Protocol for Illumina 2bRAD sample preparation

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Critical update: August 25 2014 – introduced NNRW adapters to discard PCR

duplicates

Minor update: March 20, 2015 – added NNRW to the sequence of reduced-

representation (..NG) adapter

Minor update: March 27, 2016 – replaced qPCR with picoGreen quantification in

the end

Overview

This is a modification of the protocol described in Wang et al, Nature Methods 2012 2b-RAD paper (doi:10.1038/nmeth.2023) re-designed for Illumina HiSeq and BcgI enzyme (https://www.neb.com/products/R0545-BcgI). BcgI is a relatively frequent cutter, with 75-100k sites per genome, and in our experience is one of the most efficient of the IIb-type restriction enzymes. It produces 36-base fragments with two-base overhangs and is fully heat-inactivatable, both of which facilitate ligation of adaptors. Another enzyme that can be used with this protocol is AlfI (http://www.thermoscientificbio.com/restriction-enzymes/alfi/)

The latest modifications include the use of barcoded ligated adaptors so the <u>samples can be pooled by 12</u> after ligation, and use of degenerate 5'-adaptor that makes it <u>possible to remove PCR duplicates</u>.

The protocol involves the following steps, of which steps 1-3 are performed within the same tube (or well of a 96-well plate) by consecutively adding reagents.

- 1. Restriction digest. Genomic DNA is digested with a type IIB restriction enzyme to produce restriction fragments of uniform length.
- 2. Ligation. Adaptors, barcoded for 3' end and generic for 5', are ligated to the cohesive ends generated by restriction digest. Overnight ligation at +4°C, then heat-inactivate the ligase for 10 min at 65°C.
- 3. Pool ligations with different 3' barcodes into groups of 12 (because we have 12 different 3' ligation adapters). Determine proportions from gel picture (regular pcr) or by qPCR.
- 4. Amplification and barcoding of pooled ligations.
- 5. Purification of the target 170 bp band by gel-electrophoresis (the only purification step in the whole procedure).
- 6. Quantification and mixing in equal proportions.

Digest

1. Prepare intact, high-quality genomic DNA samples each containing a total of 100-200 ng in 4 μ l. This can be accomplished using by ethanol precipitation or by drying under vacuum. Ideally, samples should have the same amount of DNA.

Note: we have successfully prepared samples from as little as 50 ng.

2. Prepare a digestion master mix. The following recipe is for a single reaction, so multiply by the number of samples plus some small amount for pipetting error.

NEB Buffer #3 0.6 μl 150 μM SAM 0.4 μl BcgI (1 U μl⁻¹) 1.0 μl

*Note: SAM [S-adenosyl-methionine] and enzyme concentrations may differ depending on the manufacturer. If they are different in your case, re-calculate the volumes, make up the difference with nuclease free water if necessary, but keep the single-reaction volume at 6μ l.

- 3. Combine 2 μ l master mix with each 4 μ l DNA sample (6 μ l total volume). Cover this preparation with a drop of mineral oil. Incubate at 37°C 1 hr. If using a heat-inactivatable enzyme (e.g. BcgI, AlfI), inactivate the enzyme at 65°C for 20 min. Hold samples on ice.
 - *Note: one way to increase digestion efficiency is to add one extra μ l of enzyme after the initial 1h incubation and incubate for one more hour before heat-inactivation.
- 4. (optional) For each sample, load 1 μ l digested DNA on a 1% agarose gel alongside a comparable amount of intact DNA from the same sample to verify the effectiveness of the digest.

The signs of an effective enzyme digest are quite subtle and include (a) a slight downward shift in the high molecular weight DNA band and (b) a subtle smear trailing downward from that band. Quite often these effects are not very clear, but the digest is happening. The key criterion for whether the library is good is the number of PCR cycles required to amplify the target band, later in the protocol – it should be 15 or less for full-representation 2bRAD.

Ligation

In this step adaptors are ligated to the restriction fragments produced above. Note that this is the stage at which reduced tag representation (RTR) must be applied by the choice of adaptor sequences.

