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Protocol status: Working
We use this protocol and it's working

Illumina library preparation and dual hybridization protocol of ERC GLOBAL

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DISCLAIMER

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Bill Baker and Donovan Bailey are thanked for drawing our attention to this option and sharing preliminary data and results.

ABSTRACT

Step by step protocol for library preparation of Annonaceae species within the ERC GLOBAL project. This protocol was followed for all libraries prepared within the project. Each step involves a purification phase (AMPure XP protocol), and is detailed at the start of this protocol. The protocol is done per series of 48 samples at a time with indexing.

The protocol also describes the steps for the dual hybridization of Daicel Arbor Biosciences Mybaits Angiosperm353 kit + Custom Annonacea baiting kit. The protocol is based on hybridisation protocols used at DIADE research unit IRD Montpellier France and on the Hybridization Capture for Targeted NGS User Manual version 5 protocol "standard" of Daicel Arbor Biosciences Mybaits.

To proceed DNA extraction of plant and in particular from herbarium and silicagel specimens of annonacea read the Annonaceae DNA extraction protocol from silicagel dried and herbarium preserved leaves

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Grant ID: 865787

ATTACHMENTS

[TAG ADAPT INDEX
sequence TX.csv](#)

IMAGE ATTRIBUTION

Photo by Thomas L.P. Couvreur

GUIDELINES

This library preparation & dual hybridisation protocol follows and is inspired by Henriksen et al (2021) using both the Annonaceae specific baits and the Angiosperm353 baits (see references). In the end we sequence both the Annonaceae specific baits and the Angiosperm353 universal ones.

Both the Annonaceae (Couvreur et al. 2019) and Angiosperm353 (Johnson et al., 2019) bait kits are available as Arbor Biosciences "myBaits" kits (Daicel Arbor Biosciences, Ann Arbor, Michigan, USA; <https://arborbiosci.com/genomics/targeted-sequencing/mybaits/>).

MATERIALS

- Pipettes and cones with "low bind" filters (1000-200-20-10µL)
- Eppendorf tubes 1.5mL-2mL-0.2mL, aluminum films, refrigerated holder, magnetic holder, centrifuge and vortex
- PLATES RT-PCR 96 (Roche)
- VWR®, Centrifugeuse de microplaques, PCR Plate Spinner
- Lightcycler 480 (Roche)
- QIAxcel "Advanced system" analyze + QIAxcel DNA screening kit (2400) or other bioanalyser
- Personal protective equipment (PPE): gown, gloves, goggles
- DNA Dilution Buffer
- Agencourt AMPure XP solution
- 70% ethanol (fresh)
- DNase/RNase Free Water
- Thermo Scientific™ ADN ligase T4 (5 U/µl) EL0013 Code produit.10548730
- Bst DNA Polymerase, Large Fragment
(NEB#: M0275)
- DNTP Mix , REF U151B 10mM U1515 or 1511 (200 or 1000µL)
- KAPA HiFi HS Real-Time Master Mix (2X) + Real-Time Standards 1 to 4
- Kit MYbaits ® "In-Solution Sequence Capture for Targeted High-Throughput Sequencing".

PROTOCOL MATERIALS

 PCR Grade Water **Catalog #AM9935** Step 2.6

 Agencourt AMPure XP **Beckman Coulter Catalog #A63880** In 9 steps

 70% EtOH In 2 steps

SAFETY WARNINGS

 Wear protections

BEFORE START INSTRUCTIONS

Clean bench and pipettes with water and then ethanol. Follow the instructions for use of the Arbor Biosciences V5 kit.

check if you have all the material needed for all the step or equivalent tools

Introduction and AMPure XP protocol

30m

- 1 Library preparation protocol is carried out after the DNA extraction detailed in :

Protocol



NAME

Annonaceae DNA extraction protocol from silicagel dried and herbarium preserved leaves

CREATED BY

Thomas LP Couvreur

[PREVIEW](#)

To start we need purified DNA diluted in water or TE buffer 50µL or more at 2 to 900ng/µL below 60ng/µL samples are considered low quantity if only diluted in 50µL samples with size less than 200pb is considered low quality/small fragments dna (use 1.8X ampure)

2 AMPURE XP DNA Purification protocol

On each steps of the protocol a purification step is need to change solution, concentrate or purify dna.

