


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


NIST
 National Institute of Standards and Technology

DNA Mixture Interpretation & Statistical Analysis

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



Steps Involved in Process of Forensic DNA Typing

1) Data Interpretation
2) Statistical Interpretation

Gathering the Data **Understanding the Data**

Collection/Storage/Characterization

Extraction/Quantitation

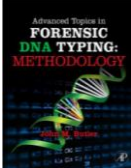
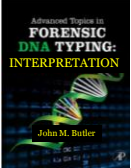
Amplification/Marker Sets

Separation/Detection

Interpretation

Report

Advanced Topics: Methodology *Advanced Topics: Interpretation*

SWGAM Website and Resources Available

<http://www.swgdam.org/resources.html>



- Home
- ByLaws
- Members
- Committees
- Meetings
- Publications

Link to <http://www.cstl.nist.gov/biotech/strbase/mixture/SWGAM-mixture-info.htm>



Additional Resources

Beginning with the development and revision of its next draft guidance document(s), SWGAM will make a "Draft for Comment" or other work product available for the purpose of receiving comments from the general public. This "Draft for Comment" solicitation will be open for a minimum of 45 days, visible through SWGAM.org. SWGAM will make all reasonable efforts to advise the forensic DNA community of the open comment period for a proposed guidance document or standard, guideline, best practice, study, or other recommendation and/or finding via as many avenues as possible to include posting notices through discipline-specific and related professional organizations. SWGAM strongly encourages all interested parties to regularly monitor SWGAM.org for the posting of such draft documents as well as public comments received by SWGAM will forwarded to the appropriate SWGAM Committee for review and consideration as a part of its formal business practice for the development of the guidance documents or other work product.

The following information resources have been produced and reviewed by members of the Mixture Committee of SWGAM and are available at <http://www.cstl.nist.gov/biotech/strbase/mixture/SWGAM-mixture-info.htm>

Mixture Training Materials

Reviewed by SWGAM Mixture Committee

SWGAM Mixture Committee Resource Page

The following information resources have been produced and reviewed by members of the Mixture Committee of the Scientific Working Group on DNA Analysis Methods (SWGAM) -- see <http://www.swgdam.org/resources.html> for additional information.


Mixture Training Examples

- Download "[Mixture 6](#)" PowerPoint show (56 Mb)
 - with voice-over by Bruce Heidebrecht (Maryland State Police), may work best if file is first saved to your computer
- Download "[Mixture IQAS2904](#)" PowerPoint show (35 Mb)
 - with voice-over by Bruce Heidebrecht (Maryland State Police), may work best if file is first saved to your computer

<http://www.cstl.nist.gov/biotech/strbase/mixture/SWGAM-mixture-info.htm>

Mixture Workshop (Promega ISHI 2010)

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



October 11, 2010

Regional workshops presented in FL, TX, MI, and AZ (April - June 2011)
 Updated mixture workshop presented at ISHI 2011

Handout >200 pages
Literature list of >100 articles

13 Modules Presented

Introduction to SWGAM (John)

Stochastic effects (Robin)

Peak height ratios (Charlotte)

Number of contributors (John)

Mixture ratios (John)

Mixture principles (Charlotte)

Statistics (Mike)

Case Example 1 (Robin)


Case Example 2 (Charlotte)

Case Example 3 (John)

NIJ Grant to Boston University funded -150 state & local lab analysts to attend

Catherine Grgicak Boston U.
Mike Coble NIST
Robin Cotton Boston U.
John Butler NIST
Charlotte Word Consultant

Promega ISHI 2012 Mixture Workshop



Forensics Amplified
 Nashville, TN • Oct. 15-18, 2012

- **John Butler**, Ph.D., NIST, Gaithersburg, MD
- **Michael Coble**, Ph.D., NIST, Gaithersburg, MD
- **Robin Cotton**, Ph.D., Boston University, Boston, MA
- **Catherine Grgicak**, Ph.D., Boston University, Boston, MA
- **Charlotte J. Word**, Ph.D., Gaithersburg, MD

This workshop is for analysts, technical reviewers and technical leaders performing and interpreting validation studies and/or interpreting and reviewing STR data, particularly more difficult mixtures. Various DNA profiles will be analyzed and interpreted using selected analytical thresholds and stochastic thresholds to demonstrate the impact of those values on the profiles amplified with low-template DNA vs. higher amounts of DNA. Different statistical approaches and conclusions suitable for the profiles will be presented.

Useful Articles on DNA Mixture Interpretation

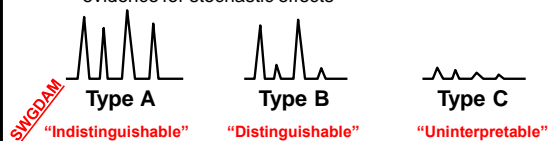
- Buckleton, J.S. and Curran, J.M. (2008) A discussion of the merits of random man not excluded and likelihood ratios. *Forensic Sci. Int. Genet.* 2: 343-348.
- Budowle, B., et al. (2009) Mixture interpretation: defining the relevant features for guidelines for the assessment of mixed DNA profiles in forensic casework. *J. Forensic Sci.* 54: 810-821.
- Clayton, T.M., et al. (1998) Analysis and interpretation of mixed forensic stains using DNA STR profiling. *Forensic Sci. Int.* 91: 55-70.
- Gill, P., et al. (2006) DNA commission of the International Society of Forensic Genetics: Recommendations on the interpretation of mixtures. *Forensic Sci. Int.* 160: 90-101.
- Gill, P., et al. (2008) National recommendations of the technical UK DNA working group on mixture interpretation for the NDNAD and for court going purposes. *FSI Genetics* 2(1): 76-82.
- Schneider, P.M., et al. (2009) The German Stain Commission: recommendations for the interpretation of mixed stains. *Int. J. Legal Med.* 123: 1-5.

German Mixture Classification Scheme

Schneider et al. (2009) *Int. J. Legal Med.* 123: 1-5

(German Stain Commission, 2006):

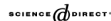
- Type A:** no obvious major contributor, no evidence of stochastic effects
- Type B:** clearly distinguishable major and minor contributors; consistent peak height ratios of **approximately 4:1** (major to minor component) for all heterozygous systems, no stochastic effects
- Type C:** mixtures without major contributor(s), evidence for stochastic effects



Available for download from the ISFG Website:
<http://www.isfg.org/Publication;Gill2006>



Available online at www.sciencedirect.com



Forensic Science International 160 (2006) 90-101



DNA commission of the International Society of Forensic Genetics:
 Recommendations on the interpretation of mixtures

P. Gill^{a,*}, C.H. Brenner^b, J.S. Buckleton^c, A. Carracedo^d, M. Krawczak^e, W.R. Mayr^f,
 N. Morling^g, M. Prinz^h, P.M. Schneiderⁱ, B.S. Weir^j

^a Forensic Science Service, Tidlex Court, 2960 Southall Parkway, Birmingham, UK
^b Forensic Science Group, School of Public Health, University of California, Berkeley, CA 94720-7391, USA
^c ESR, Private Bag 92021, Auckland, New Zealand

Our discussions have highlighted a significant need for continuing education and research into this area.

ⁱ University of Washington, Department of Biostatistics, Box 357322, Seattle, WA 98195, USA
 Received 4 April 2006; accepted 10 April 2006
 Available online 5 June 2006

Gill et al. (2006) DNA Commission of the International Society of Forensic Genetics:
 Recommendations on the interpretation of mixtures. *Forensic Sci. Int.* 160: 90-101



ISFG Recommendations on Mixture Interpretation

<http://www.isfg.org/Publication;Gill2006>

- The likelihood ratio (LR) is the preferred statistical method for mixtures over RMNE
- Scientists should be trained in and use LRs
- Methods to calculate LRs of mixtures are cited
- Follow Clayton et al. (1998) guidelines when deducing component genotypes
- Prosecution determines H₀ and defense determines H₁ and multiple propositions may be evaluated
- When minor alleles are the same size as stutters of major alleles, then they are indistinguishable
- Allele dropout to explain evidence can only be used with low signal data
- No statistical interpretation should be performed on alleles below threshold
- Stochastic effects limit usefulness of heterozygote balance and mixture proportion estimates with low level DNA

Gill et al. (2006) DNA Commission of the International Society of Forensic Genetics:
 Recommendations on the interpretation of mixtures. *Forensic Sci. Int.* 160: 90-101



Forensic Science International
 91 (1998) 55-70



Analysis and interpretation of mixed forensic stains
 using DNA STR profiling

T.M. Clayton^{a,*}, J.P. Whitaker^a, R. Sparkes^b, P. Gill^b

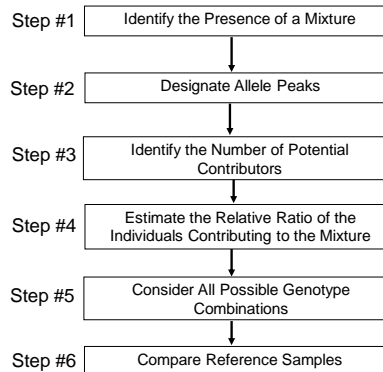
^a Forensic Science Service, Wetherby Laboratory, Sandbeck Way, Audby Lane, Wetherby, West Yorkshire LS22 4DN, UK