1. Prepare double stranded adaptors by combining each pair of primers (illBC-ii, antiBC-ii). The oligonucleotide combination used for each alternative 2b-RAD preparations and sequences of each oligo are provided at the end of this document.

For generic Adaptor 1, mix 5Ill-NNRW (10 μ M) with the same volume of Anti5ill-NNRW (10 μ M).

For barcoded Adaptor 2, mix each par: 3IllBC(1-12) (10 μ M) with the same volume of antiBC(1-12) (10 μ M).

Incubate at 42°C for 10 minutes then keep at room temperature until ligation.

2. Prepare 12 master mixes for ligations (one for each barcoded 3' primer). This recipe is for a single reaction, so scale up as needed.

10 mM ATP	0.5 µl
10X T4 ligase buffer	$2.0 \mu l$
5 μM Adaptor 1	1.0 µl
5 μM Adaptor 2(1-12)	1.0 µl
T4 DNA ligase (NEB M0202L)	1.0 µl
NFW (nuclease-free water)	14.5 µl

3. Combine 20 μ l master mix with the remaining 5 μ l of digested DNA (25 μ l total volume). Incubate at 16°C for AlfI and BcgI, at 4°C for BsaXI, for at least 2 hours.

When set up ligation in 96 well plate format we find it useful to prepare 12 master mixes, each with individual Adapter 2(BC), and aliquot it with multi channel pipette. Keep on ice while mixing.

Incubate in 4°C overnight;

Then heat at 65°C for 20 min to inactivate the ligase.

Pooling ligations into 12-plex groups

If you used the same 3' barcoded adapters for all samples skip this step, since you will not be pooling samples at the next stage.

Samples barcoded at the 3' end by different ligation Adapters 2 (1-12) can be pooled into groups of 12. Each such group (you will end up with 8 such groups in a 96 well plate) works as a single sample sample in the subsequent Amplification and Purification, so pooling saves quite a bit of effort.

However, to ensure that we are pooling equivalent amounts of each initial sample into a group, it is necessary to quantify them using one of the following procedures:

- i. Use standard pcr (see next chapter **Amplification**) with 12-14 cycles and estimate the brightness of the bands on a gel. Mix 5µl (most efficient), 10, 15 or 20µl of each ligation in groups of twelve. Label Groups 1-8.
- ii. Alternatively, set up quantitative PCR:

	(volumes given in µl)
H ₂ O	4.45
SYBR Green mix	7.5
10 mpx primer(10 μM)	0.15
any Ill-Rad-bc primer (1 μΜ	1) 1.5
IC1-P5 primer (10 μM)	0.2
IC1-P5 primer (10 μM)	0.2

a. Add 1µl of Ligation, either undiluted or diluted 10 times, to appropriate well as a template. Amplify in a qPCR machine with some NTC (no-template control) using the following profile:

95°C 5 min, (95°C 40 sec, 65°C 2 min, 72°C 30 sec) X 35 cycles

b. Arrange the data in Excel in the form of a table with four columns: sam (sample name), lane (group number), conc (DNA dilution; use 0.066 for undiluted (0.066=1/15) and 0.0066 for 10-fold diluted template), and ct (qPCR result for this sam-conc combination). There must be at least two technical replicates for each combination of sam-conc (i.e. two rows with the same sam and conc and different ct values). The order of columns and rows does not matter, but the names of the columns do matter (note that they are case sensitive).

Export the data from Excel as comma-separated values (.csv). Open script mix_illumina_qpcr.R in R and follow the instructions given in the comments within the script.

Amplification

In this step, the constructs produced by ligation are amplified using a set of four primers that introduce sample (or pooled sample) specific barcodes and the annealing sites for HiSeq amplification and sequencing primers.

