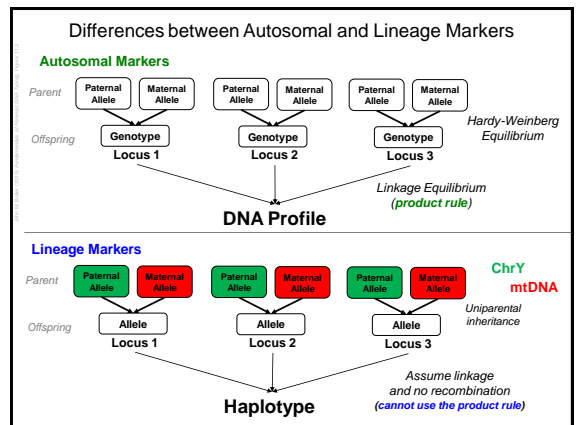
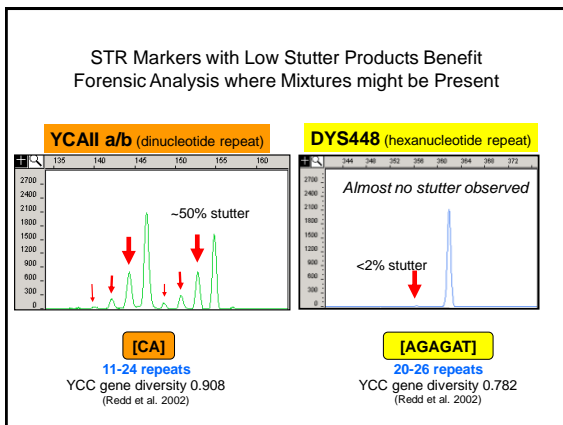
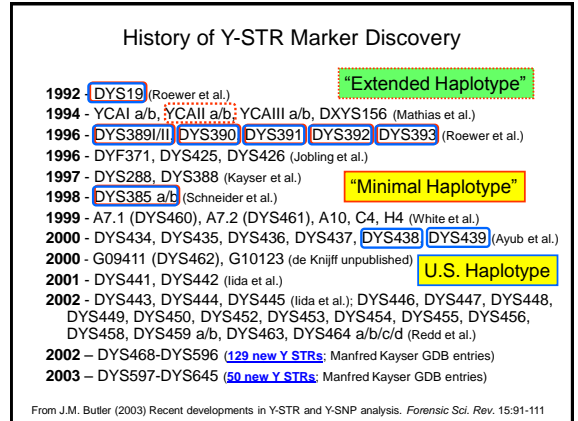


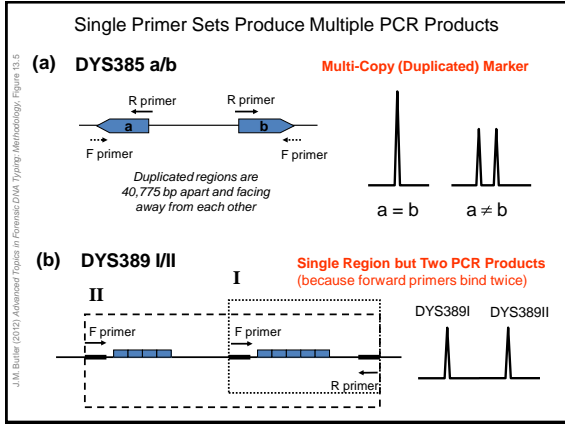
Disadvantages of the Y-Chromosome

- Loci are not independent of one another and therefore rare random match probabilities cannot be generated with the product rule; must use haplotypes (combination of alleles observed at all tested loci)
- **Paternal lineages possess the same Y-STR haplotype** (barring mutation) and thus fathers, sons, brothers, uncles, and paternal cousins cannot be distinguished from one another
- **Not as informative as autosomal STR results**
 - More like addition ($10 + 10 + 10 = 30$) than multiplication ($10 \times 10 \times 10 = 1,000$)



Y-STR Loci & Kits

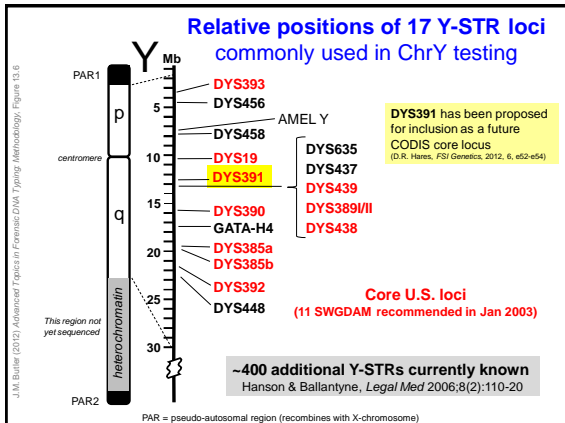




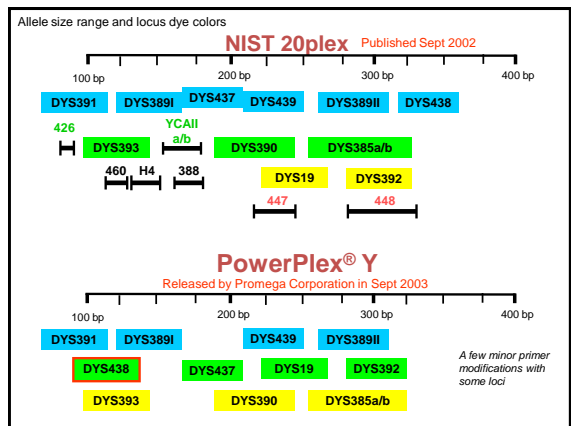
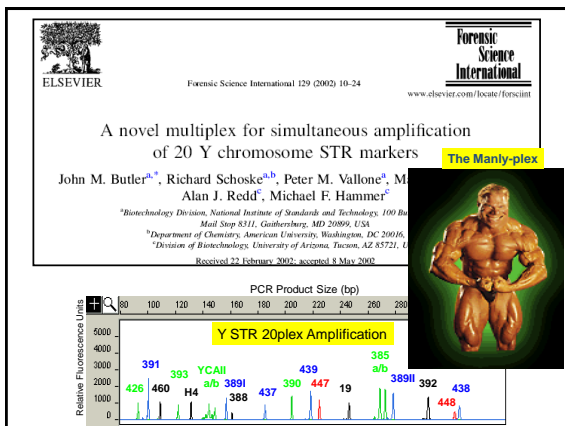
17 PCR products 15 primer sets Characteristics of the 17 Commonly Used Y-STR Loci

