shERWOOD-Ultramir Lentiviral shRNA Pooled Libraries

Format: Lentiviral Particles

I. Introduction

shERWOOD-Ultramir Lentiviral shRNA Pooled Libraries from transOMIC technologies are pooled lentiviral shRNA constructs targeting human or mouse genes to enable highly multiplexed, RNAi mediated gene knockdown. Screening libraries are available as pooled plasmid DNA or pooled high titer lentiviral particles and target gene families and pathways, whole genome or any custom gene list.

shERWOOD-UltramiR shRNA screening libraries can be used to screen hundreds to thousands of genes in a high-throughput, multiplexed assay using a standard tissue culture hood without the aid of costly automation and liquid handling required for traditional arrayed screens. Lentiviral vectors allow the user to select from the broadest range of cell lines for functional screens including primary, stem and non-dividing cells.

All shERWOOD-UltramiR shRNA (Figure 1) constructs in the pooled screening libraries are vector-based RNAi triggers with a new generation shRNA-specific design and an optimized microRNA scaffold “UltramiR” which has been shown to produce more potent and consistent knockdown performance than existing shRNA reagents. The UltramiR scaffold has been optimized for efficient primary microRNA processing (Auyeung et al., 2013) and shRNA designs are predicted using the proprietary shERWOOD algorithm developed in Dr. Gregory Hannon’s laboratory at Cold Spring Harbor Laboratory. Based on the functional testing of 270,000 shRNA sequences using a high-throughput sensor assay (Knott et al., 2014), the shERWOOD algorithm has been trained to select the rare shRNA-mir designs that are consistently potent even at single copy representation in the genome which make them ideal for pooled screens and greatly simplifies downstream analysis by decreasing both false positive and false negative results.
Lentiviral pooled shRNA screening libraries are provided standard in the ZIP-mCMV vector. They are also available in ZIP-hCMV, ZIP-hEF1-alpha, ZIP-mEF1-alpha and ZIP-SFFV vectors. **No changes to the protocols are required for use with the alternate ZIP vectors.** Detailed vector graphics are provided in Appendix D.

**Figure 1.** Schematic of shERWOOD-UltramiR shRNA. (A) Passenger (green) and Guide (orange) strand are shown with Dicer and Drosha nuclease cleavage sites are in red. (B) The final step of shRNA processing loads the Guide Strand (orange) into the RISC complex which binds to the target mRNA (blue) in a sequence specific manner.

**Figure 2.** Schematic of ZIP lentiviral shRNA vectors. All elements are identical with the exception of the promoter (LTRs are not shown). The ZIP vectors constitutively express the shRNA, fluorescent marker and puromycin selection marker from a single transcript driven by a viral promoter. This allows for direct visual confirmation of shRNA-mir expression. A promoter selection kit ([TLN0005](#)) is available for quickly determining the optimal promoter for a given cell line.
II. Safety
See Appendix A for safety considerations.

Pooled shRNA Screening Strategies
Pooled shRNA libraries are typically used to perform enrichment (positive) or dropout (negative) selection screens. Both strategies begin by transduction of cells with viral particles expressing a complex pool of shRNA such that each transduced cell expresses a single shRNA. Next, a selective pressure is applied to the heterogeneous population of cells to elicit a desired phenotypic change (e.g., survival screen as shown in 3). The shRNA is integrated into the cellular genome and its representation in the culture increases or decreases as the cells are selected for or against, respectively. Changes in the shRNA representation can be determined through sequence analysis of the shRNA sequence.

Figure 3. Schematic showing the three basic phases of the screening process: transduction, screening and analysis.

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III. Product specifications

Shipping and storage
Components are shipped on wet ice and should be stored at -20°C upon receipt.

Lentiviral particle library components

<table>
<thead>
<tr>
<th>Table 1. Components provided</th>
</tr>
</thead>
<tbody>
<tr>
<td>Component</td>
</tr>
<tr>
<td>Pooled Screening library – Viral particles</td>
</tr>
<tr>
<td>Non-targeting Control – Viral particles</td>
</tr>
<tr>
<td>Primary PCR Forward Primer</td>
</tr>
<tr>
<td>Primary PCR Reverse Primer</td>
</tr>
</tbody>
</table>

Equipment and reagents to be supplied by user

For titering:
- 1.5 ml microcentrifuge tubes
- HEK293T cells (NIH-3T3 may be used as an alternative)
- Complete media for HEK293T cells and experimental cell line
- Serum free media for each cell line
- 24-well tissue culture plate

For Puromycin kill curve:
- Complete media for the experimental cell line
- Puromycin (1.25 μg/μl stock solution)
- 24-well tissue culture plate

For Transduction:
- Polybrene
- Puromycin

For PCR:
- KOD Hot Start Polymerase (Novagen Cat# 71086-4)
- 96-well PCR plates
- Betaine
- Agarose gel
- PCR purification column (QIAquick PCR Purification Kit, Qiagen Cat# 28104)
- DNA gel extraction kit (QIAquick Gel Extraction, Qiagen Cat# 28104)
- Multiplex indexed primer kit (12-plex, TRP0002)

Equipment:
- Automatic pipetter/Pipette-aid
- Disposable or autoclaved tissue culture pipettes
- CO2 cell culture incubator at 37°C
- Fluorescent microscope with GFP filter (refer to Appendix B for alternative methods that do not require this)
- Qubit® fluorometric
- Illumina® sequencing platform
shRNA representation in pooled libraries – Quality Control

shRNA pools have been developed using strategies that maintain equimolarity of the pooled shRNA constructs – to ensure that screening results are not artificially biased to any one or some gene/s or construct/s prior to the screen. All shRNA plasmid DNA pools are evaluated by next-generation sequencing (NGS) for distribution and representation analysis. Quality metrics are outlined in the Certificate of Analysis (C of A) provided with each pool. A representative graph of shRNA distribution is shown in Figure 4. For a more detailed vector map and link to vector sequence see Appendix D.

