



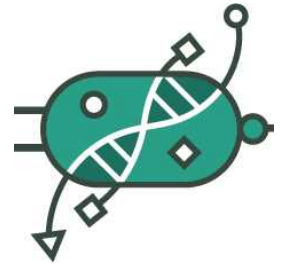
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Version 2

High-throughput workflow for the genotypic characterization of transposon insertion library variants V.2

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Protocol status: Working

We use this protocol and it's working

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Abstract

This is a workflow for the genotypic characterization of transposon library variants. It has been developed using an open-source Opentrons OT-2 robot, BLASTN for genomic annotations and modular sub-protocols (e.g., PCR sample preparation, OT-2 volume transfer, OT-2 counter selection, etc) that can be used for other tasks, thus providing a general-purpose pipeline.

All steps follow a 96-well plate format for high-throughput analysis. The protocol is described for the characterization of transposon library variants generated with SEVA-Sib pBAMD1-x and pBLAM1-x plasmid sets that follow Standard European Vector Architecture (SEVA, <https://seva-plasmids.com>) and can be amplified with the standard PS1-PS6 primers. However, it can be adapted for other transposon systems.

Changes from previous version:

- This version references updated OT-2 automation protocols which now follow the standardized LAP format (<https://www.laprepo.com>).
- Genotyping is performed with a new ad hoc python package to annotate genomic reads and identify the transposon insertion sites.

Guidelines

This workflow comprises the following sections: 1) Colony picking in selective media 2) Counter-selection and glycerol stocks pre-cultures 3) Colony selection in OT-2 liquid handler robot 4) Master 96-well plate for PCR steps 5) Control PCRs (spurious plasmid integration control and cargo insertion control) 5) Arbitrary PCRs 6) Sequencing and annotation. There is an additional section with an example on how to run the script. We recommend the use of an OT-2 protocol specially if more than 2 libraries are to be analyzed. However, we recommend to do counter-selection in the OT-2 liquid handling robot even for one plate to avoid human errors. Note that other pipettes can be used to run the workflow in the OT-2 but these were deemed the most appropriate for the overall workflow to minimize pipette changes.

Materials

Equipment:

Equipment	
Incubating mini-shaker	NAME
Incubating shaker	TYPE
Fisherbrand	BRAND
15554070	SKU
https://www.fishersci.es/shop/products/incubating-mini-shakers-3/15554070 ^{LINK}	

Equipment	
OT-2	NAME
Liquid handler	TYPE
Opentrons	BRAND
OT-2	SKU

Equipment

SPECTROstar Nano	NAME
plate reader	TYPE
BMG	BRAND
SPECTROstar Nano	SKU
https://www.bmglabtech.com/spectrostar-nano/	LINK



Equipment

Mastercycler® nexus - PCR Thermal Cycler	NAME
Thermocycler	TYPE
Eppendorf	BRAND
12304943	SKU
https://www.fishersci.es/shop/products/mastercycler-nexus-gradient-3/12304943?matchedCatNo=12304943&searchHijack=true&searchTerm=mastercycler-nexus-gradient-3&searchType=Rapid	LINK

Equipment

Centrifuge Tube Mini-Cooler

NAME

Cold Block

TYPE

BRAND

BRAND

10141921

SKU

<https://www.fishersci.es/shop/products/brandtech-scientific-brand-centrifuge-tube-mini-cooler-3/10141921>

LINK

Electrophoresis machine

Wet-lab requirements:

Material

- Non-treated flat bottom sterile 96-well plates

96-well plates flat bottom non-treated **VWR International (Avantor) Catalog #734-2781**

- Sterile breathable membrane for 96-well plate

Greiner Bio-One Sellador BREATHseal™ **Fisher Scientific Catalog #11920667**

- Storage membrane for 96-well plate

Thermo Scientific™ Láminas de papel de aluminio adhesivas para placas de PCR **Fisher Scientific Catalog #10130853**

