

For Research Use Only. Not for use in diagnostic procedures.

Amplify DNA

□ 1	Add DNA into either of the following to create a
	DNA plate:
	Midi plate: 20 µl to each DNA well
	► TCY plate: 10 µl to each DNA well
\square 2	Select MSA6 Tasks Make MSA6.
\square 3	Select the DNA plate type.
$\square 4$	Enter the Number of DNA plates .
\square 5	Place the MA1, RPM, and MSM tubes in the
	robot tube rack.
□6	Pour 15 ml NaOH into a trough and place on
	the robot bed.
\square 7	Place DNA and MSA6 plates on robot bed.
8	Select Run.
9	Enter the barcode of each DNA plate.
□ 10	Place the DNA plates on the robot bed and select OK .
□ 11	Vortex the MSA6 plate at 1600 rpm for
	1 minute.
□ 12	Centrifuge at 280 × g.
□ 13	Remove the cap mat, place the MSA6 plate
	on the robot bed, and select OK .
□ 14	When complete, select OK.
□ 15	Remove and seal the MSA6 plate.
□ 16	Centrifuge at 280 × g.

Incubate DNA

	[LIMS] Select Infinium LCG
[a Scan the barcodes.
□2	Incubate the MSA6 plate for 20–24 hours at 37°C.

Fragment DNA

\Box 1	Pulse centrifuge the MSA6 plate at $280 \times g$.
\square 2	Select MSA6 Tasks Fragment MSA6.
\square 3	Place the MSA6 plate on the robot bed.
$\square 4$	Place FMS tubes in the robot tube rack.
\Box 5	Select Run.
□6	When complete, select OK.
\square 7	Remove the plate and seal with a cap mat.
8	Vortex at 1600 rpm for 1 minute.
9	Pulse centrifuge at 280 × g.
□ 10	Place on the 37°C heat block for 1 hour.

SAFE STOPPING POINT

If you are stopping, seal the plate(s), and store at -25°C to -15°C.



☐ 22 Quickly invert the plate and drain the

☐ 23 Firmly tap until all wells are free of liquid.

supernatant.

Infinium LCG Assay Automated Workflow Checklist

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Precipitate DNA	24 Place the plate on a tube rack for 1 hour at room temperature.	Resuspend DNA
□ 1 Select MSA6 Tasks Precip MSA6. □ 2 Pulse centrifuge the sealed plate at 280 × g. □ 3 Place the MSA6 plate on the robot bed. □ 4 Place a half reservoir in the frame, and add PM1 as follows: ■ For 48 samples, add 1 tube PM1 ■ For 96 samples, add 2 tubes PM1 □ 5 Place a full reservoir in the frame, and add 2-propanol as follows: ■ For 48 samples, add 20 ml 2-propanol ■ For 96 samples, add 40 ml 2-propanol ■ Select Run. □ 7 Remove the MSA6 plate from the robot bed. Do not select OK. □ 8 Vortex at 1600 rpm for 1 minute.	□ 25 Make sure that a blue pellet is still present. SAFE STOPPING POINT If you are stopping, seal the plate(s), and store at -25°C to -15°C.	 □ 1 Select MSA6 Tasks Resuspend MSA6. □ 2 Place the MSA6 plate on the robot bed. □ 3 Place a quarter reservoir in the frame, and add RA1 as follows: ▶ For 48 samples, add 4.5 ml RA1 ▶ For 96 samples, add 9 ml RA1 □ 4 Select Run. □ 5 Remove the MSA6 plate from the robot deck. □ 6 Apply a foil seal to the MSA6 plate. □ 7 Incubate in the Illumina Hybridization Oven for 1 hour. □ 8 Vortex at 1800 rpm for 1 minute. □ 9 Make sure that the pellets are resuspended. □ 10 Pulse centrifuge at 280 × g.
9 Incubate on the heat block for 5 minutes.		SAFE STOPPING POINT
 □ 10 Centrifuge at 280 × g for 1 minute. □ 11 Set the centrifuge at 4°C. □ 12 Place the MSA6 plate on the robot bed. 		If you are stopping, store sealed MSA6 plate(s) at 2°C to 8°C for up to 24 hours. If more than 24 hours, store at -25°C to -15°C.
☐ 13 Select OK.☐ 14 Remove the MSA6 plate from the robot bed and seal.		Store sealed RA1 at -25°C to -15°C. If RA1 will be used the next day, seal it, and store it overnight at 4°C.
☐ 15 Invert 10 times to mix.		т О.
16 Incubate at 4°C for 30 minutes.		
17 Place in the centrifuge.		
18 Centrifuge at 3000 × g for 20 minutes.		
19 Remove MSA6 plate.		
20 Make sure that a blue pellet is present.		
21 Remove and discard the cap mat.		