<table>
<thead>
<tr>
<th>Pool Screening Metrics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total shRNA in Pool</td>
</tr>
<tr>
<td>shRNA Detected (by NGS)</td>
</tr>
<tr>
<td>% Recovery</td>
</tr>
<tr>
<td>Mean Read Depth (MRD)</td>
</tr>
<tr>
<td># ± 5X MRD</td>
</tr>
<tr>
<td>% ± 5X MRD</td>
</tr>
</tbody>
</table>

**Figure 4.** Example histogram showing distribution of shRNA from a plasmid DNA pool as seen by NGS analysis (left). Summary statistics of QC analysis (right). The summary includes the total shRNA constructs pooled to create the library and the number detected by NGS.
IV. Pooled shRNA Screen – Preparation and optimization

The following User’s Guide provides detailed protocols for each step performed during pooled shRNA screening. Please review Appendix A for all safety considerations prior to beginning these protocols.

1. Relative transduction efficiency and transduction optimization

The transduction step determines the fold representation of shRNA in the pool and the number of viral particles transducing a cell. Both of these parameters are integral to a successful screening process. The following protocol is designed to evaluate functional titer of the virus produced in the previous section for optimal transductions during the screen. Following transduction, puromycin will be used to remove non-transduced cells. The cell line’s natural resistance to puromycin will be evaluated using an antibiotic titrating assay to determine the optimal concentration for selection during the screen.

Increase screening quality and capacity by increasing transduction efficiency:
Optimized transduction allows for maximal shRNA fold representation and biological replicates and thus more effective screen performance. A number of variables influence transduction efficiency including components of the media, duration of transduction, cell type, cell health and plating density. It is possible to optimize many of these variables prior to the screen.

- Serum is a known inhibitor of transduction and should be minimized (0 - 2%) in transduction media. For cells sensitive to low serum conditions either reduce the transduction time in low serum media or increase the transduction time in complete media.
- Hexadimethrine bromide (Polybrene) is a cationic lipid known to enhance viral particle binding to the surface of many cell types. A range of concentration (0 - 10 µg/ml) should be tested to determine the highest transduction efficiency that can be achieved without cell toxicity.
- Extending transduction incubation times may increase efficiency. However, it may be necessary to increase the volume of media applied to the cells for transduction to limit the effects of evaporation.
- Cell density may influence transduction efficiency. Plate cells at a range of densities to determine its effect on your cell line. Rapidly dividing cells are often transduced more efficiently.
Determining Functional Titer

Functional titer must be determined using the experimental cell line to ensure optimal transduction during the screen. The functional titer is the number of viral particles, or transducing units (TU), able to transduce the target cell line per volume and is measured in TU/ml. Cell type, media components and viral production efficiency influence functional titer, and it should therefore be calculated for every batch of virus produced or when using a different cell line.

The functional titers for the viral particles provided with the screening library C of A are determined in HEK293T cells. However, many cell lines have lower or higher transduction efficiencies than HEK293T. The relative transduction efficiency is a measure of the ratio of the number of HEK293T cells transduced to the number of experimental cell line cells transduced given the same volume of viral particles. Once this ratio is determined it can be applied to remaining the tubes of viral particles. This allows the functional titer to be determined without consuming viral particles needed for the screen. Formulas and example calculations are provided below.

Once a baseline titer is known, this protocol can be used to further optimize transduction efficiency. To do so, follow this procedure and alter variables know to influence transduction efficiency.

- The following protocol evaluates titer by manually counting ZsGreen positive colonies. Alternate methods for determining titer are provided in Appendix B.
- Transduction optimization should be done with the Non-Targeting Control viral particles prior to titering the shRNA pool to ensure optimal conditions and sufficient titers for the screen. If extensive optimization is required additional non-targeting control may be purchased as viral particles (TLNVU4420).
- HEK293T cells may be used for troubleshooting. If needed, repeat the protocol with the experimental cell line and HEK293T.

Required materials

- HEK293T or NIH-3T3 cells (optional for troubleshooting)
- Complete media for cell line/s
- Serum free media for each cell line
- 24-well tissue culture plate
- Lentiviral particles
- Microcentrifuge tubes

Equipment

- Automatic pipetter /Pipette-aid
- Disposable or autoclaved tissue culture pipettes
- CO₂ cell culture incubator at 37°C
- Fluorescent microscope with GFP filter (refer to Appendix B for alternative methods that do not require this)
Titering Protocol

1. Plate experimental cell line 24 hours prior to transduction in a 24 well plate. Plate at a density of 5-8 x 10^4 cells per well in 14 wells with complete media. Cells should be seeded at a density so that the next day the cells are approximately 40% confluent at time of transduction. Incubate overnight with 5% CO2 at 37°C.

2. Prepare a serial dilution series with serum free media and viral supernatant as shown in Figure and Table 2
   a. Set up 6 sterile microcentrifuge tubes.
   b. Add 980 µl to the first tube and 800 µl of serum free medium to each remaining tube.
   c. Add 20 µl of viral stock to the first microcentrifuge tube.
   d. Mix well by gently pipetting 200 µl up and down (10 - 15 times) without creating bubbles and discard the tip.
   e. Transfer 200 µl from the first microcentrifuge tube to the second tube. Mix well and discard the tip.
   f. Repeat the procedure for 5 of the remaining tubes.

