



Topics and Techniques for Forensic DNA Analysis
Continuing Education Seminar

Capillary Electrophoresis Fundamentals and Troubleshooting

NYC OCME
Dept of Forensic
Biology



New York City, NY
March 25, 2009






Dr. John M. Butler
National Institute of
Standards and Technology
john.butler@nist.gov

Presentation Outline

- History and background on CE
- Separation
- Injection and sample preparation
- Detection
- Troubleshooting



My Goal:
To help you understand the basic chemistry behind DNA separations
and to help make CE instruments less of a "black box"

Pioneers of Capillary Electrophoresis

		
Stellan Hjertén Uppsala University	James Jorgenson University of North Carolina	Barry Karger Northeastern University
1967 First high voltage CE system (with rotating 3 mm i.d. capillaries)	1981 First "modern" CE experiments (with 75 µm i.d. capillaries)	1988/90 First DNA separations in a capillary (gel-filled/sieving polymer)

Stellan Hjertén

Uppsala University (Sweden)



In 2003 at age 75

With first fully automated capillary free zone electrophoresis apparatus in 1967

Received his PhD (1967) under Professor Arne Tiselius who had developed moving boundary zone electrophoresis in 1937 (Noble Prize in 1948)

A Brief History of Capillary Electrophoresis

- 1937 – Tiselius develops moving boundary electrophoresis
- 1967 – Hjertén uses rotating 3 mm i.d. tubes for CE
- 1981 – Jorgenson and Lukacs demonstrate first high performance CE separations with 75 µm i.d. capillary
- 1988 – Karger's group shows DNA separations of single stranded oligonucleotides with gel-filled capillaries
- 1990 – Karger's group shows DNA separations with sieving polymers on DNA restriction fragments
- 1991 – Grossman expands work with sieving polymers
- 1992 – Bruce McCord starts working on PCR product separations with STR allelic ladders

My Experience with CE, STRs, etc.

- May 1993 – began working in Bruce McCord's lab at Quantico
- Sept 1993 – developed mtDNA amplicon quantitation method (used in FBI casework from 1996 to present)
- Nov 1993 – first demonstration of STR typing by CE (using dual internal standards and TH01 ladder)
- July 1995 – defended Ph.D. dissertation entitled "Sizing and Quantitation of Polymerase Chain Reaction Products by Capillary Electrophoresis for Use in DNA Typing"
- July 1995 – ABI 310 Genetic Analyzer was released

My Experience with CE, STRs, etc. (cont.)

- 1996-1997 Developed STRBase while a postdoc at NIST
- Nov 1998 – GeneTrace Systems purchased a 310; typed several hundred samples with Profiler Plus and Cofiler kits and compared results to mass spec STR analysis
- **1999-present – Run thousands of samples with all STR kits available (except PP 1.2) and developed a number of new STR multiplex systems**
- Jan 2001 – Published “Forensic DNA Typing: Biology and Technology behind STR Markers” (2nd Edition in Feb 2005)
- April 2001-present – Use of ABI 3100 16-capillary array system

In the early 1990s the real question was how to transition from a gel to a capillary

- Cross-linked acrylamide gel filled capillaries were tried first
 - Reusable?
 - Bubble formation
 - Thermal degradation
- Alternative was to not use a gel at all
 - Refillable sieving polymers
 - However, resolution was poor early on

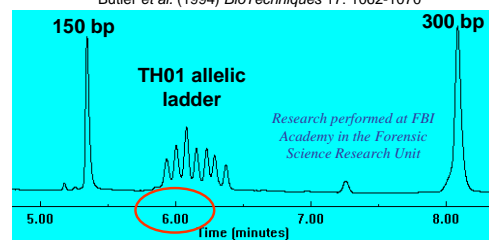
Early Work with CE and STRs

- Barry Karger’s group (1988-1990)
 - Utilized gel-filled capillaries to separate ssDNA
 - Introduced sieving polymers in the form of linear polyacrylamide to separate restriction digests
- Beckman P/ACE 2050 is introduced in 1992 as the first commercially available CE coupled to a laser to enable fluorescence detection
- John Butler and Bruce McCord (1993-1995)
 - **First STR typing with single color CE using intercalating dyes and dual bracketing internal size standards**
- Rich Mathies’ group (1995)
 - First STR typing with multi-color CE (and multi-capillary) using dye-labeled primers
- **ABI 310 is introduced in July 1995 as the first commercially available multi-color CE**

First Rapid STR Typing with Capillary Electrophoresis

Single color detection with dual internal size standards

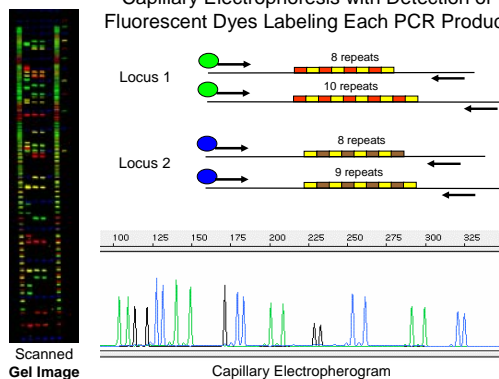
Butler et al. (1994) *BioTechniques* 17: 1062-1070



Performed in December 1993

Technology Implementation Takes Time – the FBI did not start running casework samples using STRs and CE until January 1999

STR Allele Separation Can Be Performed by Gel or Capillary Electrophoresis with Detection of Fluorescent Dyes Labeling Each PCR Product

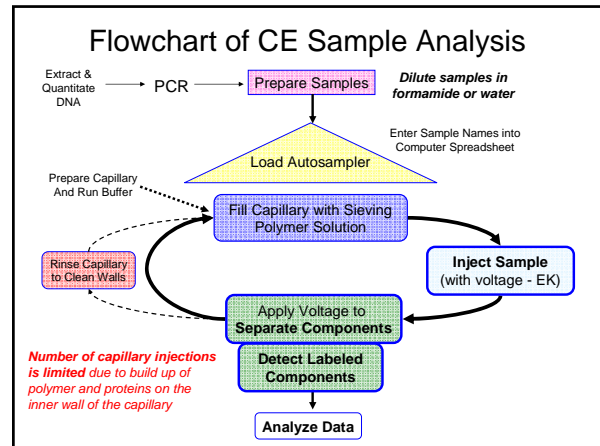
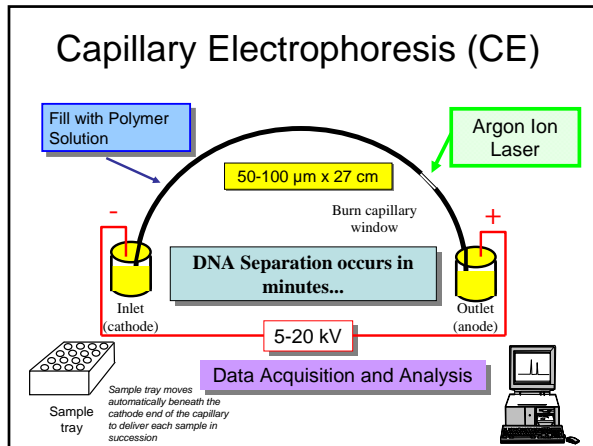


Why Use CE for DNA Analysis?

1. Injection, separation, and detection are automated.
2. Rapid separations are possible
3. Excellent sensitivity and resolution
4. The time at which any band elutes is precisely determined
5. Peak information is automatically stored for easy retrieval



Symbol first used in Oct 1994 at the Promega meeting when I had a poster introducing the use of CE for STR typing



Typical Instruments Used for STR Typing

GeneAmp 9700

Thermal Cycler for PCR Amplification

Capillary electrophoresis instruments for separating and sizing PCR products

single capillary

16-capillary array

ABI 310

ABI 3100

Review Article on STRs and CE

pdf available from <http://www.cstl.nist.gov/biotech/strbase/NISTpub.htm>

Electrophoresis 2004, 25, 1397-1412

Review

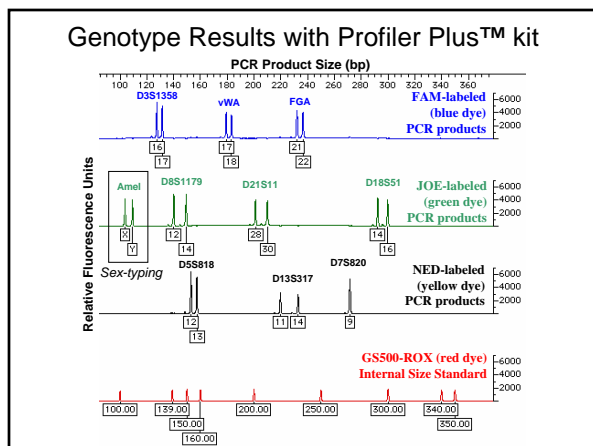
John M. Butler¹
 Eric Bue²
 Federica Crivellente^{3*}
 Bruce R. McCord⁴

Forensic DNA using the ABI for STR analysis

DNA typing with short applications including such as the ABI Prizm for many laboratories ing sample preparation results using CE systems in the context of throughput and ease

