

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



NIST National Institute of Standards and Technology

The Future of Forensic DNA Typing John M. Butler

NIST Applied Genetics Group

National Institute of Standards and Technology Gaithersburg, Maryland

National Commission on the Future of DNA Evidence



U.S. Department of Justice Office of Justice Programs National Justice of Justice



The Future of Forensic DNA Testing

Predictions of the Research and Development Working Group

http://www.ojp.usdoj.gov/nij/pubs-sum/183697.htm

A Report Fro

•Report published in Nov 2000

•Asked to estimate where DNA testing would be 2, 5, and 10 years into the future

Conclusions

STR typing is here to stay for a few years because of DNA databases that have grown to contain millions of profiles

My Thoughts on the Future

- Near-term future
 - New autosomal STR loci for expanded core loci
 - Expanded use of databases (e.g., familial searching)
 - Rapid DNA testing
- More distant future
 - Next-generation DNA sequencing?
 - Loci besides STRs for identity testing?
 - Phenotyping capabilities?

STRs vs SNPs Article

Butler et al. (2007) STRs vs SNPs: thoughts on the future of forensic DNA testing. Forensic Science, Medicine and Pathology 3:200-205.

Forensic Sci Med Pathol (2007) 3:200–205 DOI 10.1007/s12024-007-0018-1

ORIGINAL PAPER

STRs vs. SNPs: thoughts on the future of forensic DNA testing

John M. Butler · Michael D. Coble · Peter M. Vallone

- SNPs are unlikely to replace STRs for routine forensic DNA testing due to challenges with high-level multiplexing and mixture detection/interpretation
- Most likely use of SNPs will be as ancestry-informative markers (AIMs) for sample ethnicity estimation



National Academies Report on Forensic Science

Harry T. Edwards U.S. Court of Appeals (DC) Co-Chair, Forensic Science Committee

- Released February 18, 2009
- Entitled "Strengthening Forensic Science in the United States: A Path Forward"
- 13 recommendations provided to Congress
- Recommends establishing a National Institute of Forensic Science (NIFS)
- NIST will have a role in NIFS and our group has been asked to contribute expertise regarding validation and testing of DNA systems as a model for other forensic disciplines



Advisers to the Nation on Science, Engineering, and Medicine

Forensic Science Review Article

See June 15, 2009 issue of Analytical Chemistry

Anal. Chem. 2007, 79, 4365-4384

Forensic Science

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J. R. Almirall

Department of Chemistry and Biochemistry and International Forensic Research Institute, Florida International University, University Park, Miami, Florida 33199

2009 review article covers 160 DNA articles published in 2007-2008

Value of a Historical Review

"If you want to understand today, you have to search yesterday."

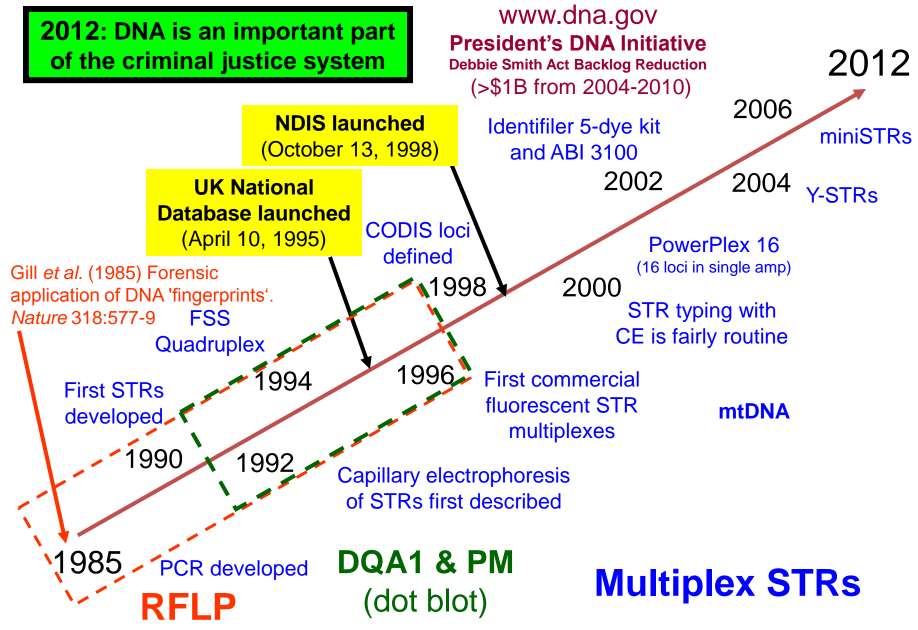
– Attributed to Pearl Buck

(http://www.quotegarden.com/history.html)



The Nobel Prize in Literature 1938

Historical Perspective on DNA Typing



Stages of Forensic DNA Progression

Stages	Time Frame	Description
Exploration	1985-1995	Beginnings, different methods tried (RFLP and early PCR)
Stabilization	1995-2005	Standardization to STRs, selection of core loci, implementation of Quality Assurance Standards
Growth	2005-2012	Rapid growth of DNA databases, extended applications pursued
Sophistication	The Future	Expanding tools available, confronting privacy concerns

Basis of DNA Profiling

The genome of **each individual is unique** (with the exception of identical twins) and **is inherited from parents**

Probe subsets of genetic variation in order to differentiate between individuals (statistical probabilities of a random match are used)

DNA typing must be **performed efficiently and reproducibly** (information must hold up in court)

Current standard DNA tests **DO NOT look at genes** – little/no information about race, predisposal to disease, or phenotypical information (eye color, height, hair color) is obtained

Short Tandem Repeat (STR) Markers

An accordion-like DNA sequence that occurs between genes

= 11 GATA repeats ("11" is all that is reported)

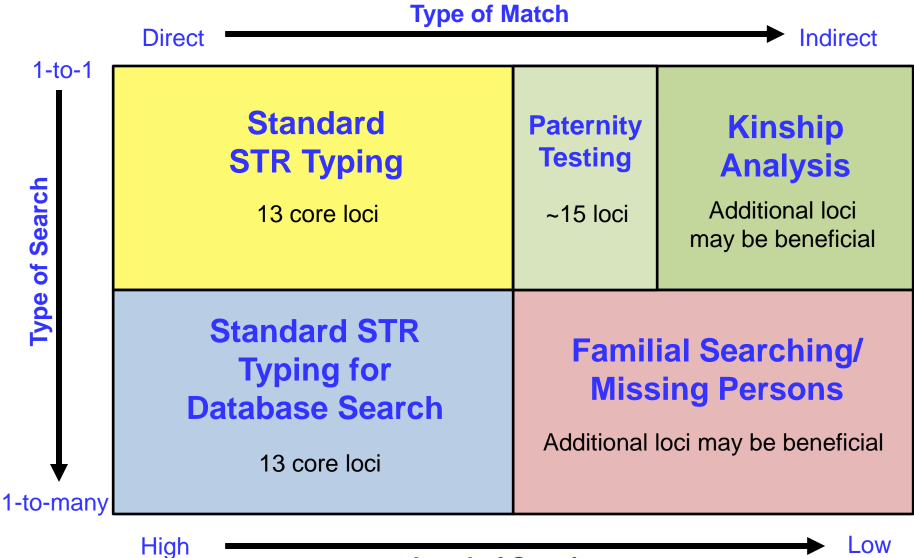
- → 7 repeats ←
- → 8 repeats ←
- → 9 repeats ←
- → 10 repeats ←
- → 11 repeats ←
- → 12 repeats ←
- → 13 repeats



The number of consecutive repeat units can vary between people

> 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

Expanding the Forensic Core Competency



Level of Certainty

Growth of DNA Databases

- Within the U.S., we have benefited from significant federal funding over the past seven years
- Expanded laws now enable more offenders to be included
- Have effectively locked technology with core STR markers used to generate DNA profiles that now number greater than 10 million profiles

Growth in Numbers of U.S. States Requiring DNA Collection for Various Offenses

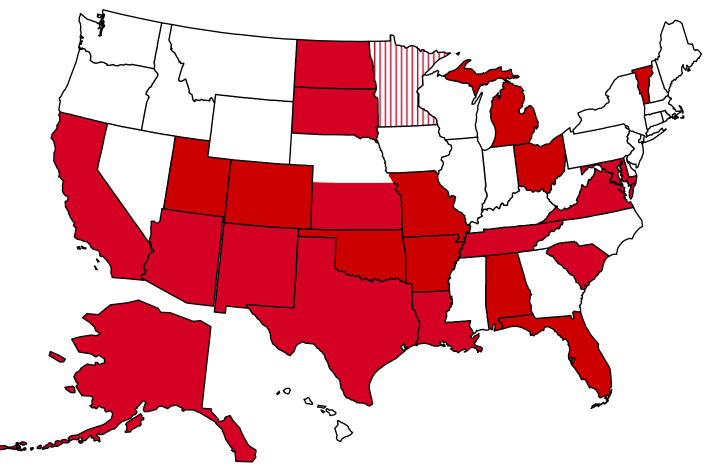
Offenses	Number of States		
	1999	2004	2008
Sex crimes	50	50	50
All violent crimes	36	48	50
Burglary	14	47	50
All felons	5	37	47
Juveniles	24	32	32
Arrestees/suspects	1	4	14

Sources: http://www.dnaresource.com and http://www.ncsl.org/programs/cj/dnadatabanks.htm

Starting initially with sex crimes, each category has grown in the past decade... burglary, all felons, arrestees...