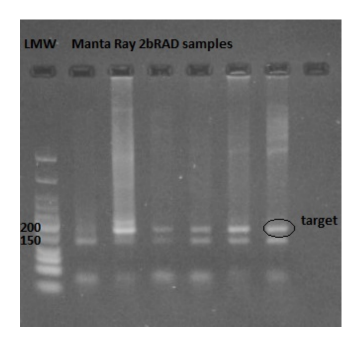
<u>Very Important</u>: if you find that it takes more than 15 cycles to amplify a visible target band, consider optimizing previous steps (DNA isolation, digestion, and ligation), since over-amplified RAD samples tend to lose heterozygosity.

Prepare a test-scale PCR to determine optimum cycle number and evaluate relative yield across samples.

1. For each reaction prepare the following master mix:

NFW	5.2 μl
2.5 mM dNTP	2.5 μl
10 μM IC1-P5	0.4 μl
10 μM IC2-P7	0.4 μΙ
10 μM Mpx primer	0.3 μΙ
5X HF buffer	4.0 μl
Phusion polymerase	0.2 μl #NEB M0530L (or M0530S)

- 2. Combine 13 μl master mix with 4 μl ligation, add 3 μl of 1 μM Ill-Rad-bc (barcode-bearing) primer to each reaction, and amplify as follows: 70°C 30 sec then (95°C 20 sec, 65°C 3 min, 72°C 30 sec) X N cycles
- 3. Sample 5 µl from each reaction at few time points. For example, at 8, 10 and 12 cycles. Load these products on a 2% agarose gel with a low molecular weight marker to confirm molecular weight of PCR product.
- 4. Select the minimum number of cycles that produces a visible, but not over-amplified, product at ~170 bp. You might also see a 130bp band, which this seems to be an artifact from the carried-over ligase (make sure to heat-inactivate the ligase before mixing the PCR reaction!).



5. Prepare the following master mix for each sample:

NFW 13 μl 2.5 mM dNTP 6 μl 10 μM Mpx primer 0.75 μl 10 μM IC1-P5 1 μl 10 μM IC2-P7 1 μl 5X HF buffer 10 μl

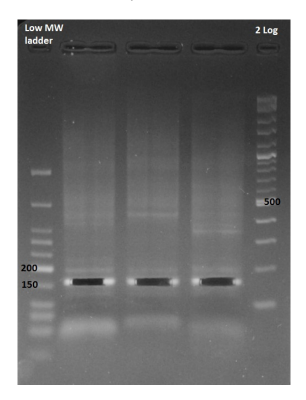
Phusion polymerase 0.5 µl #NEB M0530L (or M0530S)

6. Combine 30 μ l master mix with 10 μ l ligation (group 1-8), add 7.5 μ l of 1 μ M Ill-Rad-bc (barcode-bearing) primer individually to each reaction, and amplify using the temperature profile as in 2 above with the cycle number as determined in 4.

Purification

In this step the target band is gel-extracted to exclude high-molecular weight fragments and any chaff that may emerge during PCR (e.g., primer dimers).

- 1. Prepare a 2% agarose gel using TBE or TB. Use a wide comb that can accommodate 50 μ l sample loading plus 10 μ l loading dye, or tape together two wells if required.
- 2. Load the entire volume of each reaction alongside a low-molecular weight ladder. Run gel at low voltage for a long time until bands at 150bp and 200bp will clearly resolved. In our experience 100V for 70 minutes produce the good separation.
- 3. View the gel <u>briefly</u> (<30 seconds) on a UV or appropriate for your DNA dye black-light transilluminator set at low intensity to verify the presence of target bands and adequate separation of molecular weight standards to resolve bands at ~170 bp and ~130 bp. Typically ~5 cm run distance is sufficient. Photograph.
- 4. Cut out target band of 170 bp in a narrow gel slice, avoiding the edges of the lane (i.e., cut out the middle 70-75% of the band). Cut just inside the bottom boundary of the target band to avoid getting anything smaller than 170bp.



*Note: at this stage a commercial gel-extraction kit can be substituted for the following three steps, if you feel more confident this way. In our practice, simply soaking the gel slice in water overnight, as described below, works just fine.

