DNA Detection

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Background

In October 2023, Towson University was awarded a cooperative agreement from NIST to develop a standardized DNA training curriculum for the United States that address the components in ANSI/ASB Standard 115, *Standards for Training in Forensic Short Tandem Repeat Typing Methods Using Amplification, DNA Separation, and Allele Detection*. 2020. 1st Ed.

This presentation addresses the knowledge-based portion of the training program and covers the topic outlined in 4.2.3d in ANSI/ASB Standard 115.

Learning Objectives

- 1. history of DNA detection methods;
- 2. fluorescent dye detection;
 - i. excitation,
 - ii. emission;
- 3. dye-labeling of PCR primers;
- 4. computer software programs for DNA detection;
- 5. multicomponent analysis/spectral calibration/spatial calibration;
- 6. analytical threshold;
- 7. fragment sizing and allele calling;
- 8. bins (including virtual bins);
- 9. limitations of the technology.



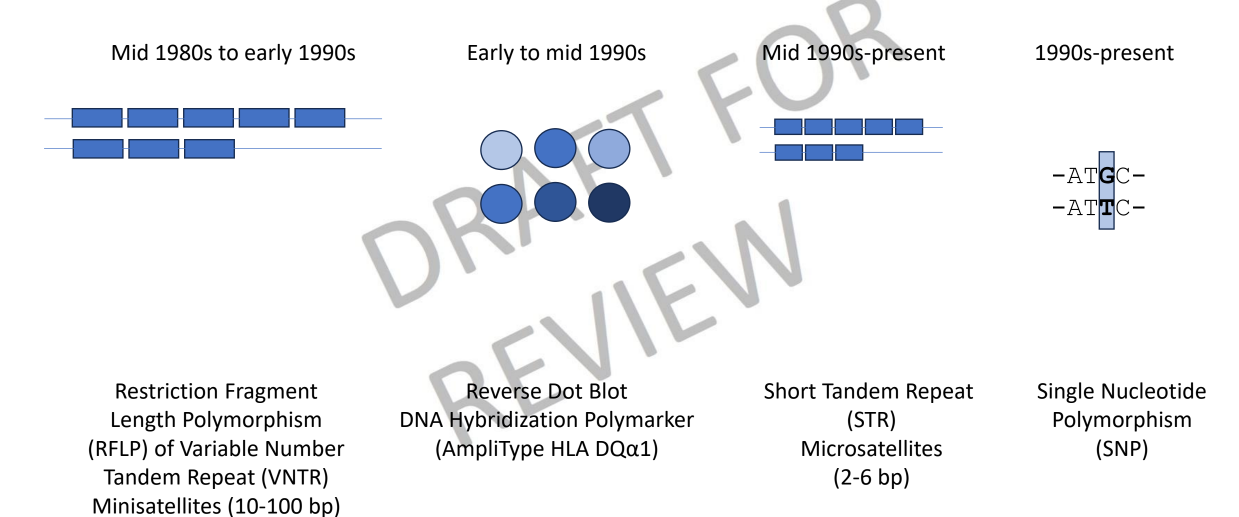
Terms & Definitions (ANSI/ASB 115)

- Analytical threshold. 1) The minimum height requirement at and above which detected peaks on a STR DNA profile electropherogram can be reliably distinguished from background noise; peaks above this threshold are generally not considered noise and are either artifacts or true alleles. 2) A "Relative Fluorescence Units" (RFU) level determined to be appropriate for use in the PCR/STR DNA typing process; a minimum threshold for data comparison is identified by the specific forensic laboratory through independent validation studies.
- Artifact. A non-allelic product of the amplification process (e.g., stutter, nontemplated nucleotide addition, or other non-specific product), an anomaly of the detection process (e.g., pull-up or spike), or a byproduct of primer synthesis (e.g., "dye blob") that may be observed on an electropherogram; some artifacts may complicate the interpretation of DNA profiles when they cannot be distinguished from the actual allele(s) from a particular sample.