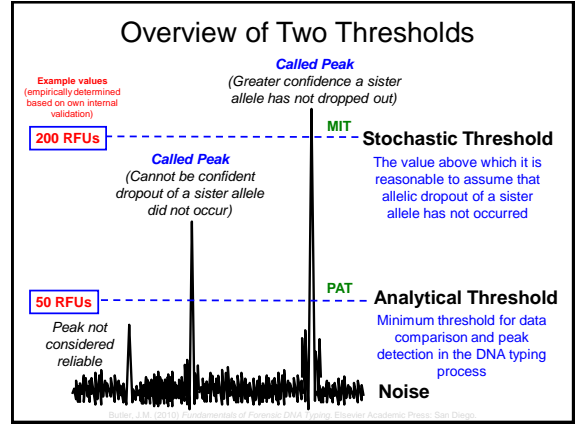
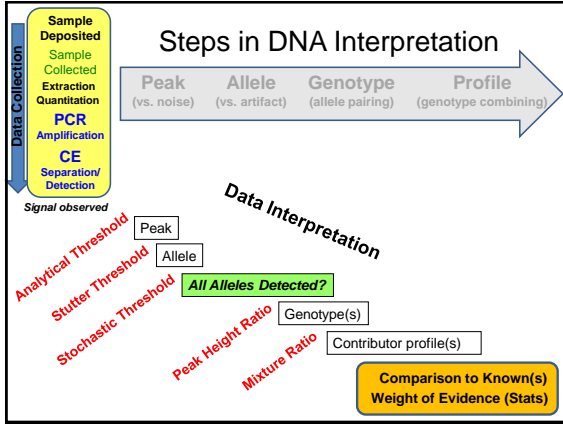
^b Forensic Science Service, Priory House, Gooch Street North, Birmingham B56QD, UK

Received 13 May 1997; received in revised form 9 October 1997; accepted 27 October 1997

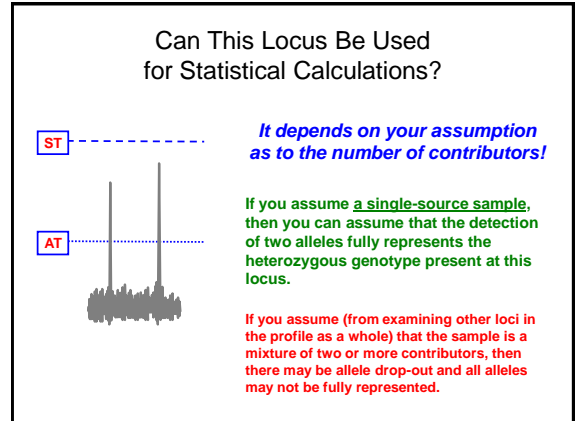
Steps in the interpretation of mixtures

(Clayton et al. *Forensic Sci. Int.* 1998; 91:55-70)

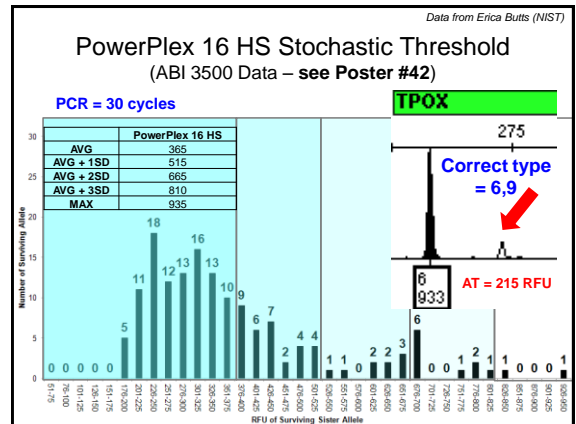




- ### Coupling of Statistics and Interpretation
- The CPE/CPI approach** for reporting an inclusionary statistic **requires that all alleles be observed** in the evidence sample
 - If allele drop-out is suspected at a locus, then any allele is possible and the probability of inclusion goes to 100% -- in other words, the locus is effectively dropped from consideration
 - If alleles are seen below the established stochastic threshold, then the locus is typically eliminated ("INC" – declared inconclusive) in many current lab SOPs



- ### Limitations of Stochastic Thresholds
- The possibility of allele sharing with a complex mixture containing many contributors may make a stochastic threshold meaningless
 - "Enhanced interrogation techniques" to increase sensitivity (e.g., increased PCR cycles) may yield false homozygotes with >1000 RFU
 - New turbo-charged kits with higher sensitivity will need to be carefully evaluated to avoid allele drop-out and false homozygotes**



Stochastic Threshold Summary

- A stochastic threshold (ST) may be established for a specific set of conditions to reflect possibility of allele drop-out, which is essential for a CPE/CPI stats approach
- ST should be re-examined with different conditions (e.g., higher injection, sample desalting, increase in PCR cycles)
- ST will be dependent on the analytical threshold set with a method and impacts the lowest expected peak height ratio
- Assumptions of the number of contributors is key to correct application of ST

Stats Required for Inclusions

SWGAM Interpretation Guideline 4.1:

“The laboratory must perform statistical analysis in support of any inclusion that is determined to be relevant in the context of a case, irrespective of the number of alleles detected and the quantitative value of the statistical analysis.”

Buckleton & Curran (2008): “There is a considerable aura to DNA evidence. Because of this aura **it is vital that weak evidence is correctly represented as weak or not presented at all.**”

Buckleton, J. and Curran, J. (2008) A discussion of the merits of random man not excluded and likelihood ratios. *Forensic Sci. Int. Genet.* 2: 343-348.

DAB Recommendations on Statistics

February 23, 2000
Forensic Sci. Comm. 2(3); available on-line at
<http://www.fbi.gov/hq/lab/fsc/backissu/july2000/dnastat.htm>

“The DAB finds either one or both PE or LR calculations acceptable and strongly recommends that one or both calculations be carried out whenever feasible and a mixture is indicated”

- Probability of exclusion (PE)
 - Devlin, B. (1993) Forensic inference from genetic markers. *Statistical Methods in Medical Research* 2: 241–262.
- Likelihood ratios (LR)
 - Evett, I. W. and Weir, B. S. (1998) *Interpreting DNA Evidence*. Sinauer, Sunderland, Massachusetts.

CPE/CPI (RMNE) Limitations

- A CPE/CPI approach assumes that all alleles are present (i.e., cannot handle allele drop-out)
- Thus, statistical analysis of low-level DNA CANNOT be correctly performed with a CPE/CPI approach because some alleles may be missing
- Charles Brenner in his AAFS 2011 talk addressed this issue
- Research is on-going to develop allele drop-out models and software to enable appropriate calculations

Notes from Charles Brenner's AAFS 2011 talk

The Mythical “Exclusion” Method for Analyzing DNA Mixtures – Does it Make Any Sense at All?

1. The claim that it requires **no assumption about number of contributors** is mostly wrong.
 2. The supposed **ease of understanding** by judge or jury is really an illusion.
 3. **Ease of use** is claimed to be an advantage particularly for complicated mixture profiles, those with many peaks of varying heights. The truth is the exact opposite. **The exclusion method is completely invalid for complicated mixtures.**
 4. The exclusion method is only **conservative** for guilty suspects.
- “Certainly no one has laid out an explicit and rigorous chain of reasoning from first principles to support the exclusion method. It is at best guesswork.”

Brenner, C.H. (2011). The mythical “exclusion” method for analyzing DNA mixtures – does it make any sense at all? *Proceedings of the American Academy of Forensic Sciences*, Feb 2011, Volume 17, p. 79

Statistical Methods in Medical Research 1993; 2: 241–262

Forensic inference from genetic markers

B Devlin Department of Epidemiology and Public Health, Yale University School of Medicine

Section 5.1 Exclusion probability

- Discussion about exclusion probabilities in **Paternity** cases.

Two types:

(1) Conditional Exclusion Probability - excluding a random man as a possible father, given the mother-child genotypes for a particular case.

(2) Average Exclusion Probability – excluding a random man as a possible father, given a randomly chosen mother-child pair.

Statistical Methods in Medical Research 1993; 2: 241-262

Forensic inference from genetic markers

B Devlin Department of Epidemiology and Public Health, Yale University School of Medicine

Section 5.1 Exclusion probability

“The theoretical concept of exclusion probabilities, however, makes no sense within the framework of normal mixture models.”

“The interpretation of conditional exclusion probability is obvious, which accounts for its value in the legal arena. Unlike [LR], however, it is not fully efficient.”

Curran and Buckleton (2010)

JOURNAL OF FORENSIC SCIENCES



J Forensic Sci, September 2010, Vol. 55, No. 5
doi: 10.1111/j.1556-4029.2010.01446.x
Available online at: interencore.wiley.com

PAPER

CRIMINALISTICS; GENERAL

James M. Curran,¹ M.Sc.(Hons.), Ph.D. and John Buckleton,² Ph.D.

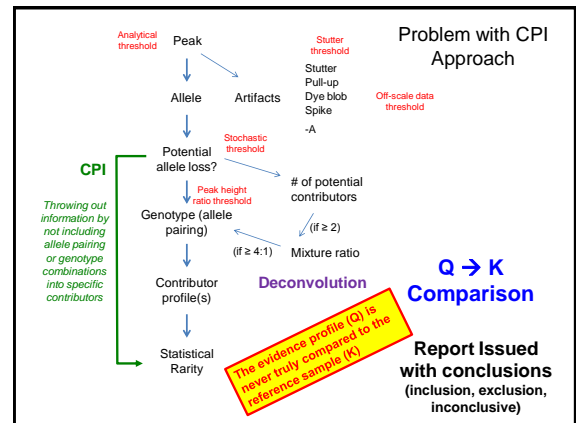
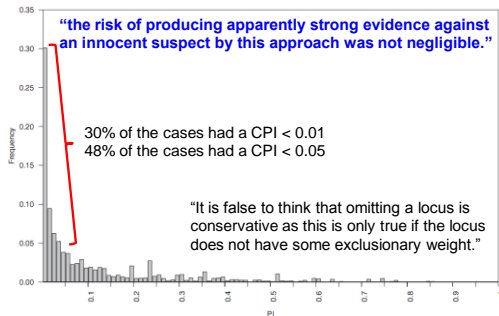
Inclusion Probabilities and Dropout

Created 1000 Two-person Mixtures (Budowle *et al.* 1999 AfAm freq.).