3. Remove media from cells in each well.
4. Add 225 µl of serum free media to each well.
5. Add 25 µl from each viral dilution to two wells (250 µl final volume) for a total of 12 wells. The remaining two wells (without viral particles) should be evaluated as negative controls.
6. Rock plate gently a few times to mix.
7. Incubate overnight with 5% CO2 at 37°C.
8. Replace the viral supernatant with complete media and allow cells to grow for 48 hours.
9. Count the number of colonies expressing ZsGreen. A colony consisting of multiple cells should be counted as a single transduction event.
   Note: Counting 50-200 colonies is sufficient to provide accurate titers. Only analyze wells that have < 20% of cells transduced to ensure none of the cells have been transduced with more than one viral particle.
10. Use the calculations below and Table 2 to determine functional titer and relative transduction efficiency. See the following page for an example calculation.

Functional titer calculations:

**Functional titer of non-targeting control (NTC) viral particles in experimental cell line:**

\[
(Number \ of \ colonies) \times (dilution \ factor) \div 0.025 \ ml = \frac{\text{TU}}{\text{ml}} \text{ functional titer in experimental cell line}
\]

**Relative transduction efficiency of experimental cell line:**

\[
\frac{(\text{NTC titer in HEK293T cells [provided in C of A]})}{(\text{functional titer of NTC in experimental cell line})} = \text{relative transduction efficiency}
\]

**Functional titer of pooled screening library viral particles in experimental cell line:**

\[
\frac{(\text{Pool titer in HEK293T cells [provided in C of A]})}{(\text{Relative transduction efficiency})} = \text{Functional titer of Pool screening library viral particles in experimental cell line}
\]
Dilution table and schematic for titration protocol

### Table 2 Dilution factors for calculating viral titer

<table>
<thead>
<tr>
<th>Tube</th>
<th>Viral particles</th>
<th>Dilution medium</th>
<th>Dilution factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>20 µl (from titer aliquot)</td>
<td>980 µl</td>
<td>50</td>
</tr>
<tr>
<td>2</td>
<td>200 µl (from Tube 1)</td>
<td>800 µl</td>
<td>250</td>
</tr>
<tr>
<td>3</td>
<td>200 µl (from Tube 2)</td>
<td>800 µl</td>
<td>1250</td>
</tr>
<tr>
<td>4</td>
<td>200 µl (from Tube 3)</td>
<td>800 µl</td>
<td>6,250</td>
</tr>
<tr>
<td>5</td>
<td>200 µl (from Tube 4)</td>
<td>800 µl</td>
<td>31,250</td>
</tr>
<tr>
<td>6</td>
<td>200 µl (from Tube 5)</td>
<td>800 µl</td>
<td>156,250</td>
</tr>
</tbody>
</table>

Figure 5 Schematic of serial dilution for viral particle titering. (--) indicates untransduced control.

Example:
If the control virus has a titer of $2 \times 10^8$ TU/ml (Refer to the C of A for the exact titer of your lot of viral particles.), then the expected number of fluorescent colonies in the titering assay using HEK293T would be:

<table>
<thead>
<tr>
<th>Tube</th>
<th>Dilution</th>
<th>Diluted titer TU/ml</th>
<th>Volume used to transduce cells (ml)</th>
<th>Fluorescent colonies expected</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1/50</td>
<td>4,000,000</td>
<td>0.025</td>
<td>100,000</td>
</tr>
<tr>
<td>2</td>
<td>1/5</td>
<td>80,000</td>
<td>0.025</td>
<td>20,000</td>
</tr>
<tr>
<td>3</td>
<td>1/5</td>
<td>160,000</td>
<td>0.025</td>
<td>4,000</td>
</tr>
<tr>
<td>4</td>
<td>1/5</td>
<td>32,000</td>
<td>0.025</td>
<td>800</td>
</tr>
<tr>
<td>5</td>
<td>1/5</td>
<td>6,400</td>
<td>0.025</td>
<td>160</td>
</tr>
<tr>
<td>6</td>
<td>1/5</td>
<td>1280</td>
<td>0.025</td>
<td>32</td>
</tr>
</tbody>
</table>

If the experimental cell line has an average of 100 colonies in well A4 and B4 then the functional titer for the experimental cell line is calculated as follows:

$$100 \text{ colonies} \times 6,250 \div 0.025 \text{ ml} = 2.5 \times 10^7 \text{ TU/ml}$$

Since the functional titer of the control in HEK293T cells = $2 \times 10^8$ (provided on the C of A) the experimental cell line is transduced 8 times less efficiently. The relative transduction efficiency is calculated as follows:

$$2 \times 10^8 \text{ TU/ml (functional titer in HEK293T)} \div 2.5 \times 10^7 \text{ TU/ml (functional titer in experimental cell line)} = 8$$

The relative transduction efficiency can be used to calculate the functional titer of the pooled shRNA viral particles in the experimental cell line. If the pool shRNA viral particles functional titer in HEK293T cells = $4 \times 10^8$ (provided in the C of A) then the functional titer in the experimental cell line is calculated as follows:

$$4 \times 10^8 \text{ TU/ml (functional titer in HEK293T)} \div 8 \text{ (Relative transduction efficiency)} = 5 \times 10^7 \text{ TU/ml (functional titer in experimental cell line)}$$
1. Puromycin selection (puromycin kill curve)
During the screen cells are transduced at an MOI of 0.3 leaving 70% of the cells untransduced. Puromycin selection should be applied to remove the untransduced cells. In the following protocol the lowest concentration of puromycin that provides adequate selection is determined for the experimental cell line.

**Required materials**
- Complete media experimental cell line
- Puromycin (1.25 µg/µl stock solution)
- 24-well tissue culture plate

**Equipment**
- Automatic pipetter /Pipette-aid
- Disposable or autoclaved tissue culture pipettes
- CO₂ cell culture incubator at 37°C

**Protocol**
1. Plate 5 x 10⁴ cells per well in 11 wells of a 24-well tissue culture plate using media without puromycin.
2. Prepare puromycin dilutions in culture media for antibiotic titration as shown in Table 3. Use a puromycin stock solution of 1.25 µg/µl stock solution.