Terms & Definitions (ANSI/ASB 115)

- **Bin.** Allele designations corresponding to the window of fragment sizes for each allele, determined by empirical testing.
- **Spectral calibration.** An examination of the contribution of overlap in the emission spectrum of fluorescent dyes used for a specific DNA test on a capillary electrophoresis instrument; permits the color deconvolution necessary for multi-color STR typing or sequencing to be performed; a poor spectral calibration may cause artifact peaks or inaccurate peak height determinations.
- **Stochastic.** 1) Chance, or random variation 2) in DNA testing, refers to random sampling error from extracts containing low levels of DNA and/or random variation in selection of alleles amplified at a particular locus.

History of DNA Detection Methods

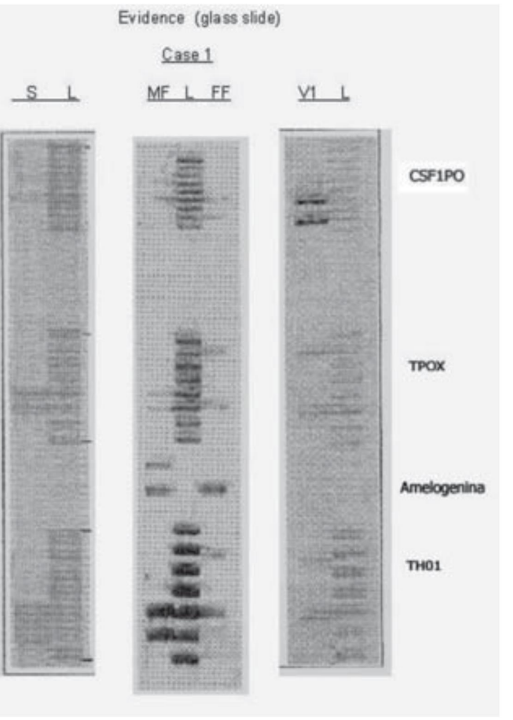


History of DNA Detection

- Followed RFLP and $\mathsf{DQ}\alpha$
- First STR kit introduced targeted TH01 in 1993 (Promega) with gel separation and silver stain detection
- CTT STR DNA typing kit (Promega, shown at right) was introduced in 1994
- Silver stain
 - Reduce silver ions (Ag⁺) from silver nitrate to metallic silver (Ag⁰) by alkaline formaldehyde
 - o LOD 2.5 ng (5x EtBr stain)
 - o Linear range 5-30 ng

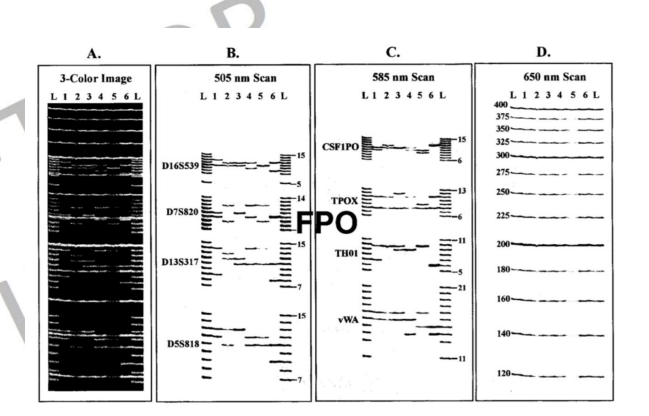
Gottlieb M, Chavko M. Silver staining of native and denatured eucaryotic DNA in agarose gels. Anal Biochem. 1987 Aug 15;165(1):33-7. doi: 10.1016/0003-2697(87)90197-7.

da Silva DA, Góes AC, de Carvalho JJ, de Carvalho EF. DNA typing from vaginal smear slides in suspected rape cases. Sao Paulo Med J. 2004 Mar 4;122(2):70-2. doi: 10.1590/s1516-31802004000200008. Figure CC BY 4.0 license



History of DNA Detection

- AB 310 CE instrument was introduced in 1995
- Detection using fluorescent dyes was introduced in 1996
 - PowerPlex[™] System (Promega)
 - 8 loci using 3 dyes
 - Fluorescein (green)
 - TMR (red)
 - CXR (blue)
 - AmpFlSTR® Blue (Applied Biosystems)
 - 3 loci using 1 dye

