Automated DNAIQ	Extraction from Bloodstains and other	Casework samples	
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Automated DNA IQTM Extraction from Bloodstains

& Other Casework Samples

1 General Information

- 1.1 WARNING: THE LYSIS BUFFER IN THE DNA IQ KITS IS CORROSIVE AND TOXIC. IT CAUSES SEVERE SKIN BURNS AND EYE DAMAGE AND IS HARMFUL IF INHALED OR SWALLOWED. If on skin: take off all contaminated clothing and rinse with water and soap. If in eyes: rinse with copious amounts of water for several minutes. Keep lysis buffer bottle tightly closed.
 - 1.1.1 The waste plates containing residual reagents can be discarded in the regular laboratory garbage.
 - 1.1.2 The remaining residual Wash buffer and Elution buffer reagents in the troughs after a run can be disposed of in the laboratory sink flushed with copious amounts of water.
 - 1.1.3 However, the remaining residual Lysis Buffer and Lysis Buffer with Resin must be collected in a properly labeled waste container and be properly discarded via a chemical waste vendor.
 - 1.1.3.1 Contact QA to collect waste containers and expired bottles for proper disposal.
- 1.2 CAUTION: DO NOT ADD BLEACH OR ACIDIC SOLUTIONS DIRECTLY TO ANYTHING CONTAINING LYSIS BUFFER INCLUDING SAMPLE WASTE. Exposure to strong acid or bleach will result in the generation of toxic gases.
 - 1.2.1 If liquid containing these buffers spill, clean with suitable laboratory detergent and water.
 - 1.2.2 If the spilt liquid contains potentially infectious agents, clean with suitable laboratory detergent and water first and then with 1% sodium hypochlorite followed by water.
- 1.3 CAUTION: DO NOT ADD BLEACH TO ANY PART OF THE HAMILTON® ROBOT INCLUDING THE DECK. Bleach will cause the robot to rust. For any spills on the robot, use ethanol and water for cleanup.
- 1.4 Sample size for the extraction should be approximately 1/3 of a swab, 3x3 mm cutting of a stain, or sample scrapings loaded in a 1.5ml flip-cap microcentrifuge tube. This extraction is applicable for all casework samples EXCEPT semen samples.
- 1.5 Do not extract evidence samples and exemplars together.

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2 Sample Preparation and Incubation

- Obtain two empty 1.5ml microcentrifuge tubes for the extraction negatives and label one as Extraction Negative 1 and the other as Extraction Negative 2.
- 2.2 Print input labels and place on spin basket units (consisting of spin basket and click fit tube) for each sample. Also print LIMS output labels and label 1.5ml screw cap tubes to be used as the elution tubes.
- 2.3 Have a **witness** verify the order of the spin basket units and the sample digestion tubes by reading the tube-top label and the entire input sample ID number for each sample. This will be your "Extraction" witness.
- 2.4 Prepare digest buffer master mix as per the calculated amounts in the Reagents tab in LIMS. Record the lot# of each reagent in LIMS.

Stock Solution	1 sample
0.05% SDS	192 μL
ProteinaseK (20 mg/mL)	8μL

- 2.5 Vortex the master mix well.
- 2.6 Add 200ul of digest buffer master mix to each of the sample tubes including the extraction negative tubes. If necessary, take a clean pipette tip and push the substrate down into the digestion liquid.
- 2.7 Incubate at 56°C for 30 minutes with shaking at 1400 rpm in the thermomixer.
- 2.8 Record the temperature of the thermomixer in LIMS. The temperature should be within $\pm 3^{\circ}$ C of 56°C.
- 2.9 Prepare 10% bleach, distilled water, and 70% alcohol in three 50ml conical tubes. Clean a pair of forceps by dipping the forceps in each of the three tubes briefly and then drying with a fresh lint free wipe.
- 2.10 While samples are incubating, proceed with the daily maintenance of the robot, if needed, in Section_5 <u>Daily Maintenance</u> below.
- 2.11 Remove the tubes from the thermomixer and briefly centrifuge the samples for a few seconds at <3000rpm.
- 2.12 For each sample, transfer the substrate or scrapings using the forceps to its associated spin basket unit. Pipette mix the sample lysate within the incubation tube a few times to disturb any pellet that may have formed, and then pipette the entire lysate volume (~200ul) over the top of the substrate

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in the spin basket. Close the tube top over the spin basket. Repeat this step for each sample, cleaning and drying the forceps between each sample.

