Quick Start Protocol Avida DNA

G9409-90500 Rev B0 For Research Use Only. Not for use in diagnostic procedures.





Nuration estimates are provided as guidelines for 8 reaction runs using 10 ng cfDNA. Your results may vary.

This Quick Start Protocol provides key protocol details for experienced users. Visit Avida DNA Reagent Kits Protocol for the full User Guide with further information including:

- Safety information
- Notices to purchaser
- List of required materials
- QC output examples

- Critical factors for system performance including component handling, mixing, and spinning specifications; pay special attention to the key critical factors such as mixing steps highlighted in this Quick Start Protocol
- Full protocol details including Notes and Caution statements
- NGS support

PCR-free Library Preparation

Note: Reagent vial cap colors are indicated by colored circles adjacent to reagent use instructions. Clear cap is indicated by "c" and white cap is indicated by "w".

in next step.				Step 2: Ligate UMI-tagged adaptors			3 0 m
ep 1: End repair a	and dA-tail fraç	gments	🚺 1 hour	Vortex thawed Ligat	ion Buffer.		
Vortex thawed End P	rep Buffer at RT till a	II particles are o	lissolved.	Prepare Ligation Ma spin > keep on ice.	ster Mix at RT (Lig	MM, see Table 3)	. Vortex > briefly
Add appropriate quar	itity of DNA sample	in strip tube. Ma	ake volume up to	Table 3: Lig MM reage	nt volumes (including	excess for suppo	rted run sizes).
50 µL with nuclease-f	ree water > keep on	ice.		Reagent	Per Rxn	8 Rxn	16 Rxn
Sample quantity rar	nge:			 Ligation Buffer 	25 µL	225 µL	450 μL
,				Ligation Enzyme	6 µL	54 µL	108 µL
Prepare End Prep Mix	(at RT (see Table 1)	Gently vortex a	t low speed >	Total	31 µL	279 µL	558 µL
brief spin. <i>The End Prep Mix is s</i>	table for up to 2 hou	ırs at 4°C.		Reagents are viscou			
Table 1: End Prep Mix re	eagent volumes (inclue	ding excess for s	upported run sizes).	📃 Add 5 µL of ● Adap	tor for ILM to each	DNA sample.	
Reagent	Per Rxn	8 Rxn	16 Rxn	Do NOT add Adaptor for ILM to the Lig MM. Avoid exposing the Adaptor			
End Prep Buffer	7 µL	63 µL	126 µL	to RT conditions.			
End Prep Enzyme	3 µL	27 µL	54 µL	Add 31 µL of Lig MN	I to coop comple N	live briefly opin	
Total	10 µL	90 µL	180 µL		i to each sample. N	nix > brieny spiri.	
Add 10 µL of End Pre	p Mix to each DNA s	ample. Mix > br	iefly spin > keep on ice.	Program the thermarun the program.	l cycler with heated	l lid turned off. Lo	bad the strip tubes
	cycler (Table 2) with	heated lid set t	o 75°C. Load the strip	Table 4: Thermal cycle	r program for end pre	ep.	
Program the thermal				Step	Tempe	rature	Time
Program the thermal tubes > run the program	program for end prep.			Step 1	20°C		30 min
		ture	Time	Step 2	4°C		Hold
tubes > run the progra	Temperat						
tubes > run the progra Table 2: Thermal cycler			30 min				
tubes > run the progra Table 2: Thermal cycler Step	Temperat		30 min 30 min	At 4°C Hold step, rer	nove from cycler >	keep at RT.	
tubes > run the progra Table 2: Thermal cycler Step Step 1	Temperat 20°C			At 4°C Hold step, rer	-	•	

Step 3: Library bead binding

🔰 30 min

Thoroughly vortex @Library Binding Beads. Allow to settle for 5 min.

Add 87 μ L (~0.9X volume) of Library Binding Beads to each adaptor-ligated DNA sample, bringing sample volume to 183 μ L. Mix > spin. Incubate at RT 10 min.

Bubbles can compromise library binding efficiency. Pipette slowly.

During 10-min incubation: Prepare Hyb Mix 1 (see Table 5). Vortex > briefly spin > keep at RT till needed.

Table 5: Hyb Mix 1 reagent volumes (including excess for supported run sizes).

	Reagent	Per Rxn	8 Rxn	16 Rxn
©	Nuclease-Free Water	17.5 µL	157.5 μL	315 µL
	Avida DNA Panel	4 µL	36 µL	72 µL
٠	Hyb Blocker	2.5 μL	22.5 µL	45 µL
	Total	24 µL	216 µL	432 µL

Collect beads using magnet (\sim 2 min) > remove and discard supernatant.