Each purification step is done with  Agencourt AMPure XP **Beckman Coulter Catalog #A63880**

The only changes between the different purification steps is the start volumes as such: volume ratio 1X to 1.8X AMPure. For example for 1.8X: for 100 µL to be purified added 180 µL Ampure

Below we provide a protocol for 1X  Agencourt AMPure XP **Beckman Coulter Catalog #A63880**

for start volume of  50 µL of DNA

resuspension (elution) volume =  20 µL.

The protocol is described for 96 well pcr plates, but can also be done in 1.5/ 2mL tubes.

2.1



Use  Agencourt AMPure XP **Beckman Coulter Catalog #A63880** at

 Room temperature , take out 30 minutes in advance.

Prepare a 70% ethanol solution.

Add  50 µL AMPure XP in each well.

2.2



Mix by pipetting or shake and wait  00:10:00 min. quickly spin down 10sec using pcr plate  10m

spinner if pcr plate or mini centrifuge for tubes

DNA will bind to the beads

2.3



Place the plate on the magnetic stand and wait  00:05:00 min.  5m

Remove liquid phase by pipetting , **avoid pipetting the beads.**

2.4



Keep on magnetic stand add  150 µL  70% EtOH **Contributed by users** .  1m

Remove liquid phase by pipetting, **avoid pipetting the beads.**

2.5



Repeat wash step on magnetic stand and add  150 µL

 70% EtOH **Contributed by users**

Remove liquid phase by pipetting, **avoid pipetting the beads.**

Allow to dry for 5 minutes at Room temperature

- 2.6** Take PCR plate out of the magnetic stand and add the needed "elution volume"; here 20 µL 10mL



of ultra pure water PCR Grade Water **Contributed by users Catalog #AM9935**
Mix by pipetting.

Wait 00:10:00 minutes at Room temperature for beads to resuspend in water.

- 2.7** The AMPure purification step is over.

Make sure to place the plate on the magnetic stand to avoid pipetting beads for next usage or quantification.

Mind that purification step reduces DNA quantity by 20 to 50% and removes small DNA fragments.

To keep small fragments use 1.8X AMPure ratio instead of 1X.

Part 1: library preparation , DNA preparation and shearing

2h

- 3** Depending on concentration of each sample, dilute the DNA in 0.2 mL microtubes Diagenode suitable for shearing with ultrasound. Target amount of DNA should, if possible, be 3-4 µg in 50 µL (to be supplemented with water). The number of shearing cycles and their duration can be modified depending on the initial degradation state of the DNA and/or the desired fragment size.
Shearing is done with the below equipment:

Equipment

Bioruptor Pico sonication device

NAME

Sonicator

TYPE

Diagenode

BRAND

B01060010

SKU

- 3.1** Pulse centrifuge the tubes.

1m

3.2 Deposit  50 µL of DNA and water if needed to dilute to match concentration to the tubes in the Pico Bioruptor (Diagenode).

3.3 Start N cycles  00:00:15 ON /  00:00:30 OFF at  4 °C.

45s

Depending on the DNA extraction control gel 0 < N < 10.

Expected result

For low quality small fragments herbarium specimen 0-3 cycles

For high quality herbarium specimen 3-8 cycles

For Silicagel specimens 5-10 cycles

3.4 Check size using  3 µL DNA + 10 µL DNA dilution buffer AL420 protocol with QIAxcel, if necessary undertake additional cycles.

Pattern must be between 150 and 600 bp, but depending on the initial degradation, it may be centred on 100 bp.

