

FEB 07, 2024

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Protocol Citation: Quadrana Leandro, pol.vendrell, basile.leduque 2024. Protocol TE Display sequencing (TED-Seq). protocols.io

https://protocols.io/view/protocolte-display-sequencing-ted-seqc7seznbe

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

Created: Jan 18, 2024

Last Modified: Feb 07, 2024

PROTOCOL integer ID: 93734

Protocol TE Display sequencing (TED-Seq)

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ABSTRACT

<u>Background:</u> Mobilization of transposable elements (TEs) can generate large effect mutations. However, because transposition is typically rare, the actual rate and landscape of new insertions remains unexplored for most TEs. Furthermore, the repetitive nature of TEs renders these sequences less amenable to be investigated by shotgun sequencing.

Results: Here, we introduce a TE Display sequencing (TED-seq) method that leverages target amplification of TE extremities to detect TE insertions with high sensitivity and specificity. By implementing TED-seq on serial dilutions of genomic DNA from Arabidopsis thaliana lines containing different repertoires of insertions, we show that our method can detect insertions that are present at a frequency as low as 1/250000 within a DNA sample. Furthermore, TED-seq can be multiplexed to simultaneously detect insertions for several TE families, including both retrotransposons and DNA transposons, increasing its versatility and cost-effectiveness to investigate complex "mobilomes". Analysis of TE insertions in large populations of A. thaliana plants undergoing a transposition burst demonstrate the power of TED-seq to investigate the rates and allele frequencies of heritable insertions. In addition, we show that TED-seq libraries can be sequenced using nanopore sequencing technologies, leading to a turn around time from DNA extraction to insertion identification of less than 24h.

<u>Conclusions:</u> TED-seq is an ultra-sensitive, specific, and cost-effective method to investigate the rate and landscape of new TE insertions, enabling systematic studies of TE mobilization.



Funders Acknowledgement:

European Research Council Grant ID: 948674

MATERIALS

- NEBNext® UltraTM II DNA Library Prep Kit for Illumina® (E7645, E7103)
- NEBNext Ultra II Q5 Master Mix (M0544S)
- AMPure XP beads (Product No: A63880)

Adapter Primer

The following primers are designed for ATCOPI93 detection, the underlined primer sequence should be specific for your TE of interest

FORKED ADAPTER

P7_adapter_up

5' GTGACTGGAGTTCAGACGTGTGCTCTTCCGATC*T 3'

P7_adapter_bottom

5' pGATCGGAAGAGCATC** 3'

Modifications: p 5'phosphate

* phosphorothioate bond

** dideoxy-C

1st PCR primer

TEDseq_outer_for: (complementary to the P7_adapter_up)

5' GTGACTGGAGTTCAGACGTG 3' (FIXED)

Reverse Primer (complementary to the target TE and downstream of the P5_linker_TE primer, in this particular case it corresponds to ATCOPIA93)

TE_outer_rev:

5' <u>GTGAGTCCTCTTCAACGGCT</u> 3' (should be changed accordingly to the specific targeted TE)

nested PCR primer

P7_primer_index4

5'CAAGCAGAAGACGGCATACGAGAT**GCCAAT**GTGACTGGAGTTCAGACGTG 3' **P7_primer_index6**

5'CAAGCAGAAGACGGCATACGAGAT**CTTGTA**GTGACTGGAGTTCAGACGTG 3' **P7_primer_index1**

5'CAAGCAGAAGACGGCATACGAGAT**CGATGT**GTGACTGGAGTTCAGACGTG 3' **P7_primer_index3**

5'CAAGCAGAAGACGGCATACGAGAT**ACAGTG**GTGACTGGAGTTCAGACGTG 3' So on so for....

Bold sequences match barcode

NEXTFLEX® PCR-Free Barcodes:

Barcode Adapter 1 CGATGT

Barcode Adapter 2 TGACCA

Barcode Adapter 3 ACAGTG

Barcode Adapter 4 GCCAAT

Barcode Adapter 5 CAGATC

So on so for...

TE Specific primer. The underlined sequence should correspond to the **(-) strand of the specific TE sequence of interest if we amplify the 5' extremity.**

Bold nucleotides are introduced to increase sequence diversity during illumina sequencing of pooled libraries generated by TED-seq.