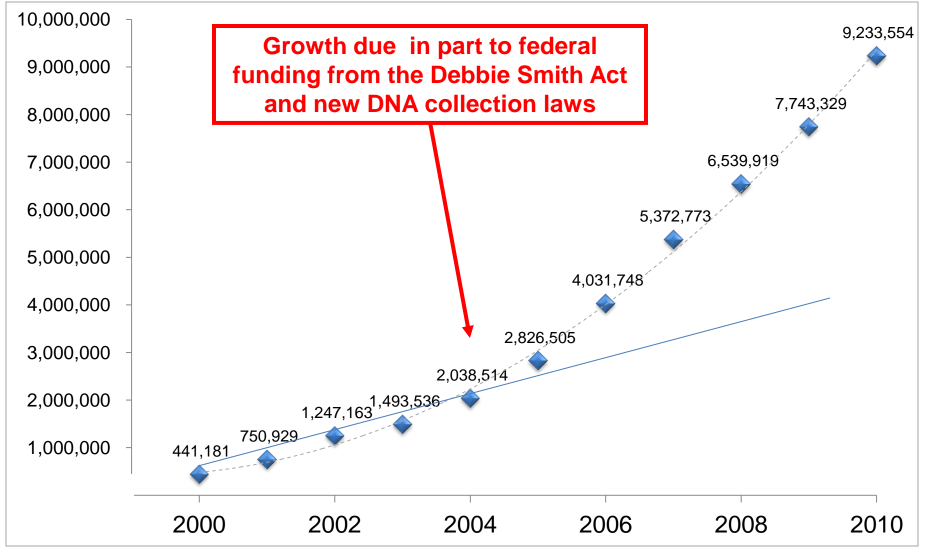
Half of the U.S. Requires Arrestee DNA Testing

+ Federal & DoD



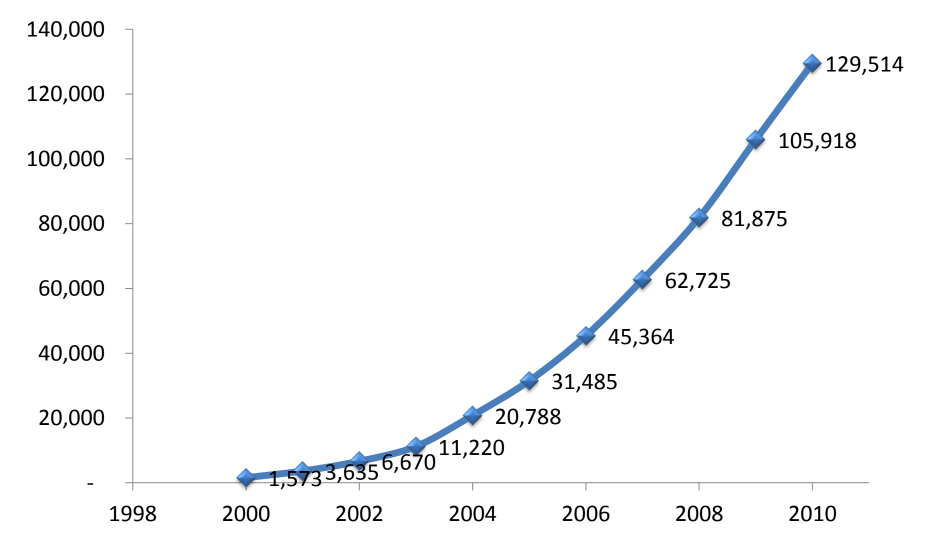
Data as of July 2010

Number of Offender DNA Profiles in the U.S. National DNA Database



Source: FBI Laboratory's CODIS Unit

Number of Investigations Aided in the U.S. National DNA Database



Source: FBI Laboratory's CODIS Unit

National DNA Index System (NDIS)





http://www.fbi.gov/hq/lab/codis/index1.htm

<u>Combined</u> <u>**DNA**</u> <u>Index</u> <u>System</u> (CODIS)



Launched in October 1998 and now links all 50 states

Used for linking serial crimes and unsolved cases with repeat offenders Convicted offender and forensic case samples along with a missing

persons index

Requires 13 core STR markers

>170,000 investigations aided nationwide as of April 2012

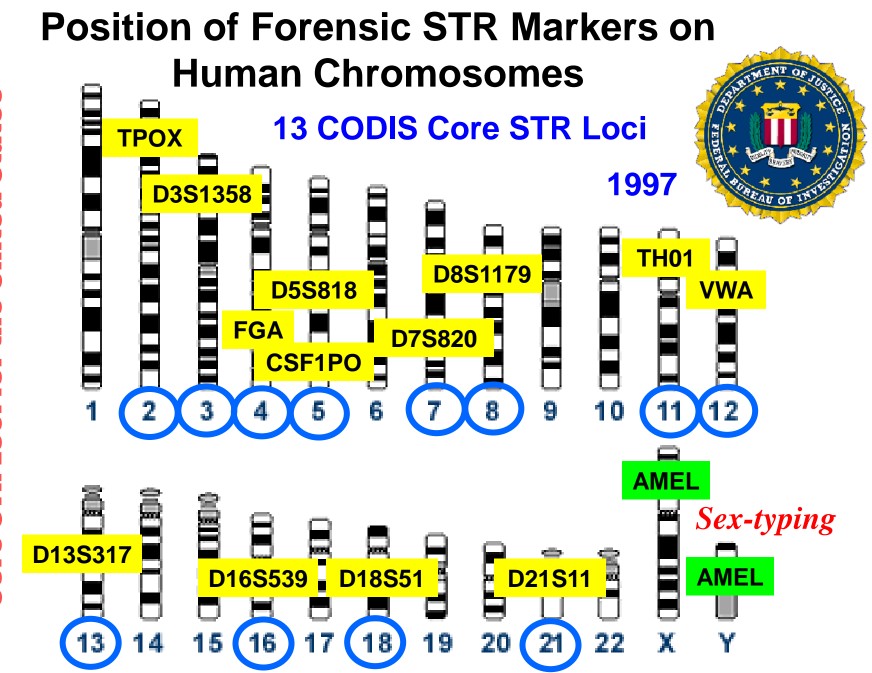
Contains more than 11 million DNA profiles

Growth in Numbers of DNA Profiles Present in Various NDIS Indices

(cumulative totals by year)

Year ending Dec 31	Forensic	Convicted Offender	Arrestee	Total Offender*
2000	21,625	441,181		441,181
2001	27,897	750,929		750,929
2002	46,177	1,247,163		1,247,163
2003				0040
2004		ist two yea	•	
2005	103,008	forensic	samples	added
2006		35 offende	-	
2007	2,000,00			
2008	248,943	6,398,874	140,719	6,539,919
2009	298,369	7,389,917	351,926	7,743,329
2010	351,951	8,559,841	668,849	9,233,554

Source: FBI Laboratory's CODIS Unit



Core STR Loci for the United States

Expanding the CODIS Core Loci

D.R. Hares (2012) Expanding the CODIS Core Loci in the United States. *Forensic Sci. Int. Genet.* 6: e52-e54 *Addendum to expanding the CODIS core loci in the United States*, Forensic Sci. Int. Genet. (2012) doi:10.1016/j.fsigen.2012.01.003



Letter to the Editor

Expanding the CODIS core loci in the United States

CODIS Core Loci Working Group

Formed in May 2010 to make recommendations to FBI CODIS Unit

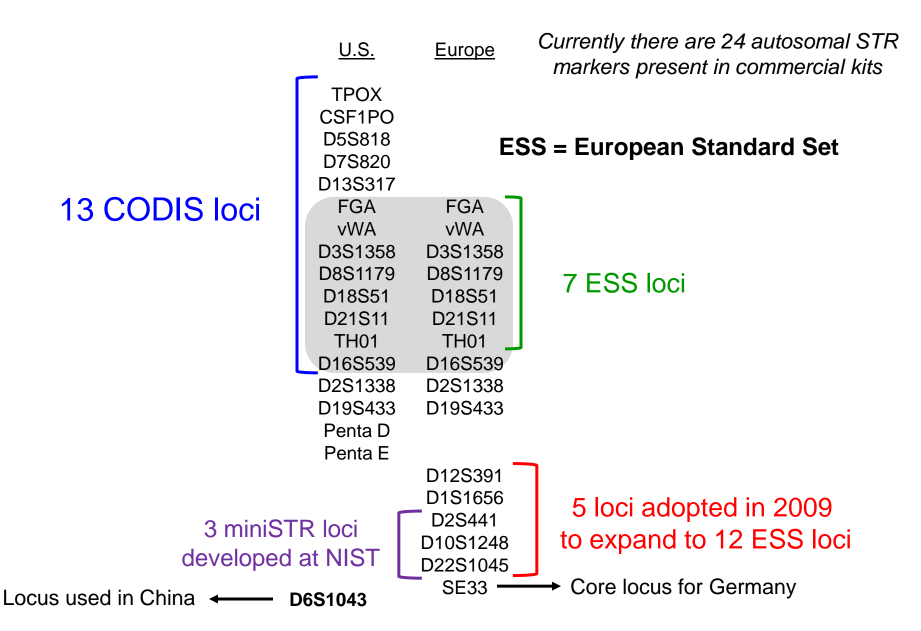
Douglas Hares (Chair) – FBI John Butler – NIST Cecelia Crouse – FL PBSO Brad Jenkins – VA DFS Ken Konzak – CA DOJ Taylor Scott – IL SP major reasons for expanding the CODIS core loci in the United States:

- (1) To reduce the likelihood of adventitious matches [7] as the number of profiles stored at NDIS continues to increase each year (expected to total over 10 million profiles by the time of this publication). There are no signs that this trend will slow down as States expand the coverage of their DNA database programs and increase laboratory efficiency and capacity.
- (2) To increase international compatibility to assist law enforcement data sharing efforts.
- (3) To increase discrimination power to aid missing persons cases.