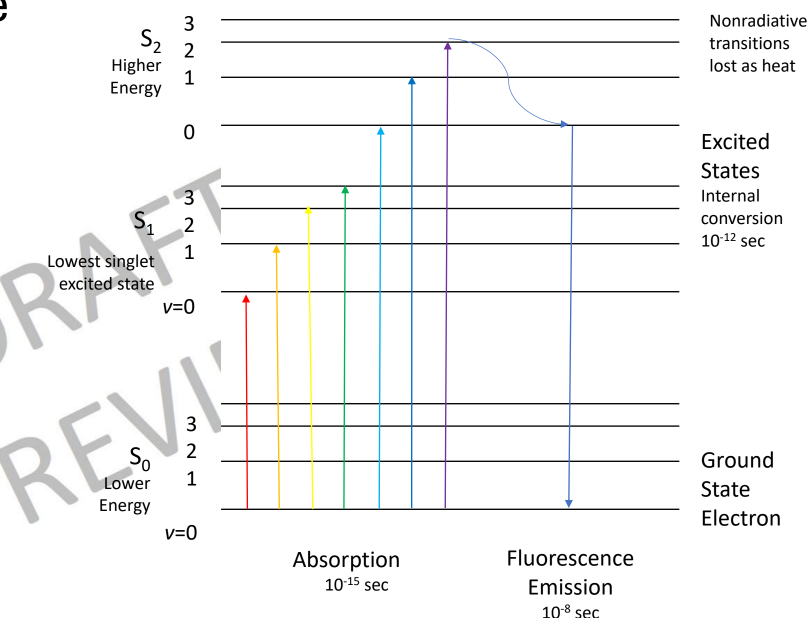


Lins, A., Micka, K., Sprecher, C., Taylor, J., Bacher, J., Rabbach, D., Bever, R., Creacy, S., and Schumm, J. "Development and Population Study of an Eight-Locus Short Tandem Repeat (STR) Multiplex System." ASTM International. *J. Forensic Sci.*. November 1998; 43(6): 1168– 1180. <u>https://doi.org/10.1520/JFS14381J</u>

Jablonski Diagram

Fluorescence Dye Detection

- **Excitation** electron in fluorophore boosted by laser or LED energy to an excited state
- Emission- energy emitted by fluorophore as electron relaxes to ground state following nonradiative transitions resulting in a longer wavelength than excitation



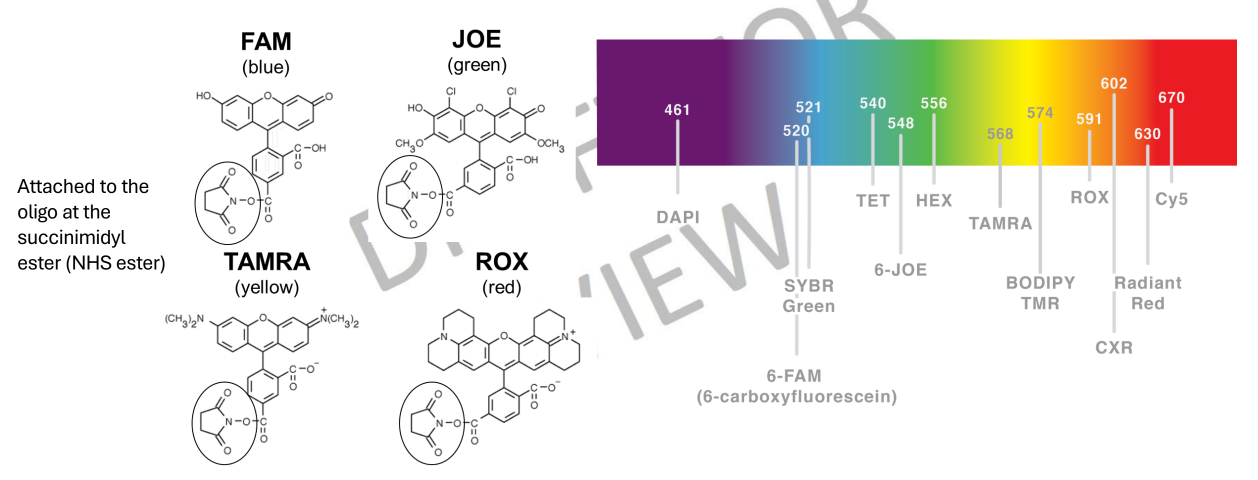
Fluorescence Detection

- Many molecules absorb UV-Vis radiation
- Some molecules fluoresce
- Fluorescent molecules (e.g., dyes) are used for DNA detection
 - o Absorptions 200-700 nm range
 - Emissions at longer wavelengths in the long UV or visible range
 - $\,\circ\,$ Peaks shift with pH changes
- Light is emitted by fluorophore