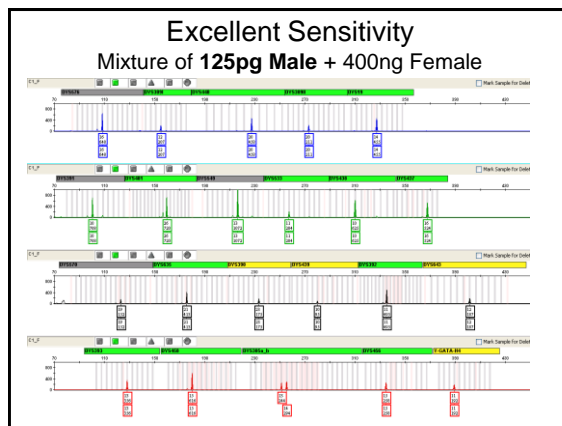
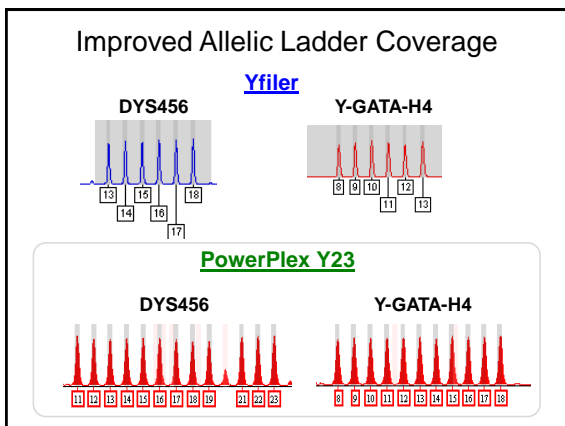
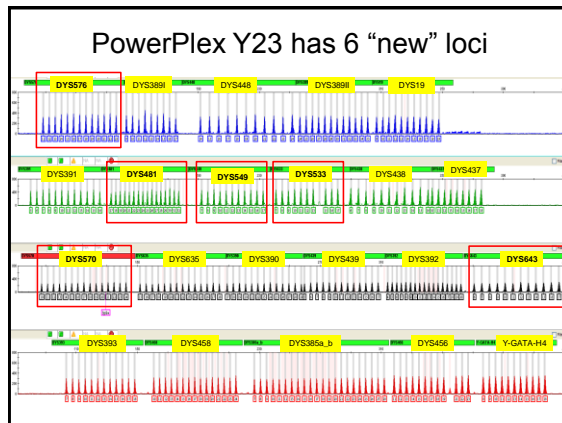
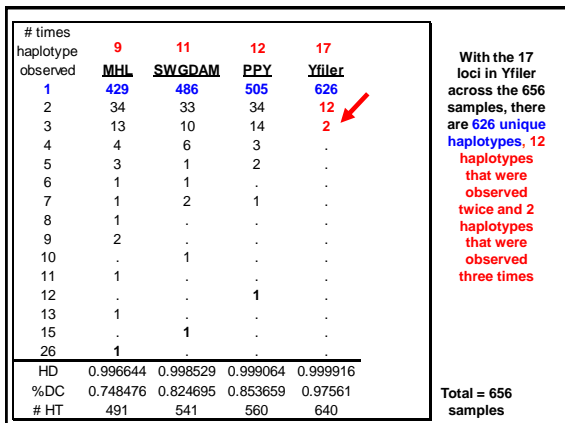
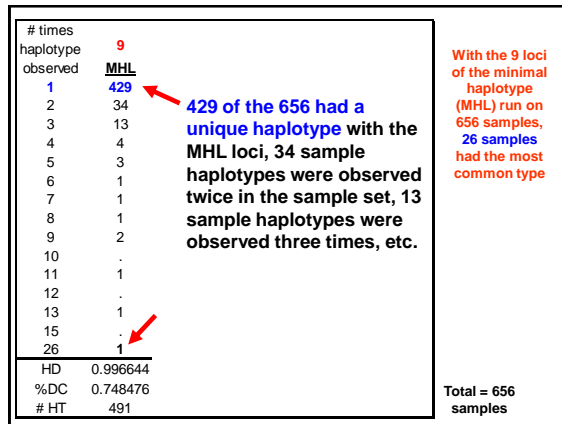
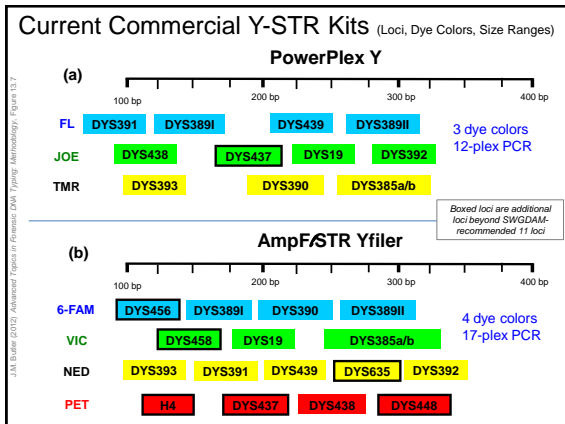
STR Marker	Position (Mb)	Repeat Motif	Allele Range	Mutation Rate*
DYS393	3.19	AGAT	8-17	0.10 %
DYS456	4.33	AGAT	13-18	0.42 %
DYS458	7.93	GAAA	14-20	0.64 %
DYS19	10.13	TAGA	10-19	0.23 %
DYS391	12.61	TCTA	6-14	0.26 %
DYS635	12.89	TSTA	17-27	0.35 %
DYS437	12.98	TCTR	13-17	0.12 %
DYS439	13.03	AGAT	8-15	0.52 %
DYS389 I/II	13.12	TCTR	9-17 / 24-34	0.25 % / 0.36 %
DYS438	13.38	TTTTC	6-14	0.03 %
DYS390	15.78	TCTR	17-28	0.21 %
GATA-H4	17.25	TAGA	8-13	0.24 %
DYS385 a/b	19.26	GAAA	7-28	0.21 %
DYS392	21.04	TAT	6-20	0.04 %
DYS448	22.78	AGAGAT	17-24	0.16 %

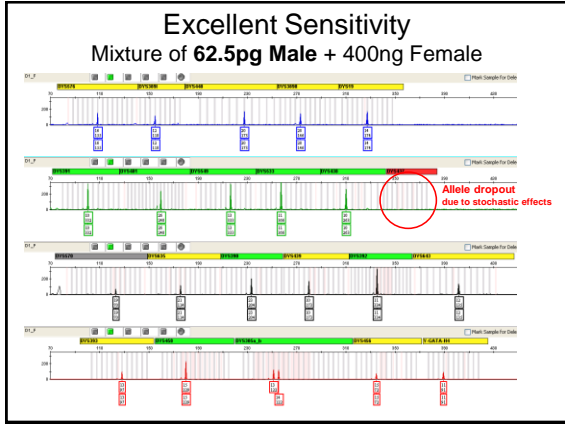
*Mutation rates are from as many as 15,000 meioses described in a YHRD summary of 23 publications in Jan 2011 (see <http://www.yhrd.org/2009/01/01/>)



- Recent Developments with Y-STR Typing**
- Promega Corporation** plans to release **PowerPlex Y23** (23 loci) in June 2012 which will enable further resolution of Y-STR haplotypes
 - Population databases will need to be developed with the new extended haplotypes
 - Manfred Kayser's group has developed a set of **rapidly mutating (RM) Y-STR loci** that have the capability to resolve fathers and sons in many instances
 - An international collaboration is currently on-going to study these RM Y-STRs in more detail (14 RM Y-STRs in 3 multiplexes)

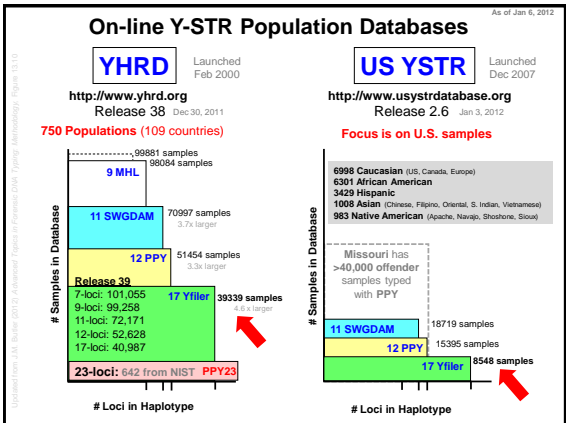






Y-STR Databases

YHRD has >100,000 haplotypes



Population Data Publications Describing Handling of Y-STR and mtDNA Haplotype Information

- The leading forensic journals **require Y-STR and mtDNA population data to be reviewed by and submitted to YHRD and EMPOP**

US YSTR Contributions

Contributor to US YSTR	# Samples	% of Database
Applied Biosystems (includes UNTHSC, NIST samples, ...)	6,159	33%
Promega	3,800	20%
ReliaGene	3,037	16%
University of Arizona	2,462	13%
NCFS (University of Central Florida)	2,440	13%
Illinois State Police	398	2.1%
Santa Clara Co. CA Crime Lab	143	0.6%
Marshall University	113	0.6%
Washington State Patrol Crime Lab	40	0.2%
San Diego Sheriff's Regional Crime Lab	39	0.2%
CA DOJ	32	0.2%
Orange County CA Coroner	30	0.2%
Richland County Sheriff's Dept.	7	0.04%
Release 2.6 (Jan 3, 2012)	18,719	8548 17-locus profiles

<http://www6.appliedbiosystems.com/yfilerdatabase/>

Population	# Haplotypes
African American	1932
Asian	330
Asian Indian	564
Caucasian	4114
Chinese	577
Filipino	105
Hispanic	1601
Japanese	1078
Malay	579
Native American	105
Sub-saharan African	59
Thai	246
Vietnamese	103
All	11393

Applied Biosystems
still maintains its
Yfiler database



Y-STR Stats & Interpretation Issues

Forensic Science International: Genetics 4 (2010) 281–291

Contents lists available at ScienceDirect

Forensic Science International: Genetics

Journal homepage: www.elsevier.com/locate/fsig

Fundamental problem of forensic mathematics—The evidential value of a rare haplotype

Charles H. Brenner^{a,b,*}

^a School of Public Health, Forensic Science Group, U.C. Berkeley, Berkeley, CA United States
^b DNA IVEV, 6807 Thornhill Drive, Oakland, CA 94611-1336, United States

“The fundamental question to decide the evidentiary significance of a trait linking suspect to crime is not one of frequency but of probability: What is the probability for such a match to happen by coincidence when the suspect is innocent?”