- 5. Transfer each gel slice into a 1.5 ml microcentrifuge tube and add 20 μ l NFW.
- 6. Make sure gel slice is in contact with water (cut or break it into a few, say 4-5, smaller pieces to make sure they sit comfortably at the bottom of the tube). Hold overnight at 4°C.
- 7. The following day transfer the eluate (~15 µl) into a new tube. This material is now ready for sequencing, pending qPCR quantification (see the writeup and scripts under "Quantifying samples for sequencing on the same Illumina HiSeq lane")

Oligonucleotide sequences for Illumina HiSeq

Name	Sequence (5' – 3')
5ILL-NNRW	CTACACGACGCTCTTCCGATCTNNRWCCNN
Anti5ill-NNRW	GGWYNNAGATCGG/3InvdT ¹ /
3ILL-NN	CAGACGTGTGCTCTTCCGATCTNN
anti-ILL	AGATCGGAAGAGC/3InvdT ¹ /
5ILL-NG ²	CTACACGACGCTCTTCCGATCTNNRWCCNG
3ILL-NG ²	CAGACGTGTGCTCTTCCGATCTNG
ILL-Mpx ³	AATGATACGGCGACCACCGAGATCTACACTCTTTCCCTACACGACGCTCTTCCGAT
ILL-RAD-bc4	${\tt CAAGCAGAAGACGGCATACGAGAT \ [barcode]^5 \ GTGACTGGAGTTCAGACGTGTGCTCTTCCGAT}$
IC1-P5	AATGATACGGCGACCACCGA
IC2-P7	CAAGCAGAAGACGGCATACGA

¹ InvdT: inverted dT to prevent extension by DNA polymerase.

- 2 These two adaptors can be used to reduce representation of the 2b-RAD tags down to approximately 1/16th of the total number. This is a useful cost-lowering trick for applications such as basic population genetics, relatedness analysis, or QTL mapping, the power of which would max out already at a few hundred polymorphic markers.
- 3 This is a standard Illumina-"universal" primer in TrueSeq v.3 (configuration 5' P5-index2-R1primingSite 3')
- 4 "bc" stands for "barcode", and is typically replaced by the barcode number (or other barcode-specific identifier) in the actual name of the primer. This primer can be substituted for a standard TruSeq v.3 barcoded oligo of the configuration 5' P7-Index1-R2primingSite 3'.
- 5 barcode: a 6-base sequence easily distinguishable from other sequences on that same sequencing run. A list of good working barcodes can be found here: https://wikis.utexas.edu/display/GSAF/Illumina+-+all+flavors

BUT NOTE that the barcode sequences listed at that site are in fact REVERSE COMPLEMENTS relative to what needs to be written in the ILL-RAD-bc oligo (we had so many fun moments because of that)

Secondary-barcoded adapters for Ligation (to be used instead of 3ILL-NN and anti-ILL)

3illBC1 CAGACGTGTGCTCTTCCGATCT ACAC NN	
3illBC2 CAGACGTGTGCTCTTCCGATCT GTCT NN	
3illBC3 CAGACGTGTGCTCTTCCGATCT TGGT NN	
3illBC4 CAGACGTGTGCTCTTCCGATCT CACT NN	
3illBC5 CAGACGTGTGCTCTTCCGATCT GATG NN	
3illBC6 CAGACGTGTGCTCTTCCGATCT TCAC NN	
3illBC7 CAGACGTGTGCTCTTCCGATCT CTGA NN	
3illBC8 CAGACGTGTGCTCTTCCGATCT AAGC NN	
3illBC9 CAGACGTGTGCTCTTCCGATCT GTAG NN	
3illBC10 CAGACGTGTGCTCTTCCGATCT GACA NN	
3illBC11 CAGACGTGTGCTCTTCCGATCT GTGA NN	
3illBC12 CAGACGTGTGCTCTTCCGATCT AGTC NN	
antiBC1 GTGT AGATCGGA/3InvdT/	
antiBC2 AGAC AGATCGGA/3InvdT/	