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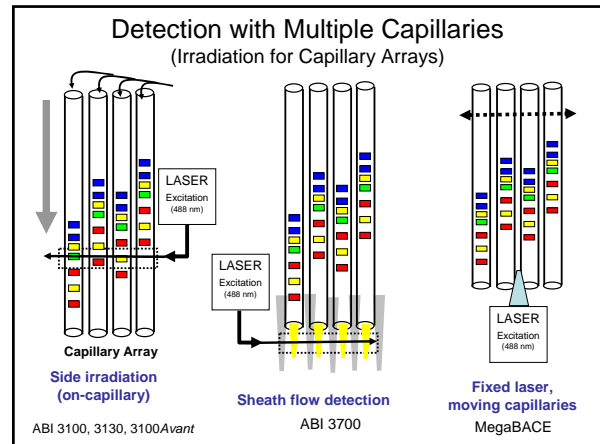
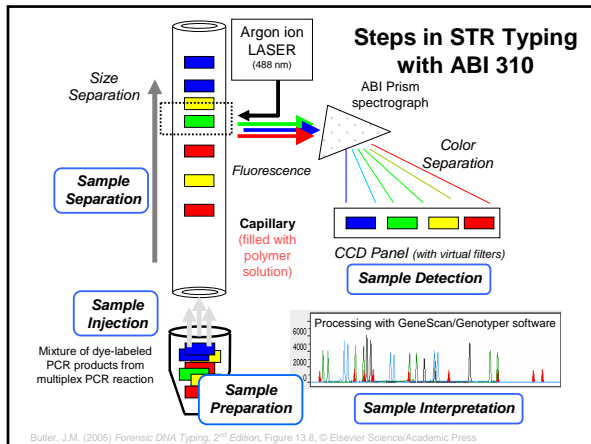
¹National Institute of Standards and Technology, Biotechnology Division, Gaithersburg, MD, USA
²Vermont Forensic Laboratory, Waterbury, VT, USA
³Ohio University, Department of Chemistry, Athens, OH, USA



Analytical Requirements for STR Typing

Butler et al. (2004) Electrophoresis 25: 1397-1412

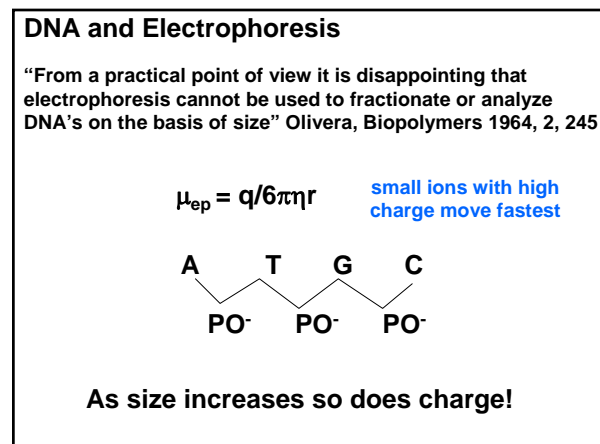
- Fluorescent dyes must be **spectrally resolved** in order to distinguish different dye labels on PCR products
- PCR products must be **spatially resolved** – desirable to have single base resolution out to >350 bp in order to distinguish variant alleles
- High **run-to-run precision** – an internal sizing standard is used to calibrate each run in order to compare data over time

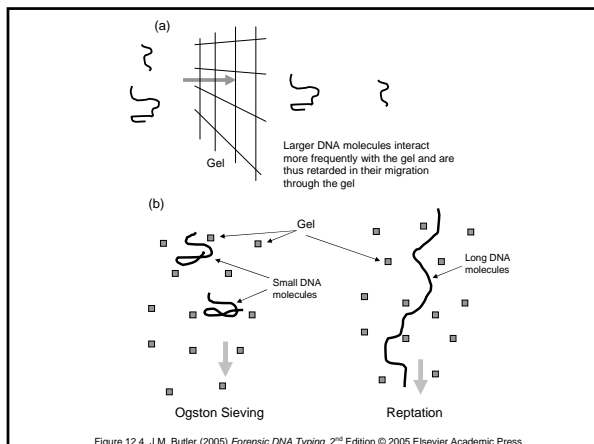


- ### Process Involved in 310/3100 Analysis
- **Separation**
 - Capillary – 50um fused silica, 43 cm length (36 cm to detector)
 - POP-4 polymer – Polydimethyl acrylamide
 - Buffer - TAPS pH 8.0
 - Denaturants – urea, pyridinone
 - **Injection**
 - electrokinetic injection process (formamide, water)
 - importance of sample stacking
 - **Detection**
 - fluorescent dyes with excitation and emission traits
 - CCD with defined virtual filters produced by assigning certain pixels

Separation

- ### Ohm's Law
- $V = IR$ (where V is voltage, I is current, and R is resistance)
 - Current, or the flow of ions, is what matters most in electrophoresis
 - CE currents are much lower than gels because of a higher resistance in the narrow capillary
 - CE can run a higher voltage because the capillary offers a higher surface area-to-volume ratio and can thus dissipate heat better from the ion flow (current)





- ### Separation Issues
- **Electrophoresis buffer** –
 - Urea for denaturing and viscosity
 - Buffer for consistent pH
 - Pyridinone for denaturing DNA
 - EDTA for stability and chelating metals
 - **Polymer solution** -- POP-4 (but others work also)
 - **Capillary wall coating** -- dynamic coating with polymer
 - Wall charges are masked by methyl acrylamide
 - **Run temperature** -- 60 °C helps reduce secondary structure on DNA and improves precision. (Temperature control affects DNA sizing)

DNA Separations in Entangled Polymer Sieving Solutions

- Size based separation due to interaction of DNA molecules with entangled polymer strands
- Polymers are **not cross-linked** (as in slab gels)
- "Gel" is **not attached** to the capillary wall
- **Pumpable** -- can be replaced after each run
- Polymer length and concentration determine the separation characteristics

Entangled Polymer Solutions

- Polymers are not cross-linked (above entanglement threshold)
- "Gel" is not attached to the capillary wall
- Pumpable -- can be replaced after each run
- Polymer length and concentration determine the separation characteristics
- Examples:
 - 1% HEC (hydroxyethyl cellulose)
 - 4% polyvinyl pyrrolidone
 - POP-4 and POP-6

POP4 Polymer
Polydimethyl acrylamide

Transient Pores Are Formed Above the Entanglement Threshold.

$C < C^*$ $C = C^*$ $C > C^*$

Ogston Sieving **Reptation** **Entanglement**

$\mu \sim \mu_0 e^{-NC}$ $\mu \sim 1/N$ $\mu \sim f(1/CN)$

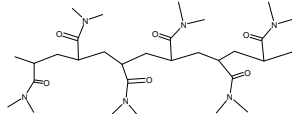
What is in POP-4 and Genetic Analyzer Buffer?

© 1997 Oxford University Press *Nucleic Acid Research*, 1997, Vol. 25, No. 19 3925-3929
Improved single-strand DNA sizing accuracy in capillary electrophoresis
 Barnett B. Rosenblum*, Frank Oaks, Steve Menchen and Ben Johnson
 PE Applied Biosystems, 850 Lincoln Centre Drive, Foster City, CA 94404, USA
 Received May 25, 1997; Revised and Accepted August 6, 1997

See also Wenz et al. (1998) *Genome Research* 8: 69-80

POP-4 (4% poly-dimethylacrylamide, 8 M urea, 5% 2-pyrrolidinone)
 US Patent 5,552,028 covers POP-4 synthesis

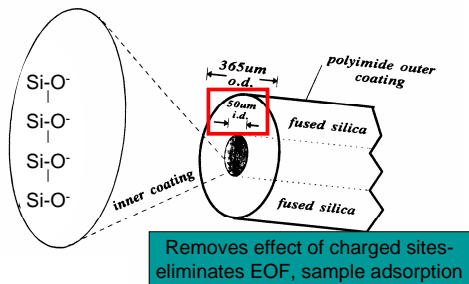
Running buffer contains 100 mM TAPS and 1 mM EDTA (adjusted to pH 8.0 with NaOH) TAPS = N-Tris-(hydroxymethyl)methyl-3-aminopropane-sulfonic acid



Why TAPS instead of Tris-borate (TBE) buffer?