Created 10,000 “third person” genotypes.

Compared “third person” to mixture data, calculated PI for included loci, ignored discordant alleles.

Curran and Buckleton (2010)



Impact of Dropping Loci

- The less data available for comparison purposes, the greater the chance of falsely including someone who is truly innocent
- Are you then being “conservative” (i.e., erring in favor of the defendant)?

Likelihood Ratio (LR)

- Provides ability to express and evaluate both the prosecution hypothesis, H_p (the suspect is the perpetrator) and the defense hypothesis, H_d (an unknown individual with a matching profile is the perpetrator)

$$LR = \frac{H_p}{H_d}$$

- The numerator, H_p , is usually 1 – since in theory the prosecution would only prosecute the suspect if they are 100% certain he/she is the perpetrator
- The denominator, H_d , is typically the profile frequency in a particular population (based on individual allele frequencies and assuming HWE) – i.e., the random match probability

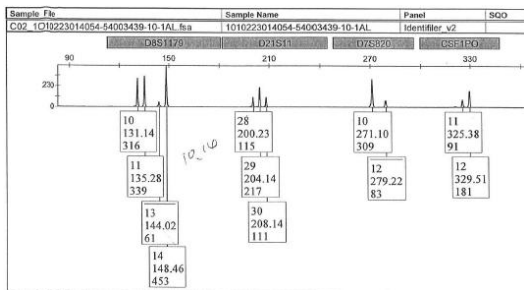
Some Important Points

- Inclusionary statements (including “cannot exclude”) need statistical support to reflect the relevant weight-of-evidence
- Stochastic thresholds are necessary if using CPI statistics to help identify possible allele dropout
- CPI is only conservative for guilty suspects as this approach does a poor job of excluding the innocent
- Uncertainty exists in scientific measurements – this fact needs to be conveyed with the statistical results
- An increasing number of poor samples are being submitted to labs – labs may benefit from developing a complexity threshold

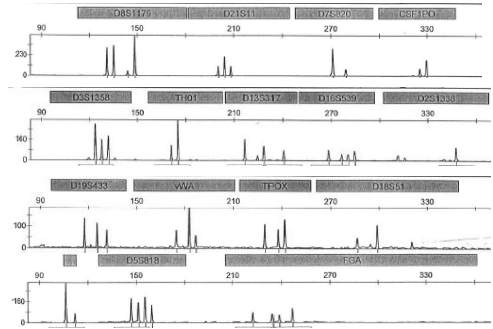
Some Mixture Examples Were Provided

- **Case 1**
 - Evidence (sexual assault victim’s underwear bra)
 - Victim
 - Suspect
- **Case 2**
 - Evidence (sexual assault victim’s panties)
- **Case 3**
 - Evidence (burglary cigarette)

Case 1 sexual assault victim’s underwear (bra)

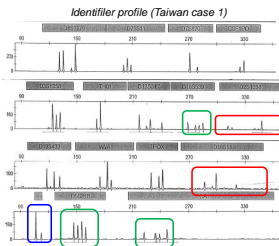


Taiwan Case 1 Evidence: Full Profile (Identifier)

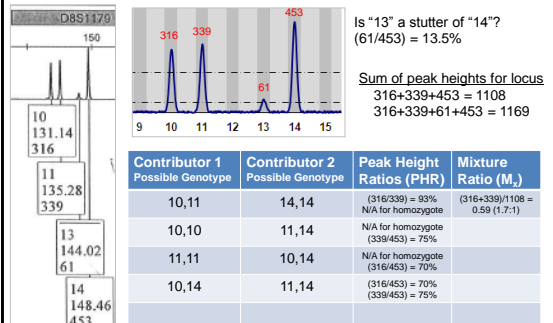


Observations from this Evidence Profile

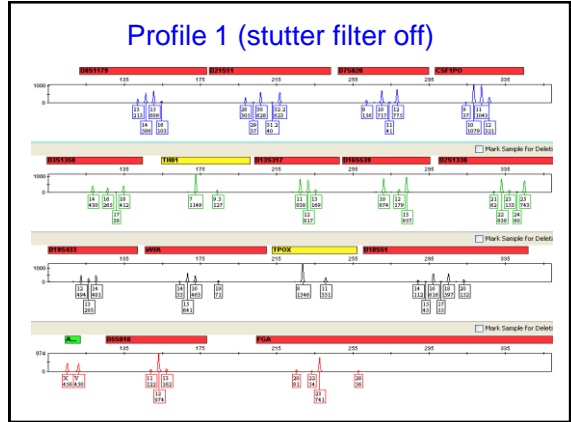
- **The sample is a mixture** since there are >2 peaks at multiple loci (at least 2 contributors)
- **Two contributors is a reasonable assumption** since there are no more than four alleles at a single locus
- **Male and female DNA are present** based on amelogenin X/Y ratio
- **A major contributor is not easily discernible** so component deconvolution is not an option
- Results at 4-allele loci (D5S818, FGA, and D16S539) suggest ≈1:1 mixture ratio
- **Overall RFU signals are low** especially for larger loci D2S1338 and D18S51 so allele drop-out is a possibility



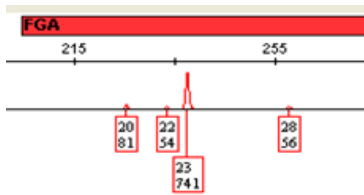
Case 1 Evidence: D8S1179



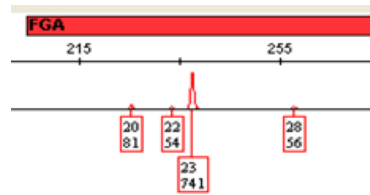
SLIDES NOT COMPLETED YET on PROVIDED MIXTURE EXAMPLES



Analytical Threshold (Peaks vs. Noise)

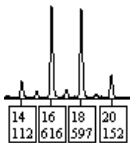


Stutter Threshold (Alleles vs. Artifacts)



Assumptions based upon # of contributors

Determination of Genotypes (PHR)

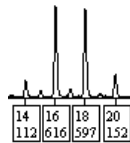


D18S51

Possible Combinations

- ~~14, 16 and 18, 20 (18%) (25%)~~
- ~~14, 18 and 16, 20 (10%) (25%)~~
- 14, 20 and 16, 18 (74%) (97%)

Determination of Mixture Ratio



Major: 16,18
Minor: 14,20

D18S51

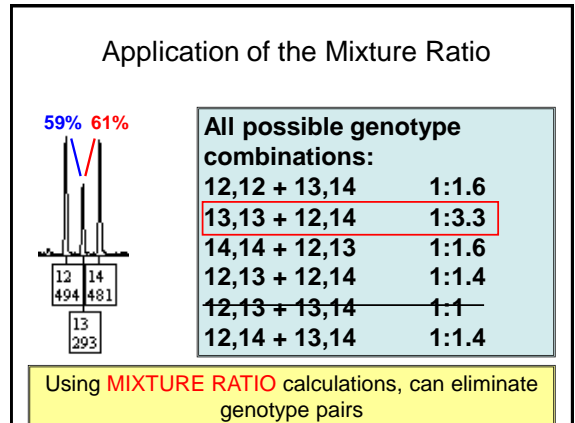
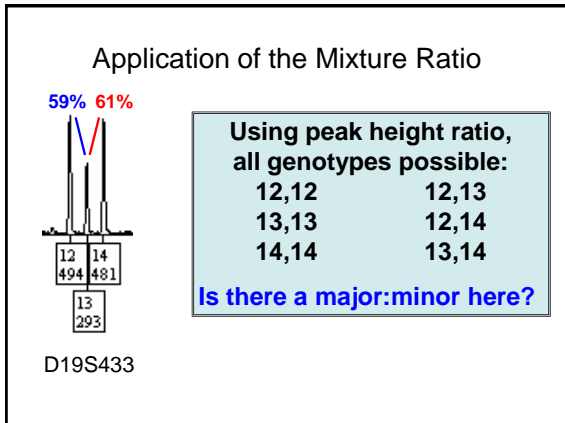
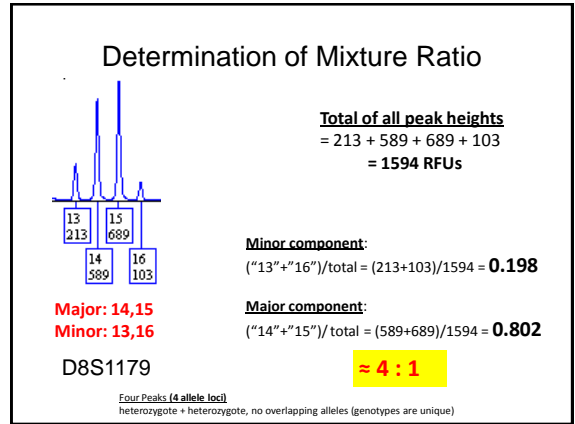
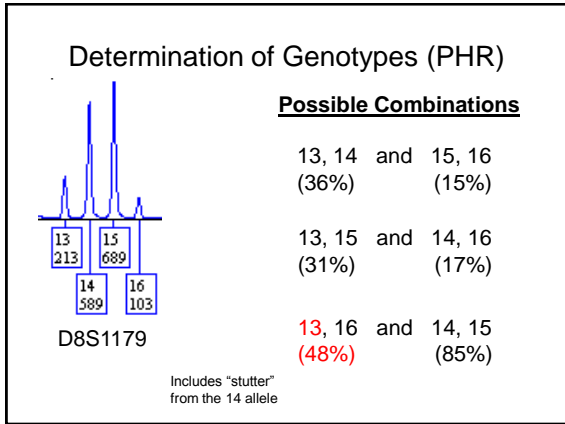
Total of all peak heights
= 112 + 616 + 597 + 152
= 1477 RFUs