- PCR plates

Thermo Scientific™ PCR Plate, 96-well, low profile, skirted, red **Fisher Scientific Catalog #10161073**

Enzymes

- DNA polymerase with green buffer

Phire Green Hot Start II PCR Master Mix **Fisher Scientific Catalog #15391732**

- DNA polymerase

Thermo Scientific™ Phire Hot Start II PCR Master Mix **Fisher Scientific Catalog #15361732**

Oligonucleotides:

- Spurious integration control: PS3, PS4, PS5, PS6 (<https://doi.org/10.3389/fbioe.2014.00046>)



- Arbitrary PCR: ARB2, ME-O-Km-Ext-F, ME-O-Km-Int-F, ME-O-Sm-Ext-F, ME-O-Sm-Int-F, ME-O-Gm-Ext-F, ME-O-Gm-Ext-R (<https://doi.org/10.3389/fbioe.2014.00046>)
- Optional insert control: PSMCS

Dry-lab requirements:

- Python 3
- Command-line BLASTN



Colony picking in selective media

1d

- 1 Dispense  100 μL of selective media (M9-citrate for *P. putida* or Luria-Bertani plus 20 ng/ μL nalidixic acid for *DH5 α* *E. coli*) plus **transposon cassette antibiotic** in a 96-well plate
- 2 Pick individual colonies into a 96-well plate with selective media
Tip: Keep tips inside of wells to keep track
- 3 Cover with a sterile breathable membrane
- 4 Grow  Overnight at  30 $^{\circ}\text{C}$ (*P. putida*) or  37 $^{\circ}\text{C}$ (*E. coli*) /  500 rpm



Counter-selection and glycerol stocks pre-cultures

1d

- 5 Measure OD_{600nm} of overnight culture grown in selective media from  plus **transposon cassette antibiotic** in a plate reader 
- 6 Inoculation of **counter-selection plate** in selective media: 10m
 - Dispense  100 μL of selective media (M9-citrate for *P. putida* or Luria-Bertani plus 20 ng/ μL nalidixic acid for *DH5 α* *E. coli*) plus **ampicillin** (backbone antibiotic) to select against spurious integration events.
 - Transfer  5 μL of overnight culture from  to counter-selection (ampicillin) plate
 - Cover with a sterile breathable membrane

Note

For steps 6 and 7, if two or more 96-well plates are used as input it is advised to use the **OT-2** protocol below to minimize human error. Dispensed volume and culture volume inoculated should be that described in these steps. Note that steps 6 and 7 could be completed together in a single run depending on the number of initial plates.



Protocol



NAME

OT-2 Media dispensing and culture inoculation protocol

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[Preview](#)

- 7 Inoculation of precultures for **glycerol stock** in rich media: 10m
- Dispense 100 μL of Luria-Bertani media plus **transposon cassette antibiotic** in a 96-well plate
 - Transfer 5 μL of overnight culture from to counter-selection (ampicillin) plate
 - Cover with a sterile breathable membrane
- 8 Grow counter-selection and glycerol stock pre-culture plates Overnight at 16h
- 30 °C (*P. putida*) or 37 °C (*E. coli*) / 500 rpm
- 9 Measure OD_{600nm} of overnight culture grown in selective media plus **ampicillin** in a plate reader

Colony selection in OT-2 liquid handler robot

- 10
- Selection of colonies to store as glycerol stocks and do further PCR reactions by running the following OT-2 protocol:



Protocol



NAME

OT-2 Counter-Selection

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Note

The OT-2 protocol will prepare three plates (2 glycerol stock plates and a "PCR plate") and perform the following:

- Dispense  75 μL of PCR-grade water to "PCR plate"
- Dispense  25 μL of 30% glycerol to two glycerol stock plates
- Transfer  25 μL of grown pre culture in Luria-Bertani media plus **transposon cassette antibiotic** from  to "PCR plate" and glycerol stock plates

- 11 Cover glycerol stock plates with a storage membrane and store at -80°C
- 12 If not proceeding to the next step right away: Store "PCR plate" at 4°C for a few days or cover with an storage membrane and store at -20°C for longer term 

Master 96-well plate for PCR steps

- 13 Transfer  50 μL of selected colonies from one or more libraries to a 96-well plate with the following OT-2 protocol: 



Protocol



NAME

OT-2 Protocol to transfer volume from several plates to a single plate

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Control PCRs

14



Safety information

Positive control (donor plasmid) and wild-type control (*P. putida* or *E. coli*) should be added to every reaction in this section.