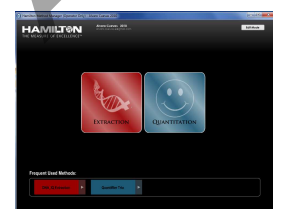
- 2.13 Centrifuge the substrates in spin baskets at 13,200 rpm to 15,000 rpm for 2 minutes.
- 2.14 Using clean and dry forceps (or a fresh lint-free wipe), remove and discard the spin baskets (including the swab remains), taking care to avoid bubbles at the rim of the open tube. Clean and dry forceps between each sample. Close the tube.
- 2.15 If the samples will be loaded immediately onto the instrument, proceed to setting up the Hamilton STAR Robot.
 - 2.15.1 Alternatively, the samples can be stored overnight in a refrigerator to be run on the robot the following day.

3 Setting up the Hamilton® STAR Robot:

- 3.1 Make sure the green power switch on the robot is turned on before logging into the computer.
- 3.2 Daily maintenance is to be performed prior to the first run of the day.
 - 3.2.1 Proceed with the daily maintenance of the robot, if needed, in Section 5 <u>Daily</u> <u>Maintenance</u> below



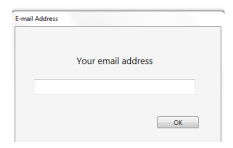
- 3.3 Once Daily Maintenance is done, click on the Method Manager icon.
- 3.4 Click on the Extraction button.



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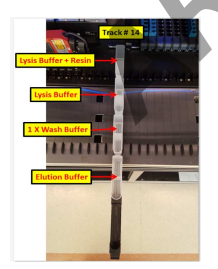
- 3.5 Then Click the Run icon.
- 3.6 The following pop up box will appear:



3.7 Enter your email address. Click OK. This will allow you to receive an email message from the robot when your run is done, or if the run is aborted. The robot can also send you a text message by entering your 10 digit phone number in an email format:

Verizon: number@vtext.com
AT&T: number@tmomail.net
Sprint: number@pm,sprint.com

- 3.8 In the next window, enter the total number of samples of your batch. Include extraction negatives. Maximum number of samples is 84. Click OK.
- 3.9 Record the lot numbers for the DNA IQ extraction reagents in LIMS.



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3.10 Load the indicated volume of each reagent into new 60ml reagent containers on track #14 as indicated by the arrows above. The calculation of the amount of reagents is calculated by the robot software. The following tables may also be used for reference.

Amount to add in mL

Buffer	8 samples	16 samples	24 samples
Lysis Buffer &	5.4	7.4	9.4
Resin**	0.2	0.2	0.3
Lysis Buffer	4.4	5.2	6.0
1X Wash Buffer	6.2	7.8	9.4
Elution Buffer	4.9	5.2	5.6

Buffer	32 samples	40 samples	48 samples
Lysis Buffer &	11.4	13.4	15.4
Resin**	0.3	0.4	0.4
Lysis Buffer	6.8	7.6	8.4
1X Wash Buffer	11.0	12.6	14.2
Elution Buffer	5.9	6.2	6.5

Buffer	56 samples	64 samples	72 samples
Lysis Buffer &	17.4	19.4	21.4
Resin**	0.5	0.5	0.6
Lysis Buffer	9.2	10.0	10.8
1X Wash Buffer	15.8	17.4	19.0
Elution Buffer	6.8	7.2	7.5

Buffer	80 samples	84 samples
Lysis Buffer & Resin**	23.4	24.4
	0.7	0.7
Lysis Buffer	11.6	12.0
1X Wash Buffer	20.6	21.4
Elution Buffer	7.8	8.0