 Nanomolar detection limit
- Detected by fluorimeter or fluorescence spectrometer charge coupled device (CCD) detector

Fluorescent Dyes

Absorption and emission maxima can be shifted to the red by additional π bonds conjugation and substituent groups



Credit Anna-Sewell Killingstad

Fluorescent Dye Detection

Dye	Colors	Excitation (nm)	Emission (nm)	
DAPI	Purple	359	461	
SYBR Green	Blue-Green	494	521	
6-FAM (6-carboxyfluorescein)	Blue-Green	495	520	
6-JOE	Green	520	548	
TET	Green	520	540	
HEX	Green-Yellow	530	556	
BODIPY TMR	Orange	542	574	
TAMRA	Yellow	542	568	
ROX	Red	567	591	
CXR	Red	580	602	
Radiant Red	Red	495	630	
Cy5	Red	650	670	
Proprietary Dyes: AQA, BTG, BTR2, E	STP. Data: https://	have his rad com (webroat (web/pd	f/ler/literature/Bulletin 2421 pdf	

Proprietary Dyes: AQA, BTG, BTR2, BTP, BTY, CCO, TOM, WEN

Data: https://www.bio-rad.com/webroot/web/pdf/lsr/literature/Bulletin_2421.pdf

DAPI, SYBR, Radiant and Cy5 are not in STR kits but are common dyes in educational settings

Fluorescent Dye Detection

100

90

80

50

40

30

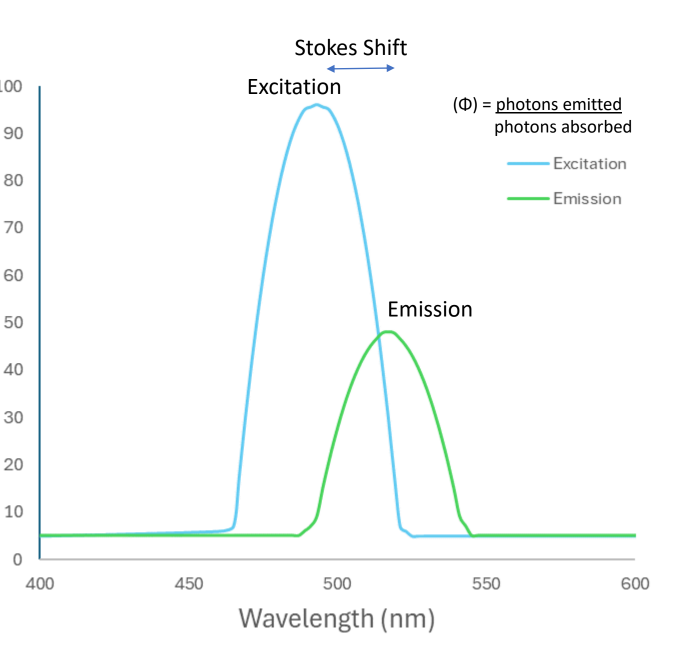
20

10

0

RFL

- Molar absorptivity (ϵ) Measure of how strongly the fluorophore absorbs light
- Quantum Yield (Φ)
 - Ratio of photons emitted/absorbed
 - Measure of the efficiency of converting absorbed photons to light emission
 - Higher quantum yield leads to higher signal intensity • Range 0 to 1.0 (100%)
- Highest sensitivity at absorption/emission maxima • LOD 125 pg