QIAxcel QIAGEN (DNA Screening Kit)

3.5 Restart N+B cycles  00:00:15 ON /  00:00:30 OFF at  4 °C.

45s



3.6 Measure DNA size again using QIAGEN QIAxcel Bioanalyser



 3 µL DNA + 10 µL DNA dilution buffer AL420 protocol analysis.

or other bioanalyser, you can also make a DNA gel

Expected result

We aim for 4µg DNA post shearing and optimal DNA size of 400pb ranging between 100-600pb.

volume after shearing step 40-50µL (40 needed for next step)

Fast End Repair & phosphorylation

1h

- 4 Prepare a buffer and enzyme mix according to the number of samples to be processed

A	B
For 1 sample:	Volume
Sheared DNA	40 µL
End Repair Reaction Buffer 10X	5 µL
End Repair Enzyme Mix	5 µL
Final volume	50 µL

Buffer mix needed per sample

NEBNext End Repair Module E6050L or E6050S

- 4.1 Distribute Δ 10 µL of enzyme and buffer mix in each well.

5m

- 4.2 Add Δ 40 µL of DNA from previous step, vortex and Spin down the sample (using pcr plate spinner).

5m

4.3 Leave  00:30:00 at  Room temperature .

30m

4.4 Stop the reaction using a purification step AMPure XP 1X ($50\mu\text{L}$) or 1.8X ($90\mu\text{L}$) if you want to retain small DNA fragments. See introduction for details of the purification step using  Agencourt AMPure XP Beckman Coulter Catalog #A63880

4.5 Elute the samples in  $30\mu\text{L}$ water and quantify using Nanoquant.



Expected result

optional if low quantity DNA elute on $16\mu\text{l}$ ($2\mu\text{L}$ for nanoquant and 14 for next step)

Ligation

3h

5 Depending on the previous dosage, calculate the volume to be taken per sample to ideally obtain  200 ng of DNA in  $13.8\mu\text{L}$.

If necessary, dilute samples that are too concentrated .

Careful not to vortex the ligase. Mix by flicking the tube.

5.1 Take the samples, place them on a 96-well plate on a refrigerated stand, and adjust all the volumes to  $13\mu\text{L}$ with water.

5.2 Then add to each sample  $2\mu\text{L}$ of TAG-P5 adapt ($4\mu\text{M}$).

5.3 Prepare a volume of Mix according to the number of samples:

A	B
1 sample (DNA = 200 ng)	
DNA + water	13.8 µL
adaptateur P5 with tag 4µM	2 µL
adaptateur P7 40µM	0.2 µL
T4 DNA ligase 5U/µL	2 µL
Buffer T4 10X	2 µL
Final Volume	20 µL

Mix volume for 1 sample

Update the thawing/freezing number of P7 P5 adapters on their storage box.

NB; Tag referses as P5 6pb barcoded adapter used as index but not read by illumina ; demultiplexed after sequencing.

adapter p7 & p5 sequences on attached file

Thermo Scientific™ ADN ligase T4 (5 U/µl) EL0013

5.4 Deposit 4.2 µL of Mix by well, vortex briefly and pulse centrifuge.

5.5 Place the plate on the thermocycler and launch the "LIG" program:

A	B	C
Step	Temperature	Duration
1	+22°C	2H30
2	+65°C	10 min
3 optional	+4°C	20 min

LIG program on thermocycler

5.6 Purify 20 μ L post ligation using



Agencourt AMPure XP **Beckman Coulter Catalog #A63880** with XP1.8X ratio
 ($\text{吸取 } 36 \mu\text{L}$ of ampure);

Then elute in $\text{吸取 } 25 \mu\text{L}$ ultra pure water.