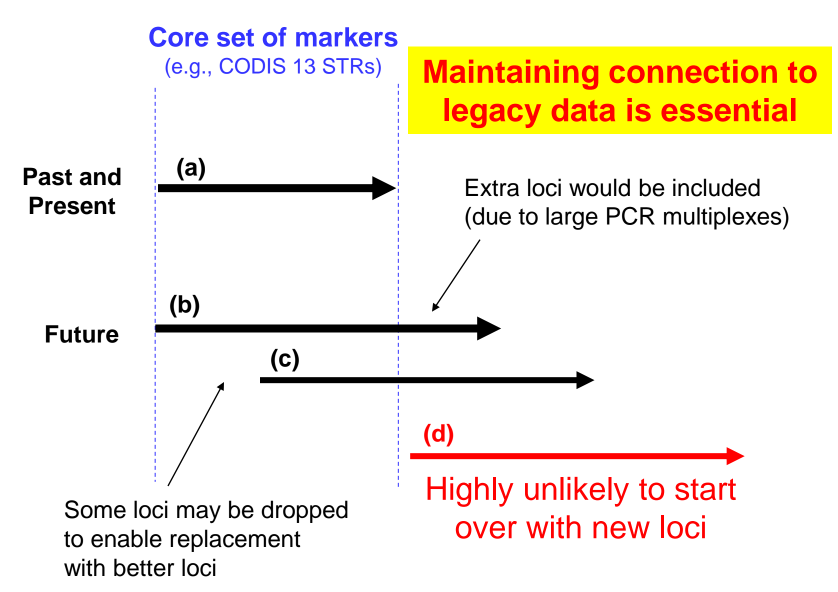
Three major reasons for expanding the CODIS core loci in the United States D.R. Hares (2012) Forensic Sci. Int. Genet. 6(1):e52-e54

- To reduce the likelihood of adventitious matches as the number of profiles stored at NDIS continues to increase each year
- To increase international compatibility to assist law enforcement data sharing efforts
- To increase discrimination power to aid missing persons cases

International Comparability

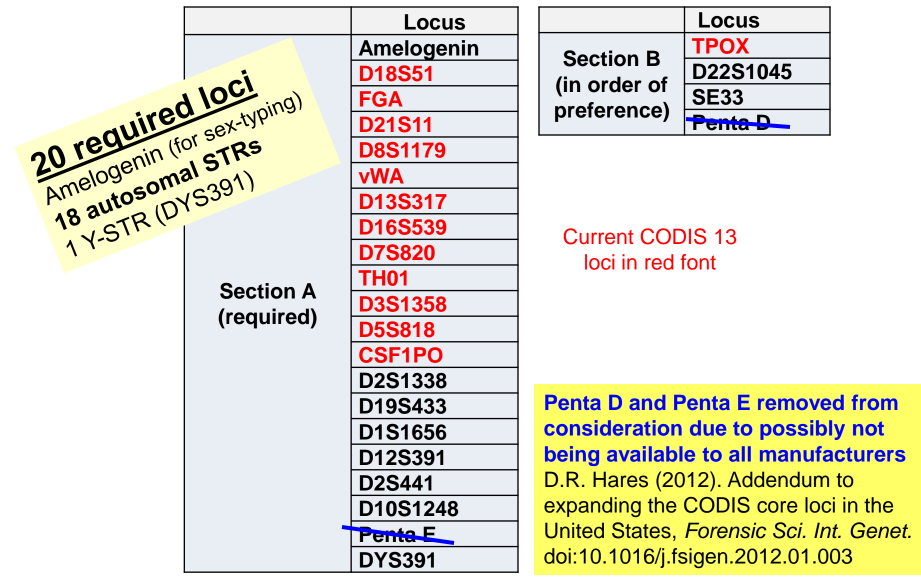


Possible scenarios for extending sets of genetic markers to be used in national DNA databases



Proposed Expanded CODIS Core Loci

D.R. Hares (2012) Forensic Sci. Int. Genet. 6(1):e52-e54



Criteria for Acceptance of Additional Loci

D.R. Hares (2012) Forensic Sci. Int. Genet. 6(1):e52-e54

Considered only short tandem repeat (STR) loci due to need for compatibility to existing database of >10 million STR profiles

STR Loci

- No known association to medical conditions or defects
- Low mutation rate
- High level of independence
- High level of discrimination
- Use by international forensic DNA community
- Number of loci vs. discrimination factor
- Compliance with Quality Assurance Standards (QAS)

Kit performance

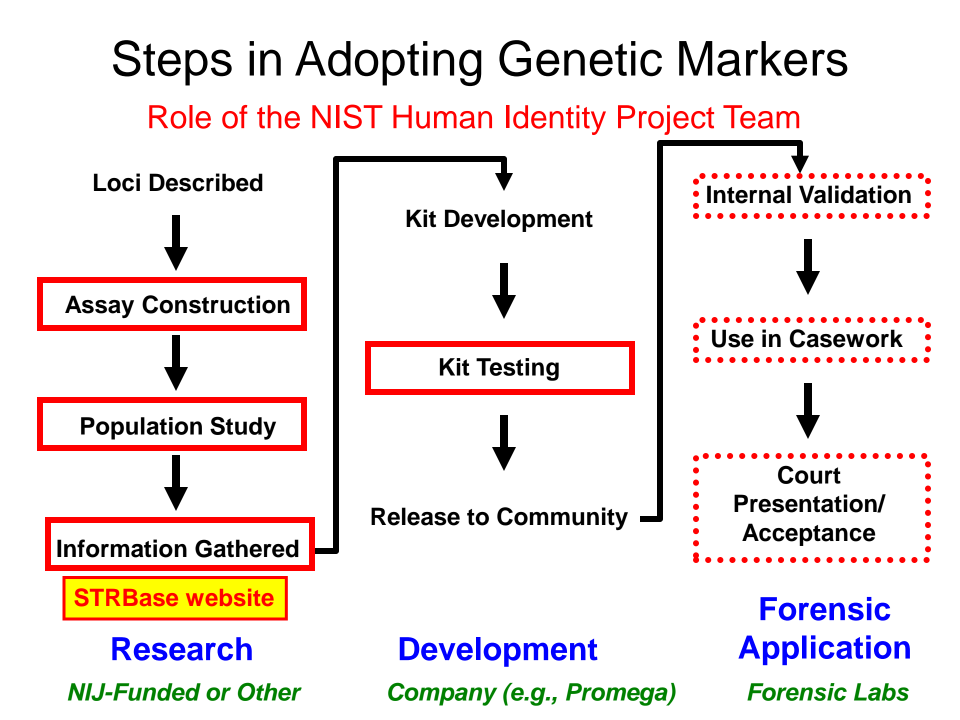
- Balance between loci
- Reliable
- Reproducible
- Sensitive
- Quality results
- Adaptable for use by NDIS laboratories (# of amplifications, ability of kit manufacturers to produce)
- QAS compliant (documentation and availability of validation requirements)

Determination of Additional CODIS Core Loci

D.R. Hares (2012) Expanding the CODIS Core Loci in the United States. *Forensic Sci. Int. Genet.* 6: e52-e54 *Addendum to expanding the CODIS core loci in the United States*, Forensic Sci. Int. Genet. (2012) doi:10.1016/j.fsigen.2012.01.003