<table>
<thead>
<tr>
<th>Volume of Puromycin Stock Solution Added (µl)</th>
<th>Total Volume of Media plus Antibiotic per 24 Well (µl)</th>
<th>Final Concentration (µg/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>500</td>
<td>0</td>
</tr>
<tr>
<td>0.2</td>
<td>500</td>
<td>0.5</td>
</tr>
<tr>
<td>0.4</td>
<td>500</td>
<td>1</td>
</tr>
<tr>
<td>0.6</td>
<td>500</td>
<td>1.5</td>
</tr>
<tr>
<td>0.8</td>
<td>500</td>
<td>2</td>
</tr>
<tr>
<td>1</td>
<td>500</td>
<td>2.5</td>
</tr>
<tr>
<td>1.2</td>
<td>500</td>
<td>3</td>
</tr>
<tr>
<td>1.6</td>
<td>500</td>
<td>4</td>
</tr>
<tr>
<td>2</td>
<td>500</td>
<td>5</td>
</tr>
<tr>
<td>3</td>
<td>500</td>
<td>7.5</td>
</tr>
<tr>
<td>4</td>
<td>500</td>
<td>10</td>
</tr>
</tbody>
</table>

3. Begin antibiotic selection the following day by replacing antibiotic free media with media containing the appropriate concentrations of puromycin.
4. Incubate cells with 5% CO₂ at 37°C, or use conditions normal for your target cells.
5. Check cells daily to estimate rate of cell death.
6. Replenish the media containing the appropriate concentrations of puromycin every 2 days for 6 days.

*Note: The optimal puromycin concentration will kill the cells rapidly (2 - 4 days). This is particularly important for screens involving essential genes that may be selected against prior to the experiment.*

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V. Primary Selection Screen: Recommendations & Considerations

Once functional titer, transduction efficiency and puromycin concentration have been evaluated in the experimental cell line, the primary selection screen can be performed. The following protocol provides recommendations and guidelines to perform the selection screen.

**Assay optimization**

The objective of a pooled screen is to selectively enrich or deplete cells in a population based on phenotypic change. Biological assays used to evaluate phenotypic changes in the pooled screen may take different forms including survival, behavior (e.g. migration or adhesion), changes in reporter expression or changes in surface marker expression. Each will have variables inherent to the assay that should be optimized such as sensitivity of the assay, concentration of selective agent, duration of treatment and method of selection.

For Example:
Selection screens identify genes affecting cell viability. If a targeted gene decreases growth by 10%, these cells will show a 40% depletion in the population on day 5, but an 80% depletion on day 15. So, it is possible to detect more subtle changes simply by extending time points. This may be of particular relevance for RNAi screens where target gene expression is reduced but not eliminated leaving the possibility of a hypomorphic phenotype (Acta Biochim Biophys Sin (2012) 44 (2): 103-112).

It is also critical to understand shRNA fold representation within the assay. The average number of cells independently transduced with constructs expressing the same shRNA is described as shRNA fold representation. Transductions with high (e.g. 500X – 1000X) and uniform representation for every shRNA provide the optimal starting point for a screen. Higher representation maintains the uniformity across the pool and increases reproducibility between biological replicates. At least two biological replicates are recommended per screen.

**Multiplicity of Infection (MOI)**

In a pooled screen it is critical that transduction is optimized to ensure that each transduced cell has a single genomic integration from the shRNA-expressing viral particle. This enables one to clearly see phenotypes associated with the knockdown of the single gene. The number of integrations correlates with multiplicity of infection (MOI) calculated as the ratio of transducing units to cells. At an MOI of 0.3 or less >95% of cells are predicted to have a single integration and is therefore recommended for pooled screening. The average number of integrations is estimated by the Poisson distribution (Figure 6). The number of cells needed at transduction is determined using the number of cells with viral integrants and the desired MOI (see calculation below).

![Figure 6. Higher MOIs result in a higher number of cells with multiple integrations. An MOI of 0.3 is recommended for pooled screening to limit the number of cells with multiple integrations (4%) and limit the number of cells required for transduction.](image-url)
Calculations for MOI

Number of viral integrants needed:

\[
\text{Number of shRNA in the pool} \times \text{Fold representation} = \text{Number of integrants needed}
\]

Number of cells needed at transduction:

\[
\frac{\text{Number of integrants needed}}{\text{MOI}} = \text{Number of cells needed at transduction}
\]

For Example:

In a pooled screen, cells should be transduced at an MOI of 0.3 to maximize the number of cells with a single integration and limit the number of cells needed at transduction. Transducing a pool of 500 shRNA at 1000-fold representation will require $5 \times 10^5$ transduction units (TU) and approximately $1.5 \times 10^6$ cells to achieve an MOI of 0.3.

Calculate as follows:

\[
500 \text{ shRNA} \times 1000 \text{ fold representation} = 5 \times 10^5 \text{ TU} \\
5 \times 10^5 \text{ TU} / 0.3 \text{ MOI} = 1.5 \times 10^6
\]

Viral particles in transduction

Prior to your screen, confirm that there is sufficient volume of viral particles for the biological replicates and representation needed for your experimental design. The total number of TU needed for the experiment and titer will be required to calculate the volume needed for the experiment. Refer to the titer calculated in previous sections.