Expected result

25 μ L of post ligation purified DNA solution 23.5 is needed for next step

optional you can conserve for 24h sample after purification step at -20°C

Nick Fill-in (Bst DNA Polymerase)

45m

6 Position the 96-well plate on a refrigerated stand.

5m

Prepare a volume of Mix according to the number of samples:

A	B
For 1 sample:	
DNA	23.5 μ L
DNTPs 5mM	1.5 μ L
Bst DNA POLYMERASE 8U/ μ L	2 μ L
Thermo Pol Buffer 10X	3 μ L
Final Volume	30 μ L

Mix for 1 sample

Bst DNA Polymerase, Large Fragment

(NEB#: M0275)

DNTP Mix , REF U151B 10mM U1515 or 1511 (200 or 1000 μ L)

6.1 Deposit $\text{吸取 } 6.5 \mu\text{L}$ of Mix per well in a AB1400 pcr plate and pulse centrifuge.

1m

6.2 Deposit $\text{PCR tube icon } 23.5 \mu\text{L}$ of DNA per well, mix with the pipette and pulse centrifuge.

6.3 Place the plate on the thermocycler and launch the "BST" program:

35m

A	B	C
Step	Temperature	Duration
1	+37°C	15 min
2	+10°C	20 min

BST program on thermocycler

6.4 Purify using PCR tube icon Agencourt AMPure XP **Beckman Coulter Catalog #A63880** XP 1X (



$\text{PCR tube icon } 30 \mu\text{L}$; see section 1.

Elute in $\text{PCR tube icon } 15 \mu\text{L}$ ultra pure water.

Expected result

15 μL of post BST purified solution, 12 μL is needed for PCR step optional you can use the remaining 3 μL for quantify or check DNA/library size and quality pre PCR

optional you can conserve for 24h sample after purification step at -20°C and done PCR step the next day

PCR1 amplification + Purification AMPure

1h

7 In a LightCycler plate® 480 Multiwell plate 96 white arranged on refrigerated stand: deposit $\text{PCR tube icon } 30 \mu\text{L}$ of each fluorescent standards ref KK2701 (1 to 4) in duplicates in 8 separate wells. Keep the plate away from

light.

7.1 Prepare a volume of Mix according to the number of samples:

A	B
For 1 sample	volume
DNA	12µL
adapter PE 34 solution 10µM	1.5µL
INDEX 10µM	1.5µL
KAPA HiFi Master Mix 2X	15µL
Final Volume =	30µL

Mix for 1 sample

Roche KAPA Hifi PCR KIT 250rx

Index and adapter sequence on attached file

Index used: AD001; AD002; AD003; AD004; AD005; AD006; AD007; AD008; AD009; AD0010; AD0011; AD0012.