Minor component:
("14"+"20")/total = (112+152)/1477 = **0.179**

Major component:
("16"+"18")/total = (616+597)/1477 = **0.821**

≈ 4.6 : 1

Four Peaks (4 allele loci)
heterozygote + heterozygote, no overlapping alleles (genotypes are unique)



Statistical Approaches with Mixtures

See Ladd et al. (2001) Croat Med J. 42:244-246

"Exclusionary" Approach	"Inferred Genotype" Approach
Random Man Not Excluded (RMNE)	Random Match Probability (RMP)
Combined Prob. of Inclusion (CPI)	Likelihood Ratio (LR)
Combined Prob. of Exclusion (CPE)	

Forensic Science International: Genetics 2 (2008) 343-348

A discussion of the merits of random man not excluded and likelihood ratios

John Buckleton^{a,*}, James Curran^b

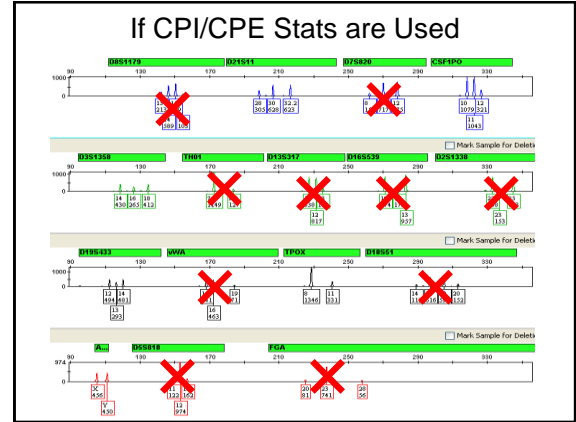
^aESR, PB 92021, Auckland, New Zealand
^bDepartment of Statistics, University of Auckland, PB 92019, Auckland, New Zealand
Received 15 January 2008; received in revised form 29 April 2008; accepted 1 May 2008

We conclude that the two matters that appear to have real force are:

- (1) LR's are more difficult to present in court and
- (2) the RMNE statistic wastes information that should be utilised.

If CPI/CPE Stats are Used

Since exclusionary statistics cannot adjust for the possibility of dropout, and does not take the number of contributors into account, any loci where alleles are below stochastic levels cannot be used in the CPI statistic.



If CPI/CPE Stats are Used

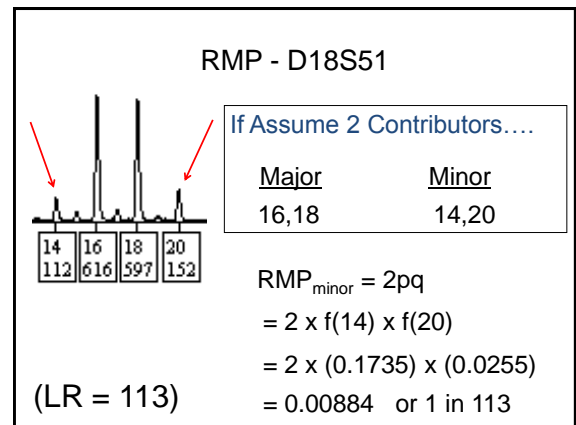
<u>Can use</u>	<u>Cannot use</u>	
D21	D8	D2
CSF	D7	vWA
D3	TH01	D18
D19	D13	D5
TPOX	D16	FGA

If CPI/CPE Stats are Used

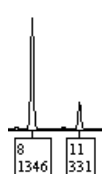
- CPI statistics using FBI Caucasian Frequencies
- 1 in 71 Caucasians included
- 98.59% Caucasians excluded

If RMP/LR Stats are Used

- Since there is an assumption to the number of contributors, it is possible to use data that falls below the ST.



RMP - TPOX

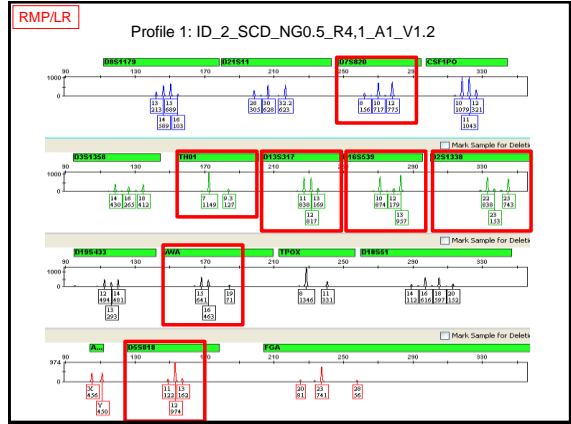


If Assume 2 Contributors....

<u>Major</u>	<u>Minor</u>
8,8	11,8 <i>OR</i> 11,11

RMP = 8,11 + 11,11
 RMP = 2pq + (q² + q(1-q)θ)

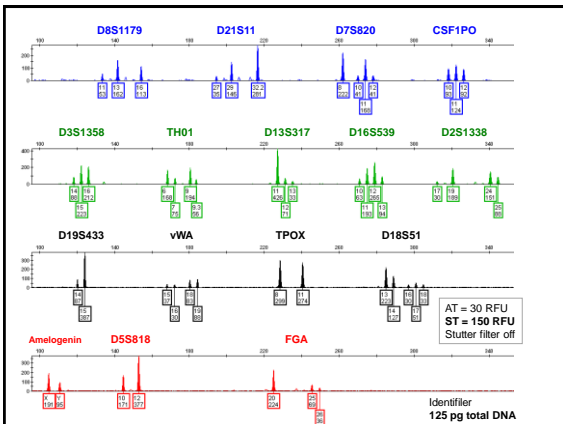
RMP = 2(0.5443)(0.2537) + (0.2537)² + (0.2537)(0.7463)(0.01)
 = 0.3424 or 1 in 2.9



If RMP/LR Stats are Used

<u>Can use</u>	<u>Loci with potential D-out</u>	
D8	D7	D2
D21	TH01	vWA
D18	D13	D5
D3	D16	
D19		
TPOX		
FGA		
CSF		

Challenges with low level, complex mixtures

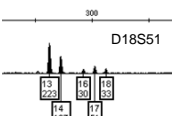


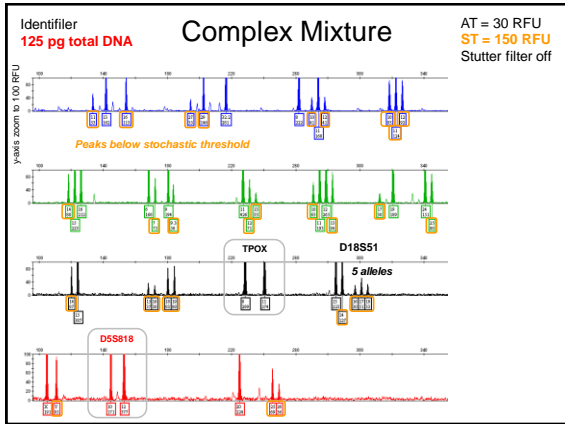
Impact of Results with Low Level DNA

Clayton et al. (1998)
ISFG (2006) Rec. #4

- Step #1** Identify the Presence of a Mixture
- Step #2** Designate Allele Peaks
- Step #3** Identify the Number of Potential Contributors
- Step #4** Estimate the Relative Ratio of Contributors
- Step #5** Consider All Possible Genotype Combinations
- Step #6** Compare Reference Samples

When amplifying low amounts of DNA (e.g., 125 pg), allele dropout is a likely possibility leading to **higher uncertainty** in the potential number of contributors and in the possible genotype combinations





What Can We Say about this Result?

- Low level DNA (only amplified 125 pg total DNA)
 - likely to exhibit stochastic effects and have allele dropout
- Mixture of at least 3 contributors
 - Based on detection of 5 alleles at D18S51
 - If at equal amounts, ~40 pg of each contributor (if not equal, then less for the minor contributors); **we expect allele dropout**
- At least one of the contributors is male
 - Based on presence of Y allele at amelogenin
- Statistics if using CPI/CPE
 - Would appear that we can only use TPOX and D5S818 results with a stochastic threshold of 150 RFU (*will explore this further*)
- **Due to potential of excessive allele dropout, we are unable to perform any meaningful Q-K comparisons**

Uncertainty in the Potential Number of Contributors with this Result

D18S51

300

13 223 16 30 18 33
14 127 17 51

5 alleles observed

- Several of the peaks are barely above the analytical threshold of 30 RFU
 - In fact, with an analytical threshold of 50 RFU or even 35 RFU, there would only be three detected alleles at D18S51
- Stochastic effects could result in a high degree of stutter off of the 17 allele making alleles 16 and 18 potential stutter products
- No other loci have >4 alleles detected

All Detected Alleles Are Above the Stochastic Threshold – Or Are They?

