STRmix v 2.7 Probabilistic Genotyping Standard Operating Instruction		perating Instructions
Status: Published Document II		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	1 OF 36

STRmixTM v 2.7 Probabilistic Genotyping Software Operating **Instructions**

1 **Guiding Principles and Scope**

- This procedure describes the use of STRmixTM v2.7 for the interpretation of PowerPlex® Fusion 1.1 DNA profiles run on 3500xL or 3130xL Genetic Analyzers within the NYC OCME Department of Forensic Biology. Readers are also referred to the STRmixTM v2.7 Users and Operation manuals for additional information.
- 1.2 For STRMixTM set-up instructions please refer to QC702a A STRmixTM v2.7 Set-Up Instructions.
- 1.3 For STRmixTM set-up instructions for the QC Monitoring Program refer to QC702b.

Preparing Data for a STRmixTM Analysis 2

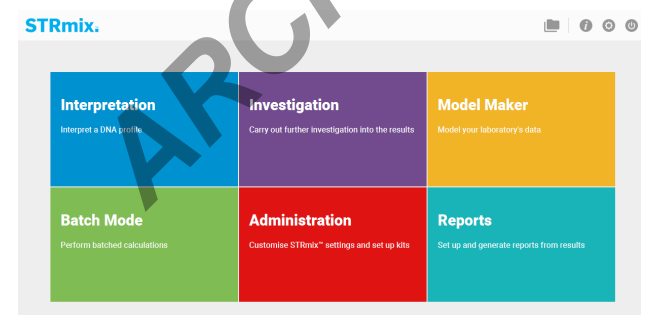
- Before performing your STRmixTM analysis, the following actions must be taken: 2.1
 - Verify that the sample is suitable for STRmixTM analysis (refer to Interpretation of PowerPlex[®] 2.1.1 Fusion data run on 3500xL).
 - Evaluate your replicates, if applicable. If there are drastic inconsistencies with the alleles 2.1.2 present between replicates and/or one has little information, only the amplification with the most information should be used, or a third amplification may be warranted.
 - 2.1.3 Determine the best described Number of Contributors to the sample (NOC). Refer to Interpretation of PowerPlex® Fusion data run on 3500xL regarding the procedure for determining the number of contributors.
 - Create folder(s) with the FB (or FBS) case number for the STRmixTM runs within the STRmix 2.1.4 Data folder: M:\STR Data\STRmix Data.
 - e.g. FB16-01234 or FBS16-05678
 - Confirm that the STR data is prepared correctly for STRmixTM analysis: 2.1.5
 - 2.1.5.1 Evidence samples must only be amplified in PowerPlex® Fusion in order to undergo STRmixTM analysis. This procedure is specifically for evidence amplified in PowerPlex® Fusion data run on the 3500xL or 3130xL Genetic Analyzers at the NYC OCME Department of Forensic Biology.

	STRmix v 2.7 Probabilistic Genotyping Standard Operating Instruction		perating Instructions
Status: Published Document ID: 5702		Document ID: 57029	
	DATE EFFECTIVE	APPROVED BY	PAGE
	10/11/2024	Nuclear DNA Technical Leader	2 OF 36

- 2.1.5.2 Sample data must be assembled into the appropriate format for STRmixTM input. The standard input for STRmixTM is a .txt file. See the <u>GeneMarker v3.0 Operation Manual</u> for instructions on exporting data for STRmixTM input.
- 2.1.5.3 Evidence samples must be edited to remove all artifacts, including pull-ups, spikes, dye blobs, etc. before being input into STRmixTM. Back and forward stutter in 3130xL data should not be removed, and back, forward, half back, and double back stutter in 3500xL data should not be removed before importing into STRmixTM. Refer to STR Results Interpretation PowerPlex® Fusion & STRmixTM, Interpretation of PowerPlex® Fusion data run on 3500xL, and the Appendix for PowerPlex® Fusion Stutter.
- 2.1.5.4 Reference samples must be edited to remove all artifacts including stutter. Incomplete or tri-allelic loci must not be imported into STRmixTM for a reference sample remove all allele(s) for that locus within the text file. If a possible drop-in peak is present in a reference sample, remove this peak from the text file before STRmixTM import.
- 2.1.5.5 Non-numeric values such as OL or OB, < or > are not permitted within the STRmixTM input files. Unambiguous alleles including those that are rare should appear in the corresponding input file as their actual allelic size designation, for example D21: 30.1. If an actual allelic size designation cannot be determined, the data for this locus should be removed completely from the text file and the locus should be ignored.
- 2.1.5.6 To modify a STRmixTM input text file: open the STRmixTM .txt file associated with the appropriate STR project (e.g. in Notepad or Microsoft Excel®). Locate the sample and locus containing the non-numeric value within the .txt file and manually replace the value with the appropriate actual allelic size designation. Save the .txt file **replacing** the original file.
- 2.1.5.7 An attempt should be made to amplify reference samples in PowerPlex® Fusion (see Case Management). If unavailable, STRmixTM allows the user to calculate a likelihood ratio when the evidence and reference samples are analyzed in different autosomal typing kits. LRs will only be calculated for those loci in common between the two kits.
- 2.1.5.7.1 If a reference sample was not amplified in Fusion, the data must be converted to a .txt file for import into STRmixTM with the locus order matching that of the evidence (PowerPlex[®] Fusion order). The reference sample data can be converted to the proper PowerPlex[®] Fusion order and appropriate .txt file format using the following macro: Identifiler to Fusion Exemplar STRmix Input Creation.
- 2.1.5.8 If a DNA donor is being used from one sample to condition or compare to another, use the following macro: Reference profile for STRmix Input Creation.

STRmix v 2.7 Probabilistic Genotyping Standard Operating Instruction		perating Instructions
Status: Published Document II		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	3 OF 36

- 2.1.6 Evaluate evidence samples to determine if a locus needs to be ignored before STRmixTM deconvolution is performed. A comment should be added to the **Case Notes** field within the STRmixTM analysis to indicate why the locus was ignored for that run.
 - 2.1.6.1 The following is a list of reasons data may need to be ignored at a particular locus. For any situation not covered in the list below, the technical leader should be consulted.
 - Tri-allelic pattern
 - Unresolved allelic or stutter peak that is visible above the AT
 - OB/OL allele or stutter peak that cannot be assigned a correct allelic designation
 - Stutter or allelic peak for an allele belonging to locus A is being called in a neighboring locus B (ignore both loci)
 - Where a conditioning sample does not have data at a locus that is present in the evidence sample
- 2.1.7 Loci should not be ignored for likelihood ratio calculations due to a partial comparison sample. Ensure that your reference sample text file has been updated appropriately as described above in 2.1.5
- 2.2 Launch the STRmixTM application and prepare the scenarios to be run in STRmixTM. Open the STRmixTM software by locating STRmixTM in the task bar or by double clicking on the STRmixTM icon on the desktop. The main menu is shown below:



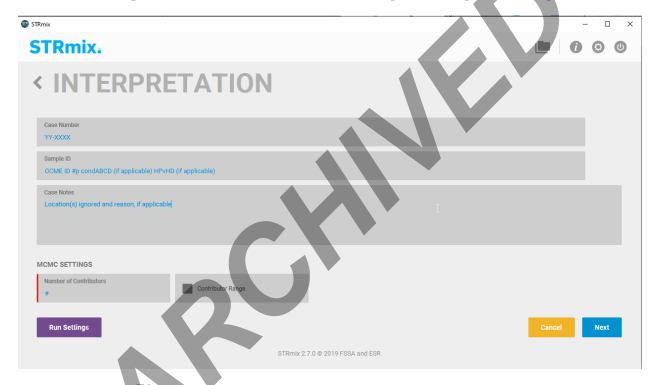
2.2.1 For deconvolutions of evidence profiles (Interpretation), go to Section 3.