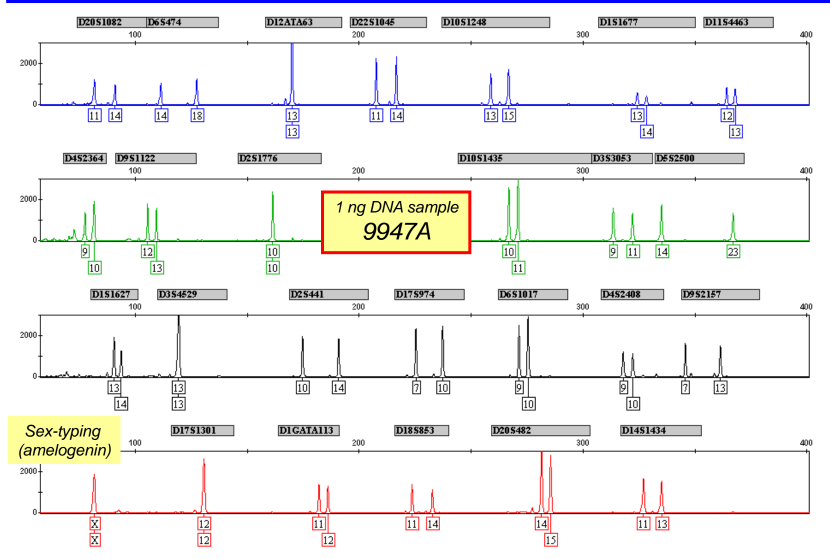
What	Why	Who/How	When
Form a Working Group (WG) to discuss initial selection	Establishes target goals	CODIS Core Loci Working Group with FBI Chair and 5 members; Web meetings	May 2010 - present
Announce proposed additional CODIS core loci	Sets desired target goals and informs manufacturers	WG Chair; Publish proposed listing of CODIS core loci	April 2011 online (published Jan 2012)
Ongoing Progress Reports	Provides updates for DNA community	WG Chair; Present updates on status of CODIS Core Loci project at meetings	2010-2012
Implementation Considerations & Strategy	Identify issues for implementation and timeline	WG	June 2011 - present
Manufacturers develop prototype kits	Creates tools to meet target goals	Manufacturers; Provide status reports to WG for timeline	2011-2012
Test and validate prototype kits	Examines if target goals can be met	Validation Laboratories; Follow QAS compliant validation plan	Beginning in 2012
Review and evaluate data from validation	Evaluates if desired performance is obtained	NIST, SWGDAM and FBI; Provide feedback, if any, to Manufacturers	In conjunction with and at the conclusion of validation
Selection of new CODIS core loci	Allows protocols to be established	FBI; seek input from DNA community and stakeholders; Notify Congress	After evaluation of validation data and kit production factors
Implementation of new CODIS core loci at the National DNA Index System	Enables target goals to be met	All NDIS-participating labs	~ 24 months after selection of new CODIS core loci

http://www.fbi.gov/about-us/lab/codis/planned-process-and-timeline-for-implementationof-additional-codis-core-loci



NIST 26plex

Gender identification + 25 autosomal STR loci in a single amplification



Hill, C.R., et al. (2009) A new STR 26plex assay for human identity testing. J. Forensic Sci. (in press)

Primer sequences and GM bins & panels available: http://www.cstl.nist.gov/biotech/strbase/str26plex.htm

NIST 26plex Demonstration

- Our group at NIST has demonstrated that 25 autosomal STRs and amelogenin (26plex with 52 PCR primers) can be co-amplified with sensitivities similar to commercial STR kits
 - Hill, C.R., Butler, J.M., Vallone, P.M. (2009) A 26plex autosomal STR assay to aid human identity testing. <u>J. Forensic Sci. 54(5): 1008-1015</u>.
 - See also <u>http://www.cstl.nist.gov/biotech/strbase/str26plex.htm</u>

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.

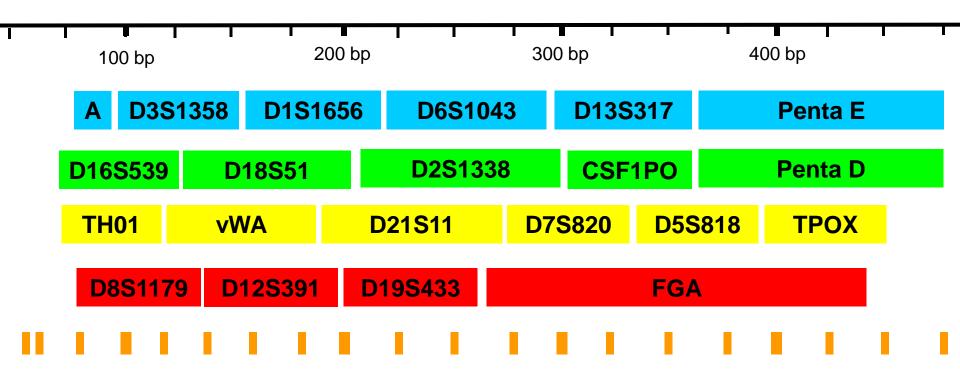
A 26plex Autosomal STR Assay to Aid Human Identity Testing*[†]

PowerPlex 21

- Promega STR kit was released in January 2012
 - NIST has been working with this kit since spring 2011 primarily for concordance testing and has permission from Promega to discuss results
- Contains 20 autosomal STRs + amelogenin
- Enables examination of performance characteristics similar to a future U.S. megaplex containing at least 20 loci

PowerPlex 21

(released by Promega in early 2012)



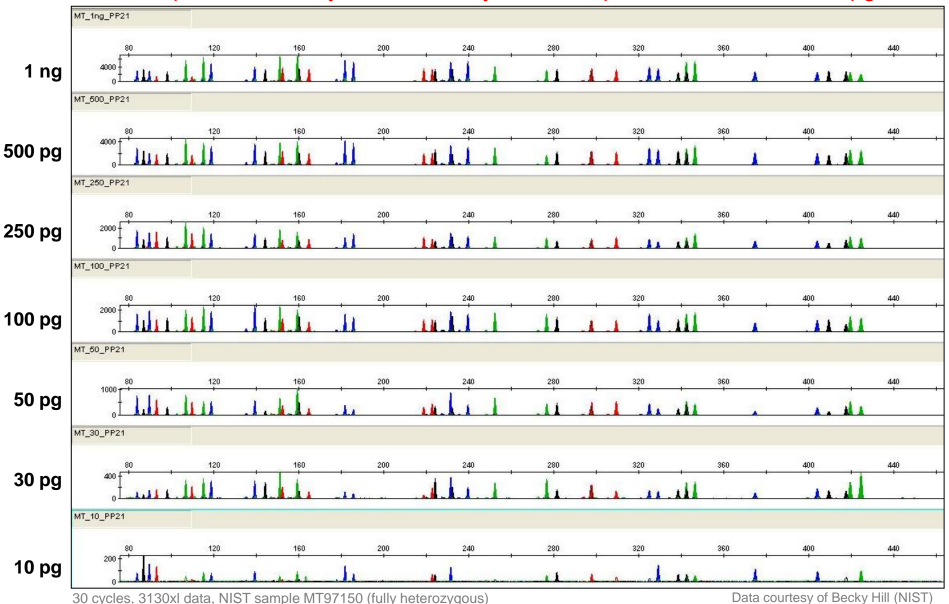
Promega 5-dye kit

13 CODIS STRs + amelogenin

Penta D & Penta E (PP16 loci) D2S1338 & D19S433 (Identifiler loci) D12S391 & D1S1656 (best new European loci) D6S1043 (previously only used in China – ABI Sinofiler kit – highly polymorphic)

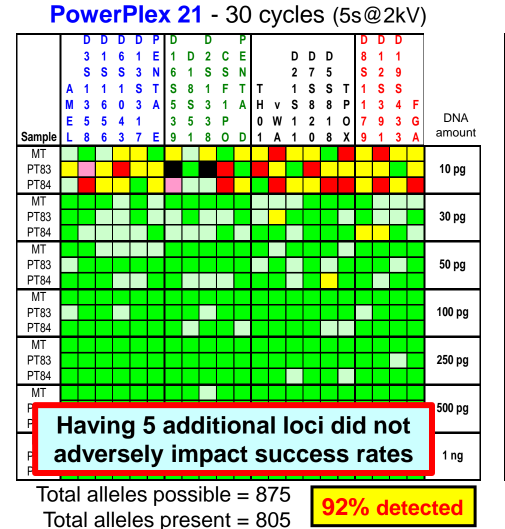
DNA Dilution Series with PowerPlex 21

As expected with any STR kit/assay, allele dropout occurs below 100 pg...

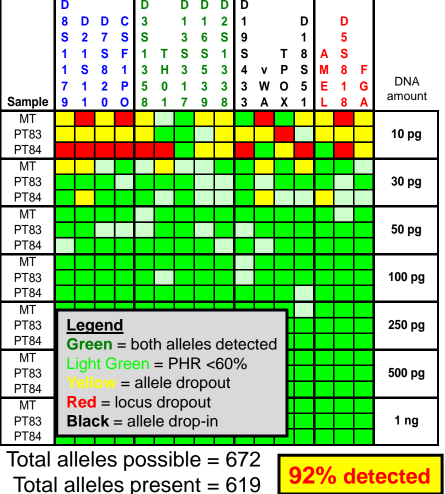


Measurement of Allele Dropout and Extreme Peak Height Imbalance for 2 STR Kits

Three fully heterozygous (except PT83 at Penta D) pristine DNA samples were examined in a dilution series with PowerPlex 21 and Identifiler Plus. Results are ordered by amplicon size and dye color.



Identifiler Plus - 28 cycles (10s@3kV)



Going Beyond the Core Competencies of Forensic DNA Testing

Core Competency

Standard STR Typing (DNA Profile)

Sufficient DNA quantity (ng)

Lower amounts of DNA being tested Direct Matching (or Parentage)

Challenging kinship search questions

Familial Searching Attempts (fishing for brothers or other relatives)

Solution: Additional Markers (Y-chromosome, more STRs) and Multiple Reference Samples

Touch DNA Attempts (poor quality, mixtures, low-level stochastic effects)

Solution: Replicate Testing

Be very cautious when outside the box...