Spurious integration control with SEVA primers pairs PS3/PS4 and PS5/PS6

Protocol



NAME

OT-2 PCR sample preparation protocol

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Preview



Note

If <48 cfu are to be analyzed both PS3/PS4 and PS5/PS6 spurious integration controls can be done in a single 96-well plate

- 15 Optional: Cargo integration control with primers PSMCS and either ME-O-Km-R/ME-O-Sm-R or ME-O-Gm-R (depending on **transposon cassette antibiotic**)



Arbitrary PCRs

- 16 **Arbitrary PCR#1** using primer pairs ARB6 and ME-O-Km-Ext-F/ME-O-Sm-Ext-F or ME-O-Gm-Ext-F depending on transposon antibiotic cassette



Protocol



NAME

OT-2 PCR sample preparation protocol

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Note

The OT-2 protocol will perform the following steps:

- Prepare a PCR master mix
- Dispense 19 μ L of PCR mastermix
- Transfer 2 μ L of pre-culture from

- 17 Seal 96-well plate, place it in thermocycler and run the following PCR program:



	A	B	C
	98°C	5 min	
	98°C	10 s	x6 cycles
	30°C	30 s	
	72°C	1 min 30 s	
	98°C	10 s	x30 cycles
	45°C	30 s	
	72°C	1 min 30 s	
	72 °C	5 min	
	4°C	hold	

- 18 Select 8-12 Arbitrary PCR#1 reactions from the 96-well plate and run them on a 1% agarose gel to verify amplification.



Note

Several bands will appear and even DNA smears even when the reaction has worked perfectly.

- 19 **Arbitrary PCR#2** using primers pairs ARB2 and ME-O-Km-Int-F/ME-O-Sm-Int-F or ME-O-Gm-Int-F, depending on transposon antibiotic cassette, run the following OT-2 protocol:

Protocol



NAME

OT-2 PCR sample preparation protocol

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**Note**

The OT-2 protocol will perform the following steps:

- Prepare a PCR master mix
- Transfer 1 μL of PCR product from Arbitrary PCR#1
- Dispense 19 μL of PCR mastermix

- 20 Seal 96-well plate, place it in thermocycler and run the following PCR program:

	A	B	C
	98°C	30 s	
	98°C	10 s	x 30 cycles
	52°C	30 s	
	72°C	1 min 30 s	
	72°C	5 min	
	4°C	hold	

- 21 Select 8-12 Arbitrary PCR#2 reactions from the 96-well plate and run them on a 1% agarose gel to verify amplification

Note

Several bands will appear and even DNA smears even when the reaction has worked perfectly.

Sequencing and annotation

1m

- 22 Prepare a PCR plate to send to sequencing by mixing  10 μL of unpurified **Arbitrary PCR#2** reaction and  10 μL of 10 μM sequencing primer (ME-O-Km-Ext-F/ME-O-Sm-Ext-F or ME-O-Gm-Ext-F depending on the transposon antibiotic cassette)



Note

These guidelines may vary depending on the sequencing service arranged for your laboratory.

23 Annotate sequencing results by running the following protocol:

1m



Protocol



NAME

Bacterial genome annotation script using BLASTN

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Note

The python package uses as input:

1. DNA sequencing results in .txt or .seq
2. Reference genome file in GENBANK format
3. Transposon sequence file in GENBANK format

Optional:

1. .ab1 files to verify sequencing quality
2. map layout from previous steps