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Hyb	oridize DNA to the BeadChip	23 [LIMS] Select Infinium LCG Prepare Hyb Chamber.
□ 1	Incubate the MSA6 plate on the heat block for	a Scan the barcodes.
	20 minutes.	☐ 24 Incubate at 48°C for 16-24 hours.
\square 2	Cool at room temperature for 30 minutes.	
\square 3	Pulse centrifuge at 280 × g.	
\square 4	Place the gasket into the hybridization	
	chamber.	
<u></u> 5	Add 400 µl PB2 into each reservoir.	
□6	Place the hybridization chamber insert into the	
	hybridization chamber.	
□7 □0	Immediately cover the chamber with the lid.	
□8	[LIMS] Select Select Infinium LCG Confirm for Hyb.	
9	[LIMS] Scan the barcodes.	
□10	Remove all BeadChips from packaging.	
□ 11	Place BeadChips into the robot BeadChip	
	alignment fixtures.	
□ 12	Select MSA6 Tasks Tasks Hyb.	
L	a Select the 24-sample BeadChip.	
L 10	b Enter the Number of MSA6 plates.	
□ 13	Place the robot BeadChip alignment fixtures onto the robot deck.	
\Box 14	Pulse centrifuge the MSA6 plate at 280 × g.	
	Place the MSA6 plate onto the robot deck.	
	Select Run.	
	Place each robot tip alignment guide on top of	
	each robot BeadChip alignment fixture.	
□18	To start the run, select OK .	
□ 19	When complete, select OK.	
□ 20	Remove the robot BeadChip alignment fixtures.	
□21	Place each BeadChip in a hybridization	
	chamber insert.	
□22	Place the lid on the chamber and secure with	
	the metal clamps.	
	į.	

Prepare for Next Day ☐1 Add 330 ml fresh 100% EtOH to the XC4 bottle. ☐2 Vigorously shake to resuspend. ☐3 Leave the bottle upright on the lab bench

☐ 4 Soak the robot tip alignment guides in 1% aqueous Alconox solution.

overnight.

☐ 5 Rinse and dry the robot tip alignment guides.



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/Va	sh BeadChips	with DI H ₂ O.	Exte	end and Stain	BeadChip	S	
1 2 3 4	Submerge the wash rack in the PB1 wash. Remove the hybridization insert. Remove the BeadChips. Remove the cover seals from the BeadChips.	WILLIAM NEWS		Fill the water circul Select Robot QC T to set to 44°C. Select XStain Task	asks Circulator		
3 5	Place the BeadChips into the submerged		$\Box 4$				
	wash rack.			Reagent	# BeadChips	Volume	
1 6	Move the wash rack up and down for 1 minute.			95% formamide/1 mM EDTA	1–8	15 ml	
	Move the wash rack to the next PB1 Wash.				9–16	17 ml	
8	Move the wash rack up and down for 1 minute.				17–24	25 ml	
]9	Confirm that you are using the correct Infinium			RA1	1–8	10 ml	
	LCG glass back plates and spacers.				9–16	20 ml	
<u> </u>	Fill the BeadChip alignment fixture with 150 ml PB1.				17–24	30 ml	
711	For each BeadChip, place one black frame			XC3	1–8	50 ml	
	into the BeadChip alignment fixture.				9–16	100 ml	
12	Place each BeadChip into a black frame.				17–24	150 ml	
13	Place a <i>clear</i> spacer onto the top of each						
_	BeadChip.		Ш5	Invert the LX1, LX2			
_14	Place the alignment bar onto the alignment fixture.			tubes to mix. Remethe robot deck.	ove the caps, ar	id place on	
715	Place a clean glass back plate on top of each		□6	Enter the number of	of BeadChips.		
_ 10	clear spacer.		\Box 7	Select Run.	•		
□16	Secure each flow-through chamber assembly with metal clamps.		□8	[Non-LIMS]Enter the stain temperature listed on the SML tube.			
□ 17	Remove the assembled flow-through chamber		□9	Place the flow-thro	ugh chambers i	nto the	
	from the alignment fixture.			chamber rack.			
☐ 18	Trim the spacers from each end of the		□ 10	Select OK .			
	assembly.		□11	Remove the flow-t	nrough chambe	rs from the	
<u> </u>	Leave assembled flow-through chambers on			chamber rack.			
	the lab bench.		∐12	Set up two top-loa PB1 and XC4.	ding wash dishe	es labeled	



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\square 13	Add 310 ml PB1 to the PB1 wash dish.
□ 14	Submerge the staining rack in the wash dish.
□ 15	Leave the staining rack in the wash dish.
□16	Disassemble each flow-through chamber.
	Place the BeadChips into the submerged staining rack.
□18	Slowly lift the staining rack 10 times.
	Soak for 5 minutes.
□ 20	Vigorously shake the XC4 bottle.
_	Add 310 ml XC4 to the XC4 wash dish and cover.
□ 22	Transfer the staining rack from the PB1 to the XC4.
□ 23	Slowly lift the staining rack 10 times.
□ 24	Soak for 5 minutes.
□ 25	Remove the staining rack and place it onto the tube rack.
□ 26	Dry each BeadChip as follows.
	a Grip the BeadChip by the barcode end. b Place onto a tube rack with the barcode
□ 27	facing up and toward you. Place the tube rack into the vacuum
LJ 21	desiccator.
□28	Dry the BeadChips for 50–55 minutes at 675 mm Hg (0.9 bar).
	[LIMS] Select Infinium LCG Coat. a Scan the barcodes.