ACCA AGATCGGA/3InvdT/
AGTG AGATCGGA/3InvdT/
CATC AGATCGGA/3InvdT/
GTGA AGATCGGA/3InvdT/
TCAG AGATCGGA/3InvdT/
GCTT AGATCGGA/3InvdT/
CTAC AGATCGGA/3InvdT/
TGTC AGATCGGA/3InvdT/
TCAC AGATCGGA/3InvdT/
GACT AGATCGGA/3InvdT/

Barcoded PCR primers (can be substituted for standard TruSeq v.3 oligos: 5' P7-**Index1**-R2primer 3')

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ILL-RAD01
            CAA GCA GAA GAC GGC ATA CGA GAT CGT GAT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD02
            CAA GCA GAA GAC GGC ATA CGA GAT ACA TCG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
            CAA GCA GAA GAC GGC ATA CGA GAT GCC TAA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD03
ILL-RAD04
            CAA GCA GAA GAC GGC ATA CGA GAT TGG TCA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD05
            CAA GCA GAA GAC GGC ATA CGA GAT CAC TGT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
            CAA GCA GAA GAC GGC ATA CGA GAT ATT GGC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD06
ILL-RAD07
            CAA GCA GAA GAC GGC ATA CGA GAT GAT CTG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD08
            CAA GCA GAA GAC GGC ATA CGA GAT TCA AGT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD09
            CAA GCA GAA GAC GGC ATA CGA GAT CTG ATC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD10
            CAA GCA GAA GAC GGC ATA CGA GAT AAG CTA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD11
            CAA GCA GAA GAC GGC ATA CGA GAT GTA GCC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
            CAA GCA GAA GAC GGC ATA CGA GAT TAC AAG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD12
            CAA GCA GAA GAC GGC ATA CGA GAT TTG ACT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD13
            CAA GCA GAA GAC GGC ATA CGA GAT GGA ACT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD14
ILL-RAD15
            CAA GCA GAA GAC GGC ATA CGA GAT TGA CAT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
            CAA GCA GAA GAC GGC ATA CGA GAT GGA CGG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD16
ILL-RAD17
            CAA GCA GAA GAC GGC ATA CGA GAT CTC TAC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD18
            CAA GCA GAA GAC GGC ATA CGA GAT GCG GAC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD19
            CAA GCA GAA GAC GGC ATA CGA GAT TTT CAC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD20
            CAA GCA GAA GAC GGC ATA CGA GAT GGC CAC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
            CAA GCA GAA GAC GGC ATA CGA GAT CGA AAC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD21
ILL-RAD22
            CAA GCA GAA GAC GGC ATA CGA GAT CGT ACG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
            CAA GCA GAA GAC GGC ATA CGA GAT CCA CTC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD23
ILL-RAD24
            CAA GCA GAA GAC GGC ATA CGA GAT GCT ACC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD25
            CAA GCA GAA GAC GGC ATA CGA GAT ATC AGT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD26
            CAA GCA GAA GAC GGC ATA CGA GAT GCT CAT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD27
            CAA GCA GAA GAC GGC ATA CGA GAT AGG AAT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
            CAA GCA GAA GAC GGC ATA CGA GAT CTT TTG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD28
ILL-RAD29
            CAA GCA GAA GAC GGC ATA CGA GAT TAG TTG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD30
            CAA GCA GAA GAC GGC ATA CGA GAT CCG GTG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD31
            CAA GCA GAA GAC GGC ATA CGA GAT ATC GTG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
            CAA GCA GAA GAC GGC ATA CGA GAT TGA GTG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD32
ILL-RAD33
            CAA GCA GAA GAC GGC ATA CGA GAT CGC CTG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD34
            CAA GCA GAA GAC GGC ATA CGA GAT GCC ATG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD35
            CAA GCA GAA GAC GGC ATA CGA GAT AAA ATG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD36
            CAA GCA GAA GAC GGC ATA CGA GAT TGT TGG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
            CAA GCA GAA GAC GGC ATA CGA GAT ATT CCG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
ILL-RAD37
ILL-RAD38