Wash 1× or 2× (optional) using Library Wash Buffer. For each wash:

- □ Add 180 µL of Library Wash Buffer to each tube without disturbing the bead pellet.
- □ Incubate at RT 2 min > Remove and discard all supernatant. Washes completed: □ 1 □ 2

Remove tubes from magnet.

Proceed immediately to Hybridization.

Hybridization/Capture/Indexing

Step 1: Hybridize libraries to probe



Add 24 µL of Hyb Mix 1 to the beads bound with adaptor-ligated library generated in the last step. Resuspend beads by gentle vortexing or pipetting > Briefly spin.

Thoroughly vortex the Hyb Buffer that has been kept at 37°C. Keep at RT until later use.

Prepare Hyb Mix 2 based on Avida DNA panel size (see Table 6). Vortex > briefly spin > keep at RT.

Table 6: Hyb Mix 2 reagent volumes for panels <500kb and \ge 500kb (including excess for supported run sizes).

	Reagent	Per Rxn		8 Rxn		16 Rxn	
		<500kb	≥500kb	<500kb	≥500kb	<500kb	≥500kb
©	Hyb Buffer	30 µL	31 µL	270 µL	279 µL	540 µL	558 µL
0	Hyb Enhancer	6 µL	5 µL	54 µL	45 µL	108 µL	90 µL
	Total	36 µL	36 µL	324 µL	324 µL	648 µL	648 µL

Add 36 μL of Hyb Mix 2 to each bead mix containing Hyb Mix 1. Mix > spin.

Program the thermal cycler (Table 7) with heated lid set to 103°C. Load the strip tubes > run the program.

Table 7: Thermal cycler program for hybridization.

Step	Temperature	Time
Step 1	98°C	2 min
Step 2	≥2.5°C/second ramp down to 60°C	
Step 3	60°C	60 min
Step 4	60°C	Hold

For panels >500 kb, a longer hybridization time (up to 16 hours) may improve capture efficiency.

Step 2: Prepare capture beads (concurrently) Step 3: Capture and wash hybridized libraries 15 min 30 min Thoroughly vortex Hyb Buffer that has been kept at RT. After hybridization, spin tubes and place on magnet (1 min). Thoroughly pipett or vortex Capture Beads stock till solution is homogeneous. Program thermal cycler for washing (Table 9) with heated lid set to 75°C. Table 9: Thermal cycler program for hybridization. Calculate volume of Capture Beads needed (8 µL/sample + overage). Step Temperature Time Transfer into 1.5-mL tube. Step 1 60°C Hold Volume calculated: Transfer each supernatant to tubes containing 8 µL aliquots of prepared Place tube on magnet (1 min) > discard supernatant > remove from magnet. Capture Beads > Mix and spin. Wash beads 2× in 1preheated Hyb Wash Buffer 1 using volume in Table 8. Incubate in thermal cycler held @60°C (heated lid 75°C) for 10 minutes. Table 8: Wash volume of Hyb Wash Buffer 1 based on number of reactions. # of rxns: 1-16 17-32 33-48 49-64 65-80 81-96 After 10 minutes @60°C, remove tubes > place on magnet > discard supernatant 100 µL 200 µL 300 µL 600 µL Volume: 400 µL 500 µL > remove from magnet. Leave the thermal cycler running. Do NOT vortex or spin the samples until instructed to do so. Handle carefully to For each wash: prevent any splashing. □ Add required volume of the preheated Hyb Wash Buffer 1 to beads. Vortex > briefly spin > place tube on magnet (1 min) > discard supernatant. Add 150 µL preheated Hyb Wash Buffer 1 to each tube. Mix well. Washes completed: $\Box 1 \quad \Box 2$ Place tubes on magnet (30 sec) > discard supernatant > remove from magnet. Resuspend Capture Beads in Hyb Buffer at original volume calculated above Add 100 µL Hyb Wash Buffer 1 to each tube > mix well. > mix well. Transfer each sample to a fresh strip tube and cap tubes. Add 8 uL resuspended Capture Beads to each tube of a fresh strip tube. Incubate in thermal cycler held @60°C (heated lid 75°C) for 3 minutes. After 3 minutes, stop the program > place tubes on magnet (30 sec) > discard supernatant > remove samples from magnet.

Add 150 µL Hyb Wash Buffer 2 kept at RT. Mix > briefly spin.