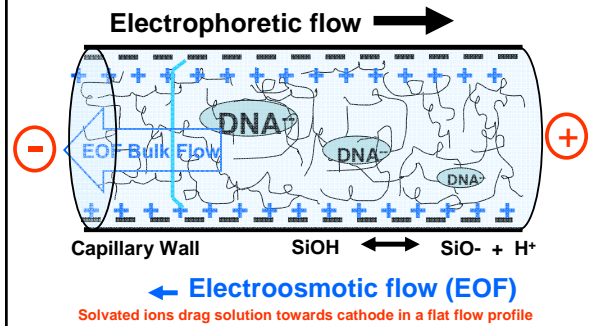
- TBE is temperature/pH sensitive
 - as temperature increases, the pH decreases (0.02 pH units with every 1 °C); this is the principle by which TaqGold activation works
- At lower pH, fluorescence emission of dyes decreases
 - see Singer and Johnson (1997) *Proceedings of the Eighth International Symposium on Human Identification*, pp. 70-77
- Thus when running at 60 °C on the ABI 310, if Tris-borate was used, fluorescent intensity of PCR products would be lower

Capillary Coating

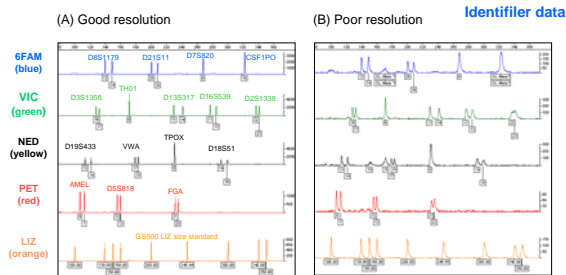


Dynamic coating of charged sites on fused silica capillary is accomplished with POP-4 polymer

Capillary Wall Coatings Impact DNA Separations

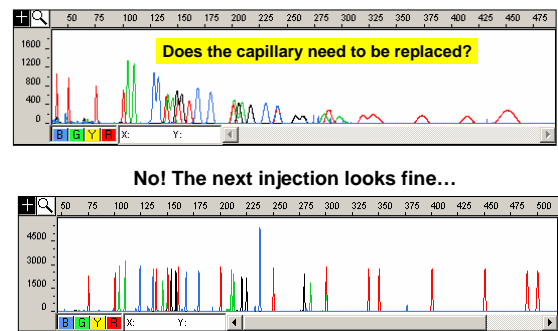


Capillary Resolution Differences



Butler, J.M., Buel, E., Crivellente, F., McCord, B.R. (2004) Forensic DNA typing by capillary electrophoresis: using the ABI Prism 310 and 3100 Genetic Analyzers for STR analysis. *Electrophoresis*, 25: 1397-1412.

What we call "melt downs" ... probably due to an incompletely filled capillary

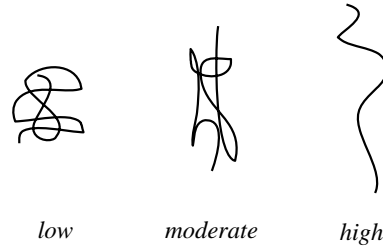


How to Improve Resolution?

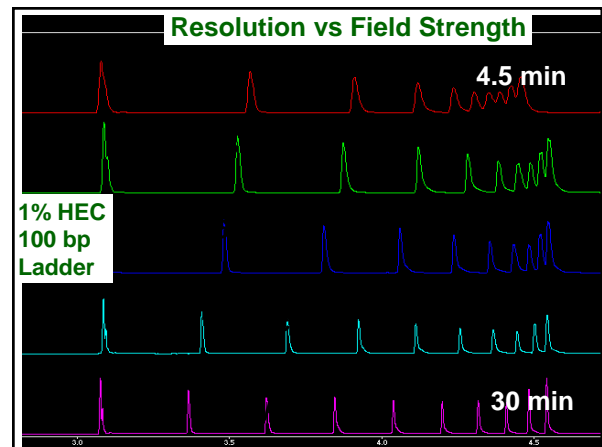
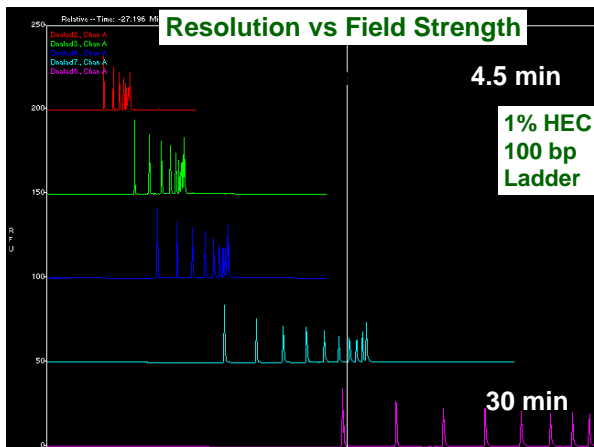
1. Lower Field Strength
2. Increase Capillary Length
3. Increase Polymer Concentration
4. Increase Polymer Length

All of these come at a cost of longer separation run times

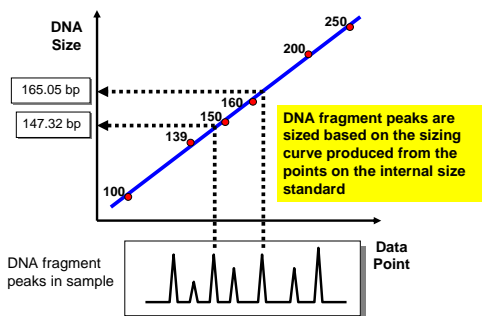
The electric field strength can influence the shape of the DNA molecule.



low moderate high
 Optimal separations usually 180-200 V/cm



Process of Sizing DNA Fragments Using an Internal Standard



Injection

CE Injection Methods

ABI 310

Hydrodynamic (pressure) Electrokinetic (voltage)

Uffelder K. J.; McCord, B. R. (1996) Capillary Electrophoresis of DNA, In *Handbook of Capillary Electrophoresis* (Landers, J., ed.), CRC Press: NY, pp. 347-378.
 Butler, J.M. (1997) Effects of sample matrix and injection on DNA separations. *Analysis of Nucleic Acids by Capillary Electrophoresis* (Heller, C., ed.), Vieweg: Germany, Chapter 5, pp. 125-134

Electrokinetic Injection Process

Amount of DNA injected is inversely proportional to the ionic strength of the solution

Salty samples result in poor injections

Sample Conductivity Impacts Amount Injected

$$[\text{DNA}_{inj}] = \frac{Et(\pi r^2)(\mu_{ep} + \mu_{eof})[\text{DNA}_{sample}](\lambda_{buffer})}{\lambda_{sample}}$$

[DNA_{inj}] is the amount of sample injected **[DNA_{sample}]** is the concentration of DNA in the sample
E is the electric field applied
t is the injection time **λ_{buffer}** is the buffer conductivity
r is the radius of the capillary **λ_{sample}** is the sample conductivity

μ_{ep} is the mobility of the sample molecules
μ_{eof} is the electroosmotic mobility

Cl⁻ ions and other buffer ions present in PCR reaction contribute to the sample conductivity and thus will compete with DNA for injection onto the capillary

Butler et al. (2004) *Electrophoresis* 25: 1397-1412

Two Major Effects of Sample Stacking

1. Sample is preconcentrated. Effect is inversely proportional to ionic strength
2. Sample is focused. Ions stop moving in low electric field
3. Mobility of sample = $\mu_{ep} = \text{velocity} / \text{electric field}$

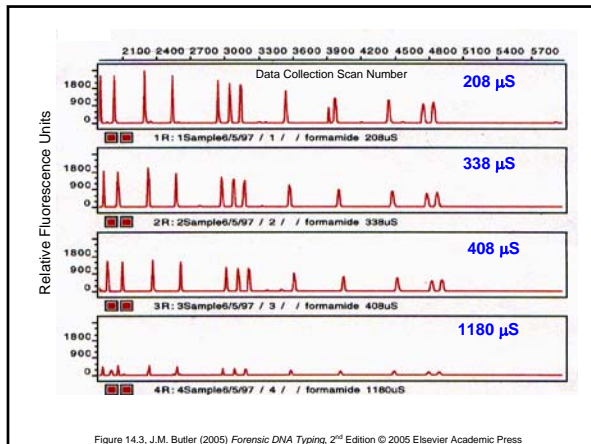
Steps Performed in Standard Module

See J.M. Butler (2005) *Forensic DNA Typing*, 2nd Edition; Chapter 14

- **Capillary fill** – polymer solution is forced into the capillary by applying a force to the syringe
- **Pre-electrophoresis** – the separation voltage is raised to 10,000 volts and run for 5 minutes;
- **Water wash of capillary** – capillary is dipped several times in deionized water to remove buffer salts that would interfere with the injection process
- **Sample injection** – the autosampler moves to position A1 (or the next sample in the sample set) and is moved up onto the capillary to perform the injection; a voltage is applied to the sample and a few nanoliters of sample are pulled onto the end of the capillary; the default injection is 15 kV (kilovolts) for 5 seconds
- **Water wash of capillary** – capillary is dipped several times in waste water to remove any contaminating solution adhering to the outside of the capillary
- **Water dip** – capillary is dipped in clean water (position 2) several times
- **Electrophoresis** – autosampler moves to inlet buffer vial (position 1) and separation voltage is applied across the capillary; the injected DNA molecules begin separating through the POP-4 polymer solution
- **Detection** – data collection begins; raw data is collected with no spectral deconvolution of the different dye colors; the matrix is applied during Genescan analysis

Typical Sample Preparation for ssDNA

1. Perform PCR with dye-labeled primers
2. Dilute 1 μL PCR product with 24 μL **deionized formamide**; add 1 μL ROX-labeled internal sizing standard
3. Denature 2 minutes at 95 °C with thermocycler
4. Cool to 4 °C in thermocycler or ice bath
5. Sample will remain denatured for at least 3 days



Comments on Sample Preparation

- Use high quality formamide (<100 μS/cm!)
 - ABI sells Hi-Di formamide
 - regular formamide can be made more pure with ion exchange resin
- Deionized water vs. formamide
 - Biega and Ducean (1999) *J. Forensic Sci.* 44: 1029-1031
 - Crivellente, *Journal of Capillary Electrophoresis* 2002, 7 (3-4), 73-80.
 - water works fine but samples are not stable as long as with formamide; water also evaporates over time...
- Denaturation with heating and snap cooling
 - use a thermal cycler for heating and cold aluminum block for snap cooling
 - heat/cool denaturation step is necessary only if water is substituted for formamide...