STRmix v 2.7 Probabilistic Genotyping Standard Operating Instruction		perating Instructions
Status: Published Document ID: 57		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	4 OF 36

- 2.2.2 For comparisons to previously deconvoluted profiles (Investigation: LR From Previous), go to Section 4.
- 2.2.3 To set up multiple STRmixTM runs to run consecutively in Batch Mode, go to Section 6.

3 Deconvolutions with STRmixTM (Interpretation)

3.1 Select **Interpretation** from the **Main Menu.** This will open the Interpretation Setup screen:



- 3.2 The STRmixTM output folder and file names are created by stringing together the values in the Case Number and Sample ID fields in the software followed by the date and time of the analysis run. The information in the file name is separated by dashes. Therefore, if other characters are entered, such as a comma, underscore, period, etc., the software will convert them into dashes.
- 3.3 Refer to Interpretation of PowerPlex® Fusion data run on 3500xL for guidance on when a conditioned contributor may be applied. A deconvolution of the evidence sample without conditioning and an LR against a potential conditioned contributor may need to be run first.
- 3.4 An LR may be run in conjunction with the deconvolution in the following scenarios:
 - Single source evidence sample that did not require a STRmixTM deconvolution for determination of a profile, needing an LR to a matching comparison sample.

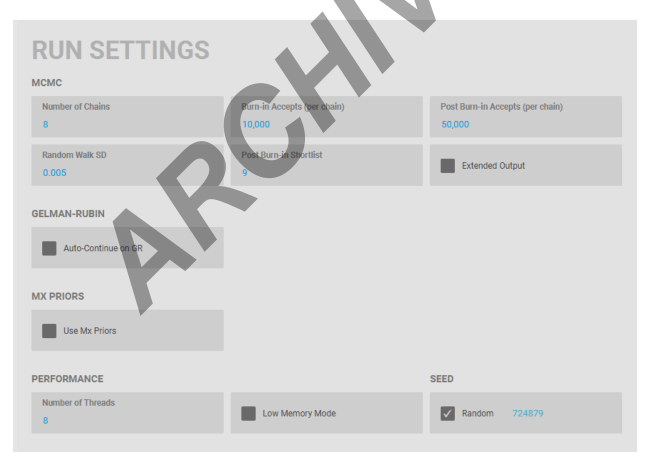
STRmix v 2.7 Probabilistic Genotyping Standard Operating Instructions		perating Instructions
Status: Published Document ID: 5702		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	5 OF 36

- Evidence mixture sample undergoing deconvolution and an LR check to determine if a reasonably expected reference sample can be used for further conditioning (ex. car owner on a swab from the steering wheel).
- To assess a probative comparison sample's presence within a mixture where that reference sample data is contained within the same evidence file (ex. victim's blood on suspect's clothing.)
- 3.5 The following naming conventions should be used for deconvolutions:
 - 3.5.1 Evidence Files
 - Case Number = YY-XXXXX (do not include "FB")
 - Sample ID = remainder of evidence sample OCME ID* #p (NOC) condElimInitials (if applicable) scenario for LR (if applicable)
 - Case notes = a comment should be added here if a locus is ignored, indicating the reason: e.g. "D2S441 was ignored due to an unresolved allelic peak"
 - *Suffixes such as '_mcon' or '_reamp' should not be included in the OCME ID
 - 3.5.2 Suspect Files
 - Case Number = SYY-XXXXX (do not include "FB")
 - Sample ID = evidence sample OCME ID* (include evidence file FB# without the "FB") #p (NOC) condElimInitials (if applicable) scenario for LR
 - Case Notes = a comment should be added here if a locus is ignored, indicating the reason: e.g. "D2S441 was ignored due to an unresolved allelic peak"
 - Suffixes such as 'mcon' or 'reamp' should not be included in the OCME ID
 - 3.5.3 For LR scenarios, the naming format should start with the comparison sample's initials, followed by any conditioned samples' initials, and then the number and "U" for unknowns, followed by a "v" to separate the numerator hypothesis from the denominator hypothesis.

Examples	Resulting STRmix [™] file name
Evidence File	
3-person, deconvolution, no conditioning, no comparisons	22-01234-567-1-1.1-trig-GS-3p
4-person deconvolution, no conditioning, comparing elimAB	22-01234-567-1-1.1-shirt-BL-4p-AB3Uv4U
3-person deconvolution, conditioning vic CD, comparing elimAB	22-01234-567-1-1.1-shirt-BL-3p-condCD-ABCD1UvCD2U
4-person deconvolution, conditioning elims CD, EF and GH, no comparisons	22-01234-567-1-1.1-trig-GS-4p-condCDEFGH
Suspect File	
1-person deconvolution, no conditioning, comparing suspTS	S22-05678-22-01234-567-2-1.1-slide-GS-1p-TSv1U

STRmix v 2.7 Probabilistic Genotyping Standard Operating Instruction		perating Instructions
Status: Published Document ID: 5		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	6 OF 36

- 3.6 Select Run Settings, and confirm the settings against the following screenshot. They should always be the same for every STRmixTM analysis unless an exception is listed below. Any changes that are made will appear in bold on the run report.
 - MCMC Settings: Burn-in Accepts (per chain) and Post Burn-in Accepts (per chain)
 must not be modified without documented approval from the technical leader (or his/her
 designee).
 - Mx Priors: this will not be modified without documented approval from the technical leader (or his/her designee). See section 3.8 for more information regarding Mx priors.
 - **Performance, Number of Threads**: it is okay to proceed if the Number of Threads does not match the screenshot below; this is specific to the computer being used.
 - **Performance, Low Memory Mode**: This setting allows the computer to minimize the memory used for the run and can be turned on in order to use less computer memory (ex. if using your computer for other tasks while running STRmixTM in the background) or if a run fails to finish due to computer memory. Using this setting will increase the run time.

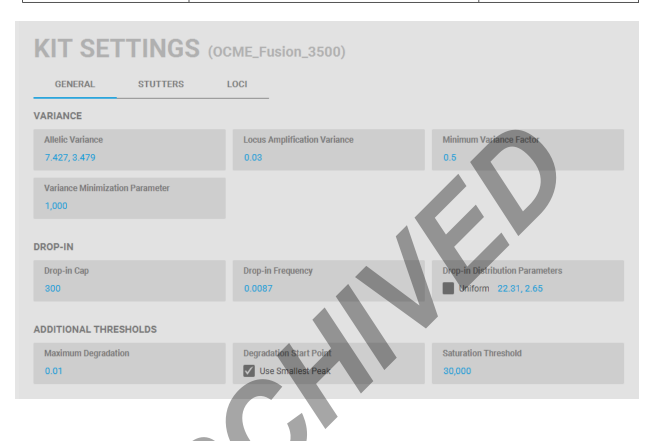


STRmix v 2.7 Probabilistic Genotyping Standard Operating Instruction		perating Instructions
Status: Published Document ID: 570		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	7 OF 36