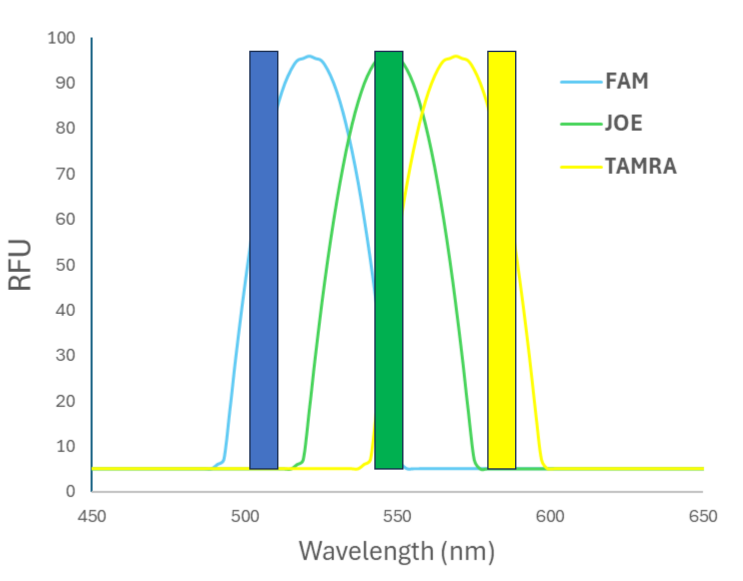


Fluorescent Dye Detection

- Dye overlap
- Excitation

 532 nm
- May choose to detect outside of emission maxima
 - \circ Detection
 - Loss of sensitivity
- Software deconvolution

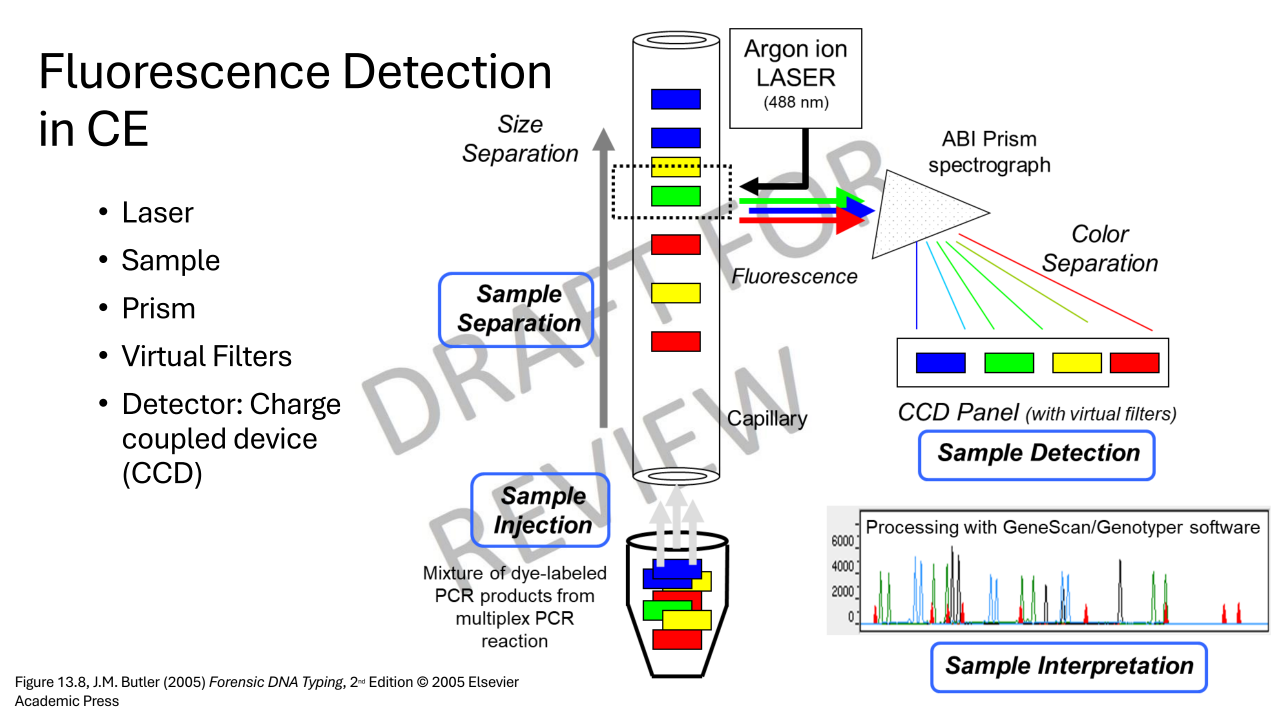
 Emission filter



Dye Labelling of PCR Primers

3'