7.2 Deposit 18 µL of Mix per well and pulse centrifuge.

7.3 Deposit 12 µL of DNA per well, mix with the pipette and pulse centrifuge.

7.4 Place the plate in the lightcycler and launch the following the "RTLib1" program.

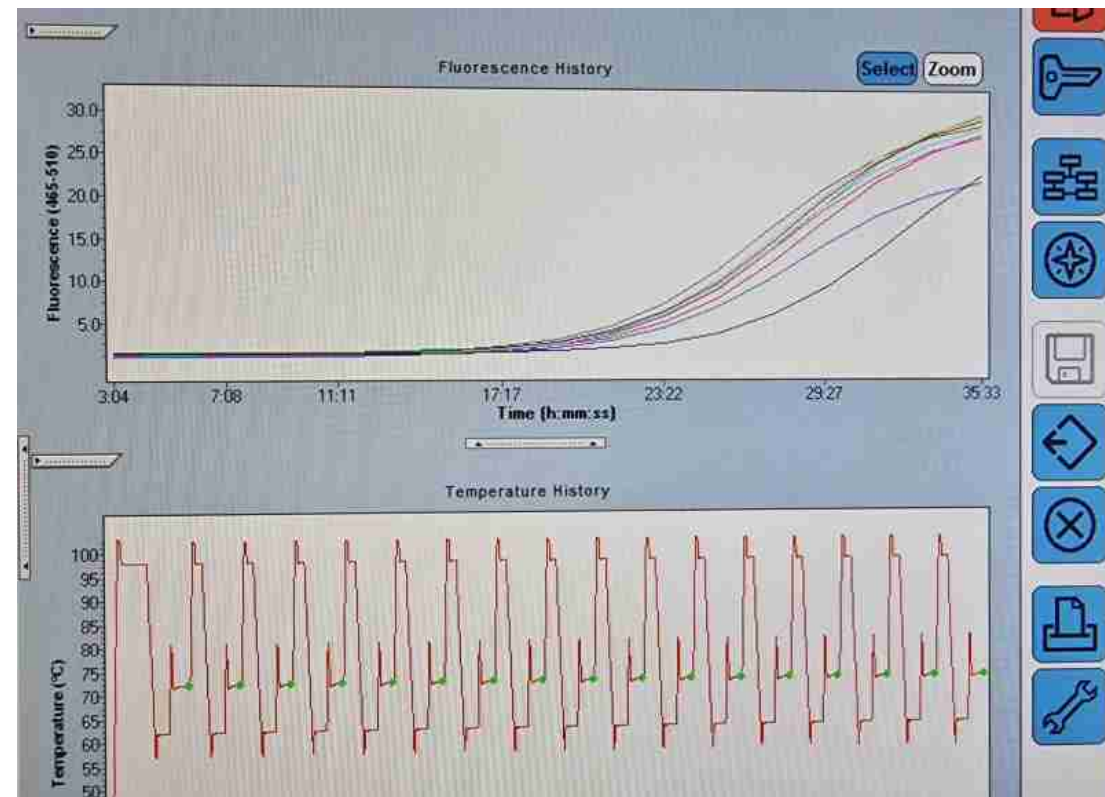


A	B	C	D	E	F
Step	temperature	acquisition	Duration	spleen	Cycles
Pre-incubation	98°C	None	45sec	4.4°C/sec	1 cycle
Amplification	98°C	None	15sec	4.4°C/sec	
	62°C	None	30sec	2.2°C/sec	
	72°C	Single	30sec	4.4°C/sec	
					20 cycles

RTLib1 program on lightcycler

PCR should be stopped when signal is between standards 1 and 3 (optimal library amplification range), generally between cycles 10-14. more for low quality or low quantity DNA

Important : Do not over amplify, it will reduce your sequencing quality, stop before reaching plateau phase of amplification



Asset URL: stop before plateau phase

- 7.5 Stop the run just after the last acquisition (green dot on the graph) when the double-stranded DNA is full length. If necessary, transfer the "passed" samples to another AB 1400 plate by piercing the film with a scalpel blade at each well concerned.
- 7.6 Repeat if necessary for samples requiring more subsequent cycles, using the same program but without pre-incubation.

- 7.7** Purify using  Agencourt AMPure XP **Beckman Coulter Catalog #A63880** XP 1X (); see section 2.
- Evaluate the samples in  of ultra pure water.

- 7.8** Perform an analysis on Qiaxcel () and dose on Nanoquant ().

7.9



Expected result

after 10-16 cycles of amplification + purification
 Mass concentration from 2 to 110ng/ μ L adn target 30ng/ μ L
 DNA size from 250 to 550pb target 400pb peak

optional you can conserve for 24h sample after purification step at -20°C and continue the next day