When sequencing together multiple TED-Seq libraries targeting different TEs, only the P5_TE_primer without linker should be used.

Example ATCOPIA93, underlined sequence (For each TE-specific family, change the TE sequence):

P5_linker_TE primer:

5'AATGATACGGCGACCACCGAGATCTACACTCTTTCCCTACACGACGCTCTTCCGATCTGCCC ACTCTCTTGTAGTACATATC

3

 $5'AATGATACGGCGACCACCGAGATCTACACTCTTTCCCTACACGACGCTCTTCCGATCT\\ \textbf{T}\underline{GCC}\\$ CACTCTCTTGTAGTACATATC

3'

5'AATGATACGGCGACCACCGAGATCTACACTCTTTCCCTACACGACGCTCTTCCGATCT**AT**<u>GC</u> <u>CCACTCTCTTGTAGTACATATC</u>

3'

5'AATGATACGGCGACCACCGAGATCTACACTCTTTCCCTACACGACGCTCTTCCGATCT**CATA**<u>GCCCACTCTCTTGTAGTACATATC</u>

3'

5'AATGATACGGCGACCACCGAGATCTACACTCTTTCCCTACACGACGCTCTTCCGATCT**GAAT T**GCCCACTCTCTTGTAGTACATATC 3'

Amplicon sequences

Amplicon after the first PCR: (TE sequence)

GTGACTGGAGTTCAGACGTG(neighboring_genomic_sequence+TSD)<u>TGCTCTTCCGATCTA</u>

<u>TTGATCAAGACTCAAATAAGAAAGGCCTAGTATTGGATATGTACTACAAGAGAGTGGGCCGAA</u>

<u>CATATGAGAAGTCTATGAAGAGGCTTCTAGAAGAGGTGAAGGCACACAAATATCTCTTGTAGCC</u>

<u>GTTGAAGAGGACTCAC</u>

Amplicon after the nested PCR (using Index 4):

ie P7adapter+(neighboring_genomic_sequence+TSD)+COPIA93 + P5adapter

Before Starting

1 Before you start, be sure to have:

P7_adapter_up

P7_adapter_bottom

P7 primer (P7_Primer_index#)

Primer for the 1st PCR (outer Primers)

Primer for the nested PCR (P7_Primer_index#, TE Specific P5adapter)

NEBNext® UltraTM II DNA Library Prep Kit for Illumina® (E7645, E7103)

PREPARING CUSTOM ADAPTER

2 Annealing adapters (final concentration 30uM):

In a 0.2ml tube, mix:

90µl of 100µM P7_adapter_up (in 10mM Tris-HCl pH8)

90µl of 100µM P7_adapter_bottom (in 10 mM Tris-HCl pH8)

30µl T4 DNA Ligase Reaction Buffer (NEB)

90µl H₂O

Heat to 95 °C and maintain the temperature for 2 min.

Cool to 25 °C over 45 min. (0.026C per seg)

Cool to 4 °C for temporary storage.

Centrifuge the PCR tube briefly to draw all moisture away from the lid.

Note: We recommend preparing aliquots of 50ul (40 reactions) of the custom adapter to avoid repeated

freeze-thawing

Long term storage at -20°C

[C] P7_Primer_index = 10µM in 10 mM Tris-HCl pH8 (stock in 100uM)

[C] P5_TE_primer = 10µM in 10mM Tris-HCl pH8 (stock in 100uM)

Sonication

3 Sonicate 100µl of 20ng/µl DNA (ie 1µg)

Diagenode Program 30"/30", 13 cycles -> 13' (Could be fragmented by any sonication system or enzymatic fragmentation. For short-reads you should aim fragments between 400-700bp).

Load in Agarose gel 1% 100 ng

Only **500ng** will be use in the following protocol

End repair and A tailing

4 Add the following components to a sterile nuclease-free tube:

(green) NEBNext Ultra II End Prep Enzyme Mix 1.5 μ l (green) NEBNext Ultra II End Prep Reaction Buffer 3.5 μ l Fragmented DNA (500ng) 25 μ l Total Volume 30 μ l

Note: For End Prep and Ligation step, NEBNext® UltraTM II DNA Library Prep Kit's volume have been adjusted to reduce the price of each library. 48 samples can be processed with one kit (E7546S) instead of 24.