- Dyes can be attached to the 3' and 5' ends of oligos
- However, PCR generates a labeled amplicon so the 5' end must be the labeled end to allow 3' extension
- Modern STR kits have up to 8 dye detection for use with CE



Computer Software Programs for DNA Detection

- Applied Biosystems Data Collection and secondary analysis software
 - \circ GeneMapper ID
 - \circ GeneMapper ID-X
- Promega Spectrum Software
- Open Source and Independent Review Interpretation System (OSIRIS)

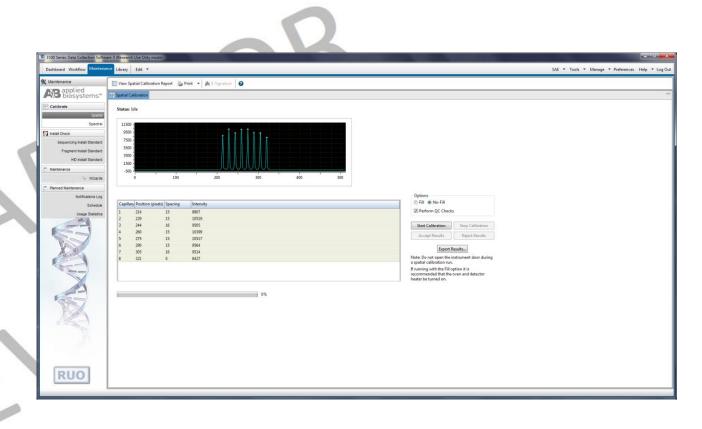
Number of Dyes Detected in CE Instruments



Capillary Electrophoresis Instrument	Number of Dyes	Number of Capillaries	Manufacturer
310	4	1	Applied Biosystems
3100	5	16	Applied Biosystems
3100-Avant	5	4	Applied Biosystems
3130	5	4	Applied Biosystems
3130xl	5	16	Applied Biosystems
3500	6	8	Applied Biosystems
3500xl	6	24	Applied Biosystems
3730	5	48	Applied Biosystems
3730xl	5	96	Applied Biosystems
SeqStudio	6	4	Applied Biosystems
SeqStudio 8 Flex	6	8	Applied Biosystems
SeqStudio [™] 24 Flex	6	24	Applied Biosystems
Spectrum Compact SE	6	4	Promega
Spectrum SE	8	8	Promega

Multicomponent Analysis

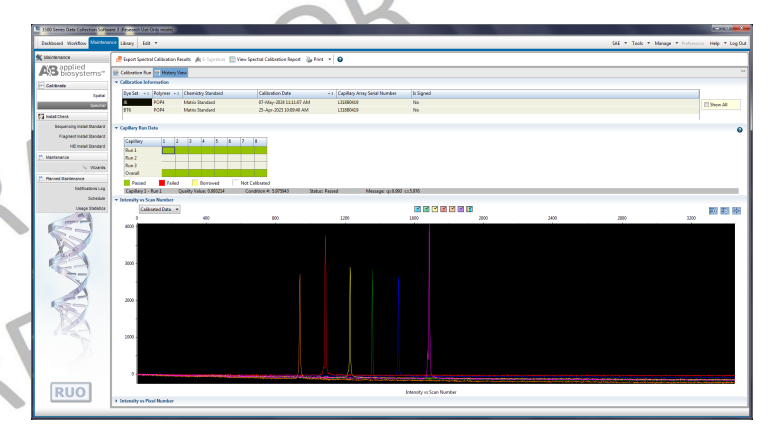
- Spectral Calibration: Color deconvolution
- Spatial Calibration: Physical space
 - Determine relationship between elution of the dye from the capillary and position of the signal as detected by the CCD camera



Van Orden A, Keller RA. Fluorescence correlation spectroscopy for rapid multicomponent analysis in a capillary electrophoresis system. Anal Chem. 1998 Nov 1;70(21):4463-71. doi: 10.1021/ac980768q.