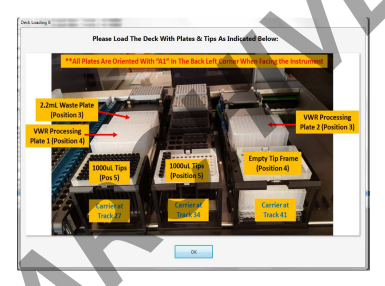
^{**}The software rounds the amount of resin to the nearest tenth of a mL

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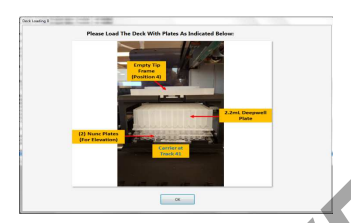
- 3.10.1 Prepare the Promega Lysis Buffer and DNA IQ[™] Resin Solution fresh before each run as per the calculations in the DNA IQ program. Before pouring the resin, thoroughly mix the resin by inversion many times before adding. Do not store solution after run.
- 3.10.2 Add Promega Lysis buffer to the second reagent container as per the calculations in the DNA IQ program. Remember to re-cap the bottle tightly to prevent crystallization!
- 3.10.3 If Promega 2X wash buffer has not already been diluted, prepare as follows and add Promega 1X wash buffer to the third reagent container as per the calculations in the DNA IQ program.
 - 3.10.3.1 Note: The wash buffer comes from the manufacturer at a concentration of 2X in each DNA IQ kit. It needs to be diluted to 1X. If you are opening a new DNA IQ kit proceed with the following:
 - 3.10.3.1.1 Add 35ml of 95–100% ethanol and 35ml of 99% isopropyl alcohol directly to the 70 ml 2X Wash Buffer bottleReplace the cap, and mix by inversion several times.
 - 3.10.3.1.2 Label as 1X Wash Buffer, indicating the addition of ethanol and isopropyl alcohol. These lot numbers do not need to be recorded in LIMS. Label the date of the dilution and your initials.
 - 3.10.3.1.3 Store at room temperature (22–25°C). Make sure bottle is closed tightly to prevent evaporation.
- 3.10.4 Add Promega Elution buffer to the fourth reagent container as per the calculations in the DNA IQ program.
- 3.10.5 Note: If graduated cylinders are used to measure the volume of the buffers, ensure to use only the labeled cylinder for each specific reagent. Clean each cylinder with detergent and rinse with distilled water after use. For volumes under a mL, use a pipette. **DO NOT ADD BLEACH OR ETHANOL TO THE CYLINDERS**.
- 3.10.6 Label the reagent containers with the reagent name.
- 3.11 Set up the Robot Platform:
 - 3.11.1 IT IS VERY IMPORTANT THAT VWR PLATES AND PROMEGA PLATES ARE ONLY USED WHERE DIRECTED. DO NOT INTERCHANGE THE DIFFERENT TYPES OF PLATES AS THIS WILL RESULT IN A Z CRASH ERROR!
 - 3.11.2 All plates should be oriented with well A1 in the top left corner.

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- 3.11.3 Load one new VWR deep well processing plate to position 4 on the carrier located on tracks 27-32.
- 3.11.4 Add another VWR plate to position 3 on carrier located on tracks 41-47.
- 3.11.5 Add one Promega 2.2ml deep well plate for waste on position 3 of carrier located on tracks 27-32.
- 3.11.6 Make sure that there is at least one full rack of 1000ul tips in position 5 on the carrier located on tracks 27-32.
- 3.11.7 Place a new empty tip frame in position 4 of carrier located on tracks 41-47.
- 3.11.8 Click OK.



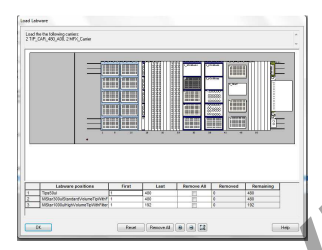
3.11.9 Add another Promega 2.2ml deep well plate under position 4 on carrier on tracks 41-47. The plate should rest on top of two Nunc plates (used for elevation only) as seen in the photo below.