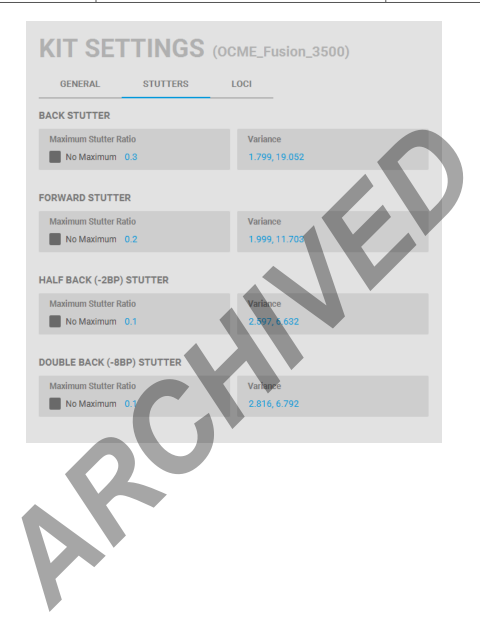
- 3.7 If you are not adjusting Mx Priors settings, skip to **3.8**; User informed Mx Priors is a function within STRmixTM that allows analysts to set approximate mixture proportion percentages for each contributor. If a proposed genotype does not fit the proportion percentage and variance set by the user, a penalty is applied to that iteration indicating an overall poor fit to the observed profile. Mx Priors settings are adjusted only with documented approval from the technical leader (or his/her designee). An analyst must show the Technical leader (or designee): 1) STRmix run data where a sample shows counterintuitive information in the output report and 2) the observed sample profile from the electropherogram. In addition, other troubleshooting techniques must be considered and tried prior to seeking approval for applying Mx Priors. See section **3.20** for set-up instructions for this function.
- 3.8 Select if all run settings are correct. If approved changes were made, Select Apply
- 3.9 Set the number of contributors and ensure that contributor range remains unchecked. Select
- 3.10 Check that the **Profiling Kit** selected is **OCME_fusion_3500** or **OCME_fusion_3130_2_7 depending on the data being analyzed**. The STRmixTM output folder and file name (based on the combined Case Number and Sample ID you previously entered) are located at the top right. The kit labeled OCME Fusion must not be used for deconvolutions in STRmixTM v2.7.
- 3.11 Click on Kit settings. There are three tabs of settings to verify against the following screenshots in this window General, Stutters, and Loci. If a locus needs to be ignored for a deconvolution, this is where you will be able to do that. See 3.133.12 for further instruction on ignoring a locus.
 - 3.11.1 For the OCME fusion 3500 Profiling Kit, ensure settings match the following:



STRmix v 2.7 Probabilistic Genotyping Standard Operating Instructions		perating Instructions
Status: Published Document ID: 570		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	8 OF 36



	STRmix v 2.7 Probabilistic Genotyping Standard Operating Instructions		perating Instructions
Status: Published		Document ID: 57029	
	DATE EFFECTIVE	APPROVED BY	PAGE
	10/11/2024	Nuclear DNA Technical Leader	9 OF 36

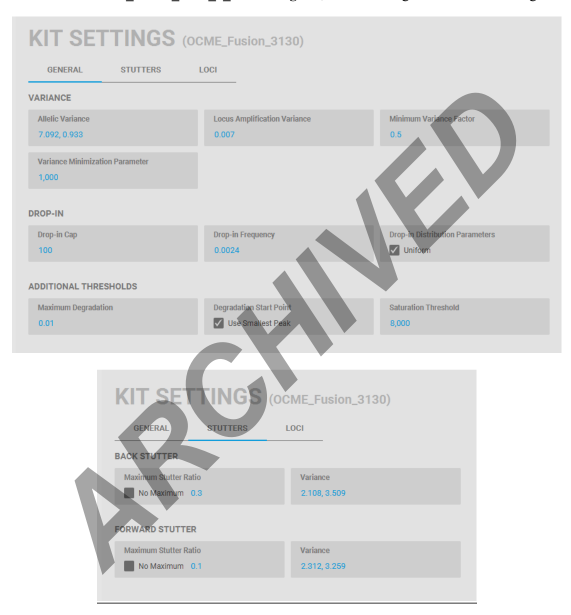


	STRmix v 2.7 Probabilistic Genotyping Standard Operating Instructions		
Status: Published Document ID:		Document ID: 57029	
	DATE EFFECTIVE	APPROVED BY	PAGE
	10/11/2024	Nuclear DNA Technical Leader	10 OF 36



STRmix v 2.7 Probabilistic Genotyping Standard Operating Instructions		
Status: Published Document ID: 57029		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	11 OF 36

3.11.2 For the OCME_fusion_3130_2_7 Profiling Kit, ensure settings match the following:



STRmix v 2.7 Probabilistic Genotyping Standard Operating Instructions		
Status: Published		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	12 OF 36

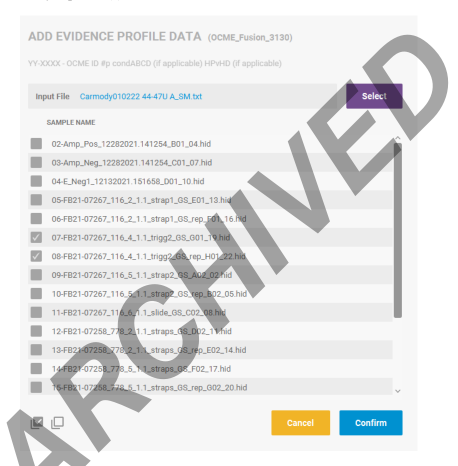


- Once all settings have been verified, click to return to the previous screen. For any locations that need to be ignored, click the checkbox in the Ignore column before clicking and make sure a note is made in the Case Notes section of the interpretation.
- 3.13 Add your **Evidence Profile Data** by either dragging and dropping the .txt file into the box or clicking on the to find your .txt file. The following window will pop up and you can select

STRmix v 2.7	7 Probabilistic Genotyping Standard Op	perating Instructions
Status: Published		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	13 OF 36

your evidence profile(s). If multiple replicates are in the same run, you can select multiple. Click to return to the file import screen.

3.13.1 Repeat for any replicate(s) from other .txt files.

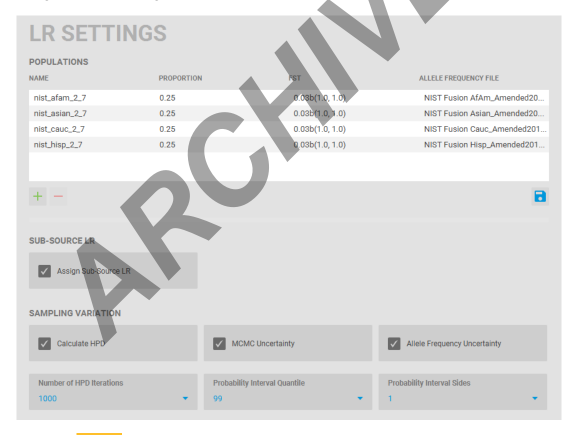


- 3.14 **To add a conditioned contributor**, add your **Reference Profile Data** by either dragging and dropping the .txt file into the box or clicking on the to find your .txt file. If you are performing a deconvolution without a conditioned contributor, proceed to step **3.15**.
 - 3.14.1 Select your reference profile(s) for conditioning using the checkboxes and click (similar to the screen for selecting your evidence profile).
 - 3.14.2 Once the profile data is added, the box for HP will be checked on the right side. To condition, you must also check the box for HD. Conditioned contributors are considered true donors in Hp and Hd.

STRmix v 2.7	7 Probabilistic Genotyping Standard Op	perating Instructions
Status: Published		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	14 OF 36



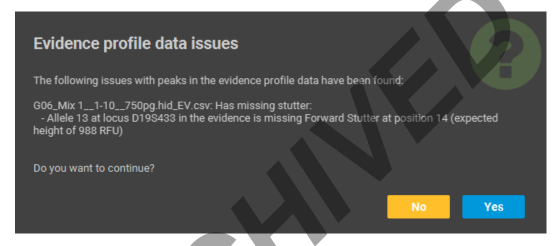
- 3.15 To perform an LR in conjunction with the deconvolution, add your Reference Profile Data by either dragging and dropping the .txt file into the box or clicking on the to find your .txt file. If you are performing a deconvolution without an LR, proceed to step 3.16.
 - 3.15.1 Select your reference profile for LRs using the checkboxes and click similar to the screen for selecting your evidence profile). The reference will be added within the numerator of the LR (i.e. assigned to HP only).
 - 3.15.2 Once the reference profile is added to your HP, you can then select against the following screenshot:



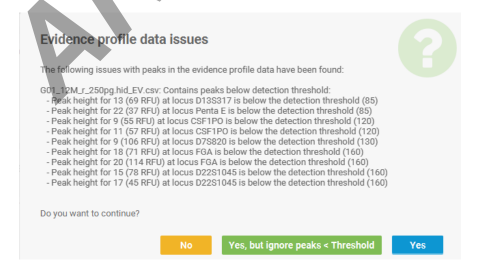
3.15.3 Select cancel to return to the file import screen.