Preparation of Bulk

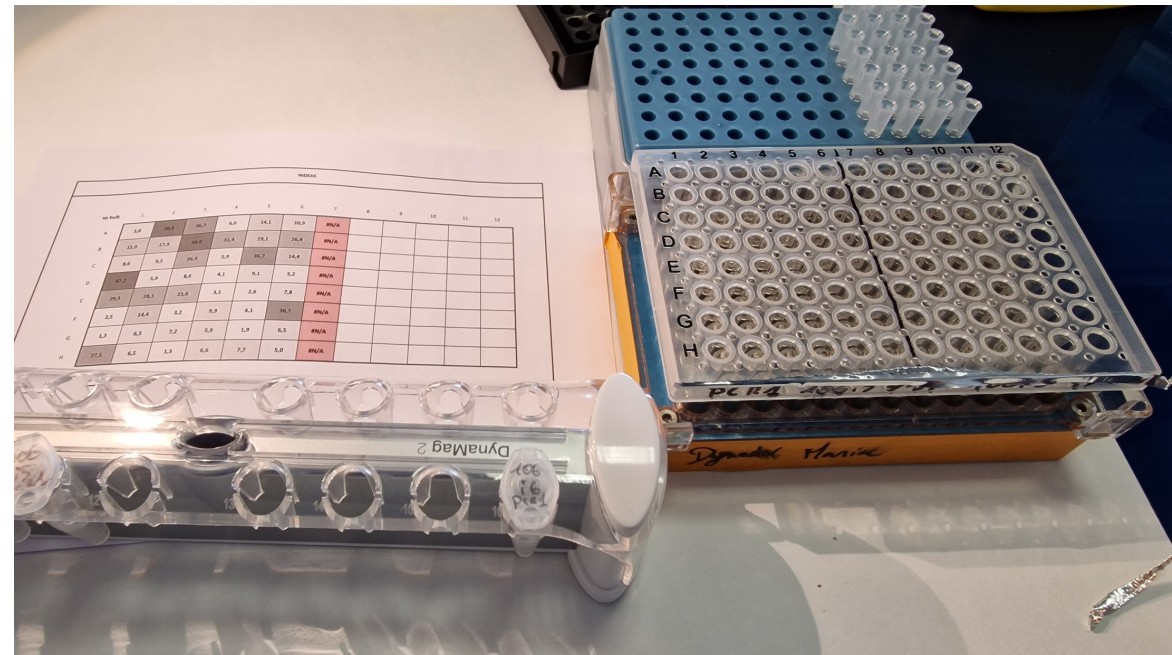
1h 30m

- 8** Prepare an equimolar mix of all 48 sample by adding  to  of each purified sample from PCR1 step. Volume will depend of concentration and size of DNA.

- 8.1** Match 0.5 pmol per samples ex :  for 400pb DNA.

1h

Each volume is added in a  DNA low bind tube, total volume between  to 



Transfer of 48 samples from PCR plate to one 1.5 mL tube with different volumes

8.2

Purify using  **Agencourt AMPure XP Beckman Coulter Catalog #A63880** XP 1.8X; see 30m
section 2. Volumes depends on the total volume of the bulk.
Eluate the samples in  $30 \mu\text{L}$ of ultra pure water.



Bulk tubes before elution step.

8.3

Expected result

30 μ L of 50-200ng/ μ L DNA bulk, size 400pb peak
1-7 μ L is needed for Hybridization step

you can conserve DNA bulks for weeks at -20°C

you can repeat library preparations protocols for multiple bulk and when you have enough N bulks of 48/specimens so you can proceed multi hybridization step in same time and make N*48 specimens per sequencing.

Part 2: Hybridization step for targeted DNA sequencing

1d

9 This part of the protocol describes dual hybridization process for targeted DNA sequencing using Daicel Arbor Biosciences Mybait KIT V5 Angiosperm353 and custom Annonaceae kit.

In this protocol we use a bulk of 48 samples for each reaction (Arbor recommend max 8 sample per reactions). This protocol describes the quantity for one reaction or 8 reactions (48 or 384 samples)

A total of $\text{pipette icon } 5.5 \mu\text{L}$ of baits is required per reaction. We used a ratio of a **3/4** Angiosperm353 baits and **1/4** Annonaceae baits per reaction following Henriks et al. (2021):

$\text{pipette icon } 4.125 \mu\text{L}$ of **Angiosperm353 baits**

$\text{pipette icon } 1.375 \mu\text{L}$ of **Annonaceae baits**

Bait mix preparation

15m

10 Program the Thermo Mixer so that it reaches $\text{thermo icon } 65^\circ\text{C}$ and a second Thermo mixer to $\text{thermo icon } 95^\circ\text{C}$. Place the Arbor Biosciences Baits on ice.