Stopping point

Samples can be stored in Hyb Wash Buffer 2 at 4°C overnight. Do NOT discard the beads.

Place tubes on magnet (1 min) > discard all supernatant > remove tubes from magnet.

Resuspend tubes in 20 μL \bullet Resuspension Buffer. Vortex > briefly spin.

Indexing/Library Purification

Step 1: Amplify and dual-index

Thaw @2X Amplification Mastermix and Avida Index Primer Pairs on ice. Vortex 2X Amplification Mastermix at low speed. Briefly centrifuge Avida Index Primer Pairs.

Set up Indexing PCR reaction: 20 μ L resuspended Capture Beads + 25 μ L 2X Amplification Mastermix + 5 μ L of the appropriate Avida Index Primer Pair. Mix > briefly spin.

Program thermal cycler for indexing PCR (Table 10) with heated lid set to 103°C. Load the strip tubes > run the program.

Table 10: Thermal cycler program for indexing PCR

Step	Number of Cycles	Temperature	Time
Initial denaturation	1	98 °C	45 sec
Amplification stage 1	5	98 °C	10 sec
		63 °C	30 sec
		72 °C	30 sec
Amplification stage 2 (based on sample input of 10 ng cfDNA)	 11 for Discovery Cancer panel 14 for Expanded Cancer panel 16 or fewer for >100 kb 	98 °C	10 sec
	 17 for >50 kb to 100 kb 18 for 10 kb to 50 kb (Focused Cancer panel) 19 for <10 kb 	72 °C	1 min
Final extension	1	72 °C	1 min
Final hold	1	4 °C	Hold

- □ High-quality gDNA samples typically require 1-2 cycles more than cfDNA samples of the same input quantity.
- Low-quality gDNA samples (e.g., FFPE-derived gDNA) may require more cycles than high-quality gDNA samples.
- □ A sample input >50 ng typically requires 1-2 fewer cycles than a sample input of 10 ng.
- At 4 °C Hold step, remove from cycler > keep on ice.

Stopping point

1 hour

Store indexed libraries at 4° C overnight or -20° C for up to 72 hours. Avoid prolonged storage >72 hours.

Prep Ahead: Put AMPure XP beads at RT for use in next step (30 min equilibration). Prepare fresh 80% ethanol (400 µL/sample) for use in next step.

ads + 25 µL 2X <primer Pair.</p> Collect beads using magnet (2 min) > remove and discard supernatant. Wash beads 2× with 200 µL freshly-prepared 80% ethanol. For each wash: Add 200 µL of 80% ethanol to beads without disturbing pellet. Incubate for 30 sec > discard supernatant. Washes completed: □1 □2 Remove residual ethanol and air-dry beads for up to 3 min.

Remove tubes from magnet. Add 23 μ L of 1X Low TE Buffer to each tube. Mix > briefly spin > incubate at RT 2 min.

Get AMPure XP beads held at RT, vortex until homogeneous.

Add 50 µl of AMPure XP beads to each amplified DNA well > mix well.

Collect beads using magnet (2 min).

Step 2: Clean-up final libraries

Incubate at RT 5 min.

Transfer 20 μL of the eluate from each tube to a new tube, being careful to avoid bead carryover.

Stopping point

30 min

Store indexed libraries at 4°C overnight or -20°C for prolonged storage.



Step 3: Quality control

Assess quality and quantity using one of these platforms:





Agilent 4200 or 4150 TapeStation System with D1000 Assay Agilent 5200, 5300 or 5400 Fragment Analyzer System with NGS Fragment Kit (1–6000 bp) or other suitable kit

Stopping point

Varies

Store indexed libraries at 4° C overnight or -20° C for prolonged storage.

Pool libraries for multiplex NGS

- Calculate the number of indexes that can be combined per lane, according to sequencer capacity and amount of sequencing data required per sample.
- Combine the libraries such that each index-tagged sample is present in equimolar amounts in the pool using one of the following methods:
- □ Dilute each sample to be pooled to the concentration of the most dilute sample using Low TE > combine equal volumes of all samples in final pool.
- Combine the appropriate volume of each sample to achieve equimolar concentration in the pool, then adjust the pool to the desired final volume using Low TE. See the assay user guide for more information on this method.

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Run Date

Operator

Probe Name or Design ID

Index Pairs

Reagent Lot Information

- DNA Reagent Box 1:
 - Avida DNA and Duo Reagent Box 2:
- Probe:
- Avida Beads Box:
- Avida Index Primer Pairs:

Sample Information

Library Pooling Information

Comments



This information is subject to change without notice.