Forensic Science International: Genetics 5 (2011) 78–83

Contents lists available at ScienceDirect

Forensic Science International: Genetics

Journal homepage: www.elsevier.com/locate/fsig

The interpretation of lineage markers in forensic DNA testing

J.S. Buckleton^a, M. Krawczak^b, B.S. Weir^{c,*}

^a FBI Lab, Private Bag 9202, Auckland, New Zealand
^b Institute of Medical Informatics and Statistics, Christian-Albrechts University, 24105 Kiel, Germany
^c Department of Biostatistics, University of Washington, Box 357232, Seattle, WA 98195-7232, USA

This article reviews and discusses a number of highly relevant topics:

- Normal vs. binomial (Clopper-Pearson) sampling distributions
- Theta corrections
- Handling rare haplotypes (Charles Brenner approach)
- Combination of lineage and autosomal markers

Current (2009) SWGDAM
Y-STR Interpretation Guidelines

- Approved July 15, 2008 by SWGDAM
- Published in *Forensic Sci. Comm.* Jan 2009 issue

Will likely be updated soon to reflect change to Clopper-Pearson...

http://www.fbi.gov/about-us/lab/forensic-science-communications/fsc/jan2009/standards2009_01_standards01.htm/

J.S. Elder (2011) Advanced topics in Forensic DNA Typing: Mitochondria, D.N.A. Box 13.1

Results of Y-STR Profile Search

The following profile was searched on 15 January 2011 against several databases:

DYS19 (14), DYS389I (13), DYS389II (29), DYS390 (24), DYS391 (11), DYS392 (13), DYS393 (13), DYS385 a/b (11,15), DYS438 (12), DYS439 (13), DYS437 (15), DYS448 (19), DYS456 (17), DYS458 (18), DYS635 (23), and GATA-H4 (12).

Database	Minimal haplotype (9 loci)	SWGDM (11 loci)	PowerPlex Y (12 loci)	Yfiler (17 loci)	3/N for zero observations
YHRD	403/89804 = 0.45 %	29/62548 = 0.046 %	14/42277 = 0.033 %	0/30300 = <0.0033 %	3/30300 = 0.0099 %
US Y-STR	6/18547 = 0.032 %	1/18547 = 0.0054 %	1/15223 = 0.0066 %	0/8376 = <0.012 %	3/8376 = 0.036 %
Yfiler database	64/11393 = 0.56 %	4/11393 = 0.035 %	4/11393 = 0.035 %	0/11393 = <0.0088 %	3/11393 = 0.026 %

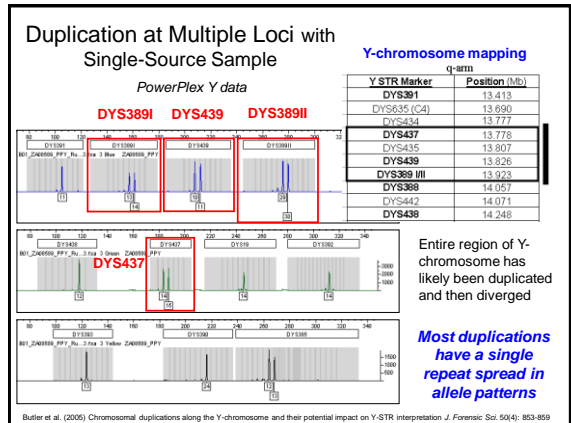
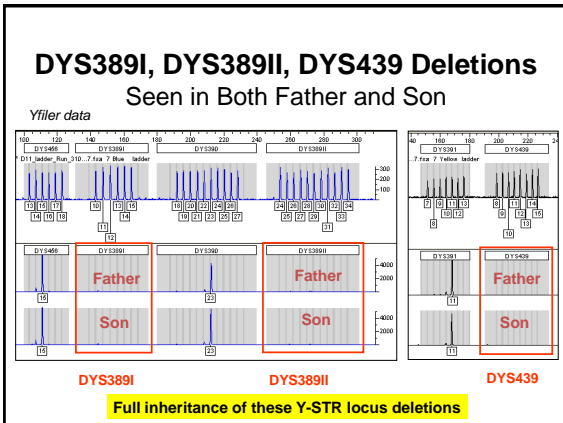
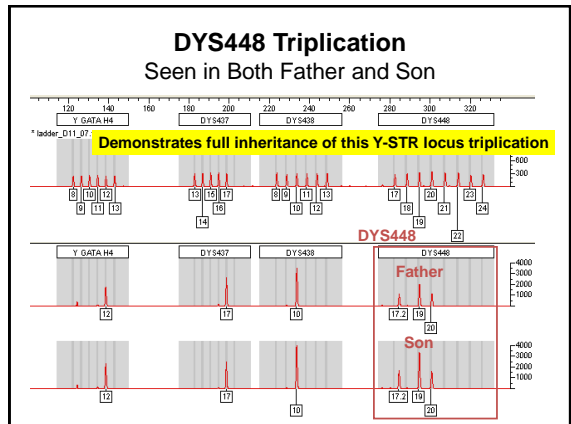
Normal vs. Clopper-Pearson

In March 2010 the US Y-STR database changed its 95 % confidence interval calculations to the Clopper-Pearson method.

Count values	Frequency $p = x/N$	Normal 95 % confidence interval	Clopper-Pearson 95 % confidence interval*
YHRD 9 loci: 403/89804	0.449 %	0.485 %	0.487 %
YHRD 12 loci: 14/42277	0.0331 %	0.0477 %	0.0518 %
US Y-STR 12 loci: 1/15223	0.0657 %	0.0174 %	0.0317 %

* Calculation performed with HaploCALC 1.0 Excel spreadsheet kindly provided by Steven P. Myers, CA DOJ

Note that with a large number of observations, such as 403 out of a database of 89804, there is almost no difference between the normal and Clopper-Pearson approaches. However, the normal method is less conservative (i.e., provides a more rare frequency) when the haplotype frequency is low, such as 1 out of 15223 or even 14 out of 42277. Although there are differences in these calculations, re-evaluation by the Clopper-Pearson method will not suddenly change a reported result by orders of magnitude or likely change the outcome of a report significantly.



Duplication and Divergence Model

Locus	# dup*	>1 repeat
DYS19	23	2
DYS389I	5	0
DYS389II	9	2
DYS390	1	0
DYS391	3	1
DYS392	0	0
DYS393	3	0
DYS385a/b	17	0

*from www.yhrd.org, literature, and our work

92% have single repeat difference

Since single-step mutations are most common, then single repeat spacing in duplicated alleles is expected

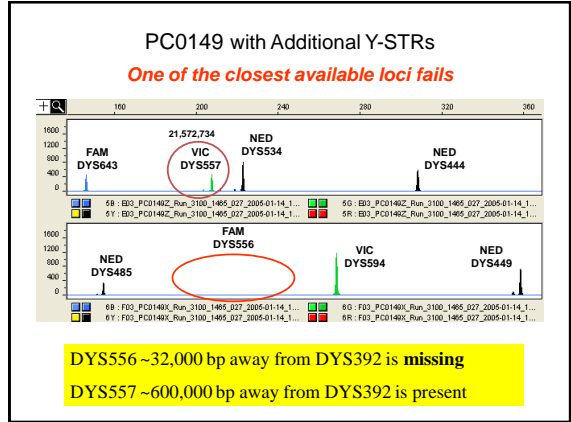
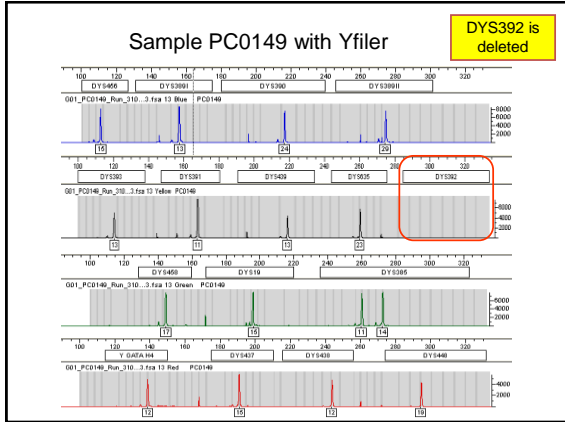
Butler et al. (2006) Chromosomal duplications along the Y-chromosome and their potential impact on Y-STR interpretation J. Forensic Sci. 50(4): 853-859

Deciphering between a Mixture of Multiple Males and Locus Duplication

- Note the number of loci containing >1 allele (other than multi-copy DYS385)
- Consider relative position on the Y-chromosome if multiple loci have two alleles
- See if repeat spread is >1 repeat unit
- Examine DYS385 for presence of >2 alleles

Locus duplication along the Y-chromosome is in many ways analogous to heteroplasmy in mitochondrial DNA, which depending on the circumstances can provide greater strength to a match between two DNA samples.