January 6, 2005 Letter from Applied Biosystems to ABI 310 Customers

- “Testing has shown that Hi-Di Formamide denatures DNA **without the need to heat samples...**”
- In other words, no heat denaturation and snap cooling needed!

Applied Biosystems Okays Use of Deionized Water for DNA Sequencing

Technical Bulletin #1 Issued August 2006 Applied Biosystems 3730/3730xl DNA Analyzer

Subject: Influence of Sequencing Injection Solution on 3730/3730xl DNA Analyzer Performance

In this Bulletin:

- Three Loading Solutions Tested on Page 1
- Loading Solution Test Data on Page 2
- Recommendations on Page 6
- Guidelines for Use on Page 6

Three Loading Solutions Tested

Loading Solution Background

Applied Biosystems presently recommends the use of Hi-Di™ Formamide as the sample-loading solution for all Applied Biosystems DNA sequencers to ensure sample preservation and resistance to evaporation. However, many users of the 3730 choose either deionized water or dilute EDTA solutions. These choices are driven largely by cost and safety/hazardous material considerations.

Detection

Detection Issues

- Fluorescent dyes
 - spectral emission overlap
 - relative levels on primers used to label PCR products
 - dye “blobs” (free dye)
- Virtual filters
 - hardware (CCD camera)
 - software (color matrix)

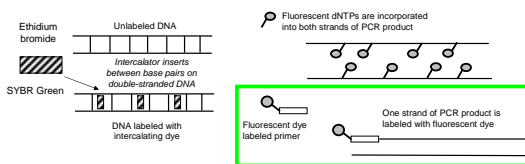
Filters determine which wavelengths of light are collected onto the CCD camera

Laser Used in ABI 310

- Argon Ion Laser
- 488 nm and 514.5 nm for excitation of dyes
- 10 mW power
- Lifetime ~5,000 hours (1 year of full-time use)
- Cost to replace ~\$5,500
- Leads to highest degree of variability between instruments and is most replaced part
- Color separation matrix is specific to laser used on the instrument

Methods for Fluorescently Labeling DNA

- Intercalating Dyes (post-PCR)
- Dye-labeled nucleotide insertion during PCR
- Dye-labeled primer insertion during PCR

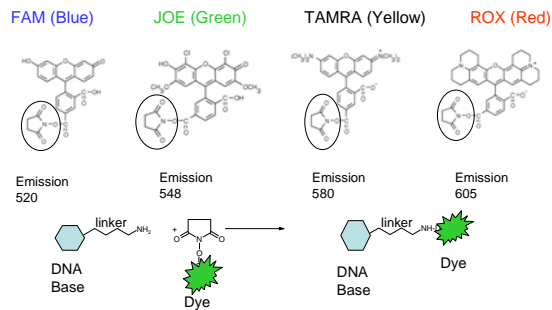


Butler, J.M. (2001) *Forensic DNA Typing*, Figure 10.2, ©Academic Press

Fluorescent Labeling of PCR Products

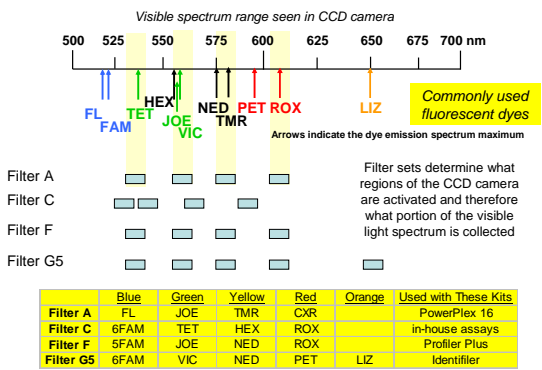
- Dyes are attached to one primer in a pair used to amplify a STR marker
- Dyes are coupled to oligonucleotides (primers) through NHS-esters and amine linkages on the 5' end of the primer: Dye-(CH₂)₆-primer
- Dye-labeled oligonucleotides are incorporated during multiplex PCR amplification giving a specific color "tag" to each PCR product
- PCR products are distinguished using CCD imaging on the 310

Amine Reactive Dyes used in Labeling DNA

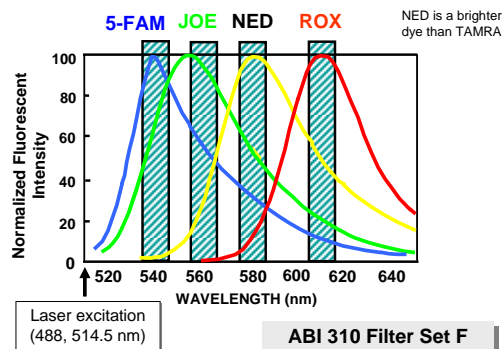


The succinimidyl ester reacts rapidly with amine linkers on DNA bases

Virtual Filters Used in ABI 310



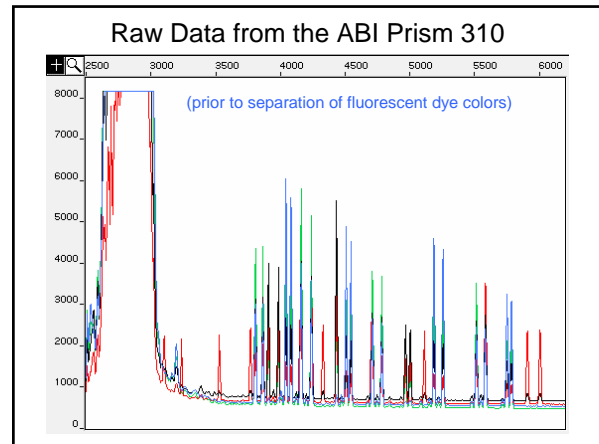
Fluorescent Emission Spectra for ABI Dyes



Butler, J.M. (2001) *Forensic DNA Typing*, Figure 10.4, ©Academic Press

Please Note!

- There are no filters in a 310
- Its just the choice of pixels in the CCD detector
- All the light from the grating is collected
- You just turn some pixels on and some off

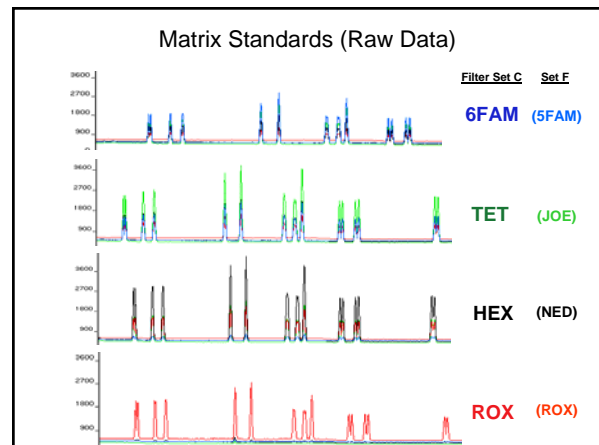


Why Make a Matrix?

The matrix is the solution to a problem:
 What's the contribution at any given wavelength
 (filter set) from each dye ?

There are 4 dyes

- Remember algebra from high school?
- To solve a problem with 4 unknowns, you need 4 equations



For Example

$$I_{540} = bx_b + gy_b + yz_b + rw_b \text{ intensity of blue}$$

$$I_{560} = bx_g + gy_g + yz_g + rw_g \text{ intensity of green}$$

$$I_{580} = bx_y + gy_y + yz_y + rw_y \text{ intensity of yellow}$$

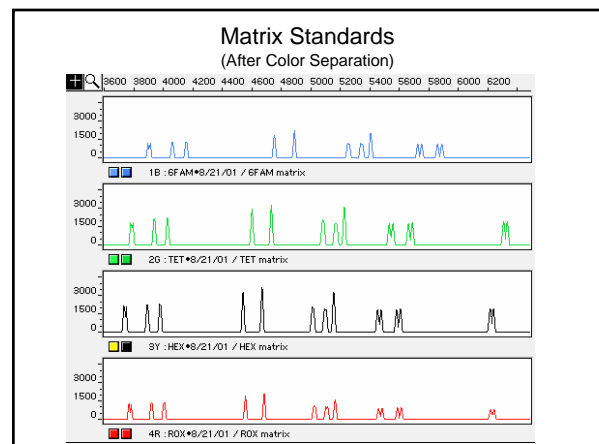
$$I_{610} = bx_r + gy_r + yz_r + rw_r \text{ intensity of red}$$

Where
b is the %blue labeled DNA
g is the %green labeled DNA, etc.

x,y,z,w are the numbers in the
 matrix (sensitivity to each color)

POP4STRM00F				
Reactions				
	B	G	Y	R
B	1.0000	0.8502	0.1380	0.0009
G	0.8300	1.0000	0.7622	0.0051
Y	0.6416	0.8324	1.0000	0.1102
R	0.4495	0.6484	0.7851	1.0000

If you solve xyzw for each dye individually
 Then you can determine dye contribution for any mixture



Matrix File Table from an ABI 310

POP4STRMODF				
Reactions				
	B	G	Y	R
B	1.0000	0.8502	0.1380	0.0009
G	0.8300	1.0000	0.7622	0.0051
Y	0.6416	0.8324	1.0000	0.1102
R	0.4493	0.6484	0.7851	1.0000

The results of the calculation are in a matrix
 (remember linear algebra?)