Heat the Hyb N and Hyb S bait reactivities at $\text{thermo icon } 60^\circ\text{C}$ and vortex to dissolve the aggregates.

Prepare a volume of Mix "**HYB**" according to the number of reactions planned, vortex and centrifuge:

A	B	C
Component	$\mu\text{L for 1 reaction}$	$\mu\text{L for 8 reactions}$
Hyb N	9.25	74
Hyb D	3.5	28
Hyb S	0.5	4
Hyb R	1.25	10

A	B	C
Baits	4.125 Angio353 + 1.375 Annonaceae	33 Angio353 + 11 Annonaceae
TOTAL	20 µL	160 µL

Mix "HYB" preparation

- 11 Incubate the **HYB** mix 00:10:00 at 60 °C and vortex to recover the condensate
For each capture reaction, aliquot 18.5 µL in 1.5 µL of DNA in a low bind tube.

15m

We refer to these reaction aliquots of hybridization Mix as "HYBs"

Let stand 00:05:00 at Room temperature

Prepare a volume of Mix "**Block**" according to the number of reactions of capture, vortex and centrifuge:

A	B	C
BLOCK	per capture	for 8 captures
Block O	5 µl	40 µl
Block X	0.5 µl	4 µl
Total volume	5.5 µl	44 µl

Mix "**Block**" preparation

For each capture reaction, aliquot 5.0 µL of "Block" in a 0.5mL tube.

Hybridization

1d

- 12 Add 7 µL of BULK DNA library 50-100 ng/µL to each 5 µL aliquot of "Block" and homogenize by 10m pipetting. We then obtain a Mix of 12 µL



These libraries with Blocker Mix aliquots are now called "LIB" (block + libraries)

Place the "LIB" on the second Thermo Mixer at 95 °C for 00:05:00.

Transfer the "LIB" and the "HYBs" mix into the Thermo Mixer at 65 °C 00:05:00 800 rpm

- 12.1 Transfer 18.5 µL of each "HYBs" to each corresponding "LIB" and homogenize by pipetting 5m times.

Incubate at hybridization temperature of  65 °C  Overnight (16h-24h).  800 rpm





eppendorf Thermomixer with Thermotop , 65°C 800RPM

Capture and washing

1h

- 13** Bring Hyb S (the reactive, not the mix), Buffer E and wash buffer to **Room temperature** in order to prepare the washing solutions about **1h30** before stopping the hybridization reaction.

Prepare the Wash Buffer X in a falcon tube of **15 mL**

A	B
HYB S	40 µL
Water	3.96 mL
Wash Buffer	1 mL
Final volume	5 mL

Wash Buffer X preparation

Wash Buffer X can be stored at **4 °C** for up to 1 month

- 13.1** Vortex and place the Wash Buffer X at hybridization temperature (+ **65 °C** in a water bath) **45m** least **00:45:00** before use.

Vortex to homogenise capture beads (kit , box at 4°C).

For each capture reaction, aliquot **30 µL** of beads (Box at **4 °C**)

*TIP: Beads can be prepared in 8 (or fewer) reaction batches (**240 µL**) in a **1.5 µL** tube.*

*Multiply all volumes by the number of reactions in the batch; i.e., for 8 reactions-worth, wash with **1.6 mL** and resuspend in **560 µL** µL Binding Buffer, then aliquot 70 µL suspension to individual tubes.*

- 13.2** Arrange tubes on a magnetic stand for **00:01:00** to **00:02:00** until the aggregate is **3m 3s** formed and remove the supernatant.
Repeat 3 consecutive times:

Add $\text{200 } \mu\text{L}$ of Binding Buffer in each tube, vortex  00:00:03, centrifuge briefly and finally eliminate the supernatant on magnetic stand.
Resuspend beads in $\text{70 } \mu\text{L}$ of Binding Buffer.

- 13.3** Place the beads at hybridization temperature on Thermo Mixer at least  00:02:00 before transfer. 7m

Quickly transfer each capture reaction ($\text{30 } \mu\text{L}$) into a tube containing the beads and homogenize by pipetting.