Arrest Made in L.A. 'Grim Sleeper' Killings

Published July 07, 2010 | Associated Press

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LOS ANGELES -- A one-time police mechanic was arrested and charged Wednesday in the serial killing of 10 people over 25 years after a DNA sample from his son was found to bear a close resemblance to DNA found on the victims.

Lonnie Franklin Jr., 57, was charged with 10 counts of murder, one count of attempted murder and special circumstance allegations of multiple murders that could make him eligible for the death penalty if convicted, District Attorney Steve Cooley said.



Lonnie David Franklin Jr.

He is charged with 10 counts of murder and one count of attempted murder for a series of killings that date back to 1985.

Victims of the Grim Sleeper

http://www.laweekly.com/2008-08-28/news/eleven-lives-stolen-and-one-lucky-survivor/

The Grim Sleeper's Victims

- 1) Debra Jackson (age 29) August 10, 1985
- 2) Henrietta Wright (age 35) August 12, 1986
- 3) Thomas Steele (age 36) August 14, 1986
- 4) Barbara Ware (age 23) January 10, 1987
- 5) Bernita Sparks (age 25) April 15, 1987
- 6) Mary Lowe (age 26) October 31, 1987
- 7) Lachrica Jefferson (age 22) January 30, 1988
- 8) Monique Alexander (age 18) September 11, 1988
- 9) Enietra Washington (raped but survived) November 1988
- 10) Princess Berthomieux (age 14) March 19, 2002
- 11) Valerie McCorvey (age 35) July 11, 2003
- 12) Janecia Peters (age 25) January 1, 2007

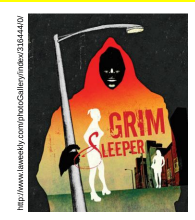


http://blogs.laweekly.com/informer/crime/grim-sleeper-son-dna-trail-led/

Ballistics on bullets recovered from the victim's bodies matched

DNA evidence recovered

Over a 13 year gap in detected crimes, hence the "Sleeper" nickname



Putative Relative Is Found

- June 30, 2010: Second familial search of the California database yielded one likely relative
- Database profile belonged to Christopher Franklin (31 years old)
 - Profile added to the database in 2009 after a felony weapons possession charge
- Grim Sleeper profile matched C. Franklin's profile with one allele at all 15 loci
- Both individuals shared the same Y-STR profile, indicating a possible paternal relationship

Identifying the Grim Sleeper

- Given that the murders spanned at least 25 years, the paternal relationship was likely father-son
- Undercover police shadowed C. Franklin's father, Lonnie David Franklin, Jr., who lived in the vicinity of the murders
- Police collected a DNA sample from Lonnie Franklin
 - Direct match between L. Franklin and the Grim Sleeper

California Familial DNA Search Team

Familial DNA Testing Scores A Win in Serial Killer Case



http://www.sciencemag.org/cgi/reprint/329/5989/262.pdf

Familial Searching in the U.S.

High-profile success in the Grim Sleeper case has led other states to consider familial searching

Experts say Texas might solve Twilight Serial Rapist cases with family DNA

July 25th, 2010 8:23 am CT

http://www.examiner.com/law-enforcement-in-wichita-falls/experts-say-texas-might-solve-twilight-serial-rapist-cases-with-family-dna

DNA DATABASE Milwaukee police on hunt for serial killer linked to 7 deaths

May 19, 2009 http://articles.cnn.com/2009-05-19/justice/wisconsin.serial.killer_1_dna-technology-dna-database-prostitutes?_s=PM:CRIME

Familial DNA hunt sought in East Coast rape case

http://www2.insidenova.com/news/2010/aug/04/familial_dna_hunt_sought_in_east_coast_rape_case-ar-428231/

March 21, 2011 Virginia announced familial searching capability Wednesday December 1, 2010

Virginia could become 3rd state to use familial DNA searches

Some concerned practice could stigmatize those related to criminals

Common PCR Thermal Cycling Times

Can we reduce PCR cycling times? What are the effects or limitations?

	Thermal Cycling	Times for	Current STR Typ	oing Kits		
Year	Run on a 9700 thermal cycler	Hot start	Time per cycle	Cycles	Post soak	Total time
1997/98	Profiler Plus/Cofiler	11 min	3 min	28	60 min	2:52
1999	SGM Plus	11 min	3 min	28	45 min	2:53
2000	PowerPlex 16	12 min	1 min 45 s	32	30 min	3:00
2001	Identifiler	11 min	3 min	28	60 min	2:58
2003	PowerPlex Y	12 min	1 min 45 s	32	30 min	3:18
2004	Yfiler	11 min	3 min	30	80 min	2:45
2007	PowerPlex S5	2 min	4 min	30	45 min	3:21
2007	minifiler	11 min	3 min 20 s	30	45 min	3:16
2009	ESI 16, 17 ESX 16,17	2 min	4 min	30	45 min	3:22
2009	PowerPlex 16 HS	2 min	1 min 45 s	32	30 min	2:42
2009	NGM	11 min	3 min 20 s	29	10 min	2:33
2009	Identifler Direct	11 min	3 min	26	25 min	2:34
2010	Idenfiler Plus	11 min	3 min 20 s	28	10 min	2:18
2011	PowerPlex 18D	2 min	1 min 10s	27	20 min	1:25

Thermal Cyclers

- 1. GeneAmp 9700 (Applied Biosystems)
- 2. Mastercycler Pro S (Eppendorf)
 - Peltier based
- 3. Rotor-Gene Q (Qiagen)
 - Air heated and cooled
- 4. SmartCycler (Cepheid)
 - Hot plates for heating, fans for cooling
- Cycling for most STR kits is run in the
- '9600 emulation mode' (1°C/s)



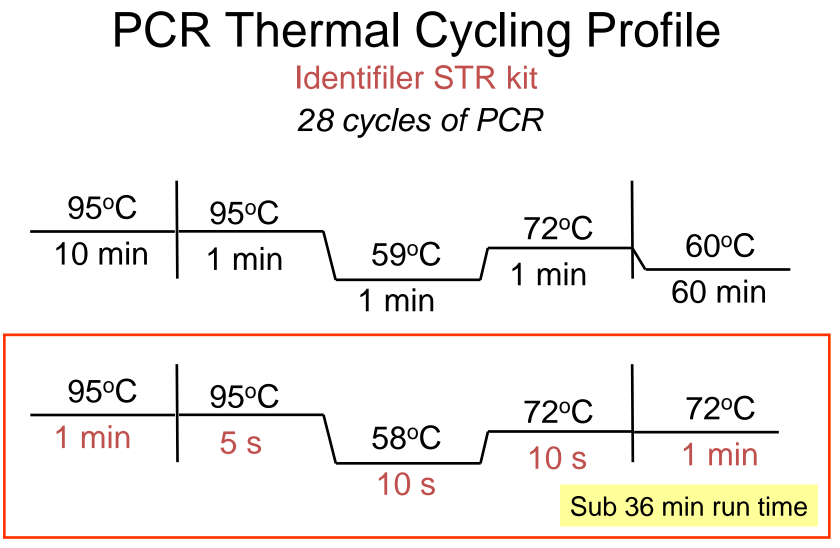


Intended for

real-time PCR





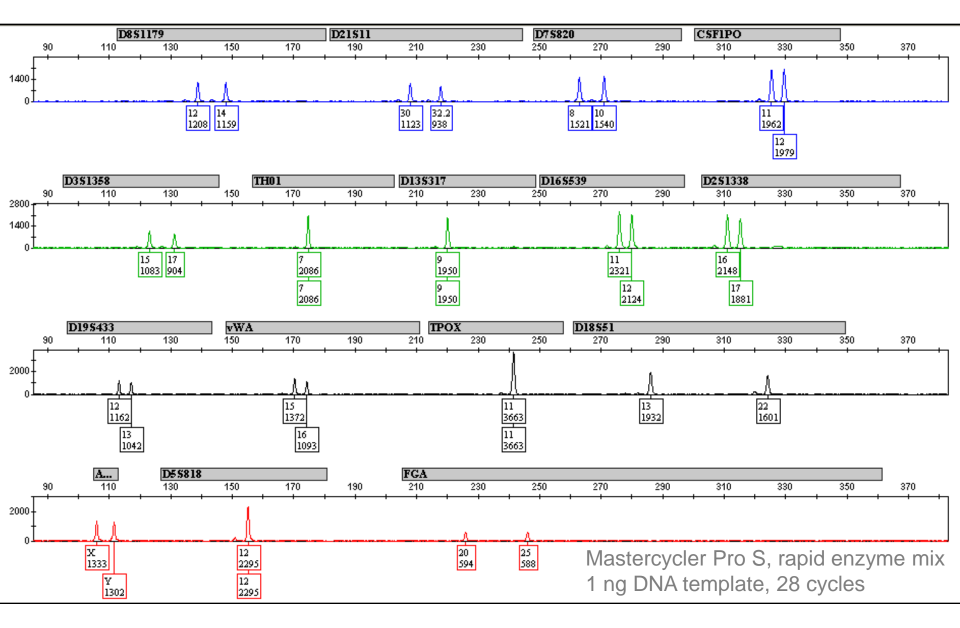


Maximum heating/cooling rate of ~2 to 6°C/s (cycler dependent)