The values represent the percent spectral overlap from each dye
 Values outside this range represent mixtures

Comments on Matrices/Spectral Calibration (Multi-Component Analysis)

- Make sure that the right filter set and matrix are applied when collecting data
- You can always apply another matrix to a sample collected on the ABI 310 but it must be run with the right filter set (matrix must be run first with ABI 3100)
- It is important to update matrices on a regular basis (depending on use) due to differences in laser power over time
- A good indication of when to run a new matrix is the observation of pull-up between dye colors when peaks are smaller than ~4,000 RFUs

Deciphering Artifacts from the True Alleles

Biological (PCR) artifacts

Stutter products

D3S1358

Incomplete adenylation

D8S1179

STR alleles

Dye blob

stutter

spike

Blue channel

Green channel

Yellow channel

Red channel

Pull-up (bleed-through)

Butler, J.M. (2000) Forensic DNA Typing, 2nd Edition, Figure 15.4, © Elsevier Science/Academic Press

Dye Blobs ("Artifacts")

- Free dye (not coupled to primer) can be injected into the CE capillary and interfere with detection of true STR alleles
- **Dye blobs are wider and usually of less intensity** than true STR alleles (amount depends on the purity of the primers used)
- Dye blobs usually appear at an apparent size that is unique for each dye (e.g., FAM ~120 bp, PET ~100 bp)

Poor primer purity

HEX dye blob

DYS437

Dye Blob Problems with Some PCR Primers Individual Y-STR Locus Amplifications

PCR product size (bp)

DYS392

DYS438

DYS437

HEX

Dye blobs

Poor primer purity

Removal of Dye Artifacts Following PCR Amplification

TH01

TPOX

CSF1PO

FGA

D21S11

D7S820

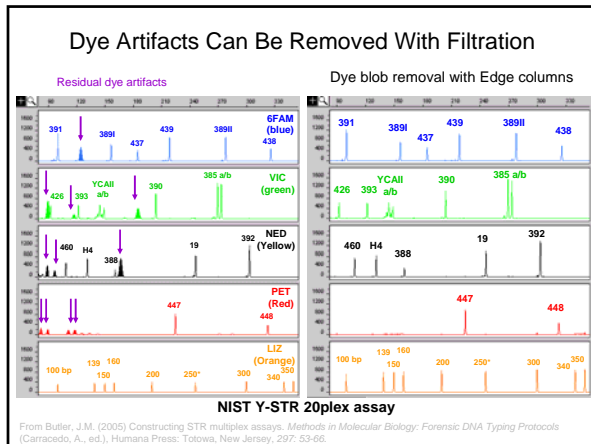
No Filtering (Straight from PCR)

Filtered with Edge columns

Note higher RFU values due to salt reduction with spin columns

EDGE GEL FILTRATION CARTRIDGES

Butler, J.M., Shen, Y., McCord, B.R. (2003) The development of reduced size STR amplicons as tools for analysis of degraded DNA, J. Forensic Sci 48(5) 1054-1064.



Conclusions

DNA typing by capillary electrophoresis involves:

- 1) The use of entangled polymer buffers
- 2) Injection by sample stacking
- 3) Multichannel laser induced fluorescence
- 4) Internal and external calibration

Practical Aspects of ABI 310/3100 Use

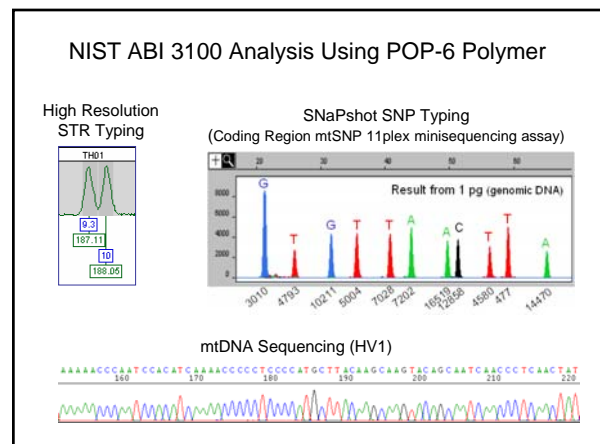
ABI Genetic Analyzer Usage at NIST

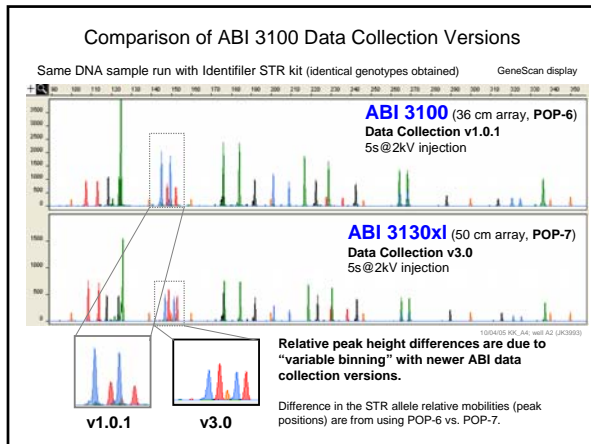
- ABI 310 x 2 (originally with Mac, then NT)
 - 1st was purchased in 1996
 - 2nd was purchased in June 2002
- ABI 3100 ([Data collection v1.0.1](#))
 - Purchased in June 2002
 - Original data collection software retained
- ABI 3130xl upgrade ([Data collection v3.0](#))
 - Purchased in April 2001 as ABI 3100
 - Upgraded to ABI 3130xl in September 2005
 - **Located in a different room**

Our Use of the ABI 3100

- Data collection software, version 1.0.1
- **POP-6** with 36 cm capillary array
- STR kits and in-house assays for autosomal STRs, Y-STRs, and miniSTRs
- SNaPshot assays for mtDNA SNPs, Y-SNPs, and autosomal SNPs
- DNA sequencing for mtDNA and STR repeat sequencing

We can routinely get more than 400 runs per capillary array by not changing the polymer between applications





Consumables for ABI 310/3100

What we use at NIST

- A.C.E.™ Sequencing Buffer 10X (Amresco)
 - \$155/L = \$0.0155/mL 1X buffer (**costs 20 times less!**)
 - <http://www.amresco-inc.com>
- 3700 POP-6 Polymer (Applied Biosystems)
 - \$530 / 200 mL = \$2.65/mL (**costs 20 times less!**)

What ABI protocols suggest

- 10X Genetic Analyzer Buffer with EDTA
 - \$78/25 mL = \$0.312/mL 1X buffer (ABI)
- 3100 POP-4 Polymer
 - \$365 / 7 mL = \$52/mL

2004 prices

Protocols Used for STR Typing

- Most forensic DNA laboratories follow PCR amplification and CE instrument protocols provided by the manufacturer
- Comments
 - Lower volume reactions may work fine and reduce costs
 - No heat denaturation/snap cooling is required prior to loading samples into ABI 310 or ABI 3100
 - Capillaries do not have to be thrown away after 100 runs
 - POP-4 polymer lasts much longer than 5 days on an ABI 310
 - Validation does not have to be an overwhelming task**

Reduced Volume PCR Amplifications

Advantages

- Lower cost** since kit contents are stretched
- Improved sensitivity perceived due to use of concentrated PCR products (since 1 uL out of a 5 uL reaction is 20% while 1 uL out of a 50 uL reaction is 2%)

Disadvantages

- Less volume of input DNA
 - Tighter control (improved precision) required in DNA quantitation**
 - If low amount of DNA, then potential for allelic dropout (LCN conditions)
 - If PCR inhibitor is present, then less opportunity for dilution of inhibitor
- Evaporation impacts PCR amplification performance

Publications:
 Gaines et al. *J Forensic Sci* 2002; 47(6):1224-1237. Reduced volume PCR amplification reactions using the AmpFISTR Profiler Plus kit.
 Leclair et al. *J Forensic Sci* 2003; 48(5):1001-1013. STR DNA typing: increased sensitivity and efficient sample consumption using reduced PCR reaction volumes.
 Fregoux et al. *J Forensic Sci* 2003; 48(5):1014-1034. AmpFISTR profiler Plus short tandem repeat DNA analysis of casework samples, mixture samples, and nonhuman DNA samples amplified under reduced PCR volume conditions (25 microL).