Butler et al. (2006) Chromosomal duplications along the Y-chromosome and their potential impact on Y-STR interpretation J. Forensic Sci. 50(4): 853-859



Practical Information on Y Deletions

- If DYS458 is deleted in Yfiler, then your sample is likely to lack an Amelogenin Y amplicon as DYS458 and AMEL Y are 1.13 Mb apart on the short arm of the human Y-chromosome – Chang *et al.* (2007) *Forensic Sci. Int.* 166: 115-120
- Many Y-chromosomes are more complicated than originally thought!

Y-STR Summary

- Mutation rates are similar to autosomal STRs (~0.2%) – based on father-son studies
- Variant alleles are observed as in autosomal STRs due to flanking region mutations, etc.
- Regions of the Y-chromosome can be duplicated or deleted causing Y-STRs to be duplicated or deleted
- Careful primer design is important to avoid X-chromosome homology or Y-chromosome duplications

Standardization is Critical for Success and Data Sharing

Needs	How/When Accomplished
Core Y-STR loci	SWGAM Y-STR Committee selected 11-loci in January 2003
Consistent allele nomenclature	NIST SRM 2395 (2003); kit allelic ladders; ISFG (2006) and NIST (2008) publications
Commercially available Y-STR kits	Early ReliaGene kits (2001-2003); PowerPlex Y (2003) and Yfiler (2004) PowerPlex Y23 (2012)
Accessible, searchable population databases for haplotype frequency estimations	YHRD (72,171 11-locus haplotypes from 750 worldwide populations) US YSTR (18,719 11-locus haplotypes from primarily U.S. population groups)
Interpretation guidelines	SWGAM Y-STR Interpretation Guidelines published in January 2009 <i>(will likely be revised soon)</i>

Predictions for the Future of Y-STR Analysis

- Continued use with casework (with excess female DNA)
- Improved frequency estimates with growing Y-STR databases
 - YHRD now at **70,997 11-locus profiles** (39,339 Yfiler)
 - USYSTR has **18,719 11-locus profiles** (8,548 Yfiler)
- Use with familial searching to eliminate false positives
 - Myers, S.P. *et al.* (2011) *FSI Genetics* 5(5): 493-500 – describes CA DOJ familial searching
- New Y-STR kits with additional loci**
 - At the ISHI meeting, Promega announced a Y-STR 23plex was being developed
 - Will take time though to grow large population databases that cover all of the new loci
- Use of fast mutating loci to help resolve paternal lineages (e.g., to separate brothers or father/son haplotypes)
 - Ballantyne, K.N. *et al.* (2010) *Am J Hum Genet* 87(3): 341-353
 - Ballantyne, K.N. *et al.* (2012) *FSI Genetics* (*in press*)
- In some cases, being able to put a lineage name to an unknown Y-STR profiles using on-line genetic genealogy information**

Expected Number of Y-STR Differences with Various Levels of Relatedness Between Tested Males

	12 loci	25 loci	37 loci	67 loci	111 loci	Interpretation
Very Tightly Related	N/A	N/A	0	0	0	Your exact match means your relatedness is extremely close. Few people achieve this close level of a match. All confidence levels are well within the time frame that surnames were adopted in Western Europe.
Tightly Related	N/A	N/A	1	1-2	1-2	Few people achieve this close level of a match. All confidence levels are well within the time frame that surnames were adopted in Western Europe.
Related	0	0-1	2-3	3-4	3-5	Your degree of matching is within the range of most well-established surname lineages in Western Europe. If you have tested with the Y-DNA12 or Y-DNA25 test, you should consider upgrading to additional STR markers. Doing so will improve your time to common ancestor calculations.
Probably Related	1	2	4	5-6	6-7	Without additional evidence, it is unlikely that you share a common ancestor in recent genealogical times (1 to 6 generations). You may have a connection in more distant genealogical times (less than 15 generations). If you have traditional genealogy records that indicate a relationship, then by testing additional individuals you will either prove or disprove the connection.
Only Possibly Related	2	3	5	7	8-10	It is unlikely that you share a common ancestor in genealogical times (1 to 15 generations). Should you have traditional genealogy records that indicate a relationship, then by testing additional individuals you will either prove or disprove the connection. A careful review of your genealogical records is also recommended.
Not Related	3	4	6	>7	>10	You are not related on your Y-chromosome lineage within recent or distant genealogical times (1 to 15 generations).

From <http://www.familytreedna.com>

If two men share a surname, how should the genetic distance at 25 Y-chromosome STR markers be interpreted?

Genetic Distance	Relationship	Interpretation
0	Related	A perfect 25/25 match between two men who share a surname (or variant) means they likely share a common male ancestor within the genealogical time frame. The probability of a close relationship is very high.
1	Related	A 24/25 match between two men who share a surname (or variant) means they likely share a common male ancestor within the genealogical time frame. For most closely related and same surnamed individuals, the mismatch markers are often DYS439, DYS385, DYS389i, DYS389ii, DYS458, DYS459, DYS449, and DYS464 which have shown themselves to move most rapidly.
2	Probably Related	A 23/25 match between two men who share a surname (or variant) means they may share a common male ancestor within the genealogical time frame. The probability of a relationship is good. However, your results show mutations and therefore more time between you and the other same surnamed person. For most closely related and same surnamed individuals, the mismatch markers are often DYS439, DYS385, DYS389i, DYS389ii, DYS458, DYS459, DYS449, and DYS464 which have shown themselves to move most rapidly.

From <http://www.familytreedna.com>

If two men share a surname, how should the genetic distance at 111 Y-chromosome STR markers be interpreted?

Genetic Distance	Relationship	Interpretation	Related in This Number of Generations or LESS			
			Confidence			
			50%	90%	95%	99%
0	Very Tightly Related	A 111/111 match indicates a very close or immediate relationship. Most exact matches are 3rd cousins or closer, and over half are related within two generations (1st cousins).	2	4	5	6
1	Tightly Related	A 110/111 match indicates a close relationship. Most one-off matches are 5th or more recent cousins, and over half are 2nd cousins or closer.	3	6	7	9
2	Tightly Related	A 109/111 match indicates a close relationship. Most matches are 7th cousins or closer, and over half are 4th or more recent cousins.	5	8	9	11
3	Related	A 108/111 match indicates a genealogical relationship. Most matches at this level are related as 9th cousins or closer, and over half will be 5th or more recent cousins. This is well within the range of traditional genealogy.	6	10	11	14

From <http://www.familytreedna.com>

Rapidly Mutating (RM) Y-STRs

Trying to separate close male relatives

Mutations Seen in 100 African American Father-Son Pairs

Ethnicity	Sample	locus	Allele (father)	Allele (child)	Comments
African American	65B	Y GATA H4	11	9	loss of 2 repeats
African American	46B	DYS389I and DYS389II	14,30	13,29	loss of 1 repeat
African American	58B	DYS389I and DYS389II	14,32	15,33	gain of 1 repeat
African American	18B	DYS390	24	23	loss of 1 repeat
African American	90B	DYS456	15	16	gain of 1 repeat
African American	16B	DYS458	18	19	gain of 1 repeat
African American	39B	DYS458	18	19	gain of 1 repeat
African American	16B	DYS635	23	22	loss of 1 repeat
African American	47B	DYS635	22	23	gain of 1 repeat
African American	72B	DYS635	22	23	gain of 1 repeat
African American	22B	DYS448	19,20	19,20	Duplication
African American	72B	DYS448	19,20	19,20	Duplication
African American	97B	DYS448	17,2,19,20	17,2,19,20	Triplication *
African American	33B	DYS389I and DYS389II			Deletion *
African American	33B	DYS439			Deletion *

