FORENSIC BIOLOGY PROTOCOLS FOR FORENSIC STR ANALYSIS

Amplification using the PowerPlex Y23 System		
Status: Published		Document ID: 59652
DATE EFFECTIVE	APPROVED BY	PAGE
08/22/2022	Nuclear DNA Technical Leader	1 OF 3

PowerPlex® Y23 Sample Preparation for Amplification

1 Procedure

PPY23 Sample Input Amount	
Optimal - 500pg of male DNA*	
Minimum – 100pg of male DNA	

^{*}The option for amplification with a greater input amount is available if determined appropriate for the sample by the analyst.

1.1 Retrieve the following reagents from the associated refrigerator and/or freezer and record the lot numbers.

PowerPlex® Y23 10X Primer Pair Mix	
PowerPlex® Y23 5X Master Mix	
Water, Amplification Grade for PPY23	
2800M Control DNA for PPY23, 10ng/μl	

- 1.2 Retrieve sample(s) needed for amplification from associated refrigerator and/or freezer.
- 1.3 Prepare dilutions (if necessary) in 1.5 mL tubes according to the values listed on the test batch data entry screen or the "FBAmplificationSheet", <u>using Promega Amplification Grade Water for PPY23</u>, for each sample, according to Table 1. Vortex and centrifuge samples prior to aliquoting for dilution.

TABLE 1: Dilutions

Dilution	Amount of DNA Template (uL)	Amount of Water (uL)
0.25	3 or (2)	9 or (6)
0.2	2	8
0.1	2	18
0.05	2	38
0.04	4 or (2)	96 or (48)
0.02	2 or (1)	98 or (49)
0.01	1 or (2)	99 or (198)
0.008	4 or (2)	496 or (248)

1.4 Label amp tubes using the values generated by LIMS. These values can be found in the test batch output samples or on the "FBAmplificationSheet".

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DATE EFFECTIVE	APPROVED BY	PAGE
08/22/2022	Nuclear DNA Technical Leader	2 OF 3

- 1.5 Centrifuge reagent tubes briefly to bring contents to the bottom and then vortex for 15 seconds before use. Do NOT re-centrifuge the Master Mix or Primer Pair Mix as this may cause the reagents to be concentrated at the bottom of the tube.
- 1.6 Consult the Reagents tab in LIMS for the exact amount of PowerPlex® Y23 10X Primer Pair Mix and PowerPlex® Y23 5X Master Mix to add.

Reagent	Per reaction
10X Primer Pair Mix	2.5 μL
5X Master Mix	5.0 μL
Mastermix total:	7.5 µL
DNA	17.5 μL

- 1.7 Vortex prepared Master Mix and all samples to be aliquoted for 5-10 seconds. After vortexing, briefly centrifuge master mix and samples.
- 1.8 Add 7.5 µL of the prepared master mix to each tube that will be utilized, changing pipette tips and remixing master mix as needed.
- 1.9 Witness Step. Have another analyst witness the sample set-up.
 - 1.9.1 For the input samples, confirm the tube label and sample ID for each sample. For the output samples, the entire amp tube label must be read for each sample.
- Positive Control for PPY23 (PPY23 2800M Control in LIMS) total input amount of 250pg. 1.10
 - 1.10.1 Perform dilution and aliquot positive control according to amplification sheet
- 1.11 Amplification Negative
 - 1.11.1 17.5 uL of Water, Amplification Grade for PPY23 (PPY23 H2O in LIMS)
- 1.12 Samples
 - 1.12.1 Aliquot samples according to amplification sheet
- 1.13 All amplification tubes should have a total final volume of 25uL.
- 1.14 Ensure that all caps are properly closed prior to sending the samples to the post-amplification laboratory.
- 1.15 Spin down samples at 1000 RPM for one minute.
- For thermal cycler usage see the Using the Mastercycler X50s manual. 1.16

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L	Status: Published		Document ID: 59652
	DATE EFFECTIVE	APPROVED BY	PAGE
	08/22/2022	Nuclear DNA Technical Leader	3 OF 3

1.17 The PPY23 PCR program is as follows:

Soak at 96°C for 2 minutes

: Denature at 94°C for 10 seconds

30 Cycles : Anneal at 61°C for 60 seconds

: Extend at 72°C for 30 seconds

20-minute incubation at 60°C.

Storage soak indefinitely at 4°C

NOTE: The 4°C storage soak step is not meant to store samples for an extended period. Samples should be removed from the instrument and placed in the 4°C refrigerator at the earliest convenience.

1.18 Place the microtube rack used to set-up the samples for PCR in the container of 10% bleach container in the Post-Amp area.