Identifiler 5 µL PCR Protocol

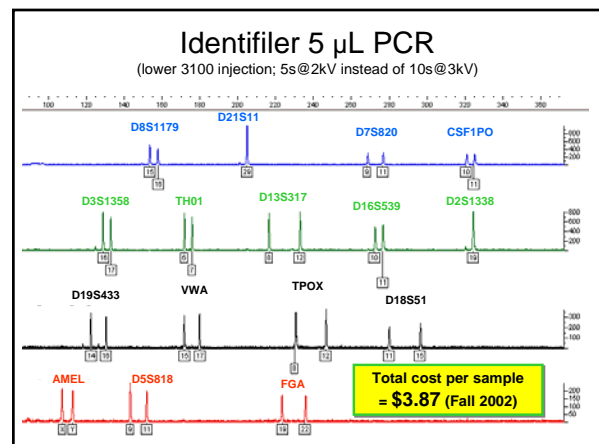
Identifiler PCR amplification was carried out on a GeneAmp® 9700 using 1 ng of DNA according to kit protocols with the exception of **reduced volume reactions** (5 µL instead of 25 µL) and **reduced cycles** (26 instead of 28).

Amplification products were diluted 1:15 in Hi-Di™ formamide and GS500-LIZ internal size standard (0.3 uL) and analyzed on the 16-capillary ABI Prism® 3100 Genetic Analyzer without prior denaturation of samples.

POP™-6 (3700 POP6) rather than POP™-4 was utilized for higher resolution separations.

Allele calls were made in Genotyper® 3.7 by comparison with kit allelic ladders using the Kazaam macro (20% filter).

Butler JM, Schoske R, Vallone PM, Redman JW, Kline MC. Allele frequencies for 15 autosomal STR loci on U.S. Caucasian, African American, and Hispanic populations. *J Forensic Sci* 2003; 48(4):908-911.



Maintenance of ABI 310/3100/3130

- Syringe – leaks cause capillary to not fill properly
- Capillary storage & wash – **it dries, it dies!**
- Pump block – cleaning helps insure good fill
- Change the running buffer regularly

YOU MUST BE CLEAN AROUND A CE!

Overall Thoughts on the ABI 310/3100/3130

- Settling on a common instrument platform has been good for the forensic DNA community in terms of data consistency (this is also true with the use of common STR kits)
- I am concerned that the community is very dependent primarily on one company...
- I really like using the instrument and can usually get nice data from it
- Like any instrument, it has its quirks...

Troubleshooting

Bruce McCord's Profiles in DNA Article

PROFILES IN DNA Volume 6 (2), Sept 2003, pp. 10-12

TECH TIPS

Troubleshooting Capillary Electrophoresis Systems

By Bruce McCord
Associate Professor of Forensic Chemistry, Ohio University, Athens, Ohio

The key to producing good DNA separations is to understand the principles underlying the injection, separation and detection of each allele.

INTRODUCTION

The development of capillary electrophoresis (CE) has played a key role in bringing about the modern application of DNA typing. Forensic laboratories are the beneficiaries of this new technology, but many practitioners are not fully aware of the underlying principles of the CE system. This article attempts to address the important issues in CE separations to aid analysts in troubleshooting problematic separations. The key to producing good DNA separations is to understand the principles underlying the injection, separation and detection of each allele. These points are addressed below.

SEPARATION

DNA analysis by CE is performed using entangled polymer buffers (Figure 1). These buffers can be easily pumped into a capillary prior to a separation and pumped out at its conclusion, providing a fresh separation matrix for each run. A typical buffer for forensic DNA separation contains 4% polydimethyl acrylamide (pDMA), buffered to pH 8

CE Troubleshooting Bruce McCord, AAFS 2006 Workshop (Seattle, WA)
February 20, 2006

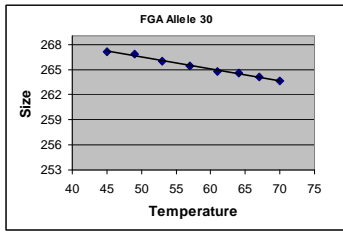
Outline for This Section

1. Chemistry/molecular biology problems – stutter, -A, degradation, inhibition, low copy #
2. Sample and buffer problems – formamide, urea, water, salt concentration, free dye ("dye blobs")
3. External factors – power supply, room temperature, cleanliness, voltage leaks
4. Instrument problems – optical system, capillary clogging, air bubbles, syringe leaks
5. Troubleshooting benchmarks/QC monitoring

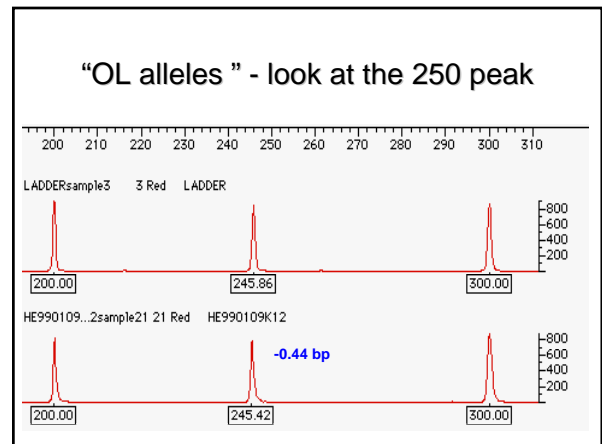
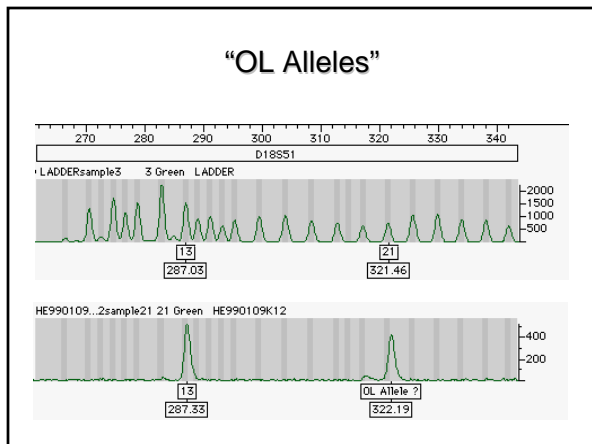
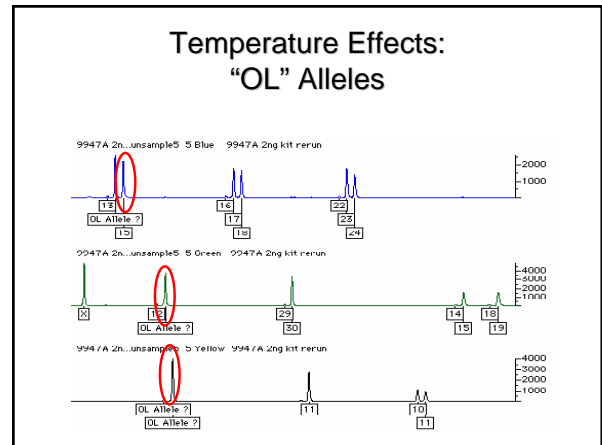
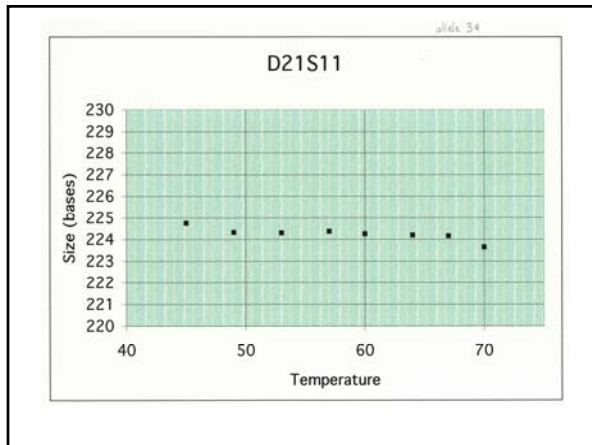
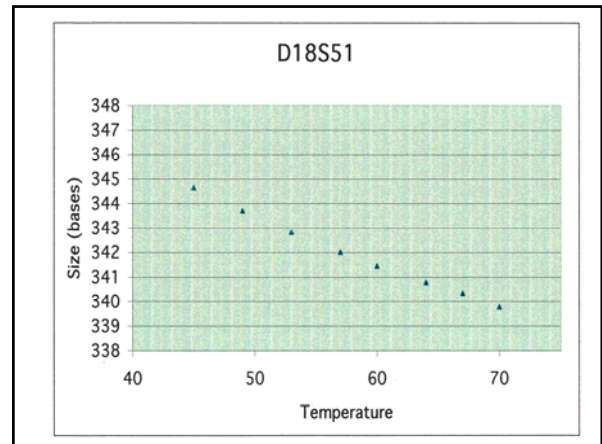
3. External Factors