Rapid PCR Conditions

- 1 X Takara PCR mastermix, 1 U SpeedStar polymerase
 Premix Ex Taq[™] (Perfect Real Time)
- 10 μ L total reaction in a thin walled tube (8-strip)
- 2 μ L of Identifiler PCR primer mix
- ~1 ng of template DNA
- Utilize maximum ramp rate on thermal cyclers
 - GeneAmp 9700 = 1.6°C/s (36 min)
 - Rotor-Gene Q = 1.6°C/s (36 min) Effective heating/cooling rates
 - SmartCycler = 5.8°C/s (20 min)
 - Mastercycler Pro S = 6.8° C/s (19 min)

Full Identifiler STR Profile with 19 min PCR



Potential Applications with Rapid PCR Capabilities

- Improve overall laboratory throughput
 - Multiplex PCR amplification is already in many situations the longest part of the DNA analysis process (depending on DNA extraction and DNA quantitation methods)
 - With increased use of robotic sample preparation and expert system data analysis, bottleneck for sample processing will shift to time for PCR amplification...
- Enable new potential DNA biometric applications (because the overall DNA analysis process is faster)
 - Permit analysis of individuals at a point of interest such as an embassy, an airport, or a country border

A "Crystal Ball" to the Future?





Compromised Sample Improvements

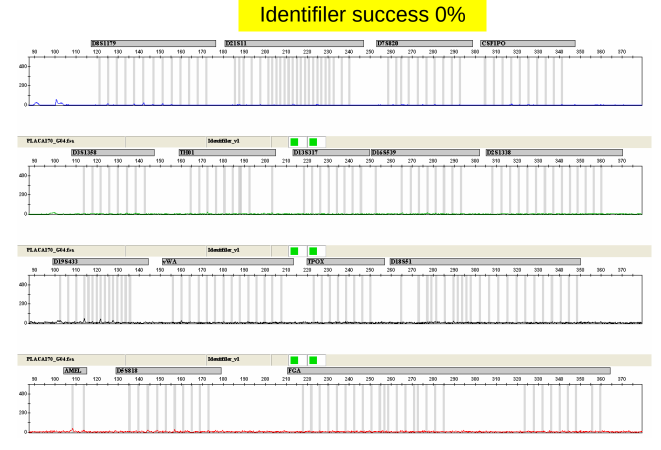
- Better DNA extraction/recovery
- Continued use of miniSTRs
 - to improve success rates for recovery of information from compromised DNA evidence
- Replicate results for reproducibility
 - to improve reliability with low-template DNA testing



Highly degraded DNA

SNP genotyping in an extreme degradation case Corpse half buried in a forest for ten years

- Uncovered by a forest fire
- Calcinated remains



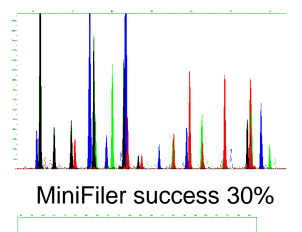


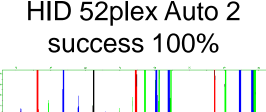
Highly degraded DNA

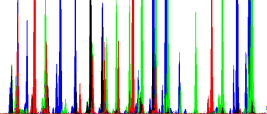
SNP genotyping in extreme degradation case Corpse half buried in a forest for ten years

- Uncovered by a forest fire
- Calcinated remains

HID 52plex Auto 1: success 100%









+SNPs

P: - 99.993

STRs

Highly degraded DNA

Carlos Vullo's group from Argentina has published similar results with both SNPforID 52plex and IPATIMUP 38plex InDel reaction on common graves from Argentinian dictatorship period

Forensic Science International: Genetics 6 (2012) 469-476



Typing short amplicon binary polymorphisms: Supplementary SNP and Indel genetic information in the analysis of highly degraded skeletal remains

C. Romanini^a, M.L. Catelli^a, A. Borosky^b, R. Pereira^d, M. Romero^a, M. Salado Puerto^c, C. Phillips^e, M. Fondevila^e, A. Freire^e, C. Santos^e, A. Carracedo^e, M.V. Lareu^e, L. Gusmao^d, C.M. Vullo^{a,b,*}

Geographical Origin Prediction

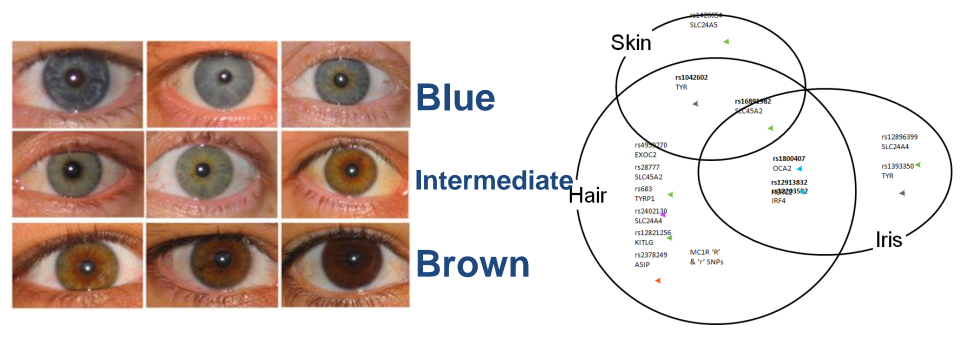


- Lao O, van Duijn K, et al. (2006) Proportioning whole-genome single-nucleotide-polymorphism diversity for the identification of geographic population structure and genetic ancestry. Am J Hum Genet 78: 680-90.
- Phillips, C., Salas, A., et al. (2007) Inferring ancestral origin using a single multiplex assay of ancestry-informative marker SNPs. FSI: Genetics 1: 273-280.
- Halder, I., Shriver, M., et al. (2008) A Panel of Ancestry Informative Markers for Estimating Individual Biogeographical Ancestry and Admixture From Four Continents: Utility and Applications. Hum Mut 29: 648-658.
- Pereira R., Phillips C., et al. (2012) Straightforward inference of ancestry and admixture proportions through ancestry-informative insertion deletion multiplexing. PLoS One;7(1):e29684.

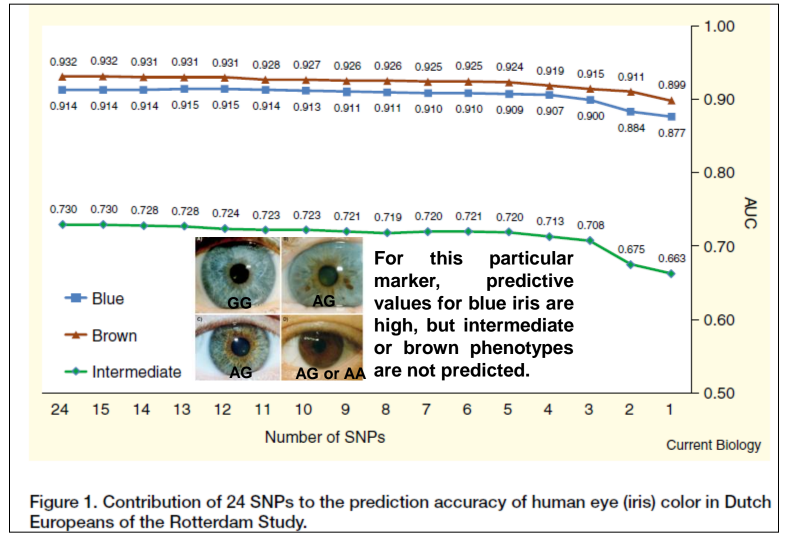
Phenotypic Trait Prediction

Traits of interest

- Traits whose variation may be classified on discreet categories.
- Regulated by a relatively low number of genes.
- Fine example: Iris and hair pigmentation.



Phenotypic trait prediction



Liu F., et al. (2009). Eye color and the prediction of complex phenotypes from genotypes, Curr. Biol. 19:R192–R193

Phenotypic trait prediction

- Currently several research groups are working on the prediction of phenotypical traits by SNP typing.
- Best predictions have been achieved on iris pigmentation.
- However the achieved predictive values are still different for each variant. Research is not yet completed.
- Branicki W, Kayser M et al. (2011). Model-based prediction of human hair color using DNA variants. *Human Genetics;* DOI 10.1007/s00439-010-0939-8.
- Walsh S., et al. (2010) IrisPlex: A sensitive DNA tool for accurate prediction of blue and brown eye colour in the absence of ancestry information. Forensic Sci. Int. Genet. (Epub)
- Kayser M., Schneider P.M. (2009) DNA-based prediction of human externally visible characteristics in forensics: motivations, scientific challenges, and ethical considerations. Forensic Sci. Int. Genet. 3(3):154-61.
- Ruiz Y., C. Phillips et al.(2012) Further development of forensic eye color predictive tests. Forensic Sci. Int. Genet. (accepted for publication).