- Set a 100 µl or 200 µl pipette to 25 µl and then gently pipette the entire volume up and down at least 10 times to mix thoroughly. Perform a quick spin to collect all liquid from the sides of the tube.

 Note: It is important to mix well. The presence of a small amount of bubbles will not interfere with performance.
- 6 Place in a thermocycler, with the heated lid set to $\geq 75^{\circ}$ C, and run the following program:

30 minutes @20°C

30 minutes @ 65°C

Hold at 4°C

If necessary, samples can be stored at -20°C; however, a slight loss in yield (~20%) may be observed. We recommend continuing with adapter ligation before stopping.

7 Incubate at 20°C for 15 minutes in a thermocycler with the heated lid off.

Adapter Ligation

8 Add the following components to a sterile nuclease-free tube:

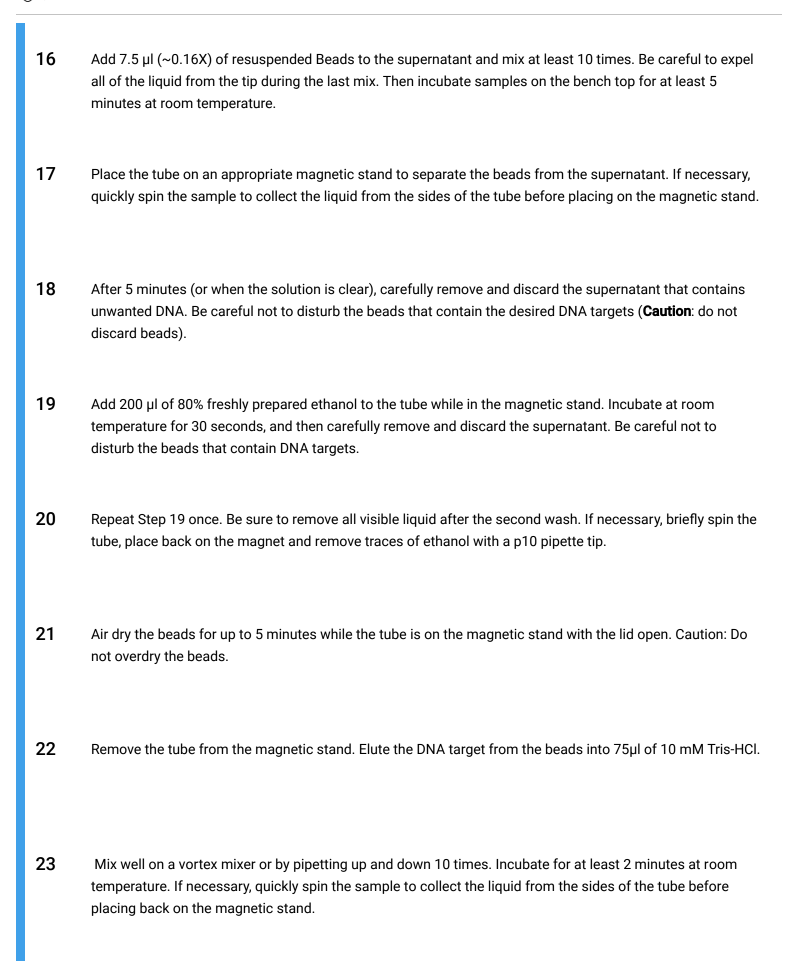
End Prep Reaction Mixture (Step 6 in End repair and A tailing Section 30 μ l (red) NEBNext Ultra II Ligation Master Mix* 15 μ l (red) NEBNext Ligation Enhancer 0.5 μ l Custom Adapter (30 μ M) 1.25 μ l

Total volume 46.75 µl

- * Mix the Ultra II Ligation Master Mix by pipetting up and down several times prior to adding to the reaction. Note: The Ligation Master Mix and Ligation Enhancer can be mixed ahead of time and is stable for at least 8 hours @ 4°C. We do not recommend adding adapter to a premix in the Adapter Ligation Step.
- 9 Set a 100 μl or 200 μl pipette to 40 μl and then pipette the entire volume up and down at least 10 times to mix thoroughly. Perform a quick spin to collect all liquid from the sides of the tube.
- 10 Incubate at 20°C for 15 minutes in a thermocycler with the heated lid off.