STRmix v 2.7	7 Probabilistic Genotyping Standard Op	perating Instructions
Status: Published		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	15 OF 36

- 3.16 Select start (or Queue for Batch Mode, Section 6). The Progress window will open showing the Pre-Burnin, Burnin progress, and MCMC Progress (or for Batch Mode, you will return to the batch screen to continue to add to the batch).
 - 3.16.1 If a flag fires about missing an expected stutter (see below), review the input data and confirm that an allele label was not inadvertently deleted during analysis and that the input .txt file is correct before proceeding.



- 3.16.1.1 If no editing changes need to be made, you can click **Yes** to continue. If changes need to be made, select **No**, correct the labeling in the electropherogram and see Section 2.1.5.6 for editing the input file. Return to the beginning of setting up the run.
- 3.16.2 If a flag fires about peaks below analytical threshold (see below). Click and do not continue. Review the input data and the GeneMarker run settings used for analysis.



STRmix v 2.7	Probabilistic Genotyping Standard Op	perating Instructions
Status: Published		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	16 OF 36

- 3.17 Once complete, you will be at the Results screen.
 - 3.17.1 You can click on the single file folder icon on the right side to be directed to that run's **Results** folder. Select to return to the Main Menu.
 - 3.17.2 Alternatively, you can manually navigate to your STRmix[™] Run Folder(s) within the **OCME_STRmix_Fileshare** folder (\\csc\ocme\OCME_STRmix_Fileshare).
- 3.18 **COPY** your run folder(s) into the previously created FB sample folder within the **STRmix Data** folder.
- 3.19 Once you have copied the folder, **CONFIRM that all files for that run have transferred over correctly** to the **STRmix Data** folder. After confirmation, the copy of the STRmixTM Run folder located in the **OCME STRmix Fileshare** folder should be **deleted**.
- 3.20 When Applying User Informed Mx Priors: **DOCUMENTED APPROVAL BY TECHNICAL LEADER NEEDED PRIOR TO APPLICATION**
 - 3.20.1 Use the stutter filtered electropherograms to determine an approximate mixture proportion percentage for each contributor. Start within the first few loci to the left in each dye channel.
 - 3.20.2 Set-up an interpretation run as per section 3.1-3.6 of this manual. Be sure your "Number of Contributors" setting is correct, and the "Contributor Range" option remains unchecked.
 - 3.20.3 Select Run Settings to open the RUN SETTINGS window and check the box marked "Use Mx Priors". A CONFIGURE MX PRIORS window will appear like the one seen below:

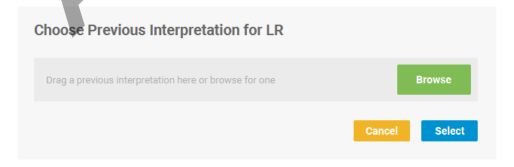


STRmix v 2.7	7 Probabilistic Genotyping Standard Op	perating Instructions
Status: Published		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	17 OF 36

- 3.20.4 Ensure contributor 1 is chosen in the drop-down menu. Using the top slider bar marked "Mean" to set the contributor proportion to the previously determined value.
- 3.20.5 Look at the range of the mixture proportion values used to determine the mean contributor as a guide for setting the bottom slider bar marked "Variance". Suggestions for variance to be set based on previous testing range between 0.125 and 6.1035E-5.
- 3.20.6 Repeat the steps in sections 3.20.4 and 3.20.5, changing the mean setting for the additional contributor(s) in the sample based on the determined values as needed. The variance setting should be set to the same value for all contributors.
- 3.20.7 Tap confirm in the CONFIGURE MX PRIORS window, then click SETTINGS window.
- 3.20.8 Click Next
- 3.20.9 Go back to section 3.10 and follow the steps needed to complete the run.

4 Likelihood Ratio Calculations with STRmixTM (Investigation: LR from Previous)

- 4.1 Note: This section is specific for running LRs using a previously run deconvolution. Samples must undergo deconvolution prior to (or in conjunction with) running an LR for a comparison sample. Refer to Section 3 for setting up deconvolutions in conjunction with the LR and the Interpretation of PowerPlex® Fusion data run on 3500xL manual for further guidance on running deconvolutions and LRs.
- 4.2 From the Main Menu, select Investigation. Within the Investigation screen, select LR from Previous. You will then select the deconvolution file for which you would like to calculate an LR for a comparison sample:



4.3 Drag and drop the entire decon run folder or select Browse to locate **the deconvolution file**, labeled as config.xml or results.xml (or settings.ini for older versions of STRmixTM) within your

STRmix v 2.7	7 Probabilistic Genotyping Standard Op	perating Instructions
Status: Published Document ID: 57029		
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	18 OF 36

run folder in the STRmix Data folder in the M drive. Click **Select** to navigate to the **LR from Previous** screen to name the run.

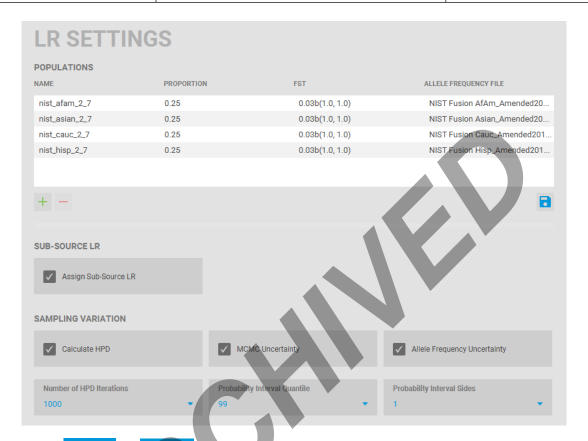
- 4.4 When naming STRmixTM Likelihood Ratio runs, the Case Number and Sample ID will autopopulate from the deconvolution file. These should be updated to the appropriate naming convention for an LR run before proceeding.
- 4.5 The following naming convention should be used for likelihood ratios:
 - 4.5.1 Evidence Files
 - Case Number = YY-XXXXX (leave off "FB")
 - Sample ID = remainder of evidence sample OCME ID* #p (NOC) condElimInitials (if applicable) scenario for LR
 - Case notes = a comment should be added here if a locus is ignored, indicating the reason: e.g. "D2S441 was ignored due to an unresolved allelic peak"
 - *Suffixes such as 'mcon' or 'reamp' should not be included in the OCME ID
 - 4.5.2 Suspect Files
 - Case Number = SYY-XXXXX (leave off "FB")
 - Sample ID = evidence sample OCME ID* (include evidence file FB# without the "FB") condElimInitials (if applicable) scenario for LR
 - Case Notes = a comment should be added here if a locus is ignored, indicating the reason: e.g. "D2S441 was ignored due to an unresolved allelic peak"
 - *Suffixes such as 'mcon' or 'reamp' should not be included in the OCME ID
 - 4.5.3 For LR scenarios, the naming format should start with the comparison sample's initials, followed by any conditioned samples' initials, and then the number and "U" for unknowns, followed by a "v" to separate the numerator hypothesis from the denominator hypothesis.