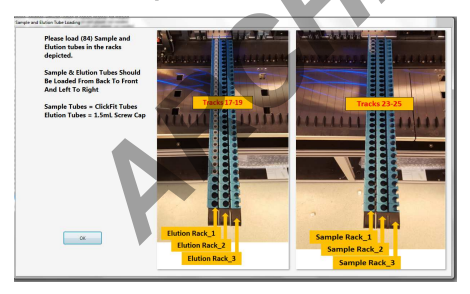
3.11.10 Refill tips as needed according to photo below. Ensure that correct size tips go in correct carrier. Ensure that the bar code on the side of the tip tray is facing to the right.



- 3.11.11 The program has a built in tip-counter that will remember which tips it used in previous runs and the number of tips used. Make sure that the number of tips on the deck corresponds to what is listed in the tip-counter window below.
 - 3.11.11.1 Click and drag the lasso tool around each tip tray to indicate which tips have not been used if there are any discrepancies.
 - 3.11.11.2 If full trays of tips have been added, click the Reset button. Then Click OK.



- 3.12 A series of prompts will ask for confirmation of the carriers located on the instrument deck. Verify the correct location of the carriers and click YES to continue.
- 3.13 Have a **witness** verify the "robot setup" by confirming the sample order while loading the sample lysates in open click-fit tubes into the blue tube adaptors on the sample carriers located in tracks 23-25. The samples should be loaded from back to front.



- 3.13.1 The witness must also confirm the elution tube order while loading the uncapped labeled elution tubes into the carriers located in tracks 17-19 from back to front.
- 3.13.2 Finally, the witness must also make sure that all carriers are pushed back into the robot, the carriers are on the correct tracks, the deck layout is correct, VWR plates and Promega plates are in the right positions, elution tubes are uncapped, input tubes are open, and the robot door is closed. This will be your "robot setup" witness in LIMS.

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- 3.14 Click Continue to begin run. Make sure to watch the run for the first few minutes to make sure it is running ok.
- 3.15 When the run is complete, an email (or text) will be sent to you. Once the Run is complete:
 - 3.15.1 Cap the elution tubes and store in a cryobox in the appropriate fridge.
 - 3.15.2 Discard input tubes, plates, reagent troughs and the empty tip frame in position 4 along with its Promega deep well plate underneath it on the carrier on tracks 41-46. If a batch of tips runs out, do not discard the tip tray. These should be saved for future runs, to be used as the tip holder at position 4 (track 41-46).
 - 3.15.3 In the LIMS system, navigate to the Data Entry page, indicate if the DNA IQ extraction passed or failed.
 - 3.15.4 Ensure that the Hamilton usage log has been filled out in LIMS.
 - 3.15.5 Close the Method Manager software and logout of the computer.
 - 3.15.6 If it is the last run for the day on Friday, or the instrument will not be used for more than 24 hours, turn off the instrument.

4 Troubleshooting

4.1 DNA IQ

Poor yield	Too much sample was used. Excessive amount of	
	sample can reduce the efficiency of DNA binding	
	to the resin. Use less sample.	
Inconsistent yield	Inconsistent amounts of resin. Vortex resin stock	
	thoroughly before adding to reagent trough. Do not	
	allow the resin to dry out.	

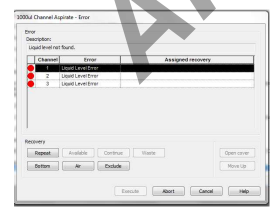
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4.2 STAR Robot

- 4.2.1 While some errors may be corrected by the user, most should be handled by QA. Error email messages are sent to ocme.nyc.gov. If an error is encountered, contact QA to check for an email. If the run is aborted, an email is sent to the email address that was entered at the beginning of the run.
- 4.2.2 Below are some common error codes that may be resolved by the analyst. For further troubleshooting errors, contact QA:

Main Error		
Code	Error	Description
5	Barcode Error	Barcode could not be read or is missing
6	Insufficient Liquid	Not enough liquid available
6	Error	
7	Tip Present Error	A tip has already been picked up
8	No Tip Error	Tip is missing use the next tip in sequence or fill in
		tips
9	No Carrier Error	No carrier present for loading
16	Cover Open Error	Cover not closed or cannot be locked
18	Wash Liquid Error	Waste is full or wash liquid is empty
100	Wrong Carrier Error	Wrong carrier barcode, a wrong carrier is loaded.
102	Liquid Level Error	Liquid surface not detected.
103	Not Detected Error	Carrier not detected at end position on deck

4.2.3 If a liquid level error occurs, for example, (like in the following pop up window), click repeat and execute to determine which trough has an insufficient volume of liquid. The error message should appear again. You can then add additional reagent and click repeat and execute again to continue the run.



4.2.4 The following describes the actions of each recovery option in the pop up window when an error is encountered. Choose the appropriate action or contact QA for assistance:

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Recovery Button		
Text	Description	
Abort	Run is aborted	
Cancel	Run is canceled and a defined user error handler is started	
Repeat	The command is repeated	
	The channel or position is excluded until a new tip pick up is	
Exclude	called.	
Waste	The tip is ejected to the default waste	
Air	Rest of missing volume is filled up with air	
Bottom	Repeats the aspiration on container bottom	
Continue	Continue without any change	
Barcode	Barcose is assigned manually	
Next	Repeat the command on next sequence position	
Available	Aspirate & dispense the available volume.	

5 Daily Maintenance

5.1 Maintenance should be performed before the first run on a calendar day. Check to see if the daily maintenance was performed by clicking on The Microlab STAR Maintenance & Verification icon on the desktop.



- 5.2 Check the Processed Date/Time to determine the date maintenance had last been completed.
 - 5.2.1 Note that the Expired Date/Time is registered based on a 24hour time frame; therefore, the system may not say required even if the Maintenance has not been performed that day.
- 5.3 If the maintenance had not been completed for the day, follow the prompts as described below.

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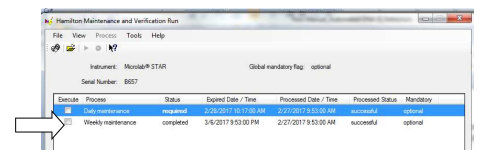
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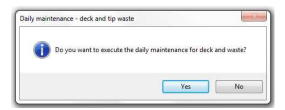
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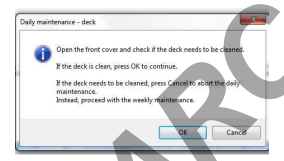
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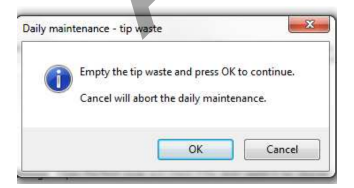
5.3.1 A window will appear after the robot initializes. Click "Yes."



5.3.2 Inspect the deck to make sure it is clean. If the deck is clean, press OK. If not, press Cancel and alert QA to run the weekly maintenance.



5.3.3 If the tip waste is full, empty it; otherwise, press OK.

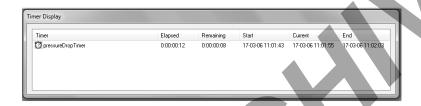


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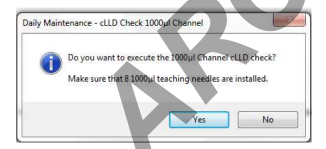
5.3.4 When prompted "Do you want to execute the 1000ul Channel tightness check?" click Yes.



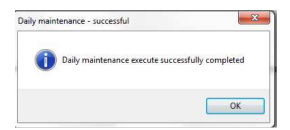
5.3.5 The window below will then appear. Wait while the channels perform a pressure check.



5.3.6 When prompted, "Do you want to execute the 1000ul Channel cLLD check?" click Yes.



5.3.7 Click OK once the maintenance has successfully completed.



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- 5.4 Update the daily maintenance log in LIMS by doing the following:
 - 5.4.1 Navigate to the LIMS tramstop "Other", then instrument → *instrument name* → select instrument → click [maintenance log] → Add → select "daily maintenance performed" in the Process Performed from the dropdown menu and click Save Entry.