Size Selection of Adapter-ligated DNA

- 11 Vortex AMPure XP Beads to resuspend.
- Add 7.5 μ I (\sim 0.16X) of resuspended beads to the 46.75 μ I ligation reaction. Mix well by pipetting up and down at least 10 times. Be careful to expel all of the liquid out of the tip during the last mix. Vortexing for 3-5 seconds on high can also be used. If centrifuging samples after mixing, be sure to stop the centrifugation before the beads start to settle out.
- 13 Incubate samples on bench top for at least 5 minutes at room temperature.
- Place the tube on an appropriate magnetic stand to separate the beads from the supernatant. If necessary, quickly spin the sample to collect the liquid from the sides of the tube before placing on the magnetic stand.
- After 5 minutes (or when the solution is clear), carefully transfer the supernatant containing your DNA to a new tube (**Caution**: do not discard the supernatant). Discard the beads that contain the unwanted large fragments.



Place the tube on the magnetic stand. After 5 minutes (or when the solution is clear), transfer 75 μl to a new PCR tube.

Samples can be stored at -20°C.

1st PCR: Enrichment of Adapter-ligated DNA

25 Add the following components to a sterile strip tube:

Adapter Ligated DNA Fragments (Step 24)	7.5 µl
(blue) NEBNext Ultra II Q5 Master Mix	12.5 µl
(blue) TEDseq_outer_for (10μM)	2.5 µl
(blue) TE_outer_rev (10μM)	2.5 µl
Total volume	25 µl

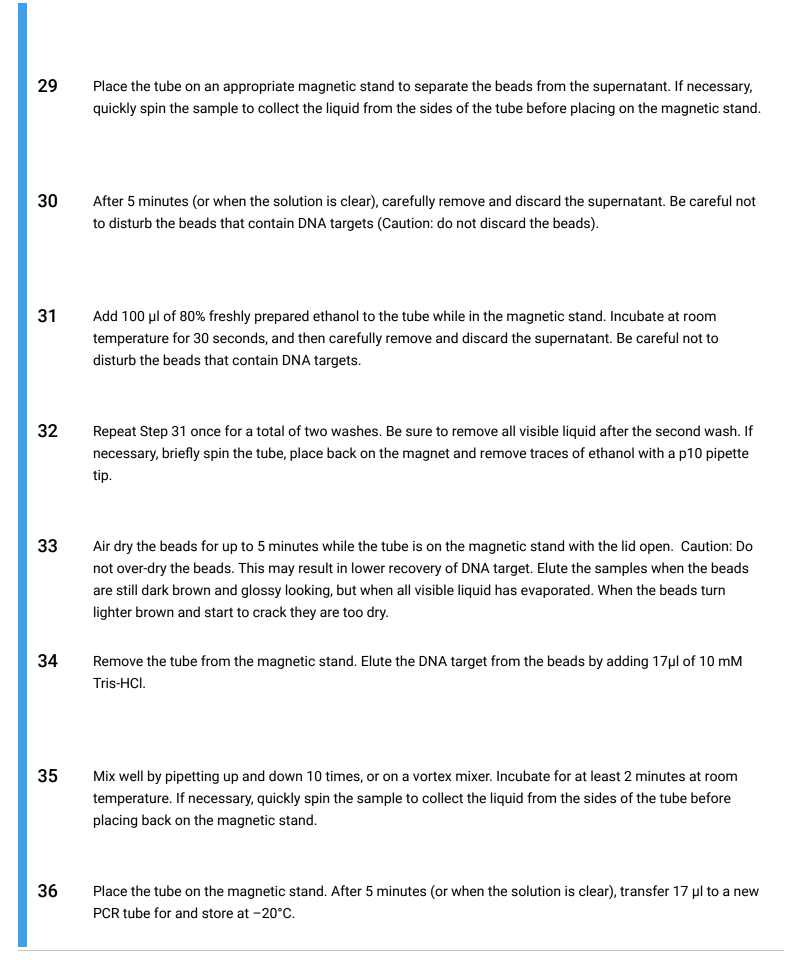
Set a 100 μ l or 200 μ l pipette to 20 μ l and then pipette the entire volume up and down at least 10 times to mix thoroughly. Perform a guick spin to collect all liquid from the sides of the tube.