Mutations in both DYS458 and DYS635 were observed in father and son 16B

Rapidly Mutating Y-STRs for Separating Male Relatives

LOCUS (average mutation rate)

- DYS449 (1.2%)
- DYS518 (1.8%)
- DYS547 (2.4%)
- DYS570 (1.2%)
- DYS576 (1.4%)
- DYS612 (1.4%)
- DYS626 (1.2%)
- DYS627 (1.2%)

DYS458 (0.64%) is highest in Yfiler loci where average is ~0.2%

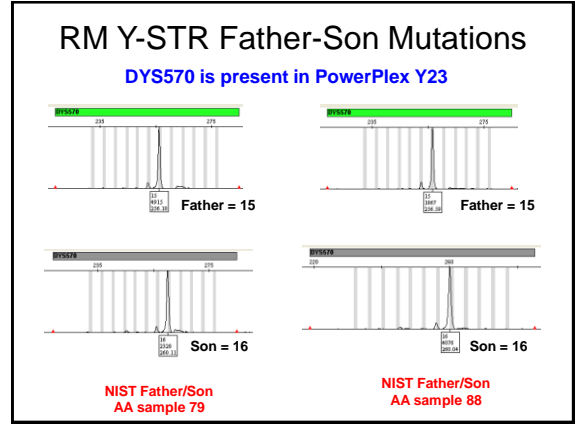
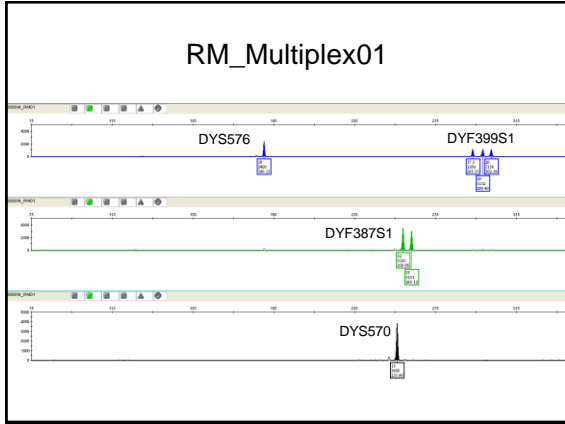
ARTICLE IN PRESS
Forensic Science International: Genetics
journal homepage: www.elsevier.com/locate/bsfig

A new future of forensic Y-chromosome analysis: Rapidly mutating Y-STRs for differentiating male relatives and paternal lineages
Kajre N. Ballantyne^{a,1,2}, Victoria Keel^{a,1,3}, Andreas Wollstein^{a,8}, Ying Choi⁴, Sofia B. Zamiga⁵, Arwin Ralf⁴, Mark Vermeulen⁶, Peter de Knijff⁷, Manfred Kayser^{a,4}

DYF387S1 (1.6%)
DYF399S1 (7.7%)
DYF403S1 a/b (3.1/1.2%)
DYF404S1 (1.3%)
DYS26 a/b (1.3%)

Mutability of Y-Chromosomal Microsatellites: Rates, Characteristics, Molecular Bases, and Forensic Implications

ARTICLE



Statistical Calculations

Statistical Calculations on Y-STR Data

- **Locus (gene) Diversity** = $(n/n-1)(1 - \sum p_i^2)$ where n is the number of samples in the dataset and p_i is the frequency of the i^{th} allele
- **Haplotype Diversity (HD)** = $(n/n-1)(1 - \sum p_i^2)$ where n is the number of samples in the dataset and p_i is the frequency of the i^{th} haplotype
- **Random Match Probability (RMP)** = $1 - HD$
- **Discrimination Capacity (DC)** – total number of observed haplotypes divided by the total number of individuals in the dataset
- **Unique Haplotypes (UH)** – number of haplotypes that occur only once in the dataset

Calculating Gene (STR) Diversity

Locus	Allele	Size Range (bp)	Count	Combined Freq (N = 661)
DYS463	17	222.45	1	0.0015
	18	227.34-227.44	27	0.0408
	19	232.30-232.39	7	0.0106
	20	237.24-237.44	151	0.2284
	21	242.21-242.41	67	0.1014
	22	247.12-247.40	74	0.1120
	23	252.13-252.33	35	0.0530
	24	257.05-257.49	256	0.3873
	25	262.01-262.26	37	0.0560
26	267.05-267.21	5	0.0076	
	277.22	1	0.0015	
	failure	2		
	TOTAL	661		STR diversity 0.7684

$D = (n/n-1)(1 - \sum x_i^2)$

Haplotype Diversity

- is a measure of the uniqueness of a particular haplotype in a given population

$$H = \frac{N}{N-1} (1 - \sum_i x_i^2)$$

Population size

Relative frequency

Discrimination Capacity

- is a measure of the number of unique haplotypes in a given population

$$DC = \frac{\#H}{N}$$

← # of Haplotypes
↑
 Population size

N = 100

$$H = \frac{N}{N-1} \left(1 - \sum_i x_i^2\right)$$

0

DC = 1/100 = 0.01

N = 100

$$H = \frac{N}{N-1} \left(1 - \sum_i x_i^2\right)$$

0.758

DC = 4/100 = 0.04

N = 100

$$H = \frac{N}{N-1} \left(1 - \sum_i x_i^2\right)$$

0.989

DC = 100/100 = 1.0

# times haplotype observed	MHL	SWGAM	PPY	Yfiler	ALL 37
1	429	486	505	626	652
2	34	33	34	12	2
3	13	10	14	2	.
4	4	6	3	.	.
5	3	1	2	.	.
6	1	1	.	.	.
7	1	2	1	.	.
8	1
9	2
10	.	1	.	.	.
11	1
12	.	.	1	.	.
13	1
14
15	.	1	.	.	.
16
17
18
19
20
21
22
23
24
25
26	1
HD	0.996644	0.998529	0.999064	0.999916	0.999991
DC	0.748476	0.824695	0.853659	0.97561	0.996951
# HT	491	541	560	640	654

N = 656

Haplotype Diversity (HD) vs. Discrimination Capacity (DC)

$$HD = \frac{N}{N-1} (1 - \sum x^2)$$

x = frequency of each haplotype

$$DC = \frac{\#HT}{N}$$

Acknowledgments

Funding from interagency agreements 2008-IJ-R-029 (and previously 1999 & 2003) between the National Institute of Justice and the NIST Office of Law Enforcement Standards

NIST Past and Present Team Members:
 Amy Decker, Richard Schoske, Christian Ruitberg, Jill Appleby, Mike Coble, Becky Hill, Margaret Kline, Peter Vallone, Dave Duewer