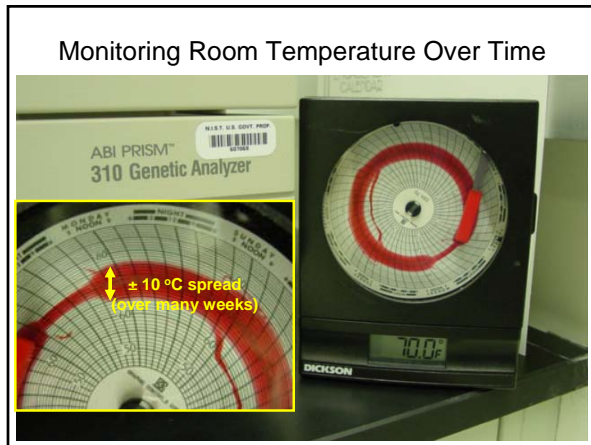
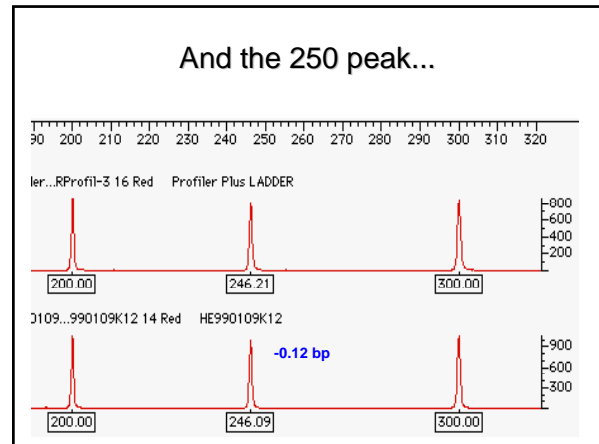
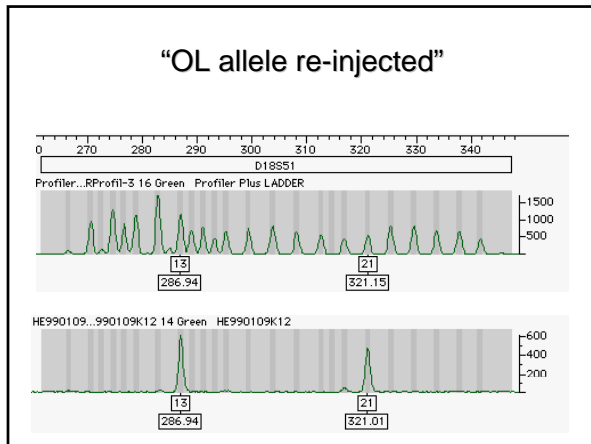
- Room temperature
 - Variations in room temperature can cause mobility shifts with band shifts and loss of calibration
 - Temperature is also important due to effects of high humidity on electrical conductance
- Cleanliness
 - Urea left in sample block can crystallize and catalyze further crystal formation causing spikes, clogs and other problems.
 - Best bet is to keep polymer in system and not remove or change block until polymer is used up.

Effect of Temperature on allele size



Slope is 0.14 bases/degree centigrade
 Therefore a small change in temperature has a big effect
 (A 1-2 degree shift in temperature of the heat plate can produce an OL allele)





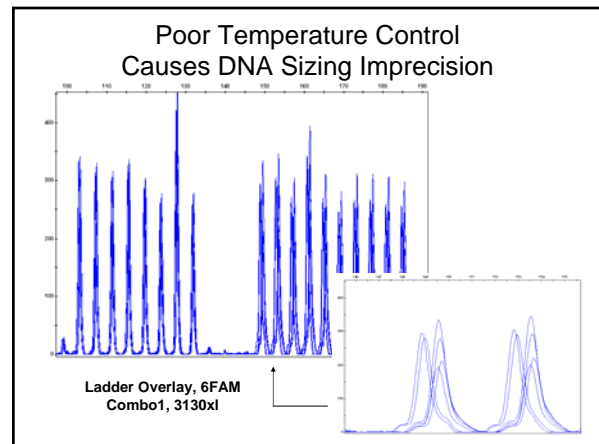
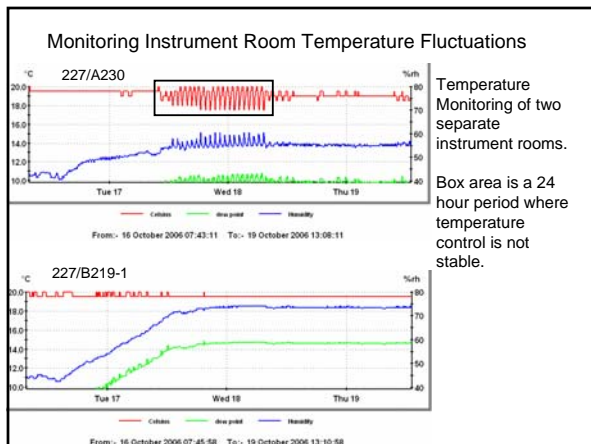
Temperature Probes

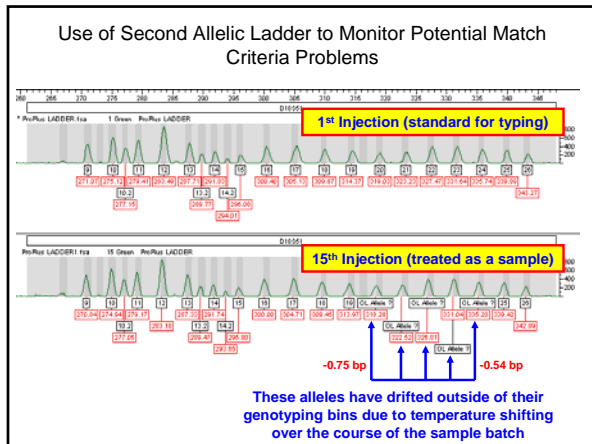
Refrigerator and freezer monitoring

Frig/Freeze Monitors \$240
#DT-23-33-80 – USB Temperature Datalogger
PLUS Software \$79.00 (#DT-23-33-60)
Room Monitors, # DT-23039-52 – USB Temperature-Humidity Datalogger \$91.00
(Cole Parmer, Vernon Hills IL)

Room temperature monitoring

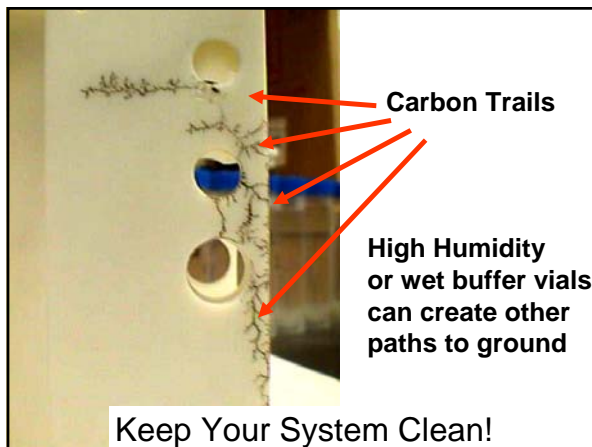
3130xl Genetic Analyzer





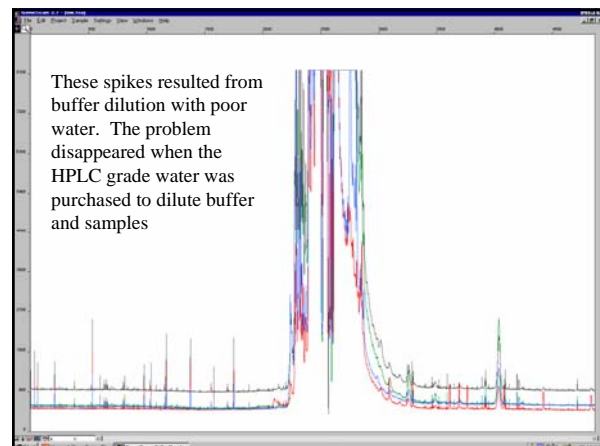
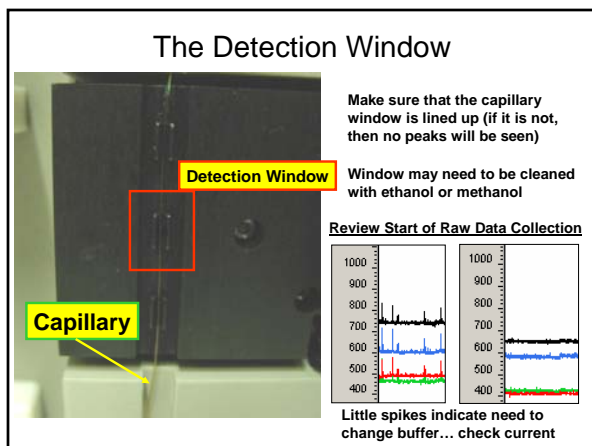
Cleanliness

- Urea sublimates and breaks down to ionic components - these find a path to ground
- Similarly wet buffer under a vial creates paths to ground
- Capillary windows must be clear or matrix effects will occur
- Laser will often assist in this process
- Vial caps will transfer low levels of DNA to capillary



4. Instrumental Factors

- Optical System
 - Sensitivity changes with age, capillary diameter, capillary cleanliness, instrument calibration
- Fluidic System
 - Effects of bubbles, dust, urea crystals, leaks in syringe and capillary ferrule
- Matrix Calculations
 - Changes in buffer, optics, sample dye can alter the software calibrations
- Capillary Problems
 - Chemisorbed materials on capillary surface can produce osmotic flow, DNA band broadening and inconsistent resolution (meltdowns)