Examples	Resulting STRmix file name
Evidence File	
comparing elimAB to 4p decon, no cond	22-01234-567-1-1.1-shirt-BL-AB3Uv4U
comparing elimAB to 2p decon, cond vicCD	22-01234-567-1-1.1-shirt-BL-ABCDvCD1U
comparing elimAB to 3p decon, cond vicCD and elimEF	22-01234-567-1-1.1-shirt-BL-ABCDEFvCDEF1U
Suspect File	
comparing suspTS to 1p decon, no cond	S22-05678-22-01234-567-2-1.1-slide-GS-TSv1U
comparing suspTS to 2p decon, no cond	S22-05678-22-01234-567-2-1.1-slide-GS-TS1Uv2U
comparing suspTS to 3p decon, cond vicCD	S22-05678-22-01234-567-2-1.1-slide-GS-TSCD1UvCD2U
comparing suspTS to 4p decon, cond vicCD and	S22-05678-22-01234-567-2-1.1-slide-GS-
elimEF	TSCDEF1UvCDEF2U

	7 Probabilistic Genotyping Standard Op	_
Status: Published		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	19 OF 36

- The run settings will be pulled from the deconvolution and should not be changed. To check, click on Run Settings and see 3.6 for Run Settings screenshot. Click Cancel if all settings are correct and click Next to proceed.
 - **Performance, Number of Threads**: it is okay to proceed if the Number of Threads is different; this is specific to the computer being used.
 - Performance, Low Memory Mode: This setting allows the computer to minimize the
 memory used for the run and can be turned on in order to use less computer memory (ex.
 if using your computer for other tasks while running STRmixTM in the background) or if
 a run fails to finish due to computer memory. Using this setting will increase the run
 time.
 - if a run fails to finish due to computer memory.
 - **Performance**, **Seed**: This number may be different; there is a Seed for each run, deconvolution or LR.
- 4.7 The kit settings will also be pulled from the deconvolution. The kit will be pulled in as appropriate based on the STRmix deconvolution file:
 - 4.7.1 OCME_Fusion_3500 for 3500xL evidence data.
 - 4.7.2 OCME_Fusion_3130_2_7 for 3130xL evidence data deconvoluted using STRmix v2.7.
 - 4.7.3 **OCME Fusion** for 3130xL evidence data deconvoluted using STRmix v2.4.
- 4.8 If a locus needs to be ignored, click on the Kit settings.
 - 4.8.1 Rarely, a locus may be ignored at this step. For example, in the case of a tri-allelic pattern that matches your reference sample, which was not recognized at the deconvolution stage. For any locations that need to be ignored, click the checkbox in the Ignore column before clicking
- 4.9 Your Evidence Profile Data and Reference Profile Data for any conditioned contributors (HP & HD selected) will be populated from the deconvolution.
- 4.10 Add the **profile(s)** for comparison to the **Reference Profile Data** by either dragging and dropping the .txt file into the box or clicking on the to find your .txt file(s).
 - 4.10.1 Select your reference profile(s) for the LR using the checkboxes and click confirm. The reference will be added within the numerator of the LR (i.e. assigned to HP only).
 - 4.10.2 Once the reference profile(s) is added to your HP, you can then select them against the following screenshot:

STRmix v 2.7	7 Probabilistic Genotyping Standard Op	perating Instructions
Status: Published Document ID: 57029		
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	20 OF 36



- 4.11 Select start (or Gueue for Batch Mode, Section 6). The Progress window will open (or for Batch Mode, you will return to the batch screen to continue to add to the batch).
- 4.12 Once complete, you will be at the Results screen.
 - 4.12.1 You can click on the single file folder icon on the right side to be directed to that run's **Results** folder. Select finish to return to the Investigation Menu.
 - 4.12.2 Alternatively, you can manually navigate to your STRmix Run Folder within the **OCME_STRmix_Fileshare** folder (\\csc\ocme\OCME_STRmix_Fileshare).
- 4.13 **COPY** your run folder into the previously created FB sample folder within the **STRmix Data** folder.
- 4.14 Once you have copied the folder, **CONFIRM that all files for that run have transferred over correctly** to the **STRmix Data** folder. After confirmation, the copy of the STRmix Run folder located in the **OCME STRmix Fileshare** folder should be **deleted**.

STRmix v 2.7	7 Probabilistic Genotyping Standard Op	perating Instructions
Status: Published Document ID: 57029		
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	21 OF 36

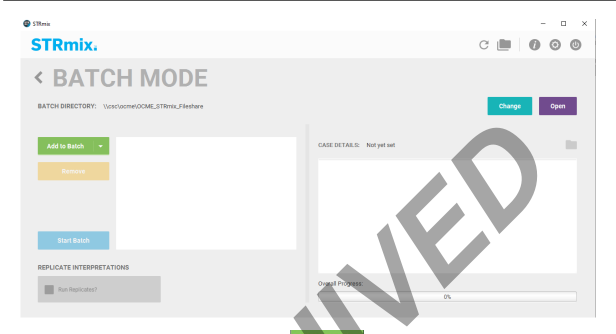
5 LRs for Convicted Offender Match Testimony

- 5.1 For testimony in relation to a convicted offender match, where a statistic is needed, an LR must be calculated through STRmixTM.
 - 5.1.1 Obtain the Convicted Offender profile through the CODIS software:
 - 5.1.1.1 Go to the Specimen Manager window → Click the filter button at the top of the Specimen ID column → Click "custom" → Enter your specimen ID from your evidence sample in the top line in the dialog box → Hit "search"
 - 5.1.1.2 Once your specimen ID pops up, right click on the line and click "view matches" → Right-click and choose "print reports", choose "match details short report"
 - 5.1.2 Add this to your case file and use the convicted offender profile from this report in order to create a comparison sample profile for STRmixTM LR calculation using the form: Reference Profile for STRmix Input Creation.
 - 5.1.3 Once the STRmixTM LR report is generated, it will need to undergo technical review (documented using case contacts) and recertification prior to testimony.

6 Running STRmixTM using Batch Mode

- 6.1 Several STRmixTM runs can be set up and queued to run sequentially. To set up a queued analysis for multiple runs, select **Batch Mode** from the STRmixTM main window. Batch mode should only be used for samples from one individual case within a single batch.
- 6.2 Before setting up a batch, navigate to the OCME_STRmix_Fileshare folder (\\csc\ocme\OCME_STRmix_Fileshare). Create a new folder within the STRmix Fileshare folder with the name of "Batch Mode [date] [your initials]."
- 6.3 Select change to change the Batch Directory. Navigate to and select the created folder inside the OCME_STRmix_Fileshare. This new folder will now appear at the top of the screen:

STRmix v 2.7	Probabilistic Genotyping Standard Op	perating Instructions
Status: Published Document ID: 57029		
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	22 OF 36



- Use the down arrow next to Add to Batch and Add Interpretation or Add LR from Previous to set up each individual run. If you do not click the arrow and click Add to Batch, it will automatically open whichever run type was used last.
 - 6.4.1 These batch details will be saved even if you close STRmixTM until you set up a new run; the batch can be stopped and restarted at a later date.
- 6.5 Complete the analysis set up for the first sample following the corresponding setup instructions: <u>Section 3 for Deconvolutions (Investigation)</u> or <u>Section 4 for LRs (Investigation: LR from Previous)</u>.
- 6.6 After setting up your run, click queue to return to the Batch Mode setup window.
- 6.7 Repeat steps **6.4-6.6** to add additional runs.
 - 6.7.1 If you need to edit a run, you will need to remove it and redo the setup. To remove a sample from the batch mode, highlight the case/sample in the **Calculations in Batch** part of the window and select **Remove**.
- 6.8 Select Start Batch to start the batch run.
- After completion of analyses, use the left arrow at the top left next to Batch Mode to return to the STRmixTM Main Menu. You can click open to be directed to the Batch Mode folder you created.
 - 6.9.1 Alternatively, if you have additional runs to perform within the same batch, you can add additional runs and continue the batch.

STRmix v 2.7 Probabilistic Genotyping Standard Operating Instructions				
Status: Published		Document ID: 57029		
DATE EFFECTIVE	APPROVED BY	PAGE		
10/11/2024	Nuclear DNA Technical Leader	23 OF 36		

- 6.9.2 The **Batch Log** will show if any of the individual runs failed and why. If an individual run failed, the run folder will not contain a Results folder, and a .txt file called BATCH CALCULATION FAILED will be made instead.
- 6.10 Results folders from Batch Mode will be saved in the folder you created within the OCME_STRmix_Fileshare folder (\\csc\ocme\OCME_STRmix_Fileshare).
- 6.11 **COPY your run folders** into the previously created FB sample folder(s) within the **STRmix Data** folder.
- 6.12 Once you have copied the folders, **CONFIRM that all files have transferred over correctly** to the **STRmix Data** folders. After confirmation, the copy of the STRmix Run folder and the Batch folder with your initials and date located in the **OCME_STRmix_Fileshare** folder should be **deleted**.