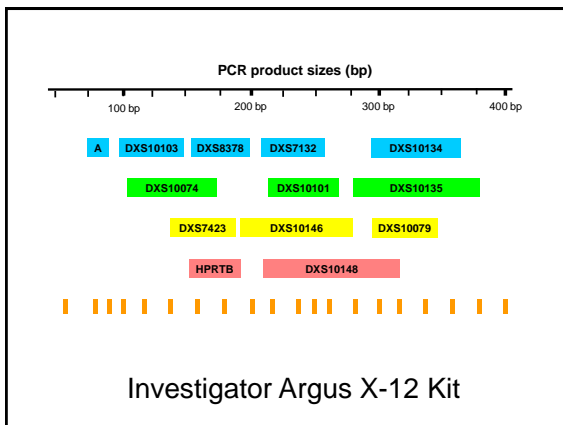
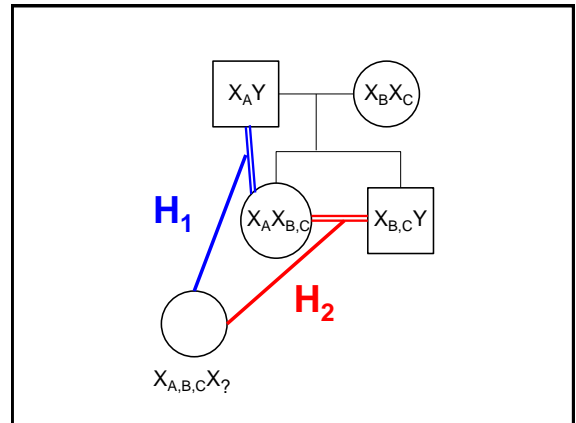
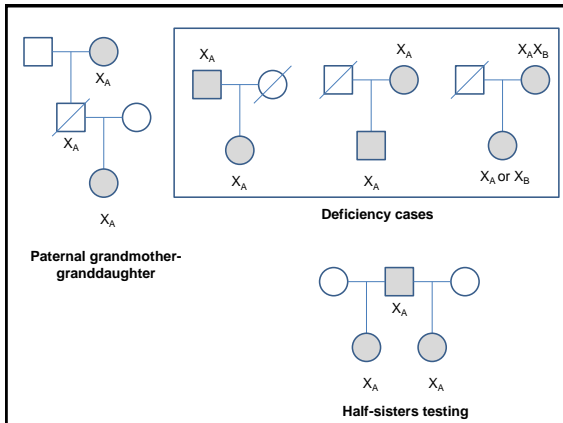
Past Collaborators:
 Mike Hammer, Alan Redd, Tom Reid, ISFG DNA Commission, SWGDAM Y-STR Committee

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

X-Chromosome Markers

Applications of X-Chromosome Analysis

- Complex kinship cases involving at least one female
- Disputed paternity to a daughter (especially in motherless cases)
- Half-sister testing where the father is the common relative
- Grandparent—grandchild comparisons
- Paternity testing in incest cases

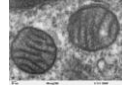


X-STR Summary

- ChrX analysis has potential forensic and human identity testing applications due to its inheritance pattern compared to other genetic markers
- As with the rest of the human genome, STR markers are prevalent along the X-chromosome with comparable density to autosomal STRs
- A number of X-STR assays and kits are available
- Population studies are regularly published with X-STR data

Mitochondrial DNA (mtDNA)

Why Mitochondrial DNA?



- Mitochondria are organelles within cells
 - Produce energy via Krebs Cycle
- Separate genome from the nucleus (≈ 16,569 bp)
- Human cells have hundreds of mitochondria
- Between 2 – 10 genome copies per mitochondrion ≈ 1000 genome copies per cell
- A single cell's mtDNA can be amplified by PCR
 - 6 pg of DNA = 1 nuclear genome = 1000 mtDNA copies
 - When nuclear DNA fails to amplify, can often obtain mtDNA results
- In forensic samples quantity of evidence is sometimes a limitation
 - Trace evidence (hair, blood, bone)

<http://neel.dartmouth.edu/magajournal/magajournal/EMTsource8.htm>

Primary mtDNA Characteristics

- High copy number of mtDNA
- Maternal inheritance of mtDNA
- Lack of recombination
- High mutation rate compared to single copy nucDNA

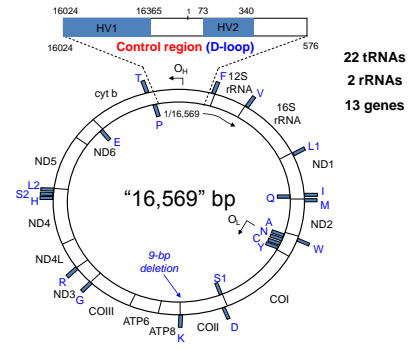
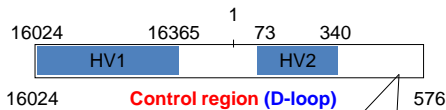


Figure 10.1, J.M. Butler (2005) Forensic DNA Typing, 2nd Edition © 2005 Elsevier Academic Press

Control Region (16024-576)

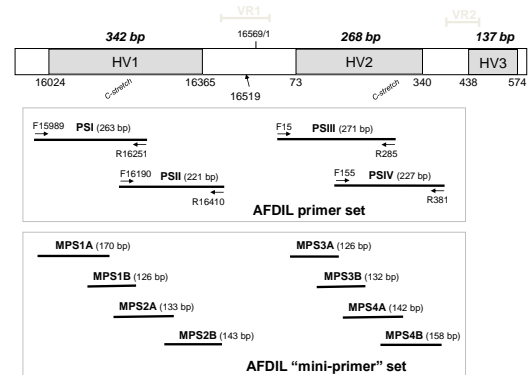
1,122 nucleotide positions



Forensic Focus

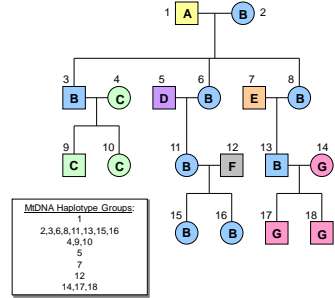
Typically only **610 bases** examined
 – (HVI: 16024-16365; HVII: 73-340)

- (AC)₃
- (AC)₄
- (AC)₅
- (AC)₆
- (AC)₇



Maternal Inheritance of mtDNA

- Fertilizing sperm contributes only nuclear DNA
- Cellular components including the mitochondria in the cytoplasm come from the mother's ovum
- Any sperm mitochondria that may enter a fertilized egg are selectively destroyed due to a ubiquitin tag added during spermatogenesis
- Barring mutation, a mother passes her mtDNA type on to her children



Candidates for mtDNA Testing

- Shed hairs lacking root bulb or attached tissue
- Fragments of hair shafts
- Aged bones or teeth that have been subjected to long periods of exposure
- Crime scene stains or swabs that were unsuccessful for nuclear DNA testing
- Tissues (muscle, organ, skin) that were unsuccessful for nuclear DNA testing

Terry Melton – International Symposium on the Application of DNA Technologies in Analytical Sciences

Process for Evaluation of mtDNA Samples

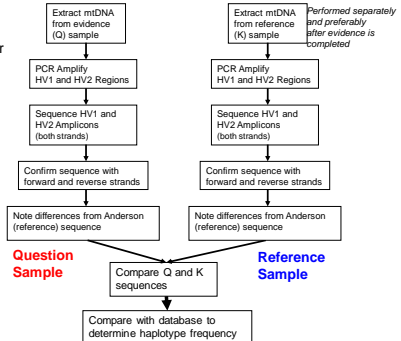
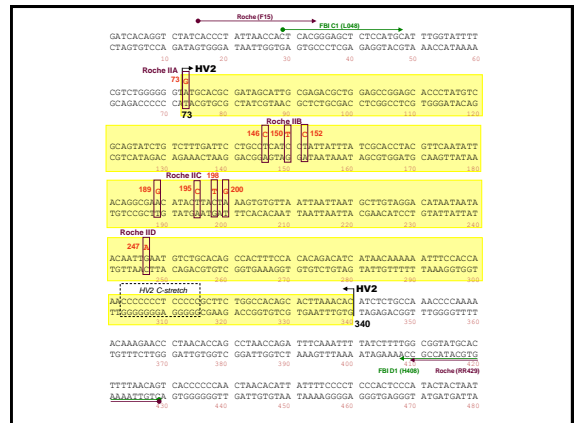
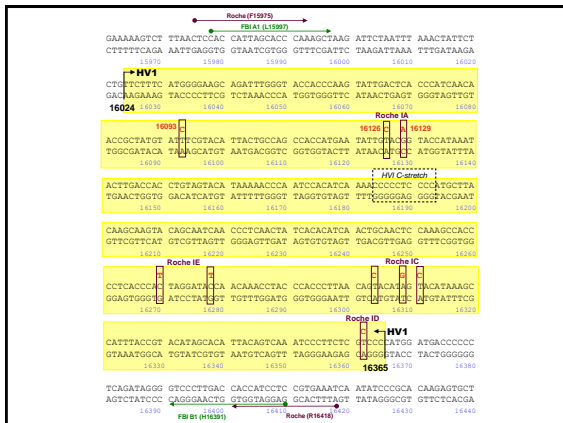
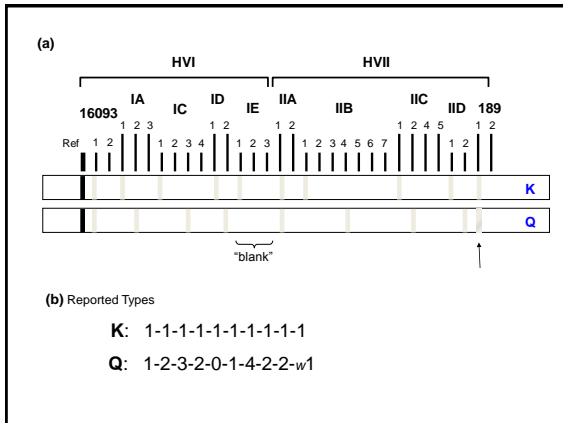
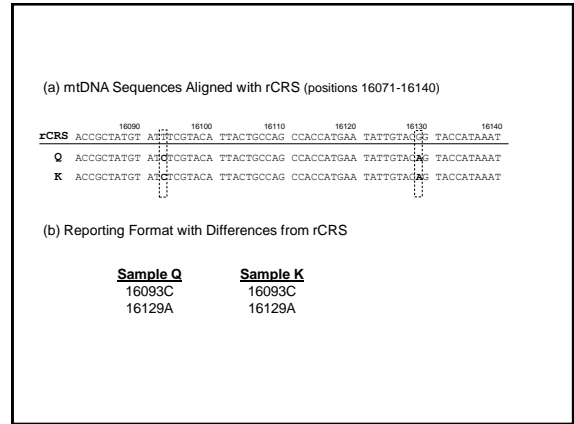
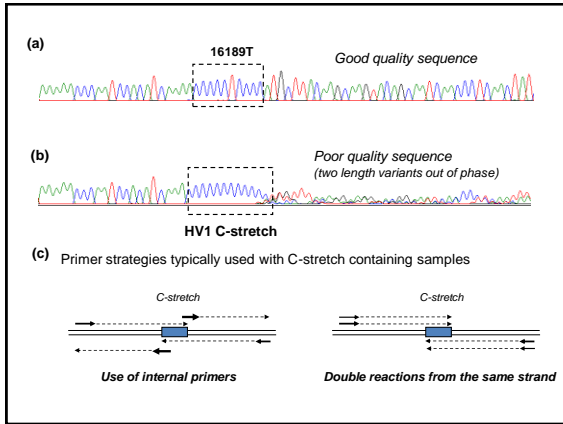