Place the tube on a thermocycler and perform PCR amplification using the following PCR cycling conditions:

- 1. 98°C 30"
- 2. 98°C 10"
- 3. 61°C 75"
- 4. GO TO .2 20x
- 5. 61C 5'
- 6. Hold at 4C

Cleanup of PCR Reaction

- Vortex AMPure XP to resuspend.
- 27 Add 22.5 µl (0.9X) resuspended beads to the PCR reaction. Mix well by pipetting up and down at least 10 times. Be careful to expel all of the liquid out of the tip during the last mix. Vortexing for 3-5 seconds on high can also be used. If centrifuging samples after mixing, be sure to stop the centrifugation before the beads start to settle out.
- Incubate samples on bench top for at least 5 minutes at room temperature.



Samples can be stored at -20°C.

Nested PCR: Adding P5 and P7 sequences

37 Add the following components to a sterile strip tube:

Adapter Ligated DNA Fragments (Step 36)	7.5 µl
(blue) NEBNext Ultra II Q5 Master Mix	12.5 µl
(blue) P7_Primer_index# (10μM)	2.5 µl
(blue) P5_linker_TE_primer (10µM)	2.5 µl
Total volume	25 µl

Set a 100 μ l or 200 μ l pipette to 20 μ l and then pipette the entire volume up and down at least 10 times to mix thoroughly. Perform a quick spin to collect all liquid from the sides of the tube.

Place the tube on a thermocycler and perform PCR amplification using the following PCR cycling conditions:

- 1. 98C 30"
- 2. 98C 10"
- 3. 61C 75"
- 4. GO TO .2 6X
- 5. 98C 10"
- 6. 72C 75"
- 7. GO TO .5 10X
- 8. 72C 5'
- 9. Hold a 4C

Cleanup of PCR Reaction

- **38** Vortex AMPure XP to resuspend.
- Add 22.5 μ I (0.9X) resuspended beads to the PCR reaction. Mix well by pipetting up and down at least 10 times. Be careful to expel all of the liquid out of the tip during the last mix. Vortexing for 3-5 seconds on high can also be used. If centrifuging samples after mixing, be sure to stop the centrifugation before the beads start to settle out.
- 40 Incubate samples on bench top for at least 5 minutes at room temperature.

- Place the tube on an appropriate magnetic stand to separate the beads from the supernatant. If necessary, quickly spin the sample to collect the liquid from the sides of the tube wells before placing on the magnetic stand.
 After 5 minutes (or when the solution is clear), carefully remove and discard the supernatant. Be careful not
- Add 100 µl of 80% freshly prepared ethanol to the tube while in the magnetic stand. Incubate at room temperature for 30 seconds, and then carefully remove and discard the supernatant. Be careful not to disturb the beads that contain DNA targets.

to disturb the beads that contain DNA targets (Caution: do not discard the beads).

- Repeat Step 43 once for a total of two washes. Be sure to remove all visible liquid after the second wash. If necessary, briefly spin the tube, place back on the magnet and remove traces of ethanol with a p10 pipette tip.
- Air dry the beads for up to 5 minutes while the tube is on the magnetic stand with the lid open. Caution: Do not over-dry the beads. This may result in lower recovery of DNA target. Elute the samples when the beads are still dark brown and glossy looking, but when all visible liquid has evaporated. When the beads turn lighter brown and start to crack they are too dry.
- Remove the tube from the magnetic stand. Elute the DNA target from the beads by adding 33 μ l of 10 mM Tris-HCl.
- Mix well by pipetting up and down 10 times, or on a vortex mixer. Incubate for at least 2 minutes at room temperature. If necessary, quickly spin the sample to collect the liquid from the sides of the tube wells before placing back on the magnetic stand.
- Place the tube on the magnetic stand. After 5 minutes (or when the solution is clear), transfer 30 μ l to a new PCR tube for and store at -20° C.

Library can be stored at -20°C.