7 Evaluation of the STRmixTM Analysis Setup

- 7.1 Verify the evidence and reference input sections of the STRmixTM printout against the associated electropherograms. Ensure that:
 - 7.1.1 All appropriate edits were made; no artifact peaks were left labeled and no allelic peak labels were removed.
 - 7.1.1.1 If an Evidence profile issue flag was generated at the beginning of your run and checked for any issues against the electropherogram prior to running, this will also show up in the EVIDENCE PEAK ISSUES section at the end of the report. The information listed here should be aligned with the issue flag.
 - 7.1.2 Correct input file(s) have been selected.
 - 7.1.3 The correct file was imported into an LR from previous analysis, if applicable.
 - 7.1.4 All suitable replicates have been utilized.
 - 7.1.4.1 Check the **Inter replicate efficiency** that is generated by STRmixTM. If there are drastic inconsistencies between the two efficiencies that are consistent with the amount of data present in the replicates, the STRmix analysis may be marked as 'not reported' and a new analysis may be performed with the amplification containing the most information. A third amplification may be warranted.
- 7.2 The number of contributors that best describes the data was chosen, as applicable.
- 7.3 The correct assumptions (i.e. conditioning) have been made, if applicable.
- 7.4 The appropriate proposition has been selected (i.e. LR calculation), if applicable.

STRmix v 2.7 Probabilistic Genotyping Standard Operating Instructions			
Status: Published		Document ID: 57029	
DATE EFFECTIVE	APPROVED BY	PAGE	
10/11/2024	Nuclear DNA Technical Leader	24 OF 36	

- 7.5 The **SEED** value listed at the beginning of the report, with the CASE NUMBER, SAMPLE NAME, and COMMENTS, is the starting number used within the random number generator.
- 7.6 Check the **SETTINGS** at the end of the report to verify that the STRmixTM run was set up properly; note that any edited settings values are bolded by the program.
 - 7.6.1 The **CASE SETTINGS** will be specific to your case and should be used to check for correct setup with the run name.
 - 7.6.2 The MCMC SETTINGS are only for runs that include a deconvolution.
 - 7.6.2.1 Check the remaining settings against the following screenshot keeping in mind the numbers of accepts may be different (and bolded) if this setting has been approved for use.

MCMC SETTINGS	
Number of contributors	4
Use Mx priors	N
Number of chains	 8
Burn-in accepts per chain	10,000
Post burn-in accepts per chain	50,000
Random walk SD	0.005
Post burn-in shortlist	 9.0
Auto-continue on Gelman-Rubin	N

7.6.3 The **KIT SETTINGS** will be included in deconvolutions and LRs. If any loci were ignored, they will be listed here.

STRmix v 2.7	7 Probabilistic Genotyping Standard Op	perating Instructions		
Status: Published	Status: Published Document ID: 57029			
DATE EFFECTIVE	APPROVED BY	PAGE		
10/11/2024	Nuclear DNA Technical Leader	25 OF 36		

OCME_fusion_3500 Kit

KIT SETTINGS

Ignored loci	DYS391	
Detection thresholds	D3S1358	85
	D1S1656	85
	D2S441	85
	D10S1248	85
	D13S317	85
·	Penta E	85
	D16S539	120
	D18S51	120
	D2S1338	120
	CSF1PO 🔺	120
	Penta D	120
	TH01	130
	vWA	130
	D21S11	130
	D7S820	130
	D5S818	130
	TPOX	130
	D8S1179	160
	D12S391	160
· ·	D19S433	160
	FGA	160
	D22S1045	160
Saturation	30,000	
Degradation starts at	-1.0	
Degradation max	0.01	
Drop-in cap	300	
Drop-in rate parameter	0.0087	
Drop-in parameters (α, β)	22.31, 2.65	
Min variance factor	0.5	
Variance minimization parameter	1,000	
LSAE variance parameter (1/ λ)	0.03	
Allelic variance parameters (α, β)	7.427, 3.479	

STRmix v 2.7 Probabilistic Genotyping Standard Operating Instructions			
Status: Published		Document ID: 57029	
DATE EFFECTIVE	APPROVED BY	PAGE	
10/11/2024	Nuclear DNA Technical Leader	26 OF 36	

Back Stutter	
Maximum stutter ratio	0.3
Stutter variance parameters (α, β)	1.799, 19.052
Forward Stutter	
Maximum stutter ratio	0.2
Stutter variance parameters (α, β)	1.999, 11.703
Half Back (-2bp) Stutter	
Maximum stutter ratio	0.1
Stutter variance parameters (α, β)	2.597, 6.632
Double Back (-8bp) Stutter	
Maximum stutter ratio	0.1
Stutter variance parameters (α, β)	2.816, 6.792

STRmix v 2.7 Probabilistic Genotyping Standard Operating Instructions				
Status: Published		Document ID: 57029		
DATE EFFECTIVE	APPROVED BY	PAGE		
10/11/2024	Nuclear DNA Technical Leader	27 OF 36		

OCME_fusion_3130_2_7 Kit KIT SETTINGS

Ignored loci	DYS391	
Detection thresholds	D3S1358 50	
	D1S1656 50	
	D2S441 50	
	D10S1248 50	
	D13S317 50	
	Penta E 50	
	D16S539 50	
	D18S51 50	
	D2S1338 50	
	CSF1PO 50	
	Penta D 50	
	TH01 50	
	vWA 50	
	D21S11 50	
	D7S820 50	
	D55818 50	
	TPOX 50	
	D8S1179 50	
	D12S391 50	
	D19S433 50	
	FGA 50	
	D22S1045 50	
Saturation	8,000	
Degradation starts at	-1.0	
Degradation max	0.01	
Drop-in cap	100	
Drop-in rate parameter	0.0024	
Drop-in parameters (α, β)	0.0, 0.0	
Min variance factor	0.5	
Variance minimization parameter	1,000	
LSAE variance parameter (1/λ)	0.007	
Allelic variance parameters (α, β)	7.092, 0.933	
Back Stutter		
Maximum stutter ratio	0.3	
Stutter variance parameters (α, β)	2.108, 3.509	
Statter variance parameters (u, p)	2.100, 3.303	
Forward Stutter		
Maximum stutter ratio	0.1	
Stutter variance parameters (α, β)	2.312, 3.259	

STRmix v 2.7 Probabilistic Genotyping Standard Operating Instructions			
Status: Published Document ID: 57029			
DATE EFFECTIVE	APPROVED BY	PAGE	
10/11/2024	Nuclear DNA Technical Leader	28 OF 36	

OCME_Fusion Kit (for likelihood ratios on 3130xL STRmixTM v2.4 data only)

KIT SETTINGS		
Ignored loci	DYS391	
Detection thresholds	D3S1358	50
	D1S1656	50
	D2S441	50
	D10S1248	50
	D13S317	50
	Penta E	50
	D16S539	50
	D18S51	50
	D2S1338	50
	CSF1PO	50
	Penta D	50
	TH01	50
	vWA	50
	D21511	50
	D75820	50
	D5S818	50
	TPOX	50
	D8S1179	50
	D125391	50
	D19\$433	50
	FGA	50
	D22S1045	50
Saturation	8,000	
Degradation starts at	-1.0	
Degradation max	0.01	
Drop-In cap	100	
Drop-In rate parameter	0.0024	
Drop-In parameters (α, β)	0.0, 0.0	
Min variance factor	0.5	
Variance minimization parameter	1,000	
LSAE variance parameter (1/\(\lambda\)	0.0065	
Allelic variance parameters (a, β)	9.1374, 0.7472	
Back Stutter		
Maximum stutter ratio	0.3	
Stutter variance parameters (α, β)	1.5007, 12.9748	
Forward Stutter		
Maximum stutter ratio	0.1	
Stutter variance parameters (α, β)	9.1374, 0.7472	

7.6.4 The **PROFILE SETTINGS** can be used to check input file names and that the run was set up properly. If a stutter flag was generated as seen in step 3.16, "Ignore peaks below detection threshold" may be present as seen below; however this <u>does not</u> indicate that there are peaks below threshold within the run.