TPOX

220

8 298 11 274

Stochastic threshold = 150 RFU

Does this result guarantee no allele drop-out?

We have assumed three contributors. If result is from an equal contribution of 3 individuals...

Then some alleles from individual contributors would be below the stochastic threshold and we could not assume that all alleles are being observed!

Assuming Three Contributors... Some Possible Contributions to This Result

1:1:1

3:1:1

Stochastic alert!

Stochastic alert!

Stochastic alert!

Stochastic alert!

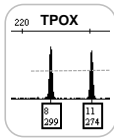
Stochastic alert!

Stochastic alert!

All Loci Are Not Created Equal when it comes to mixture interpretation

- In the case of less polymorphic loci, such as TPOX, there are fewer alleles and these occur at higher frequency. Thus, there is a greater chance of allele sharing (peak height stacking) in mixtures.
- **Higher locus heterozygosity is advantageous for mixture interpretation** – we would expect to see more alleles (within and between contributors) and thus have a better chance of estimating the true number of contributors to the mixture

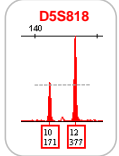
Even if you did attempt to calculate a CPI/CPE statistic using loci with all observed alleles above the stochastic threshold on this result...



TPOX Allele Frequencies (NIST Caucasian, Butler et al. 2003)
 8 = 0.53
 11 = 0.24
 $CPI = (0.53 + 0.24)^2 = 0.59$ or **59%**

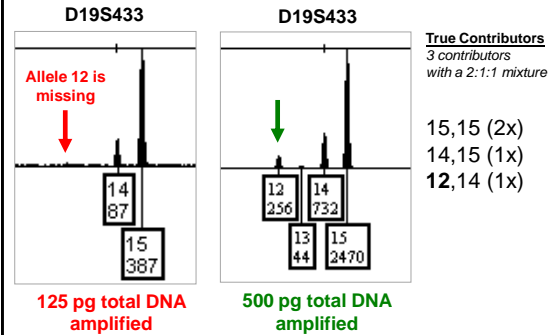
Combine loci = $0.59 \times 0.18 = 0.11$ or **11%**

Approximately 1 in every 9 Caucasians could be included in this mixture



D5S818 Allele Frequencies (NIST Caucasian, Butler et al. 2003)
 10 = 0.05
 12 = 0.38
 $CPI = (0.05 + 0.38)^2 = 0.18$ or **18%**

Impact of Amplifying More DNA



How should you handle the suspect comparison(s) with this case result?

- **No suspect comparisons should be made as the mixture result has too much uncertainty** with stochastic effects that may not account for all alleles being detected
- **Declare the result "inconclusive"**

How not to handle this result

- "To heck with the analytical and stochastic thresholds", **I am just going to see if the suspect profile(s) can fit into the mixture allele pattern observed** – and then if an allele is not present in the evidentiary sample try to explain it with possible allele dropout due to stochastic effects
- This is what Bill Thompson calls "painting the target around the arrow (matching profile)..."

Thompson, W.C. (2009) Painting the target around the matching profile: the Texas sharpshooter fallacy in forensic DNA interpretation. *Law, Probability and Risk* 8: 257-276

What to do with low level DNA mixtures?

- **German Stain Commission "Category C"** (Schneider et al. 2006, 2009)
 - Cannot perform stats because stochastic effects make it uncertain that all alleles are accounted for
- **ISFG Recommendations #8 & #9** (Gill et al. 2006)
 - Stochastic effects limit usefulness
- **Fundamentals of Forensic DNA Typing (2010)** Butler 3rd edition (volume 1), chapter 18
 - Don't go "outside the box" without supporting validation



ISFG Recommendations on Mixture Interpretation

<http://www.isfg.org/Publication;Gill2006>

1. The likelihood ratio (LR) is the preferred statistical method for mixtures over RMNE
2. Scientists should be trained in and use LRs
3. Methods to calculate LR's of mixtures are cited
4. Follow Clayton et al. (1998) guidelines when deducing component genotypes
5. Prosecution determines H_p and defense determines H_d and multiple propositions may be evaluated
6. When minor alleles are the same size as stutters of major alleles, then they are indistinguishable
7. Allele dropout to explain evidence can only be used with low signal data
8. No statistical interpretation should be performed on alleles below threshold
9. Stochastic effects limit usefulness of heterozygote balance and mixture proportion estimates with low level DNA

Gill et al. (2006) DNA Commission of the International Society of Forensic Genetics: Recommendations on the interpretation of mixtures. *Forensic Sci. Int.* 160: 90-101



A Complexity/Uncertainty Threshold

New Scientist article (August 2010)

- **How DNA evidence creates victims of chance**
 - 18 August 2010 by Linda Geddes
- From the last paragraph:
 - **In really complex cases, analysts need to be able to draw a line** and say "This is just too complex, I can't make the call on it," says Butler. "Part of the challenge now, is that every lab has that line set at a different place. But the honest thing to do as a scientist is to say: **I'm not going to try to get something that won't be reliable.**"

<http://www.newscientist.com/article/mg20727743.300-how-dna-evidence-creates-victims-of-chance.html>

Is there a way forward?

"On the Threshold of a Dilemma"

- Gill and Buckleton (2010)
- Although most labs use thresholds of some description, this philosophy has always been problematic because there is an inherent illogicality which we call the falling off the cliff effect.

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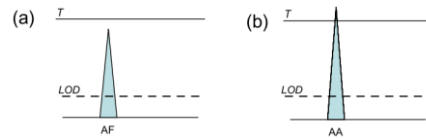


Commentary on: Budowle B, Onorato AJ, Callaghan TF, Della Manna A, Gross AM, Guerrieri RA, Luttman JC, McClure DL. Mixture interpretation: defining the relevant features for guidelines for the assessment of mixed DNA profiles in forensic casework. *J Forensic Sci* 2009;54(4):810–21.

J Forensic Sci, January 2010, Vol. 55, No. 1
doi: 10.1111/j.1556-4029.2009.01257.x
Available online at: intercience.wiley.com

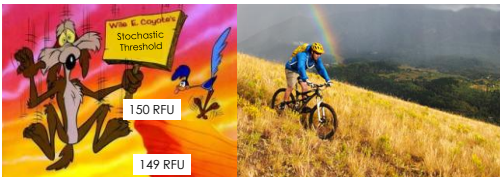
"Falling off the Cliff Effect"

- If T = an arbitrary level (e.g., 150 rfu), an allele of 149 rfu is subject to a different set of guidelines compared with one that is 150 rfu even though they differ by just 1 rfu (Fig. 1).



Gill and Buckleton *JFS* 55: 265-268 (2010)

Falling off the Cliff vs. Gradual Decline




<http://blog.stoneconsulting.com/a/6a00204c1e1c3a01118cc93970c-2e>

<http://ultrafocusecceptic.blogspot.com/2010/08/mourntank.html>

Gill and Buckleton *JFS* 55: 265-268 (2010)

- "The purpose of the ISFG DNA commission document was to provide a way forward to demonstrate the use of **probabilistic models to circumvent the requirement for a threshold** and to safeguard the legitimate interests of defendants."




PAPER

CRIMINALISTICS

Mark W. Perlin,¹ M.D., Ph.D.; Matthew M. Legler,¹ B.S.; Cara E. Spencer,¹ M.S.; Jessica L. Smith,¹ M.S.; William P. Allan,¹ M.S.; Jamie L. Belrose,² M.S.; and Barry W. Duceman,³ Ph.D.


 Validating TrueAllele® DNA Mixture Interpretation*[†]

- Quantitative computer interpretation using Markov Chain Monte Carlo testing
 - Models peak uncertainty and infers possible genotypes
 - Results are presented as the Combined LR



True Allele Software (Cybergenetics)

- We purchased the software in September 2010.
- Three day training at Cybergenetics (Pittsburgh, PA) in October.
- Software runs on a Linux Server with a Mac interface.