Beware of Urea Crystals



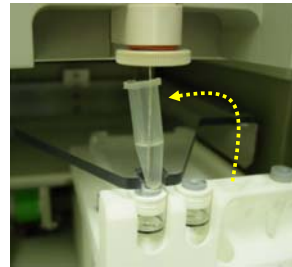
Urea crystals have formed due to a small leak where the capillary comes into the pump block

Urea sublimates and can evaporate to appear elsewhere

Use a small balloon to better grip the ferrule and keep it tight

Pump block should be well cleaned to avoid problems with urea crystal formation

Storage when ABI 310 is not in use



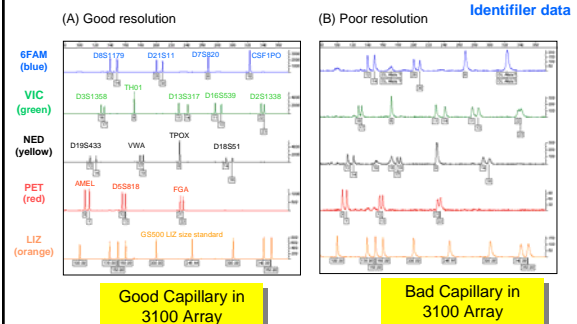
- Keep inlet of capillary in water...if it dries out then urea crystals from the polymer will clog the opening
- The waste vial (normally in position 3) can be moved into position
- A special device can be purchased from Suppelco to rinse the capillary off-line
- Store in distilled water
- Note that the laser is on when the instrument is on

Remember that the water in the open tube will evaporate over time...

Buffer Issues

- The buffer and polymer affect the background fluorescence- affecting the matrix
- Urea crystals and dust may produce spikes
- High salt concentrations may produce reannealing of DNA
- High salt concentrations affect current
- Low polymer concentrations affect peak resolution

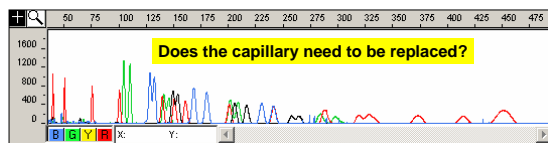
Capillary Meltdowns



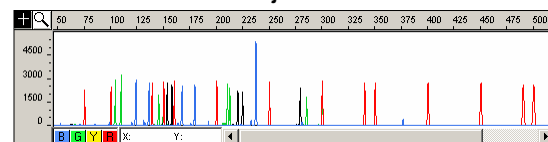
Butler, J.M., Buel, E., Crivellente, F., McCord, B.R. (2004) Forensic DNA typing by capillary electrophoresis: using the ABI Prism 310 and 3100 Genetic Analyzers for STR analysis. *Electrophoresis* 25: 1337-1412

Meltdowns can be permanent or transitory

as we have seen these may result from sample contamination effects



No! The next injection looks fine...



Meltdowns may be the result of

- Bad formamide
- Excess salt in sample/renaturation
- Water in the polymer buffer
- **Syringe leak** or bottom out
- Poisoned capillary
- Conductive polymer buffer due to urea degradation
- Crack/shift in capillary window
- Detergents and metal ions

5. Troubleshooting benchmarks

- **Monitor run current**
- Observe syringe position and movement during a batch
- Examine ILS (ROX) peak height with no sample
- Observe "250 bp" peak in GS500 size standard
- Monitor resolution of TH01 9.3/10 in allelic ladder and size standard peak shapes
- **Keep an eye on the baseline signal/noise**
- Measure formamide conductivity
- Reagent blank – **are any dye blobs present?**
- See if positive control DNA is producing typical peak heights (along with the correct genotype)

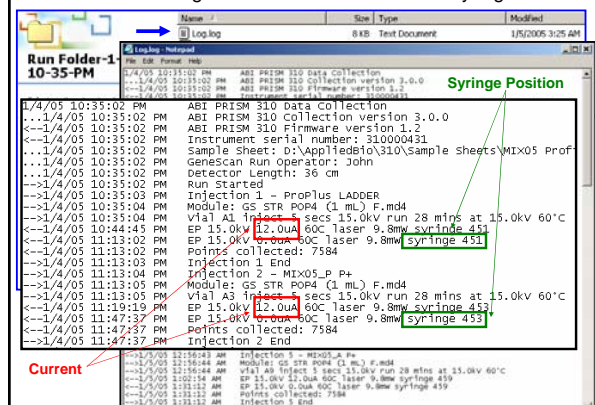
Measurement of Current

- $V/I = R$ where R is a function of capillary diameter, [buffer], and buffer viscosity
- In a CE system the voltage is fixed, thus changes in resistance in the capillary will be reflected in the current observed
- Air bubbles, syringe leaks, alternate paths to ground, changes in temperature, changes in zeta potential, and contamination, will be reflected in the current
- A typical current for a CE system with POP4 buffer is **8-12 μ A** (microamps)

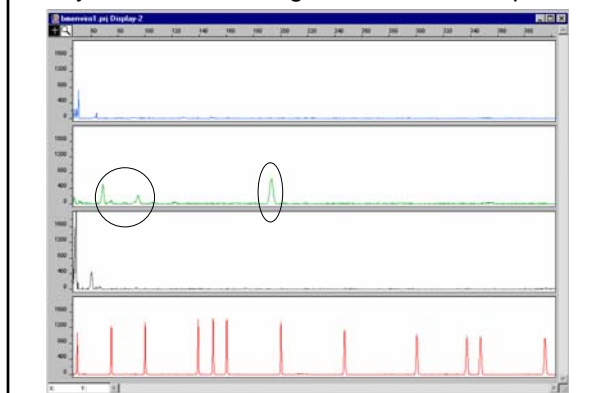
Syringe Travel

- The ABI 310 instrument also keeps track of the position of the syringe (in the log file)
- Depending on the resistance to flow, the syringe will travel different lengths
- Syringe leaks may be reflected in a longer distance traveled prior to each injection
- These leaks occur around the barrel of the syringe and at the connection to the capillary block

Use of ABI 310 Log File to Monitor Current and Syringe Travel



Dye Blobs in the Negative Control Sample



Measuring Formamide Conductivity



(not this way)

The key is to measure the bottle when it comes in or buy the good stuff and immediately pipette it into small tubes with or without ROX already added. Then freeze the tubes.

Do not ever open a cold bottle of formamide. Water will condense inside and aid in the formation of conductive formic acid.



Conclusion:
Troubleshooting is more than following the protocols

It means keeping watch on all aspects of the operation

1. Monitoring conductivity of sample and formamide
2. Keeping track of current and syringe position in log.
3. Watching the laser current
4. Watching and listening for voltage spikes
5. Monitoring room temperature and humidity

Multiplex_QA Article Published

Electrophoresis 2006, 27, 3735-3746 October 2006 issue of *Electrophoresis* 3735

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Received March 3, 2006
Revised April 21, 2006
Accepted May 11, 2006

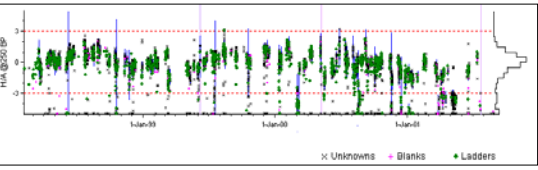
Research Article
Multiplex_QA: An exploratory quality assessment tool for multiplexed electrophoretic assays

Multiplex_QA is a data analysis tool for visualizing short- and long-term changes in the performance of multiplexed electrophoretic assays, particularly the commercial short tandem repeat (STR) kits used by the human forensic identity community. A number of quality metrics are calculated from the signal collected for the internal size standard included in nearly all multiplex assays. These quality metrics are related to the signal intensity, symmetry, retention, resolution, and noise of data collected by capillary electrophoresis systems. Interlocking graphical displays enable the identification of changes in the quality metrics with time, evaluation of relationships among the metrics, and detailed examination of electrophoretic features of particularly interesting analyses. While primarily intended for exploring which metrics are most useful for documenting data quality, the current version of the tool is sufficiently robust for use by forensic scientists with an interest in data analysis and access to a fast desktop computer.

Keywords: Electropherograms / Exploratory data analysis / Quality assessment / Resolution
DOI 10.1002/elps.200600116

User manual (127 pages) available for download from STRBase

Multiplex_QA Overview




- **Research tool** that provides quality metrics to review instrument performance over time (e.g., examines resolution and sensitivity using internal size standard peaks)
- Runs with Microsoft Excel macros. Requires STR data to be converted with NCBI's BatchExtract program into numerical form.

Available for download from STRBase:
<http://www.cstl.nist.gov/biotech/strbase/software.htm>

Acknowledgments

NIST Human Identity Project Team
Leading the Way in Forensic DNA...



John Butler (Leader) Margaret Kline Pete Vallone Jan Redman Amy Decker Becky Hill Dave Dwyer

Funding from interagency agreement 2003-IJ-R-029 between the National Institute of Justice and the NIST Office of Law Enforcement Standards

Many wonderful collaborators from industry, university, and government laboratories.

Bruce McCord (Florida International University) for many of the slides

Thank you for your attention...

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

Questions?



See also <http://www.dna.gov/research/nist>
<http://www.cstl.nist.gov/biotech/strbase>
john.butler@nist.gov