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STRmix v 2.7	Probabilistic Gen	otyping Standard Op	perating Instructions Document ID: 57029
DATE EFFECTIVE	APPRO	OVED BY	PAGE
10/11/2024	Nuclear DNA	Technical Leader	29 OF 36
PROFILE SETTINGS Number of evidence profiles		1	
Evidence profile filenames		63-FB21-07367_658_1.4.1_1.1_DS1_ST_G08_20.hid_EV.csv	
Number of Hp knowns		0	
Number of Hd knowns		0	
Ignore peaks below detection threshold		N	

7.6.5 The **LR SETTINGS** will only be present when an LR was run (whether separately or along with the deconvolution).

LR SETTINGS

Number of populations	4
Assign sub-source LR	Υ
Calculate HPD	X
HPD Iterations	1,000
Use MCMC uncertainty	Y
Use Allele Frequency uncertainty	У
HPD quantile	99%
HPD sides	1
NIST_AFAM_2.7	
Proportion	0.25
FST	0.03b(1.0, 1.0)
Allele frequency file	NIST Fusion AfAm_Amended2017.csv
NIST_ASIAN_2.7	
Proportion	0.25
FST	0.03b(1.0, 1.0)
Allele frequency file	NIST Fusion Asian_Amended2017.csv
NIST_CAUC_2.7	
Proportion	0.25
FST	0.03b(1.0, 1.0)
Allele frequency file	NIST Fusion Cauc_Amended2017.csv
NIST_HISP_2.7	
Proportion	0.25
FST	0.03b(1.0, 1.0)
Allele frequency file	NIST Fusion Hisp_Amended2017.csv

- 7.6.6 The **PERFORMANCE SETTINGS** shows the number of threads (specific to the computer used to run) and whether Low Memory Mode was used.
- 7.6.7 The **ADDITIONAL KIT DETAILS** lists the stutter files used in a deconvolution analysis.

STRmix v 2.7 Probabilistic Genotyping Standard Operating Instructions				
Status: Published		Document ID: 57029		
DATE EFFECTIVE	APPROVED BY	PAGE		
10/11/2024	Nuclear DNA Technical Leader	30 OF 36		

OCME_fusion_3500 Kit ADDITIONAL KIT DETAILS

Size Regression File	fusion_sizeregression.csv
Size regression file	TUSION_SIZE (ER) (ESSIONICS)
Back Stutter	
Position Relative to Parent	-1,0
Inversely Proportional To	Observed Height of Parent Allele
Stutter Regression File	OCME_BackStutterFile3500_022321.txt
Stutter Exceptions File	OCME_BSExceptionsFile_020321.csv
Forward Stutter	
Position Relative to Parent	1,0
Inversely Proportional To	Expected Height of Stutter Peak
Stutter Regression File	OCME_ForwardStutterFile3500_022321.txt
Stutter Exceptions File	OCME_FSExceptions File_092020.csv
Half Back (-2bp) Stutter	
Position Relative to Parent	0,-2
Inversely Proportional To	Expected Height of Stutter Peak
Stutter Regression File	OCME_HalfBackStutterFile3500_022321.txt
Stutter Exceptions File	OCME_HBSExceptionsFile_092020.csv
Double Back (-8bp) Stutter	
Position Relative to Parent	-2, 0
Inversely Proportional To	Expected Height of Stutter Peak
Stutter Regression File	OCME_DoubleBackStutterFile3500_022321.txt
Stutter Exceptions File	OCME_DBSExceptionsFile_092020.csv

STRmix v 2.7 Probabilistic Genotyping Standard Operating Instructions				
Status: Published		Document ID: 57029		
DATE EFFECTIVE	APPROVED BY	PAGE		
10/11/2024	Nuclear DNA Technical Leader	31 OF 36		

OCME_fusion_3130_2_7 Kit ADDITIONAL KIT DETAILS

Size Regression File	Fusion_SizeRegression.csv
Back Stutter	
Position Relative to Parent	-1,0
Inversely Proportional To	Observed Height of Parent Allele
Stutter Regression File	OCME_Fusion_Stutter.txt
Stutter Exceptions File	OCME_Fusion_Exceptions.csv
Forward Stutter	
Position Relative to Parent	1,0
Inversely Proportional To	Expected Height of Stutter Peak
Stutter Regression File	OCME_Fusion_Forward Stutter.txt
Stutter Exceptions File	

8 Evaluation of the STRmixTM Analysis Diagnostics

- 8.1 The presence of a single sub-optimal diagnostic is not always an indication that rework is required. In some instances, a sub-optimal diagnostic(s) may be due to the nature of the sample (ex. low amounts of input DNA and/or stochastic effects) and not due to an issue with the STRmixTM run. Refer to the Troubleshooting Guide (Section 8) for further steps that may be taken to improve a sub-optimal diagnostic result.
- 8.2 For deconvolutions, verify that the following (**primary**) diagnostics conform to your qualitative expectations when compared to the electropherogram(s):
 - 8.2.1 The mixture proportions and template amounts assigned to the contributor(s).
 - 8.2.2 The **weights** assigned to the genotypes for each contributor listed in the **COMPONENT INTERPRETATION**.
 - 8.2.3 The **degradation** values and degradation plots (at the beginning of the report). The per contributor plot can be helpful for recognizing extreme degradation its potential influence over a single contributor's genotype weights across a profile.
 - 8.2.4 The Locus Efficiencies (LSAE), following the variance charts.
- 8.3 For deconvolutions, evaluate the following (**secondary**) diagnostics for the run information listed in the **POST BURN-IN SUMMARY** at the beginning of the report.

STRmix v 2.7	7 Probabilistic Genotyping Standard Op	perating Instructions
Status: Published		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	32 OF 36

- 8.3.1 **Total iterations:** If the total iterations exceeds 2.14 billion (2.14 x 10⁹), this may lead to incorrect genotype weightings being assigned.
- 8.3.2 **Acceptance rate:** A very low acceptance rate (e.g. 1 in thousands to millions) may, in combination with the other diagnostics, indicate that the analysis needs to be run with additional accepts.
- 8.3.3 **Effective sample size (ESS):** A low ESS in relation to the total number of iterations suggests that the MCMC has not moved very far with each step or has had a low acceptance rate. A low ESS value (tens or hundreds) means that there is potential for a large difference in weights if the analysis is run again. For an ESS = 8, see the following troubleshooting guide.
- 8.3.4 **(Log)likelihood:** The larger this value, the better STRmixTM has been able to describe the observed data. A negative value suggests that STRmixTM has not been able to describe the data very well given the information it has been provided. A low or negative value for the log(likelihood) may indicate to users that the analysis requires additional scrutiny.
- 8.3.5 **Gelman-Rubin diagnostic:** If this value is above 1.2 then it is possible that the analysis has not converged, and the analysis requires additional scrutiny.
- 8.3.6 **LSAE variance**: The LSAE variance probability density chart in the report may be used in conjunction with the LSAE efficiency plot to identify unusual amplification within a profile (ex. extreme inhibition).
- 8.3.7 **Allele variance and stutter variance:** These variances should be compared to the modes and variance charts included in the reports. If the numbers are significantly elevated, the analysis may require additional scrutiny.
- 8.4 For LR comparisons, the overall **category of support** (inclusion, uninformative, exclusion) should conform to your qualitative expectations in comparison to the data.
 - 8.4.1 Evaluate the **Per Locus Likelihood Ratio** table per locus and per sample, as well as the range of LR's **between population subgroups**; pay special attention to outliers and/or zero values.
- 8.5 For LR comparisons that result in support for an inclusion, check to ensure that the comparison sample falls in the appropriate contributor order. See report section titled **CONTRIBUTOR ORDER GIVING HIGHEST LR** at the beginning of the report.
 - 8.5.1 If multiple comparison samples are positively associated with the same sample, results should be evaluated to ensure that they do not align with the same contributor. See Section 9 for troubleshooting when more than one comparison aligns with the same contributor.