Figure 10-4. J.M. Butler (2005) Forensic DNA Typing, 2nd Edition © 2005 Elsevier Academic Press





Interpretational Issues - Heteroplasmy

- Heteroplasmy – the presence of more than one mtDNA type in an individual
- Once thought to be rare, heteroplasmy exists (at some level) in all tissues
- Especially important in forensic mtDNA analysis of hair

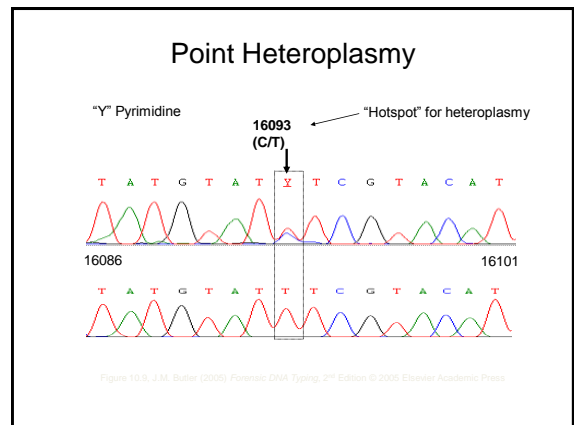
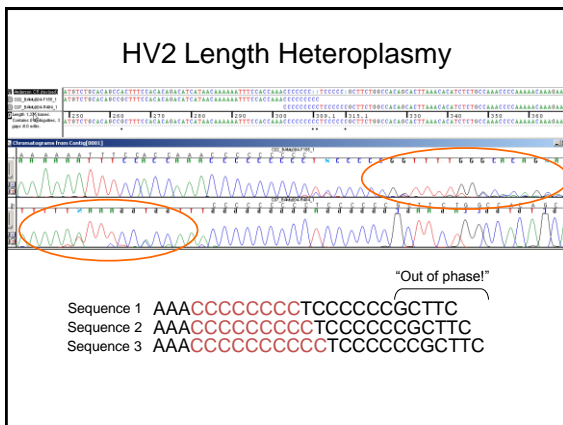


Figure 10.6. J.M. Butler (2005) Forensic DNA Typing, 2nd Edition © 2005 Elsevier Academic Press.

Origination of Heteroplasmy

Ovum – 100K mitochondria

↓

Very little mito growth until implantation

↓

Females – produce ~7 million ova during fetal development only a few hundred become mature oocytes

FIGURE 2. The mitochondrial genetic bottleneck

Chinnery et al. (2000) Trends in Genetics

Int J Legal Med (2008) 122:189–197
DOI 10.1007/s00414-007-0190-6

ORIGINAL ARTICLE

Single lymphocytes from two healthy individuals with mitochondrial point heteroplasmy are mainly homoplasmic

Sabine Lutz-Bonengel · Timo Sanger · Walther Parson · Helena Muller · Joachim W. Elwart · Marie Follo · Bernhard Bonengel · Harald Niederstatter · Marielle Heinrich · Ulrike Schmidt

OPEN ACCESS Freely available online

PLOS ONE

Detection of Heteroplasmic Mitochondrial DNA in Single Mitochondria

Joseph E. Reiner^{1*}, Rani B. Kishore¹, Barbara C. Levin², Thomas Albanetti³, Nicholas Boire³, Ashley Knipe³, Kristian Helmersen³, Koren Holland Deckman³

¹Physical Measurement Laboratory, National Institute of Standards and Technology, Gaithersburg, Maryland, United States of America, ²Material Measurement Laboratory, National Institute of Standards and Technology, Gaithersburg, Maryland, United States of America, ³Department of Chemistry, Gettysburg College, Gettysburg, Pennsylvania, United States of America

December 2010 | Volume 5 | Issue 12 | e14359

mtDNA Base Composition Analysis by Mass Spectrometry

Kevin Kiesler provided slides and performed NIST work

Sequencing vs. Mass Spec

Sanger sequencing

DNA extraction → PCR amplification → Clean-up → Cycle sequencing reaction

Clean-up → Capillary Electrophoresis → Sequence assembly → Sequence analysis

Cost: **\$300** per sample Time: **10** hours

Plex-ID system

DNA extraction → PCR amplification → Automated Clean-up → Automated Mass analysis

Data Q.C.

Cost: **\$185** per sample Time: **5** hours

Significant savings in time, labor, and cost!

Key: Manual step (dark blue), Automated step (light blue)

Base Composition by Mass Spectrometry

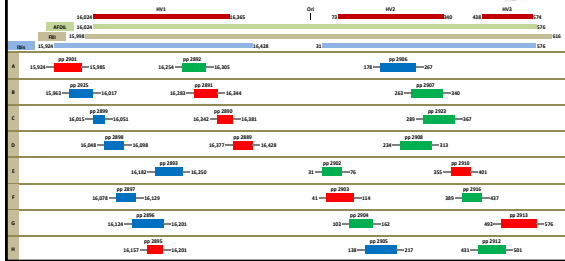
- Electrospray is a soft ionization method
 - Does not fragment molecules
 - Dissociates strands of DNA (PCR product)
 - 5 kV ionization negatively charges DNA (multiple charge states)
- Masses of forward and reverse strands measured
 - Time of flight analyzer
 - Mass/charge ratio (m/z) is result

Plex-ID Instrument

- Automated DNA cleanup
- Closed system

mtDNA 2.0 Triplex PCR Reactions

- The control region is amplified by 24 PCR primer pairs in eight triplex PCR reactions
- Tiled over HV1, HV2, and HV3
- Each nucleotide position is assayed at least once



mtDNA 2.0 Assay from Ibis Biosciences

	1	2	3	4	5	6	7	8	9	10	11	12
A	2906	2906	2906	2906	2906	2906	2906	2906	2906	2906	2906	2906
B	2901	2901	2901	2901	2901	2901	2901	2901	2901	2901	2901	2901
C	2902	2902	2902	2902	2902	2902	2902	2902	2902	2902	2902	2902
D	2907	2907	2907	2907	2907	2907	2907	2907	2907	2907	2907	2907
E	2908	2908	2908	2908	2908	2908	2908	2908	2908	2908	2908	2908
F	2909	2909	2909	2909	2909	2909	2909	2909	2909	2909	2909	2909
G	2910	2910	2910	2910	2910	2910	2910	2910	2910	2910	2910	2910
H	2911	2911	2911	2911	2911	2911	2911	2911	2911	2911	2911	2911
Sample	Sample	Sample	Sample	Sample	Sample	Sample	Sample	Sample	Sample	Sample	Negative Control	Positive Control
	1	2	3	4	5	6	7	8	9	10		