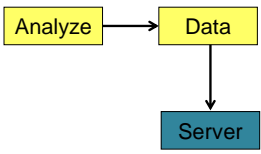
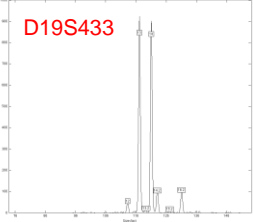


True Allele Casework Workflow 5 Modules

Analyze

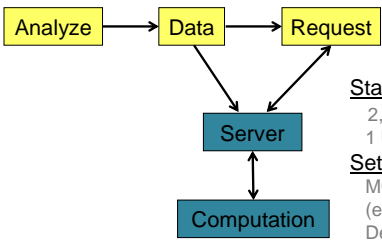
.fsa files imported
 Size Standard check
 Allelic Ladder check
 Alleles are called

True Allele Casework Workflow 5 Modules

All Peaks above 10 RFU are considered

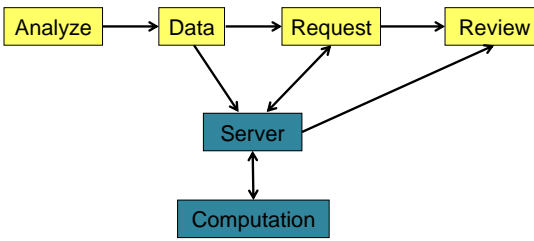
True Allele Casework Workflow 5 Modules

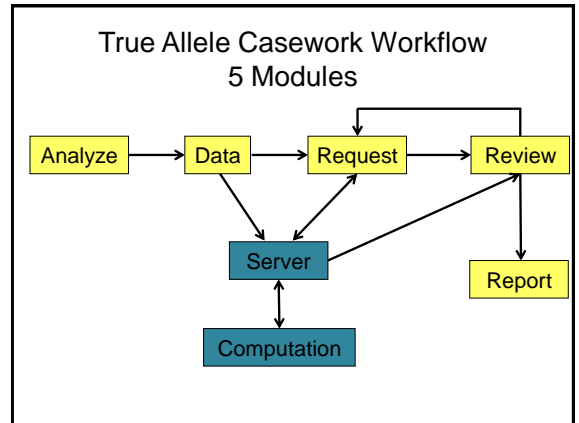
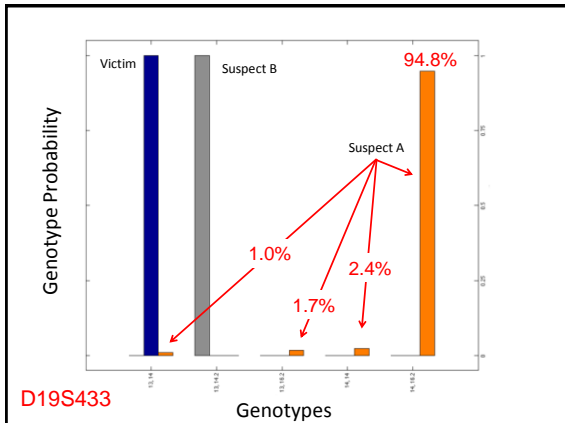
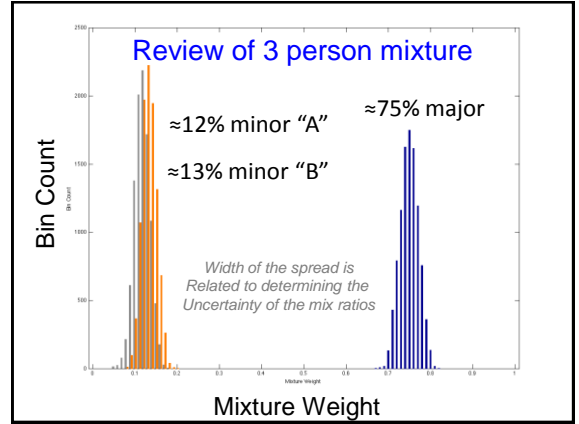
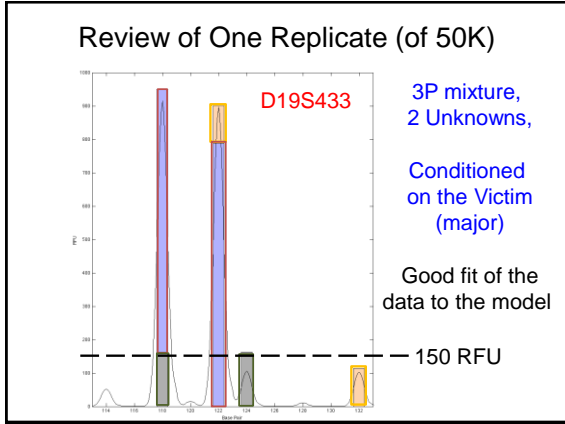


State Assumptions
 2, 3, 4 unknowns
 1 Unk with Victim?

Set Parameters
 MCMC modeling (e.g. 50K)
 Degradation?

True Allele Casework Workflow 5 Modules





Determining the LR for D19S433

Suspect A = 14, 16.2 $H_p = 0.967$

Allele Pair	Probability Before Conditioning
14, 16.2	0.967
14, 14	0.003
13, 16.2	0.026
13, 14	0.001

$LR = \frac{0.967}{0.0122} = 79.26$ H_D

Determining the LR for D19S433

Suspect A = 14, 16.2 $H_p = 0.967$

Allele Pair	Probability Before Conditioning	Genotype Frequency	Probability * Genotype Freq
14, 16.2	0.967	0.0120	0.01164
14, 14	0.003	0.0498	0.00013
13, 16.2	0.026	0.0131	0.00034
13, 14	0.001	0.1082	0.00009
sum			0.0122

$LR = \frac{0.967}{0.0122} = 79.26$ H_D

Combined LR = 5.6 Quintillion

locus	allele pair x	Likelihood l(x)	Genotype Probability Distribution		Reference r(x)	Suspect s(x)	Weighted Likelihood		Likelihood Ratio LR	log(LR)
			Questioned q(x)	Denominator d(x)*r(x)			Numerator l(x)*s(x)	Denominator l(x)*r(x)		
CSF1PO	11, 12	0.686	0.778	0.1448	1	0.68615	0.1292	5.31	0.725	
D13S317	9, 12	1	1	0.0291	1	0.99952	0.02913	34.301	1.535	
D16S539	9, 11	0.985	0.995	0.1238	1	0.98451	0.12188	8.036	0.905	
D18S51	13, 17	0.999	1	0.0154	1	0.99915	0.01543	64.677	1.811	
D19S433	14, 16.2	0.967	0.948	0.012	1	0.96715	0.01222	79.143	1.898	
D21S11	28, 30	0.968	0.98	0.0872	1	0.96809	0.08648	11.194	1.049	
D2S1338	23, 24	0.998	1	0.0179	1	0.99831	0.01787	55.866	1.747	
D3S1358	15, 17	0.988	0.994	0.1224	1	0.98759	0.12084	8.14	0.911	
D5S818	11, 11	0.451	0.394	0.0537	1	0.45103	0.07309	6.17	0.79	
D7S820	11, 12	0.984	0.978	0.0356	1	0.98383	0.03617	27.198	1.435	
D8S1179	13, 14	0.203	0.9	0.1293	1	0.20267	0.02993	6.771	0.831	
FGA	21, 25	0.32	0.356	0.028	1	0.31986	0.01906	16.783	1.225	
TH01	7, 7	0.887	0.985	0.1739	1	0.88661	0.15588	5.687	0.755	
TPOX	8, 8	1	1	0.1375	1	1	0.13746	7.275	0.862	
VWA	15, 20	0.998	0.996	0.0057	1	0.99808	0.00569	174.834	2.243	

Results

- Results are expressed as logLR values

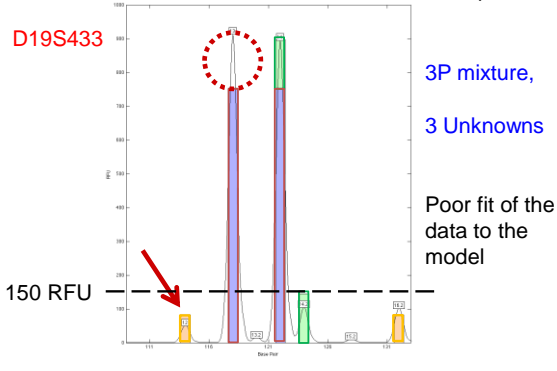
$$LR = 1,000,000 = 10^6$$

$$\log(LR) = \log 10^6$$

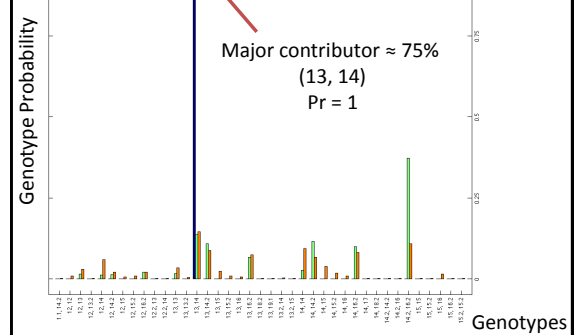
$$\log(LR) = 6 * \log 10 (1)$$

$$\log(LR) = 6$$

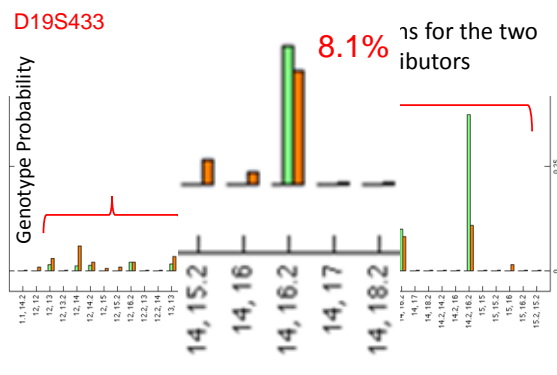
Review of One Replicate (of 50K)



No Conditioning (3 Unknowns)