STRmix v 2.7	Probabilistic Genotyping Standard Op	perating Instructions
Status: Published		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	33 OF 36

9 Troubleshooting Guide

9.1 The purpose of this guide is to address commonly seen scenarios which arise in casework. These guidelines are based on validation studies, literature references, and casework experience. However, not every situation can be covered by the Troubleshooting Guide. If a diagnostic issue arises that is not covered here, please discuss the issue with your supervisor, manager or the Technical Lead Team as needed.

Observations	Actions
EVALUATION OF DECONVOLUTION DIAG	NOSTICS
The mixture proportions or template amount do not reflect what is observed	Re-evaluate the number of contributors; consider another STRmix TM analysis with one higher or one lower number of contributors.
AND/OR the degradation does not reflect what is observed	Consider amplifying a replicate if one has not already been done, with increased input amount when available and appropriate.
AND/OR the interpreted contributor genotypes do not conform to your qualitative expectations	Inhibition has occurred—microcon to clean and reamplify sample.
AND/OR the category of support for an LR comparison does not conform to your qualitative expectations	Consider another STRmix TM analysis at a greater number of accepts (typically, 100,000 burn-in accepts and 500,000 total accepts per chain). Note: this requires approval by the Technical Leader (or his/her designee).
The mixture proportions and genotype weights do not conform to your qualitative expectations based on the electropherograms, and other troubleshooting options (including additional iterations) have been exhausted. Most common in mixtures with extreme ratios (ex. 98:2, 50:48:2,85:10:4:1) and/or many shared alleles between contributors)	Consider another STRmix analysis utilizing user informed Mx priors. This requires documented approval by the Technical Leader (or his/her designee).
The total iterations exceeds 2.14 billion (2.14 x 10 ⁹).	This could indicate the genotype weightings have been incorrectly assigned. Please contact the Technical Lead Team. An additional STRmix TM analysis may be required.
Effective sample size (ESS) = 8	Consider another STRmix TM analysis at a greater number of accepts (typically, 100,000 burn-in accepts and 500,000 total accepts per chain). Note: this requires approval by the Technical Leader (or their designee). You may want to use a computer with higher processing power.

STRmix v 2.7 Probabilistic Genotyping Standard Operating Instructions			
Status: Published		Document ID: 57029	
DATE EFFECTIVE	APPROVED BY	PAGE	
10/11/2024	Nuclear DNA Technical Leader	34 OF 36	

10/11/2024	Nuclear DIVA	Technical Leader	34 OF 30
A low or negative(log) likelihood		Re-evaluate number of contributors; consider another STRmix TM analysis with one higher or one lower number of contributors. Data has been removed that is allelic and/or stutter;	
		data must be re-imported.	,
		Artifact peaks have been le removed.	eft labeled and must be
Gelman-Rubin value is greate	er than 1.2	Consider another STRmix number of accepts (typical accepts and 500,000 total amay sometimes reduce the this requires approval by this/her designee).	lly, 100,000 burn-in accepts per chain). This e GR to below 1.2. Note:
Stutter and/or allele variance elevated usually at or beyond t asymptote (may be in conjunct (log) likelihood)	he horizontal	STRmix TM analysis with o number of contributors.	ntributors; consider another one higher or one lower licate if one has not already
EVALUATION OF LR DIAG	ENOSTICS		
Differences in Per Locus LRs obtained for each locus where profile is qualitatively included evidentiary profile, with one locus = 0	the comparison d in the	locus and perform analysis supervisor as needed. Inhibition has occurred—reamplify sample. Consider amplifying a rep been done, with increased available and appropriate.	ed at a given locus – ignore s again. Note: discuss with microcon to clean and licate if one has not already input amount when arributors; consider another one higher or one lower TM analysis at greater lly, 100,000 burn-in accepts per chain). Note:
Multiple elimination and/or co gave LRs supporting inclusion sample and are aligning with t contributor and/or there is an	to an evidence the same	Consult a supervisor and t	onal deconvolution and/or

relatedness.

STRmix v 2.7 Probabilistic Genotyping Standard Operating Instructions		
Status: Published		Document ID: 57029
DATE EFFECTIVE	APPROVED BY	PAGE
10/11/2024	Nuclear DNA Technical Leader	35 OF 36

The 99% 1-sided HPD and Unified LR result(s) for one or more **population subgroup(s)** is several orders of magnitude lower than the other population subgroups in comparison to the point estimate LR, and the lowest LR of the four population subgroups does not conform to your qualitative expectations of the comparison.

Consult your supervisor or manager and the Technical Lead Team. A stratified likelihood ratio using census data may be calculated and reported; this must be approved by the Technical Leader (or their designee).

ERROR MESSAGES

Upon opening the STRmix software, an error that indicates "All connections to the floating license are currently in use"

Multiple licenses of STRmixTM are open on your computer. Close out the error message and determine if you have another active window of STRmixTM running.

Your previous use of the software was closed incorrectly. Re-start your computer, and if this does not resolve the issue, file an IT ticket to close your open STRmixTM license connections to the server.

All licenses are in use by other members of the laboratory.

Evidence profile data flag during pre-checks (as seen in step 3.16)

This is an example of a **missing stutter** flag. You will be notified if there's an expected stutter peak that is not included based on the heights of the allelic peaks in the input file. See Section 3.16.1. Elevated stutter variance may occur in conjunction with this flag. The data should be evaluated to determine if the elevated variance makes intuitive sense with the peaks present in the profile. Consult the Tech Lead Team if further assistance is needed.

Another profile data issue that you may see is if there are peaks within the input file that are below the set analytical threshold. The original data should be reevaluated in GeneMarker to determine if incorrect settings were applied.

Pre-Burnin failed:
Determine Acceptable Genotypes failed: Locus # (locus name) in the evidence cannot be explained given the parameters you have chosen.

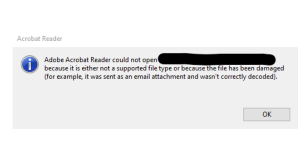
Check your input value for NOC. Re-evaluate number of contributors; consider another STRmixTM analysis with one higher number of contributors.

Consider amplifying a replicate if one has not already been done, with increased input amount when available and appropriate.

(OutofMemoryError) GC overhead limit exceeded

The run could not be completed with the computer power supplied. The run should be set up again using the Low Memory Mode setting and/or on a computer with more processing power.

STRmix v 2.7 Probabilistic Genotyping Standard Operating Instructions			
Status: Published		Document ID: 57029	
DATE EFFECTIVE	APPROVED BY	PAGE	
10/11/2024	Nuclear DNA Technical Leader	36 OF 36	



The results .pdf may be corrupted. Check that your run folder contains all other folders and files. The Reports folder may be missing the PNG and Excel files. You can regenerate your .pdf in STRmix by clicking on the Reports module from the main menu. Drag your full folder into the Previous Calculation field. Click the second Browse button that appears and add "_regen" to the end of your file name to not overwrite your original corrupted file. Include all files in the case folder. Add a case contact stating that a new .pdf was generated due to a corrupted file.

10 References:

- 10.1 STRmixTM v.2.7 Operation Manual and previous versions
- 10.2 STRmixTM v. 2.7 User's Manual and previous versions
- 10.3 NYC OCME Internal Validation of STRmix[™] v2.7 for Fusion 5C/3500xL (September 2021)
- 10.4 NYC OCME Internal Validation of STRmixTM v2.7 for Fusion 5C/3500xL Data STRmixTM Parameters (August 2021)
- 10.5 NYC OCME Stutter Study for GeneMarker® HID 3.0.0 and STRmixTM Version 2.7- PowerPlex® Fusion Data run on 3500xl Genetic Analyzers (September 2021)
- 10.6 NYC OCME Internal Validation of STRmix[™] V2.4 for Fusion (January 2017)
- 10.7 NYC OCME STRmixTM V2.4.08 Performance Check. (July 2018)