Number of transducing units:

\[
\text{Representation} \times \text{shRNA per pool} \times \text{Biological replicates} = \text{Total TU needed for experiment}
\]

Total volume of virus needed for experiment:

\[
\frac{\text{TU for experiment}}{\text{Functional Titer in Experimental Cell Line (TU/ml)}} = \text{Volume (ml) of virus needed for the experiment}
\]
VI. Primary Selection Screen: Guidelines & Protocols

The primary selection screen protocol provides a basic outline of the transduction process. The following should be optimized before proceeding to the primary selection screen:

- Transduction media: % Serum, Polybrene µg/ml
- Time exposed to transduction media: hours or overnight
- Selection media: µg/ml puromycin
- Number of live cells that should be plated when splitting
- Number of cells that should be harvested for each sample

Required materials

- Complete media for experimental cell line
- Complete media for experimental cell line with puromycin
- Pooled screening library as viral particles
- Assay specific controls and reagents
- Qubit® dsDNA BR Assay or Quant-iT™ PicoGreen ® dsDNA Kits (or other fluorometric system specific for quantitation of dsDNA)

Equipment

- Automatic pipetter /Pipette-aid
- Disposable or autoclaved tissue culture pipettes
- CO₂ cell culture incubator at 37°C
- Assay specific equipment
- Qubit® fluorometer or standard spectrofluorometer with fluorescein excitation and emission wavelengths

Protocol:

Prepare cells

1. Plate cells such that they are actively dividing and 40 - 60% confluent at the time of transduction.
2. Feed cells with complete media 3 - 4 hours prior to transduction.

Transduce cells

3. Exchange media with transduction media.
   
   Note: media should be serum free for maximum transduction efficiency.
4. Incubate cells 12 - 24 hours in transduction media.
5. Replace transduction media with complete media (no puromycin).

Puromycin selection

6. Allow cells to grow for 48 hours.
7. Replace media with selection media.
8. Continue feeding cells selection media until untransduced cells have been removed.
Sampling and passaging of cells during screen

9. Split the culture into a control sample and an experimental sample. Each sample should be derived from enough cells to maintain representation of the pool (500-1000 times the number of shRNA in the pool).

10. Start assay selection and take samples as needed. Each sample should include enough cells to maintain representation of the pool (500-1000 times the number of shRNA in the pool).

Genomic DNA (gDNA) extraction

11. Extract gDNA from samples. Ensure that gDNA is extracted from enough cells to maintain representation of the pool (500-1000 times the number of shRNA in the pool).

\[ \text{Note: Subsequent primary PCR should be performed on all extracted gDNA. To ensure that representation is not lost during the genomic DNA purification step the maximum column capacity should not be exceeded as this can decrease yields. In addition, a second elution from the column preparation may be necessary to maximize yields.} \]

12. Quantitate gDNA isolated from each sample and ensure that 260/280 ratios are greater than or equal to 1.8.

For example:
A pool containing 1500 shRNA used for screening with coverage of 1000 viral integrants per shRNA would require a total of $1.5 \times 10^6$ transduced cells. To maintain the 1000-fold coverage of shRNA used during the screen, at least $1.5 \times 10^6$ cells are needed for gDNA isolation from each sample.
VII. shRNA Amplification and Next Generation Sequencing

The representation of each shRNA is detected by next generation sequencing (NGS). The shRNA sequence integrated into the targeted cells genome can be amplified using common sequences flanking the hairpin. Indexes are then added to each sample using the indexed primer kit (TRP0002) and the pooled PCR product from all samples may be analyzed in parallel using NGS analysis. Index primer kits are available from transOMIC technologies.

Individual shRNA are amplified from gDNA with two rounds of PCR. The primary PCR amplifies the shRNA and the flanking region. The secondary PCR uses nested primers to enrich for the primary PCR amplicons using modified primers adapted for NGS on an Illumina® sequencer.

Figure 7. Schematic of PCR primers and sequencing primers as aligned with viral insert.
1. Primary PCR

Primary PCR reactions should be carried out with no more than ~850 ng of gDNA per 50 µl reaction, therefore multiple PCR reactions will be required per sample. The combined PCR reactions for each sample should include all extracted gDNA to maintain representative of shRNA coverage used during the screen of the pool (500-1000 times the number of shRNA in the pool). See the example below for calculations.

For example:
A pool containing 1500 shRNA used for screening with coverage of 1000 viral integrants per shRNA would require 9.9 µg of gDNA per sample be used for amplification.

- 1500(shRNA) x 1000(coverage) x 6.6\(^{12}\) (g/diploid genome)= 9.9\(^6\) grams gDNA (or 9.9µg)

Each 50 µl PCR reaction will contain 850 ng of gDNA.

- 9.9 µg (or 9900 ng) / 850 ng DNA = 11.6 reactions

Twelve PCR reactions per sample will be required for gDNA amplification from this pool.

Required materials

- KOD Hot Start Polymerase (Novagen Cat# 71086-4)
- 96 well PCR plates
- Agarose gel
- Qubit® dsDNA BR Assay or Quant-iT™ PicoGreen ® dsDNA Kits (recommended for their specificity to quantitate dsDNA in solution)
- PCR purification column (QIAquick PCR Purification Kit, Qiagen Cat# 28104)
- DNA gel extraction kit (QIAquick Gel Extraction, Qiagen Cat# 28104)
- Multiplex indexed primer kit (12-plex, TRP0002)

Required equipment

- Qubit® fluorometer or standard spectrofluorometer with fluorescein excitation and emission wavelengths
- NGS platform
Primary PCR Protocol

Note: KOD Hot Start Polymerase, Novagen Cat# 71086-4 is recommended for PCR.

1. Based on the gDNA quantitation determine the total number of reactions required to amplify gDNA using no more than 850 ng of gDNA per 50 µl PCR reaction.
2. Set up PCR reactions using components and volumes outlined below for each sample. A 96-well PCR plate is recommended.