Next Generation Sequencing

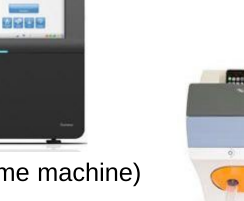
- High throughput or ultra-high throughput sequencing
- Thousands or millions of sequencing reads in parallel
- DNA sequencing, RNA expression, clinical diagnostics, microbial forensics, …
- 1. Library preparation (genomic DNA or PCR amplicons)
- 2. Sequencing
- 3. Data analysis (assembly of reads)

NGS Platforms

GS Junior Roche

- Roche
 - 454 FLX
 - 454 GS Junic
- PacificBio
 - Pacbio RS
- Illumina
 - GAIIx
 - HiSeq
 - HiScanSQ
 - MiSeq
- Life Tech
 - 5500 series
 - Ion torrent Proton
 - Ion torrent PGM (personal genome machine)









Nature Biotechnology 2012

Performance comparison of benchtop high-throughput sequencing platforms

Nicholas J Loman¹, Raju V Misra², Timothy J Dallman², Chrystala Constantinidou¹, Saheer E Gharbia², John Wain^{2,3} & Mark J Pallen¹

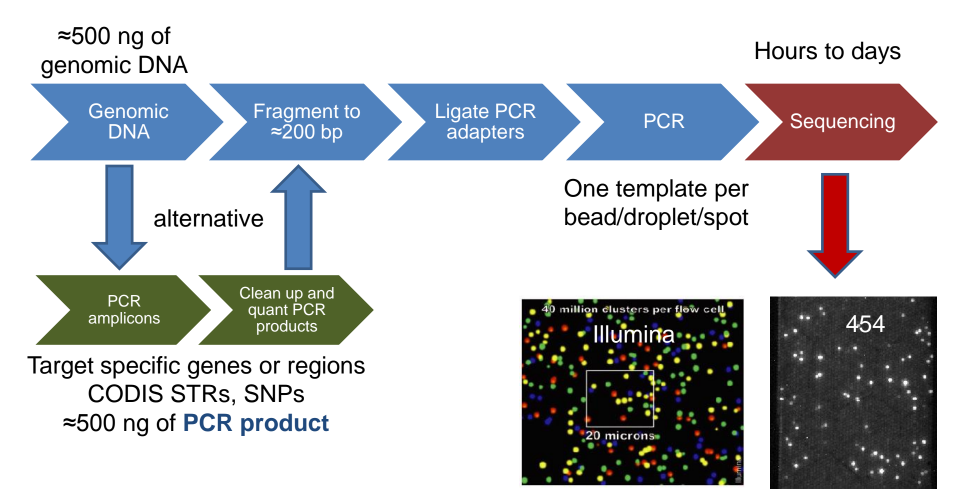
- 1. 454 GS Junior (Roche)
- 2. MiSeq (Illumina)
- 3. Ion Torrent PGM (Life tech)

Table 1 Price comparison of benchtop instruments and sequencing runs

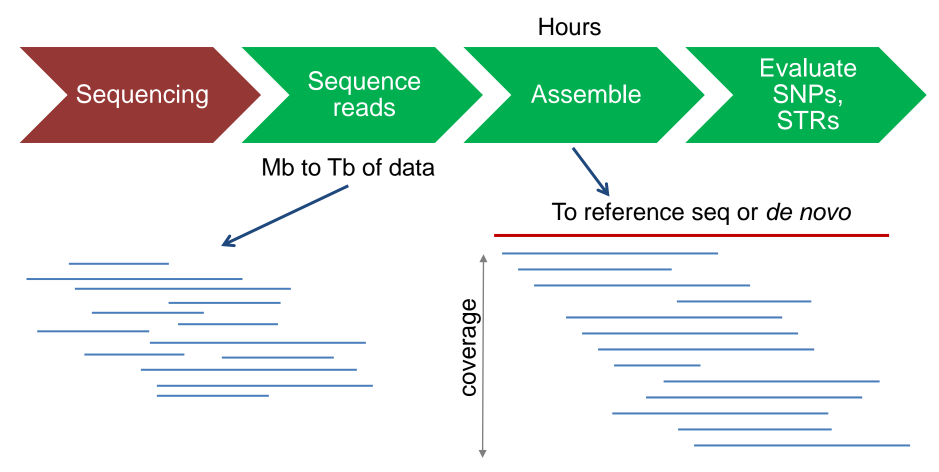
Platform	List price	Approximate cost per run	Minimum throughput (read length)	Run time	Cost/Mb	Mb/h
454 GS Junior Ion Torrent PGM	\$108,000	\$1,100	35 Mb (400 bases)	8 h	\$31	4.4
(314 chip)	\$80,490 ^{a,b}	\$225 ^c	10 Mb (100 bases)	3 h	\$22.5	3.3
(316 chip)		\$425	100 Mb ^d (100 bases)	3 h	\$4.25	33.3
(318 chip)		\$625	1,000 Mb (100 bases)	3 h	\$0.63	333.3
MiSeq	\$125,000	\$750	1,500 Mb (2 $ imes$ 150 bases)	27 h	\$0.5	55.5

Note pricing may vary between countries and/or sales territories. Instrument prices do not include service contracts. Sample prices do not include the cost of generating the initial fragmented genomic DNA library with adaptors (an additional cost of between \$75–200 depending on method used). Cost per megabase assumes one sample and one sample sequencing kit per run. Unless stated, pricing information is from the online supplement of ref. 3. ^aIon Torrent PGM pricing from Invitrogen US territory website (http://www.invitrogen.com/, accessed 21 February 2012). ^bPrice includes Ion Torrent PGM, server, OneTouch and OneTouch ES sample automation systems. ^cIon Torrent PGM prices include chip and sample preparation kit. ^dConfiguration used in this study.

Generalized NGS Workflow



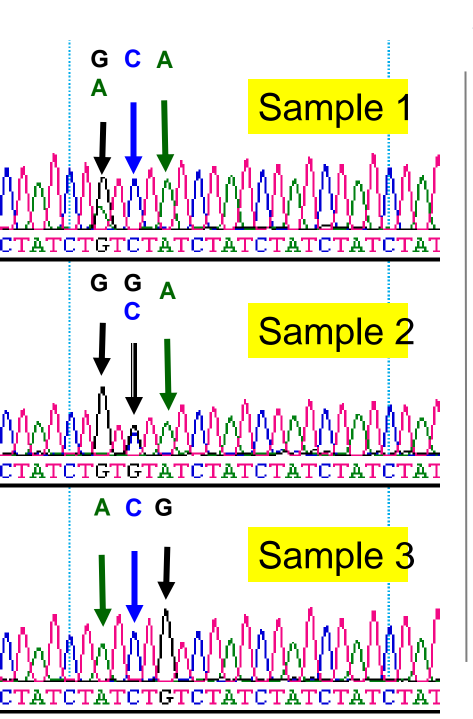
Generalized NGS Workflow



Next Generation Sequencing Forensic Applications

- Going in depth into STR loci and beyond
 - STRs are useful for legacy (databases)
 - SNPs within STRs identify 'sub-alleles'
 - Millions of bases of sequence variants (SNPs)
- Opens up new human identity applications: biogeographical ancestry, externally visible traits, complex kinship, degraded samples, mixtures, other applications

Applications are currently being addressed by the forensic genetics community (Kayser and deKnijff 2011)



SNPs within the D8S1179 repeat All 3 samples '13,13' [TCTA]₁₃

Allele A - $[TCTA]_{13}$ Allele B - TCTA TCT<u>G</u> $[TCTA]_{11}$

Allele B - TCTA TCTG [TCTA]₁₁

Allele C - TCTA TCTG TGTA [TCTA]₁₀

There are **4** different '13' alleles in these 3 samples.

Allele D - $[TCTA]_2 TCTG [TCTA]_{10}$ Allele D - $[TCTA]_2 TCTG [TCTA]_{10}$

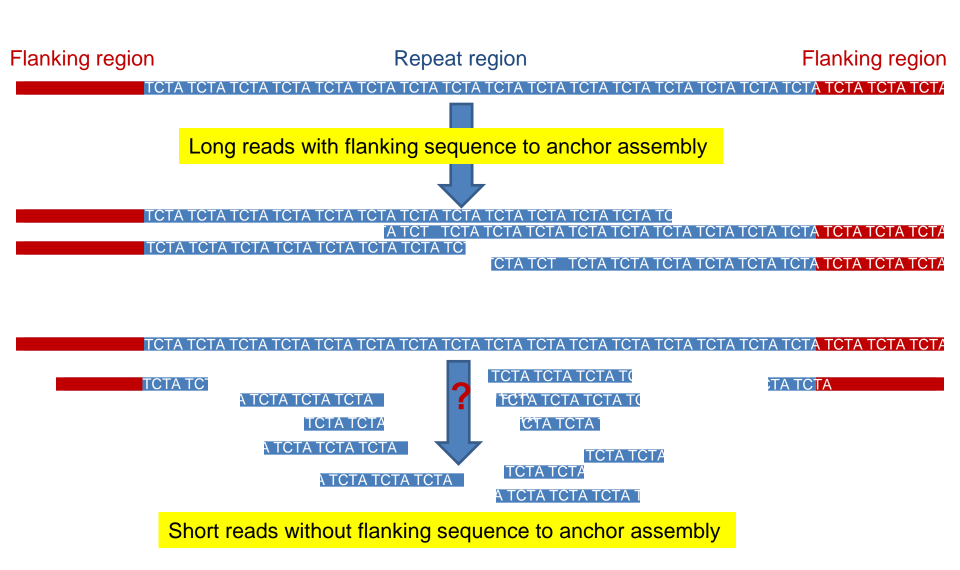
Data provided by Margaret Kline and Becky Hill