No Conditioning (3 Unknowns)



locus	allele pair	l	q	r	s	l*r	l*s	LR	log(LR)
D19S433	13, 14	0.002	0.146	0.002	0.0000	0.0000	0.0000		
	14, 2, 16, 2	0.270	0.189	0.0044	0.00118	0.0000	0.0000		
	14, 14	0.002	0.003	0.0000	0.0000	0.0000	0.0000		
	13, 14, 2	0.017	0.004	0.0002	0.0000	0.0000	0.0000		
	14, 16, 2	0.015	0.001	0.0120	1	0.0120	0.00016		
	13, 16, 2	0.018	0.014	0.0031	0.00013	0.0000	0.0000		
	14, 14, 2	0.009	0.007	0.0001	0.0000	0.0000	0.0000		
	12, 14	0.002	0.009	0.0000	0.0000	0.0000	0.0000		
	12, 13	0.002	0.029	0.0041	0.00010	0.0000	0.0000		
	13, 15	0.001	0.024	0.0017	0.00002	0.0000	0.0000		
	12, 16, 2	0.017	0.011	0.0008	0.0000	0.0000	0.0000		
	12, 14, 2	0.015	0.009	0.0130	0.00015	0.0000	0.0000		
	14, 15, 2	0.001	0.018	0.0075	0.00001	0.0000	0.0000		
	15, 16	0.002	0.015	0.0006	0.00000	0.0000	0.0000		
	13, 15, 2	0.001	0.009	0.0039	0.00003	0.0000	0.0000		
	12, 15, 2	0.003	0.009	0.0137	0.00004	0.0000	0.0000		
	14, 16	0.000	0.009	0.0017	0.00000	0.0000	0.0000		
	12, 12	0.004	0.009	0.0121	0.00004	0.0000	0.0000		
	12, 15	0.001	0.006	0.0172	0.00001	0.0000	0.0000		
	13, 16	0.000	0.006	0.0019	0.00000	0.0000	0.0000		
	13, 13, 2	0.001	0.004	0.0061	0.00003	0.0000	0.0000		
	13, 2, 14	0.001	0.003	0.0010	0.00002	0.0000	0.0000		
	12, 2, 15	0.001	0.002	0.0003	0.00001	0.0000	0.0000		
	14, 18, 2	0.002	0.002	0.0017	0.00000	0.0000	0.0000		
	13, 15, 1	0.013	0.002	0.0008	0.00000	0.0000	0.0000		
	12, 13, 2	0.002	0.002	0.0120	0.00003	0.0000	0.0000		
	16, 2, 2	0.001	0.002	0.0006	0.00000	0.0000	0.0000		
	12, 2, 13	0.001	0.002	0.0168	0.00002	0.0000	0.0000		
	13, 2, 16, 2	0.002	0.001	0.0019	0.00000	0.0000	0.0000		
	12, 2, 14	0.001	0.001	0.0135	0.00001	0.0000	0.0000		
	14, 2, 14, 2	0.004	0.001	0.0003	0.00003	0.0000	0.0000		
	15, 13	0.000	0.001	0.0010	0.00000	0.0000	0.0000		
	15, 15, 2	0.000	0.001	0.0005	0.00000	0.0000	0.0000		
	14, 17	0.001	0.001	0.0000	0.00000	0.0000	0.0000		
	15, 16, 2	0.000	0.001	0.0042	0.00000	0.0000	0.0000		
	15, 2, 13, 1	0.001	0.001	0.0010	0.00000	0.0000	0.0000		
	1, 1, 14, 2	0.072	0.001	0.0007	0.00003	0.0000	0.0000		

Suspect "A" Genotype

39 probable genotypes

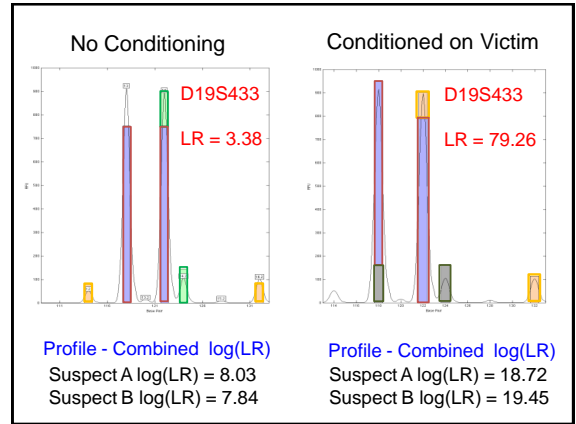
D19S433

Suspect A = 14, 16.2 $H_p = 0.013$

Allele Pair	Probability	Genotype Frequency	Prob* GenFreq
13,14	0.002	0.1082	0.00020
14.2, 16.2	0.270	0.0044	0.00118
14, 14	0.002	0.0498	0.00008
13, 14.2	0.017	0.0392	0.00068
14, 16.2	0.013	0.0120	0.00016
13, 16.2	0.018	0.0131	0.00023
etc...	etc...	etc...	etc...
			Sum 0.00385

$LR = \frac{0.013}{0.00385} = 3.38$ H_D

No Conditioning (3 Unknowns) **D19S433**

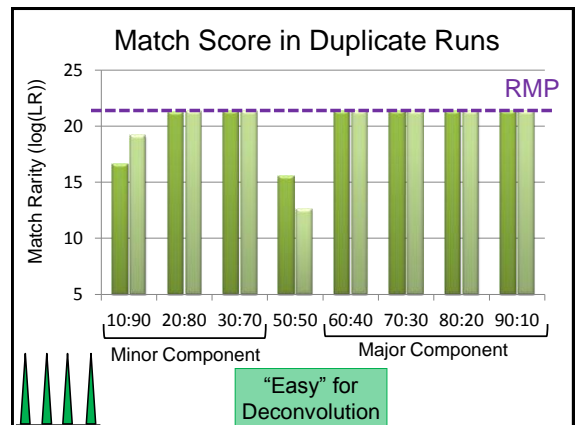
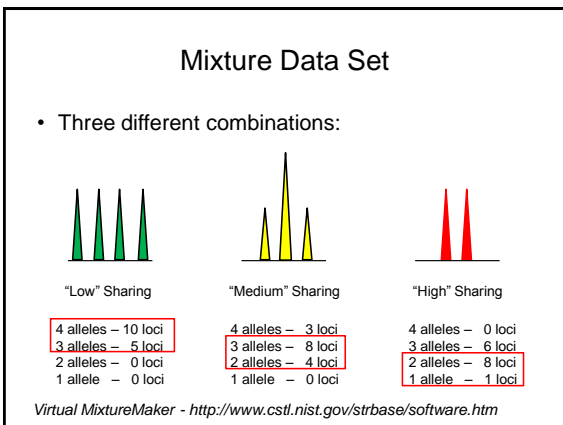


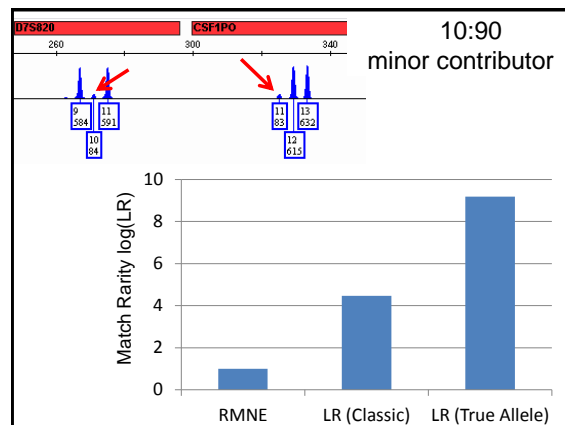
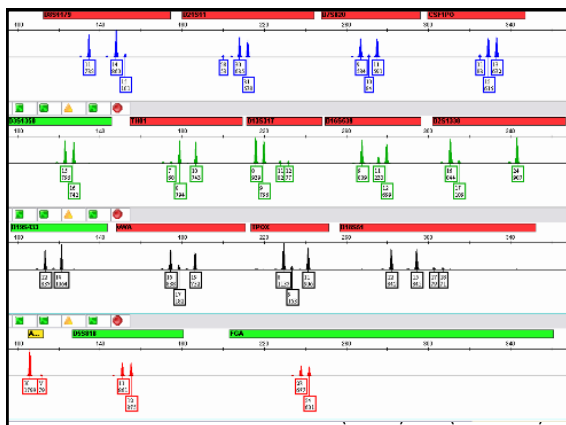
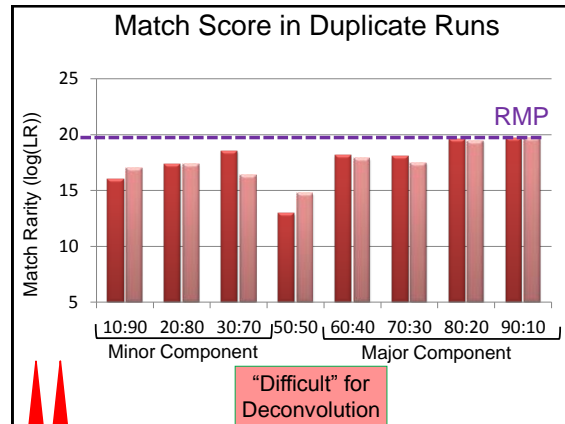
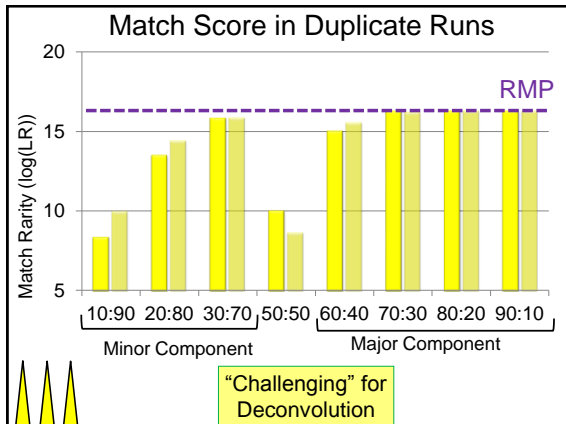
Exploring the Capabilities