A master mix of all components (excluding gDNA) should be made and then added to wells containing the same volume and concentration of gDNA.

<table>
<thead>
<tr>
<th>Components</th>
<th>µl per reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>10X KOD Buffer</td>
<td>5</td>
</tr>
<tr>
<td>dNTP Mix (2 mM each)</td>
<td>5</td>
</tr>
<tr>
<td>MgSO₄ (25 mM)</td>
<td>4</td>
</tr>
<tr>
<td>Betaine (5 M)</td>
<td>5</td>
</tr>
<tr>
<td>Primary PCR Forward (10 µM)</td>
<td>1.7</td>
</tr>
<tr>
<td>Primary PCR Reverse (10 µM)</td>
<td>1.7</td>
</tr>
<tr>
<td>KOD Hot Start Polymerase</td>
<td>1.5</td>
</tr>
<tr>
<td>Genomic DNA (850 ng)</td>
<td>--</td>
</tr>
<tr>
<td>Nuclease free water</td>
<td>--</td>
</tr>
<tr>
<td>Total Volume</td>
<td>50</td>
</tr>
</tbody>
</table>

3. Set up the following PCR cycling program on a thermal cycler with a heated lid.

<table>
<thead>
<tr>
<th>Cycles</th>
<th>Temperature (°C)</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>98</td>
<td>5 min</td>
</tr>
<tr>
<td></td>
<td>95</td>
<td>30 sec</td>
</tr>
<tr>
<td></td>
<td>57</td>
<td>30 sec</td>
</tr>
<tr>
<td></td>
<td>72</td>
<td>30 sec</td>
</tr>
<tr>
<td>1</td>
<td>72</td>
<td>5 min</td>
</tr>
<tr>
<td>1</td>
<td>4</td>
<td>Hold</td>
</tr>
</tbody>
</table>

4. Place samples in the thermal cycler with heated lid on and run the PCR program outlined above.
5. After cycling is complete, briefly centrifuge the 96 well plate containing PCR reactions.
6. Select at least 4 PCR reactions per sample and run 10 µl from each on an agarose gel to verify the 370 bp amplicon.
7. Pool PCR reactions from each sample and purify following manufacturer’s protocol ensuring that column capacity is not exceeded (it may be necessary to use multiple columns per sample for purification). QIAquick PCR Purification Kit, Qiagen Cat# 28104 is recommended.
8. Quantitate purified PCR reactions using a fluorometric assay specific for double stranded DNA (dsDNA).
2. Secondary PCR

Note: Each sample should be amplified using a uniquely indexed (6 base pair index) Secondary PCR Reverse primer if multiple samples are run in parallel on the sequencer. Indexed primer kit is available in 12-plex format (TRP0002) from transOMIC technologies.

Protocol

1. 2.0 µg of primary PCR product should be amplified across 4 reactions (500 ng DNA per 50 µl PCR reaction) for each sample.

2. Set up PCR reactions using components and volumes outlined below for each sample. A 96 well plate is recommended. 

A master mix of all components should be made for each Secondary PCR Reverse- Indexed primer used (excluding DNA) which can then be added to wells containing the same volume and concentration of DNA.

<table>
<thead>
<tr>
<th>Component</th>
<th>µl per reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>10X KOD Buffer</td>
<td>5</td>
</tr>
<tr>
<td>dNTP Mix</td>
<td>5</td>
</tr>
<tr>
<td>MgSO₄ (25mM)</td>
<td>4</td>
</tr>
<tr>
<td>Betaine (5M)</td>
<td>2</td>
</tr>
<tr>
<td>Secondary PCR Forward (10 µM)</td>
<td>7.5</td>
</tr>
<tr>
<td>Secondary PCR Reverse (10 µM)- Indexed</td>
<td>7.5</td>
</tr>
<tr>
<td>KOD Hot Start Polymerase</td>
<td>1.5</td>
</tr>
<tr>
<td>Pooled Primary PCR DNA (500 ng)</td>
<td>--</td>
</tr>
<tr>
<td>Nuclease-free water</td>
<td>--</td>
</tr>
<tr>
<td>Total Volume</td>
<td>50</td>
</tr>
</tbody>
</table>

3. Set up the following PCR cycling program on a thermal cycler with a heated lid.

<table>
<thead>
<tr>
<th>Cycles</th>
<th>Temperature (°C)</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>98</td>
<td>5 min</td>
</tr>
<tr>
<td>15</td>
<td>94</td>
<td>30 sec</td>
</tr>
<tr>
<td></td>
<td>52</td>
<td>30 sec</td>
</tr>
<tr>
<td></td>
<td>72</td>
<td>30 sec</td>
</tr>
<tr>
<td>1</td>
<td>72</td>
<td>5 min</td>
</tr>
<tr>
<td>1</td>
<td>4</td>
<td>Hold</td>
</tr>
</tbody>
</table>

4. Place samples in the thermal cycler with heated lid on and run the PCR program outlined above.

5. After cycling is complete, centrifuge the 96 well plate containing PCR reactions.

6. Pool PCR reactions from each sample.

7. Gel purify the 408 bp product following manufacturer’s recommendations from the kit used. Elute in 30 µl EB (or molecular-grade water).

We recommend using QIAquick Gel Extraction, Qiagen Cat# 28104.
3. Quality Control of NGS-adapted shRNA amplicon libraries
After secondary PCR and purification, it is necessary to accurately quantitate each sample prior to NGS analysis.