Specific issues with STRs

- Typically comprised of tetra nucleotide repeats
- Range 70 450+ bp regions
- Longer STRs can be difficult to assemble based on read length
- Illumina GAIIx (read length 150 bp)
 - Generated 1000-2500 bp amplicons (13 core loci)
 - Problems detecting D21S11 32.2 and 34.2 alleles
 - Issues detecting D18S51
 - Custom informatics tools for assembling STRs

Bornman et al., 2012 Biotechniques Rapid Dispatch: 1-6

Assembling STRs



Next Generation Sequencing

- Challenges
 - Repeating sequences (STRs) and read lengths
 - Sample amount requirements (10 ng to 5 µg)
 - Cost and time per unit of information
 - Data analysis (storage, assembly, interpretation)
 - Policy, privacy, disease related markers
 - Validation
 - Standards/reference materials
 - Nomenclature
 - Accuracy of sequence information
 - Errors, platform and bioinformatics-based bias

Next Generation Sequencing Workshop

- Interagency Workshop on the use of Next-Generation DNA Sequencing for Human Identification and Characterization (Jan 31 2012)
- Discussion of forensic applications of NGS (NIST, DoD, FBI, DHS) – materials can be found at:
 - <u>http://www.nist.gov/mml/biochemical/genetics/ngs_hid_workshop.cfm</u>
- We are in the process of looking at platforms to characterize forensic markers (mitochondrial, STRs, SNPs)
- Evaluate accuracy, reproducibility, identify initial requirements for a NGS forensic reference material

Improvements in Forensic DNA Analysis

- Biology
 - Improved DNA extraction with automation
 - New capabilities for recovery of information from degraded DNA samples (e.g., miniSTRs)
- Technology
 - Parallel processing of DNA with capillary arrays
 - Expert systems for automated data interpretation
- Genetics
 - Ethnicity estimations (with STRs and/or SNPs)
 - Larger Y-STR and mtDNA population databases

Effective Training is Needed in All Areas

Some Thoughts on the Future...

PCR amplification

- Faster enzymes to enable rapid PCR
- More robust enzymes and master mixes to overcome inhibition

Instrumentation

- More dye colors to aid higher levels of multiplexing
- Rapid, integrated devices
- Alternatives to capillary electrophoresis: PLEX-ID and NGS

Quantitative information

qPCR and digital PCR

Marker systems

- Expanding sets of STR loci for growing DNA databases
- Other marker systems: SNPs, InDels, X-STRs, RM Y-STRs
- Body fluid identification with mRNA, miRNA, and DNA methylation
- Phenotyping for external visible characteristics
- Challenges with potential whole genome information

Data interpretation

- Probabilistic genotyping for low-level DNA and mixture interpretation
- Probability of dropout

AAFS 2009 Topics Regarding Forensic DNA

From abstracts of presentations at AAFS meeting in Denver, CO (Feb 2009)

- Improved DNA extraction
- Predicting hair color and ancestry with SNPs
- X-chromosome STRs
- Familial searching
- Y-STRs and mixtures
- Low level DNA samples
- miniSTRs
- DNA screening assays
- Optimizing database labs
- Microfluidic biochip systems

- Use with property crimes
- Recovery from handguns
- DNA from IEDs
- Expert systems
- Automation with robotics
- DNA quantitation qPCR
- PCR directly from blood
- mtDNA
- RNA
- Non-human DNA (dogs & cows)
- Mixture interpretation

www.DNA.gov Website

ne Grant Funding Training Statistics Res	earch Publications State Profiles	Search
roviding funding, training and assistance) solve crimes, protect the innocent and	e to ensure that forensic DNA reaches its full potential identify missing persons.	Training Course Catalog
Highlights	Highlights	Register for online courses
About Forensic DNA	VIDEO	Login and view your courses
DNA Databases	NamUs Behind the Scenes: How It Works, Why It Matter (6 minutes)	Reset your password
Reducing the Backlog	TRAINING	Request your username
Solving Crimes	Advanced DNA Technologies and Forensic Automation	Browse by Audience
Identifying Persons and Victims	Analyst Training Program	Officers/ Investigators
Postconviction Testing	PUBLICATIONS <u>Research articles from NIJ-funded projects updated</u>	Forensic Scientists
Tools for Forensic Scientists	through 2008	Officers of the Court
	AVAILABLE FUNDING	Crime Lab Managers
Services for Laboratories	Forensic DNA Unit Efficiency Improvement	Onnie Lab Managers

Research on	Statistics on	Grant funding for	
Human DNA quantitation	Profiles in the database	Forensic backlog reduction	
Y Chromosome	DNA crime labs	Convicted offender/arrestee	
Compromised DNA ovidence	Backleg of complex	 backlog reduction 	

Summary of NIJ-Funded Research



The DNA Field Moves Forward...

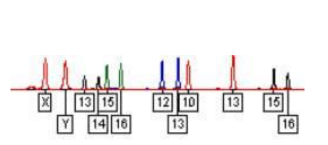
The Past

Sexual Assault Case

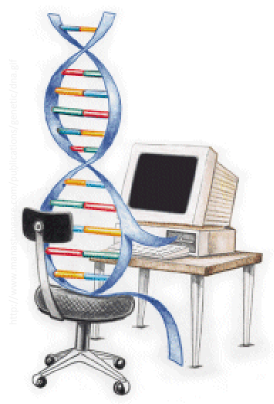
RFLP

The Present

The Future



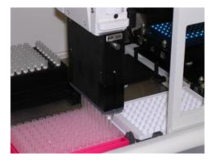
STRs





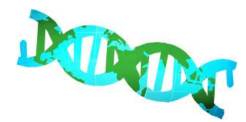
The Future

- More Robotics
- Expert Systems
- Animal & Plant DNA





- Physical Characteristics
- Ethnicity Estimation





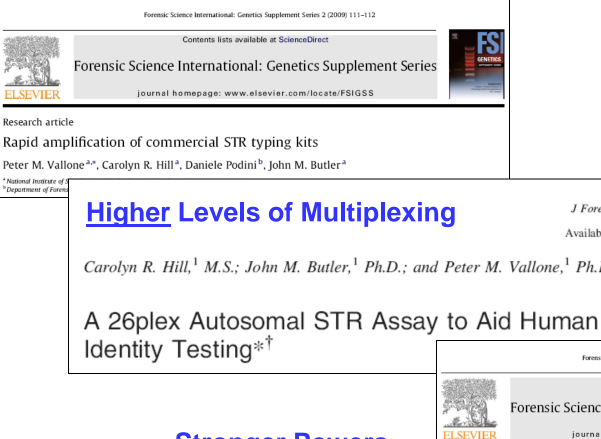
The Future of Forensic DNA

is Similar to the Olympic Motto of "Swifter, Higher, Stronger"



Recent NIST Publications Demonstrating "Swifter, Higher, Stronger" DNA Analysis

Swifter PCR Amplification

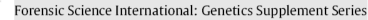


J Forensic Sci, September 2009, Vol. 54, No. 5 doi: 10.1111/j.1556-4029.2009.01110.x Available online at: www.blackwell-synergy.com

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

Forensic Science International: Genetics Supplement Series 2 (2009) 23-24

Contents lists available at ScienceDirect





journal homepage: www.elsevier.com/locate/FSIGSS

Research article

The single most polymorphic STR Locus: SE33 performance in U.S. populations

John M. Butler^{a,*}, Carolyn R. Hill^a, Margaret C. Kline^a, David L. Duewer^a, Cynthia J. Sprecher^b, Robert S. McLaren^b, Dawn R. Rabbach^b, Benjamin E. Krenke^b, Douglas R. Storts^b

^a National Institute of Standards and Technology, Gaithersburg, MD 20899-8312, USA ^b Promega Corporation, Madison, WI 53711, USA



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- Slides from Pete Vallone (NIST) and Manuel Fondevila (NIST, USC)
- Funding from National Institute of Justice and FBI Biometrics Center of Excellence for work performed within the NIST Applied Genetics Group

Thank you for your attention

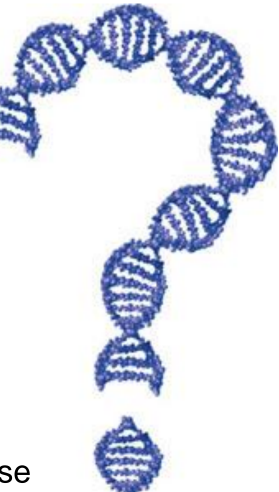
Acknowledgments: A great team of scientists within our NIST Applied Genetics Group and funding from the National Institute of Justice and the FBI

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



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