- Degree of Allele Sharing
- Mixture Ratios
- DNA Quantity

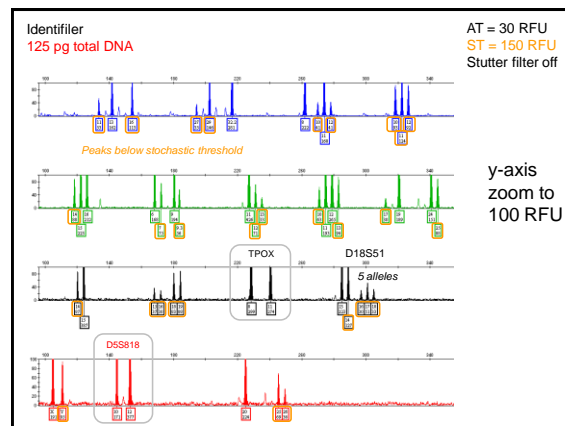
Mixture Data Set

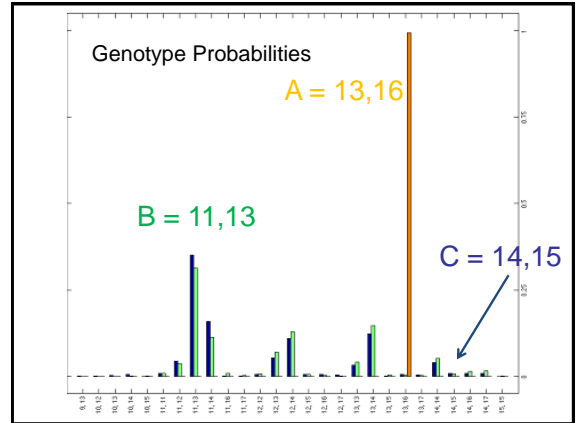
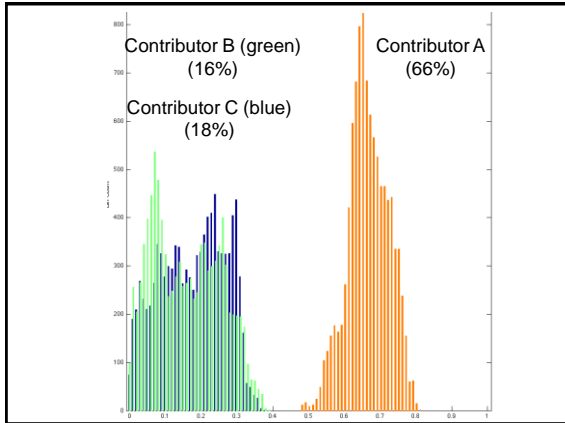
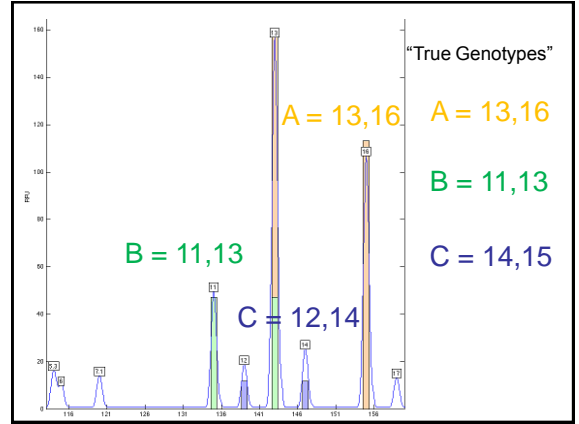
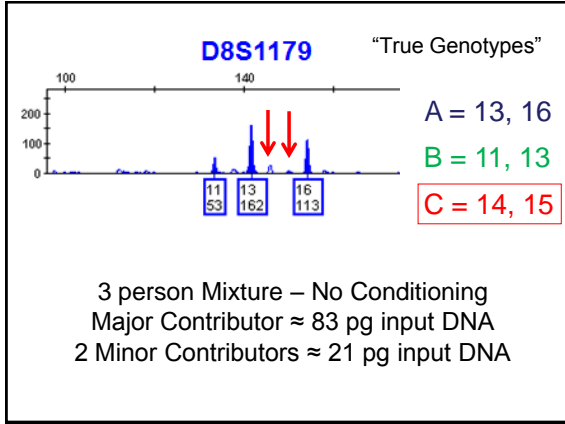
- Mixtures of pristine male and female DNA amplified at a total concentration of 1.0 ng/μL using Identifiler (standard conditions).
- Mixture ratios ranged from 90:10, 80:20, 70:30, 60:40, 50:50, 40:60, 30:70, 20:80, and 10:90
- Each sample was amplified twice.





- ### Exploring the Capabilities
- Degree of Allele Sharing
 - Mixture Ratios
 - DNA Quantity





Results for Contributor A (male)

Locus	Allele Pair	Probability		Genotype Suspect	H _p		H _d		LR
		Likelihood	Frequency		Numerator	Denominator	Numerator	Denominator	
CSF1PO	10, 11	0.572	0.1292				0.07395		
	11, 12	0.306	0.2133	1	0.30563	0.0652			
	10, 12		0.12	0.1547			0.01861		
						0.30563	0.15791	1.935	
D13S317	11, 11	1	0.1149	1	1	0.11488	8.704		
D8S1179	13, 16	0.998	0.0199	1	0.99786	0.0199	49.668		

The match rarity between the evidence and suspect is 1.21 quintillion

Results for Contributor B (female)

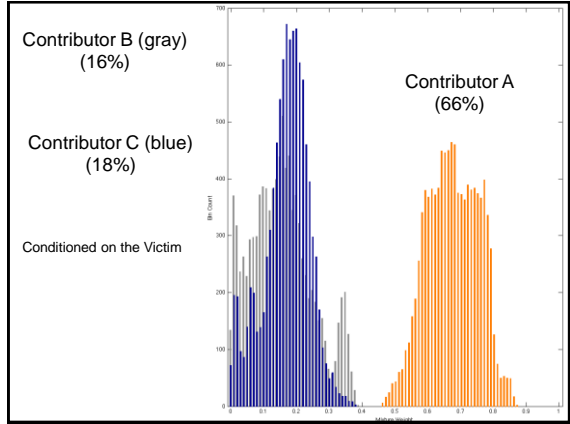
Locus	Allele Pair	Probability		Genotype Suspect	H _p		H _d		LR
		Likelihood	Frequency		Numerator	Denominator	Numerator	Denominator	
D8S1179	11, 13	0.073	0.0498	1	0.07338	0.00366			
	11, 14	0.034	0.0271			0.00092			
	13, 14	0.006	0.0996			0.00065			
	12, 14	0.011	0.0606			0.00068			
	12, 13	0.005	0.1115			0.0006			
	11, 12	0.018	0.0303			0.00054			
	14, 14	0.004	0.0271			0.00012			
	13, 13	0.003	0.0916			0.00031			
	14, 16	0.003	0.0108			0.00003			
	14, 15	0.001	0.0379			0.00003			
etc...									9.197

The match rarity between the evidence and suspect is 1.43 million

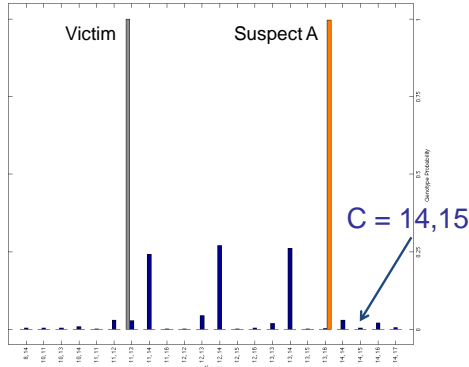
Results for Contributor C (male)

Locus	Allele Pair	Probability		Suspect	H _p		H _d		LR
		Likelihood	Genotype Frequency		Numerator	Denominator			
D8S1179	11, 13	0.056	0.0498					0.00279	
	13, 14	0.007	0.0996					0.00066	
	12, 14	0.011	0.0606					0.00068	
	11, 14	0.021	0.0271					0.00056	
	12, 13	0.006	0.1115					0.00066	
	14, 14	0.005	0.0271					0.00013	
	etc...	etc...	etc...					etc...	
	14, 15	0.001	0.0379	1	0.00056			0.00002	
	12, 15	0.001	0.0424					0.00003	
	etc...	etc...	etc...					etc...	
	10, 15	0	0.0227					0.00001	
						0.00056		0.00665	0.084

The match rarity between the evidence and suspect is 9.16 thousand



The Power of Conditioning



The Power of Conditioning

	LR (no conditioning, 3unk)
Contributor A	1.21 Quintillion
Contributor B (victim)	1.43 Million
Contributor C	9.16 Thousand

	LR (conditioned on victim + 2unk)
Contributor A	1.32 Quintillion
Contributor B (victim)	2.19 Million
Contributor C	59.8 Thousand

↑
Ranged from 1.13 to 800K

Summary

- True Allele utilizes probabilistic genotyping and makes better use of the data than the RMNE approach.
- However, the software is computer intensive. On our 4 processor system, it can take 12-16 hours to run up to four 3-person mixture samples.

Summary

- **Allele Sharing:** Stacking of alleles due to sharing creates more uncertainty.
- **Mixture Ratio:** With "distance" between the two contributors, there is greater certainty. Generally, True Allele performs better than RMNE and the classic LR with low level contributors.

Summary

- **DNA Quantity:** Generally, with high DNA signal, replicates runs on True Allele are very reproducible.
- However, with low DNA signal, higher levels of uncertainty are observed (as expected).
- There is a need to determine an appropriate threshold for an inclusion log(LR).

Thank you for your attention

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