Protocol
1. Quantitate shRNA amplicon libraries using a fluorometric assay specific for double stranded DNA (dsDNA).
2. Determine molarity of DNA and dilute each sample to 15 nM using the calculations below.
   - \( \text{(base pairs in DNA amplicon)} \times (660 \text{ grams/mol/base pair}) = \text{molecular weight (MW) of DNA in sample} \)
   - \( \frac{(DNA \ ng/\mu l)}{(MW \ of \ amplicons)} \times 10^6 = DNA \ nM \)
   - \( \frac{(DNA \ nM)}{15} \) = dilution factor for 15 nM final concentration
   - \( \text{(Dilution factor)} \times (Sample \ \mu l) = \text{Total Volume of Sample and dilution buffer} \)

   For example:
   A sample with concentration of 140 ng/µl and amplicon size of 380 bp would require the following dilution factor to achieve a 15 nM final concentration:
   - \( (380 \text{ bp}) \times (660 \text{ grams/mol/bp}) = 247000 \text{ grams/mol} \)
   - \( (140 \text{ ng/µl}) \div (247000 \text{ grams/mol}) \times 10^6 = 566.8 \text{ nM} \)
   - \( (566.8 \text{ nM}) \div 15 = \text{dilution factor of 37.8} \)
   - \( 37.8 \times (3 \text{ µl sample}) = 113.4 \mu l \text{ total volume} \)
   - \( (3 \mu l \text{ sample}) + (110.4 \mu l \text{ EB}) = 15 \text{ nM sample} \)

3. Samples diluted to 15 nM, should be analyzed by qPCR to verify concentrations of Illumina® adapted amplicons. NGS qPCR kits that are designed for quantitation with Illumina adapted libraries will work on all samples amplified with Secondary PCR primers (typically, NGS core facilities will provide this service prior to sequencing).
4. Dilute shRNA amplicon libraries to 10 nM based on qPCR quantitation.
5. Pool equimolar (uniquely indexed) amplicon libraries. It is not necessary to use the entire volume from each library when pooling, 5-10 µl from each should be sufficient.

4. Sequencing shRNA amplicon libraries
An NGS core facility is recommended if assistance is needed for NGS sequencing and analysis. The instructions below describe changes (to standard Illumina NGS protocols) that are required to prepare and sequence shRNA amplicon libraries.

A 50 base pair, single read sequencing run is sufficient to sequence through the unique 22 bases of the guide/sense and identify shRNA. Follow manufacturer protocols for preparing libraries (including denaturing, diluting and loading) and sequencer runs on an Illumina sequencer. We recommend loading ~12 - 18 pM for the pooled libraries, however optimization may be necessary to obtain cluster densities that are recommended by the manufacturer.
Note the following when using Illumina kits:

- The custom shRNA Loop Read 1 Primer is provided as 100 µM, dilute accordingly (typically 1:200 in HT1 buffer).
- Illumina® provided Read 1 and Read 2 Primer Mixes should not be used/selected for sequencing shRNA amplicon libraries.
- PhiX control should not be run with shRNA amplicon libraries as the custom shRNA Loop Read 1 primer is not compatible with PhiX control.
- If samples were amplified using Secondary Indexed Reverse Primers, Illumina provided Index Primer Mix should be used/selected for index reads (indexes correspond to Illumina® TruSeq indices 1 - 12)

5. Data Analysis

The custom shRNA Loop Read 1 Primer binds to the loop of the shRNA (as shown in Figure 6) therefore, the first base read is the 5’ end of the guide/sense sequence. The 22 bases of the guide/sense sequence can be used to identify shRNA. Multiplexed samples should be separated and analyzed according to their index reads.

Analysis of NGS data from a shRNA screen is complex and can be difficult to evaluate without the proper experience or guidance. There are many web-based tools that can be used for alignment and analysis of NGS data; however, for optimal analysis with viable hit identification, it is recommended to have screen data analyzed by an expert in the field of NGS and bioinformatics. Once data has been analyzed and individual shRNA representation has been identified as statistically significant between control and experimental samples, secondary validation of these hits by shRNA should be performed for confirmation of hits. Individual shRNA constructs can be used for hit validation. In many cases, further validation can be performed from over expression as well.
V. Appendix

Appendix A. Safety

Recombinant lentivirus is considered a Biosafety Level 2 organism by the National Institutes of Health and the Center for Disease Control and Prevention. However, local health and safety regulations should be determined for each institution.

For more information on Biosafety Level 2 agents and practices, download Biosafety in Microbiological and Biomedical Laboratories (BMBL), Fifth Edition (Revised December 2009) published by the U.S. Department of Health and Human Services Centers for Disease Control and Prevention and NIH. The publication can be found here: http://www.cdc.gov/biosafety/publications/bmbl5/.

If additional measures are needed, review biosafety guidance documents such as the NIH’s “Biosafety Considerations for Research with Lentiviral Vectors” which refers to “enhanced BL2 containment”. More information can be found through the NIH Office of Biotechnology Activities web site (http://oba.od.nih.gov/rdna_rac/rac_guidance_lentivirus.html)

Summary of Biosafety Level 2 Practices
The following is meant to be a summary of Biosafety Level 2 practices and should not be considered comprehensive. A full account of required practices should be determined for each institute and/or department.