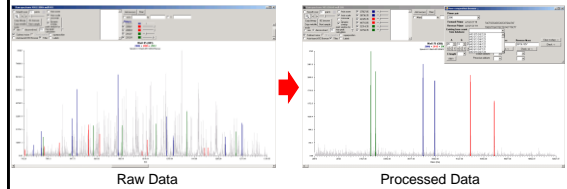
mtDNA 2.0 Assay from Ibis Biosciences

	1	2	3	4	5	6	7	8	9	10	11	12
A	2906	2906	2906	2906	2906	2906	2906	2906	2906	2906	2906	2906
B	2901	2901	2901	2901	2901	2901	2901	2901	2901	2901	2901	2901
C	2902	2902	2902	2902	2902	2902	2902	2902	2902	2902	2902	2902
D	2907	2907	2907	2907	2907	2907	2907	2907	2907	2907	2907	2907
E	2908	2908	2908	2908	2908	2908	2908	2908	2908	2908	2908	2908
F	2909	2909	2909	2909	2909	2909	2909	2909	2909	2909	2909	2909
G	2910	2910	2910	2910	2910	2910	2910	2910	2910	2910	2910	2910
H	2911	2911	2911	2911	2911	2911	2911	2911	2911	2911	2911	2911
Sample	Sample	Sample	Sample	Sample	Sample	Sample	Sample	Sample	Sample	Negative Control	Positive Control	
	1	2	3	4	5	6	7	8	9	10		

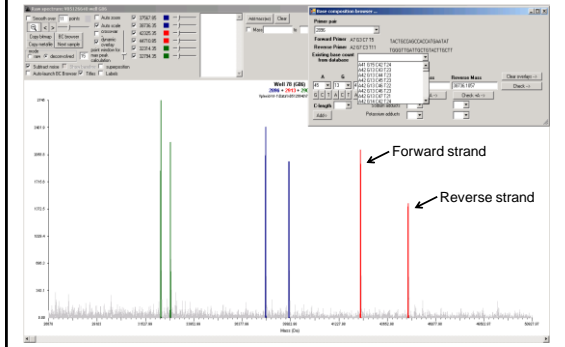
- Prefabricated 96-well plate contains all reagents for amplification
- Each well in plate contains 3 PCR primer pairs (24 amplicons total)
- For each sample, 5 µL is placed in 8 wells in a column (A - H)
- PCR is cycled off-platform then loaded onto the PlexID input stacker
- PCR products are desalted with magnetic bead-based purification
- Automated cleanup and injection is performed by the instrument
- Up to 15 plates can be processed in a single, fully automated run

Results – Mass Spectra

- Complex spectrum of multiple charge states of DNA are deconvolved into simplified spectrum

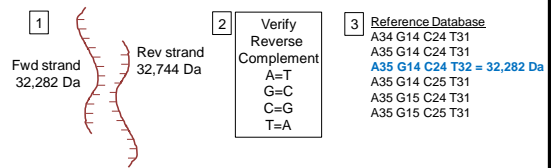


Results – Mass Spectra



Assigning Base Compositions to Mass Measurements

- Forward and reverse strands of PCR amplicons are measured independently (1)
- Watson-Crick reverse complement rules are applied to confirm that the two measured strands are from the same PCR product (2)
- Masses are correlated to a reference database of known base compositions in order to arrive at a measured base composition (3)
 - Combined base compositions of 24 amplicons are the "profile"
 - Can be used to search databases for matching profiles



DNA Mass Spectrometry Limitation: Masses of Natural Nucleotides

- When A -> G & T -> C polymorphisms are present within an amplicon, the mass difference is **+1 Dalton** compared to the reference sequence
 - Cannot be differentiated by mass spec
 - A -> G (329.2 - 313.2 = +16 Da)
 - T -> C (289.2 - 304.2 = - 15 Da)
 - tagctagctgacgatcga**tgctag** mass = 7455 Da
 - tagctagctg**C**gatcga**C**gctag mass = 7456 Da
- To resolve this limitation, the assay uses a G nucleotide labeled with heavy carbon isotope, ^{13}C
 - Adds 10 Da to the mass of the nucleotide
 - Eliminates the ambiguity in combinations of nucleotide masses
 - tagctagctgacgatcga**g**ctagctag mass = **7525 Da**
 - tagctagctg**g**gatcga**C**gctagctag mass = **7536 Da**

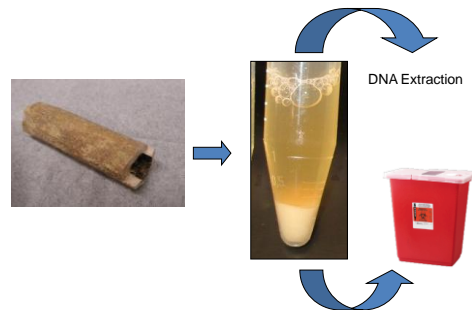
Base Composition vs. Sequence

- Sequencing results in an ordered string of bases
 - AAGAGGTTTCACCTGGTT
- Base composition yields an empirical formula of bases without knowing the order
 - $\text{A}_4\text{G}_6\text{C}_4\text{T}_6$
- The signature of 24 amplicons' base comp signature is almost as unique as sequence
 - Difference: cannot resolve reciprocal base changes within one amplicon
 - Example: C -> T + T -> C = **no change in mass**

Improved extraction protocols for mtDNA testing

Slides from Mike Coble
and work performed at AFDIL

Current Extraction Protocols – Forensic mtDNA Labs



Available online at www.sciencedirect.com

ScienceDirect

Forensic Science International: Genetics 1 (2007) 191–195

ELSEVIER

FSI GENETICS

www.elsevier.com/locate/bscig

Short communication

High efficiency DNA extraction from bone by total demineralization^{*}

Odile M. Loreille^{*}, Toni M. Diegoli, Jodi A. Irwin, Michael D. Coble, Thomas J. Parsons¹

Armed Forces DNA Identification Laboratory, 141F Research Bldg., Bldg. 301, Rockville, MD 20850, United States

Received 24 January 2007; accepted 3 February 2007

Demineralization protocol





- EDTA 0.5M, pH 8.5
- Detergent
- Proteinase K
- 1g powder**

15ml extraction buffer

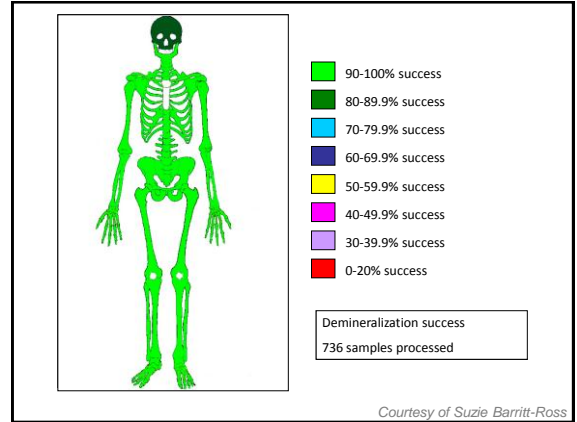
- *Organic extraction (phenol-chloroform)
- *Concentration and washes in filtration devices.

Casework SOP

Deminerlization protocol

10mM Tris, pH 8.0, 100mM NaCl, 50mM EDTA, pH 8.0, 0.5% SDS; ProK



Acknowledgments

- Thanks to Mike Coble (NIST) and Kevin Kiesler (NIST) for many of the slides
- For more information, see *Advanced Topics in Forensic DNA Typing (2012)*
 - Chapter 13 Y-Chromosome DNA Testing
 - Chapter 14 Mitochondrial DNA Analysis
 - Chapter 15 X-Chromosome Analysis