Spectral Calibration: 6 Dye

- Spectral calibration
 - Injection of dyes into capillary
 - Separate emission files are created for each capillary in an array
 - Must be performed for each dye set



J6 on AB 3500 CE

Spectral Calibration: 8 Dye

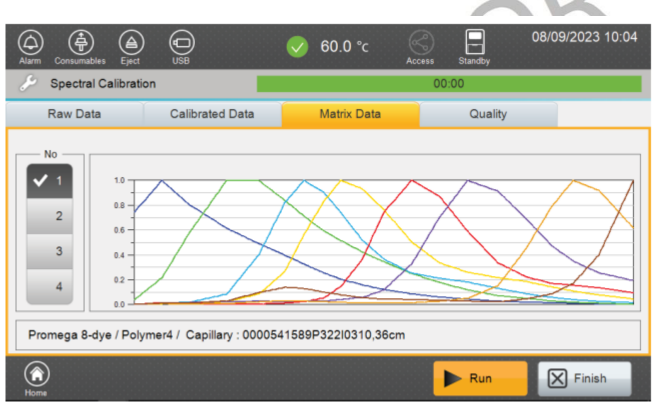
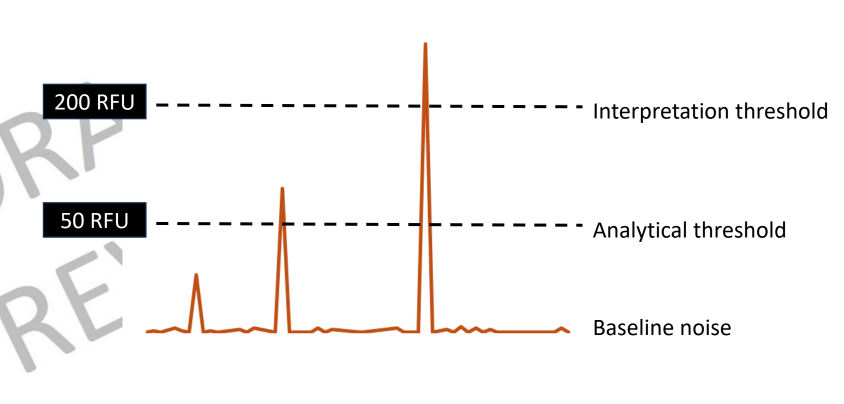


Figure 12. Spectral Calibration 'Matrix Data' tab screen.

https://www.promega.com/-/media/files/resources/protocols/technical-manuals/tmd/tmd062-powerplex-matrix-standards-for-use-on-the-spectrum-compact-ce-system-technical-manual.pdf?rev=2aba86082a804beaa73fe2d32b252ffb&sc_lang=en

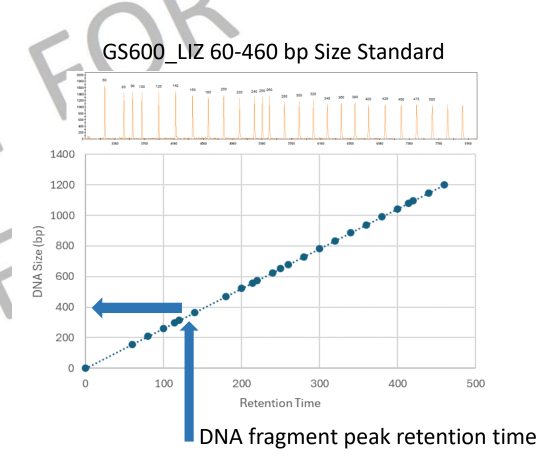
Analytical Threshold

- Peaks above the interpretation threshold reproducible and reliable and are called
- The analytical threshold is the limit of detection (LOD) and peaks above can be used for exclusion
- A peak or baseline noise below the analytical threshold is stochastic and not reliable for interpretation



Fragment Sizing using an Internal Standard

- Internal size standard is added to the sample prior to separation
 - $\,\circ\,$ DNA fragments of known size
- Sizing curve is constructed
 - Plot of DNA size vs. migration time for the standard fragments
 - A regression line is constructed and fit is determined
 - In the Global Southern Method, the slope is used to compute size of sample fragments using their migration times
 - Forensic STR size calculations use Local Southern Method sizing in which the peaks in the size standard above and below the peak of interest are used to compute the size of the sample peak