Standard microbiological practices
- Limit access to work area
- Post biohazard warning signs
- Minimize production of aerosols
- Decontaminate potentially infectious wastes before disposal
- Use precautions with sharps (e.g., syringes, blades)
- Review biosafety manual defining any needed waste decontamination or medical surveillance policies

Safety equipment
- Biological Safety Cabinet, preferably a Class II BSC/laminar flow hood (with a HEPA microfilter) used for all manipulations of agents that cause splashes or aerosols of infectious materials; exhaust air is not recirculated
- Protective personal equipment includes: protective laboratory coats, gloves, face protection if needed

Facilities
- Autoclave available for waste decontamination
- Chemical disinfectants available for spills
Appendix B. Alternate methods for titering

1. Puromycin titering by selection and colony counting:
   a. Begin the antibiotic selection by replacing the media with complete media supplemented using the optimal puromycin concentration determined in “puromycin kill curve”
   b. Continue feeding and observe the cells for approximately 7 days until you see single colonies surviving the selection. The negative control should have no surviving cells.
   c. Use a microscope to count the number of surviving colonies.
   d. Calculate the functional titer using the number of colonies visible at the largest dilution that has colonies.

\[
\text{Functional titer} = \frac{\text{Number of colonies} \times \text{dilution factor} \div 0.025 \text{ ml}}{0.025 \text{ ml}}
\]

2. ZsGreen titering by FACS analysis
   a. When calculating the percentage of transduced cells use the number of cells present on the day of transduction as the denominator.

\[
\text{Functional titer} = \frac{\text{Number of cells at transduction} \times \text{dilution factor} \div 0.025 \text{ ml}}{\text{Number of ZsGreen positive cells in well}} = \frac{\text{TU}}{\text{ml}}
\]
Appendix C. **PCR and NGS sequencing primer sequences**

### Primary PCR Forward (5’->3’)
CAGAATCGTGTGCCGCACTCT TGGAAAC

### Primary PCR Reverse (5’->3’)
CGTATCCACATAAGGTAAAGGAGCAAC

### Secondary PCR Forward primer (5’->3’)
AATGATACGGCGACCACACAGATCTTTCCCTACACGACGCTCTCCGATCTCTAGTGAAGCCACAGATGTA

### Secondary PCR Reverse Indexed primer (5’->3’)
CAAGCAGAAGACGGCATACAGATNNNNNNGTGACTGGAGTTCAAGACGTGCTCTCCGATCTCCTGCCATACATAGGTAAAGG

<table>
<thead>
<tr>
<th>Secondary PCR Reverse Index</th>
<th>Secondary PCR Primer sequences with 6 base pair index (5’-&gt;3’)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>CAAGCAGAAGACGGCATACAGATCGTGTGACTGGAGTTCAAGACGTGCTCTCCGATCTCCTGCCATACATAGGTAAAGG</td>
</tr>
<tr>
<td>2</td>
<td>CAAGCAGAAGACGGCATACAGATACATCGTGTGACTGGAGTTCAAGACGTGCTCTCCGATCTCCTGCCATACATAGGTAAAGG</td>
</tr>
<tr>
<td>3</td>
<td>CAAGCAGAAGACGGCATACAGATGCCTAAATGACTGGAGTTCAAGACGTGCTCTCCGATCTCCTGCCATACATAGGTAAAGG</td>
</tr>
<tr>
<td>4</td>
<td>CAAGCAGAAGACGGCATACAGATGTGACTGGAGTTCAAGACGTGCTCTCCGATCTCCTGCCATACATAGGTAAAGG</td>
</tr>
<tr>
<td>5</td>
<td>CAAGCAGAAGACGGCATACAGATCGTGTGACTGGAGTTCAAGACGTGCTCTCCGATCTCCTGCCATACATAGGTAAAGG</td>
</tr>
<tr>
<td>6</td>
<td>CAAGCAGAAGACGGCATACAGATATTGGC</td>
</tr>
<tr>
<td>7</td>
<td>CAAGCAGAAGACGGCATACAGATGCCTAAATGACTGGAGTTCAAGACGTGCTCTCCGATCTCCTGCCATACATAGGTAAAGG</td>
</tr>
<tr>
<td>8</td>
<td>CAAGCAGAAGACGGCATACAGATGTGACTGGAGTTCAAGACGTGCTCTCCGATCTCCTGCCATACATAGGTAAAGG</td>
</tr>
<tr>
<td>9</td>
<td>CAAGCAGAAGACGGCATACAGATCGTGTGACTGGAGTTCAAGACGTGCTCTCCGATCTCCTGCCATACATAGGTAAAGG</td>
</tr>
<tr>
<td>10</td>
<td>CAAGCAGAAGACGGCATACAGATAGCTATGACTGGAGTTCAAGACGTGCTCTCCGATCTCCTGCCATACATAGGTAAAGG</td>
</tr>
<tr>
<td>11</td>
<td>CAAGCAGAAGACGGCATACAGATCGTGTGACTGGAGTTCAAGACGTGCTCTCCGATCTCCTGCCATACATAGGTAAAGG</td>
</tr>
<tr>
<td>12</td>
<td>CAAGCAGAAGACGGCATACAGATATAAGCTATGACTGGAGTTCAAGACGTGCTCTCCGATCTCCTGCCATACATAGGTAAAGG</td>
</tr>
</tbody>
</table>

**Note:** Since the sequences shown in the above table are reverse primers in 5’->3’ orientation, the highlighted indexes are shown as reverse complements. Indexes 1-12 above correspond to Illumina TruSeq indexes 1-12.

**shRNA Loop Sequencing Primer (5’->3’)**
ACGACGCTCTTTCCGATCTTTAGTAAGCGCACAGATGTA

If low level multiplexing is needed (6 samples or less), please refer to *Multiplexing Sample preparation Guide (Illumina® Part# 1005361)* for recommendations on which sets of the indexes above can be used together.
Appendix D. Vector map

Sequence information and detailed maps for all ZIP vectors can be found on the Pooled Screening Documents Page.
References and recommended reading


Additional Reading


Tran et al. Knockdown of specific host factors protects against influenza virus-induced cell death. 2013, Cell Death and Disease 4, e769.

Sarah Cooper and Neil Brockdorff. Genome-wide shRNA screening to identify factors mediating Gata6 repression in mouse embryonic stem cells. 2013, Development 140, 4110-4115.


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