Incubate the library and beads mixture at hybridization temperature of  65 °C on Thermo Mixer for  00:05:00 at 1200 rpm (program "CAPT").

Eliminate the supernatant on magnetic stand.

- 13.4** Repeat 3 consecutive times: 5m

Add $\text{375 } \mu\text{L}$ of Wash Buffer X (heated) in each tube, vortex, centrifuge briefly, incubate at hybridization temperature on Thermo Mixer for  00:05:00 minutes at 1200 rpm and finally eliminate the supernatant on magnetic stand.

Elution

15m

- 14** Add $\text{25 } \mu\text{L}$ of Buffer E box -20° (bring back to  Room temperature before hand) 5m
Incubate at  98 °C for  00:05:00 on ThermoMixer or Heating Block (use lid).
Place the tubes on a magnetic stand and quickly recover the supernatant containing the enriched libraries and transfer it into a new PCR tube / plate.

RT-PCR 2 (final)

30m

- 15** In a LightCycler plate® 480 Multiwell plate with 96 arranged on refrigerated plate.



Deposit $\text{50 } \mu\text{L}$ of each fluorescent marker 1 to 4 repeated twice in 8 separate wells on the plate. Keep the plate away from the light.

Prepare a Mix for each INDEX used. There are 12 Illumina INDEX that can be used.

A	B
By well:	
Bulk DNA	20μL
adapter PE 34 solution 10 μM	2.5μL
INDEX 10 μM	2.5μL

A	B
KAPA HiFi Master Mix 2X	25µL
Final volume	50µL

Mix for each INDEX per well

index and adapter sequence on attached file

Index used: AD001; AD002; AD003; AD004 AD005 AD006 AD007 AD008 AD009 AD0010 AD0011 AD0012

Deposit **30 µL** of Mix (index+PE34+kapa pcr mix) per well and pulse centrifuge.

Deposit **20 µL** of DNA per well, mix with the pipette and pulse centrifuge.

Place the plate at the lightcycler and launch the following program:

A	B	C	D	E	F
Step	Temperature	acquisition	Duration	spleen	Cycles
Pre-incubation	98°C	None	45sec	4.4°C/sec	1 cycle
Amplification	98°C	None	15sec	4.4°C/sec	
	62°C	None	30sec	2.2°C/sec	
	72°C	Single	30sec	4.4°C/sec	
					55 cycles

Lightcycler program

Stop the run before the plateau phase. Generally 10 to 16 cycles. If necessary, add some extra cycles for some samples. Important : Do not over amplify, it will reduce your sequencing quality.

Carry out a purification step using  Agencourt AMPure XP **Beckman Coulter Catalog #A63880** XP 1X on half the volume **25µL**, keep the leftover volume (approx. **25 µL**) at **-20 °C** as backup.

15.1 Proceed with the purification with **1.5 mL** DNA low bind tube as follows:

5m

1. Prepare the sample: Mix the 25µL DNA sample with 25µL AMPure.
2. Bind: Incubate the mixture for 10 minutes at room temperature to allow the DNA to bind to the beads and put on magnetic stand;
3. Wash: after 5 minutes on magnetic stand remove supernatant;
4. Wash the beads twice with 70% ethanol to remove impurities;
5. Let dry for 5 minutes to remove residual ethanol and dry the purified DNA;
6. Remove from magnetic stand and resuspend the dried DNA by adding **20 µL** water.

Let DNA elute at least  00:05:00 min.

Perform a quantification with Nanodrop and quality analysis on Qiaxcel.

Preparation of bulk for illumina sequencing

30m

- 16** Prepare an equimolar mix of all samples.

30m

We end up with a single tube with all indexes mixed, and ready to sequence.

Refer to your sequencing platform for final volume and concentration.

In our case we needed a minimum of 50 µL and DNA fragments should be at 400pb.

All samples were sequenced with Novogen using a NovaSeq PE150.