Allele Calling using a Ladder

- The allele ladder is a mixture of DNA fragments of known size and sequence.
- It includes known and most common alleles for each locus.
- An internal size standard (bottom panel, orange, known sizes) is included to correlate the migration time of the ladder peaks to the sample peaks



GlobalFiler Ladder and Internal Size Standard

Bins and Virtual Bins

- Bin (allele bin)
 - $\,\circ\,$ Region that defines an allele within a locus
 - \circ Sensitivity to within ±1 bp
 - $\,\circ\,$ Resolution to ±0.1 nt

• Physical Bin

• Region physically defined by the allele ladder

Virtual Bin

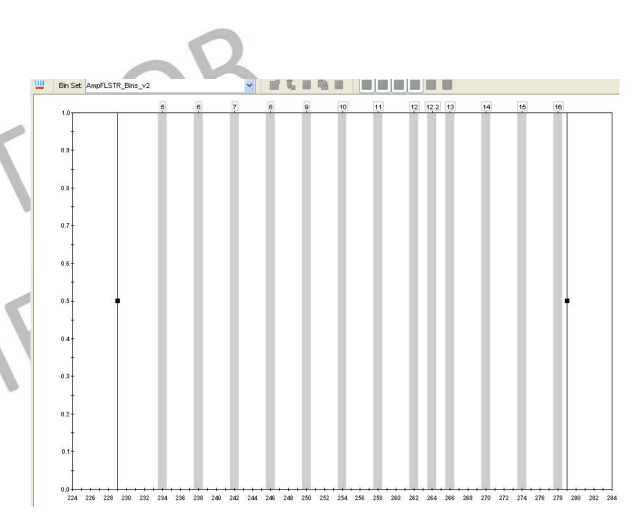
 Alleles not present in the ladder but that have been reported in the literature

• Bin Offset

 $\,\circ\,$ Size difference between the physical bin and allele ladder fragments

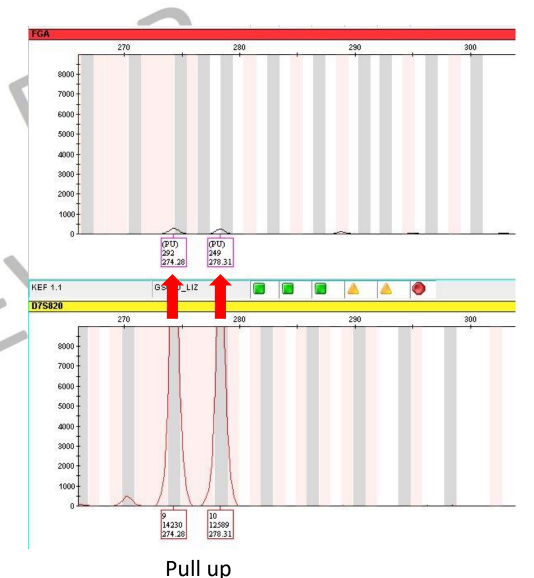
Bin Overlap

 Rerun the ladder or replace with one that did not produce overlap



Limitations of the Technology

- Range of visible spectrum wavelengths available limits the range of available colors
- Spectral overlap is not fully removed by matrix deconvolution
- Spectral overlap can lead to pull-up in dye channels
- Size but not sequence is determined



Study Questions

- Define fluorescence.
- What methods have been used to detect DNA?
- How can DNA be detected using fluorescence?
- What is a spectral calibration and why is it performed?
- Define analytical threshold.
- Define interpretation threshold.
- What is the allele ladder used for?
- How are alleles detected and called?
- Define a bin and virtual bin.
- Using fluorescent STR allele detection technology, how many reaction primers are labeled and what are the dye labels and colors?
- What are limitations of using fluorescence to detect DNA?

Suggested Readings

- ANSI/ASB Standard 115, Standard for Training in Forensic Short Tandem Repeat Typing Methods using Amplification, DNA Separation, and Allele Detection. 2020.
 1st Ed. <u>https://www.aafs.org/sites/default/files/media/documents/115_Std_e1.pdf</u>
- Butler, J.M. <u>Advanced Topics in Forensic DNA Typing: Methodology, Elsevier,</u> 2011.
- FBI, Quality Assurance Standards for Forensic DNA Testing Laboratories (QAS), effective September 1, 2011.