            CAA GCA GAA GAC GGC ATA CGA GAT AGC TAG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT
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ILL-RAD39 CAA GCA GAA GAC GGC ATA CGA GAT GTA TAG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD40 CAA GCA GAA GAC GGC ATA CGA GAT TCT GAG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT CAA GCA GAA GAC GGC ATA CGA GAT GTC GTC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD41 ILL-RAD42 CAA GCA GAA GAC GGC ATA CGA GAT TAG CGC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD43 CAA GCA GAA GAC GGC ATA CGA GAT CGA TTA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD44 CAA GCA GAA GAC GGC ATA CGA GAT GCT GTA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD45 CAA GCA GAA GAC GGC ATA CGA GAT ATT ATA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD46 CAA GCA GAA GAC GGC ATA CGA GAT GAA TGA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD47 CAA GCA GAA GAC GGC ATA CGA GAT TCG GGA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD48 CAA GCA GAA GAC GGC ATA CGA GAT CTT CGA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD49 CAA GCA GAA GAC GGC ATA CGA GAT TGC CGA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD50 CAA GCA GAA GAC GGC ATA CGA GAT GTG TTT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT CAA GCA GAA GAC GGC ATA CGA GAT CCT TCA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD51 ILL-RAD52 CAA GCA GAA GAC GGC ATA CGA GAT TAT GTT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD53 CAA GCA GAA GAC GGC ATA CGA GAT GAC GCG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD54 CAA GCA GAA GAC GGC ATA CGA GAT TGT ATC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT CAA GCA GAA GAC GGC ATA CGA GAT CAC ACC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD55 ILL-RAD56 CAA GCA GAA GAC GGC ATA CGA GAT TTC TTA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT CAA GCA GAA GAC GGC ATA CGA GAT CTC GCT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD57 ILL-RAD58 CAA GCA GAA GAC GGC ATA CGA GAT TAA CCG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD59 CAA GCA GAA GAC GGC ATA CGA GAT AAA GCT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD60 CAA GCA GAA GAC GGC ATA CGA GAT AGA CCA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT CAA GCA GAA GAC GGC ATA CGA GAT GGG ATA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD61 ILL-RAD62 CAA GCA GAA GAC GGC ATA CGA GAT ACG ACA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD63 CAA GCA GAA GAC GGC ATA CGA GAT GTG GGG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD64 CAA GCA GAA GAC GGC ATA CGA GAT TCG TAT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD65 CAA GCA GAA GAC GGC ATA CGA GAT CAA GGG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD66 CAA GCA GAA GAC GGC ATA CGA GAT GCC GGT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD67 CAA GCA GAA GAC GGC ATA CGA GAT CAG TAA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT CAA GCA GAA GAC GGC ATA CGA GAT AGT TCC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD68 ILL-RAD69 CAA GCA GAA GAC GGC ATA CGA GAT AAT AAC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD70 CAA GCA GAA GAC GGC ATA CGA GAT ACT TTT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT CAA GCA GAA GAC GGC ATA CGA GAT TCC CTT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD71 CAA GCA GAA GAC GGC ATA CGA GAT ATA CTT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD72 ILL-RAD73 CAA GCA GAA GAC GGC ATA CGA GAT AGA TGT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD74 CAA GCA GAA GAC GGC ATA CGA GAT AAT CGT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT CAA GCA GAA GAC GGC ATA CGA GAT CGG CGT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD75 ILL-RAD76 CAA GCA GAA GAC GGC ATA CGA GAT GAG AGT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD77 CAA GCA GAA GAC GGC ATA CGA GAT GAT TCT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD78 CAA GCA GAA GAC GGC ATA CGA GAT CCC AAT GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD79 CAA GCA GAA GAC GGC ATA CGA GAT ACG CGG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD80 CAA GCA GAA GAC GGC ATA CGA GAT AGG GCG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD81 CAA GCA GAA GAC GGC ATA CGA GAT CTG CAG GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD82 CAA GCA GAA GAC GGC ATA CGA GAT AAC TTC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD83 CAA GCA GAA GAC GGC ATA CGA GAT GGG TGC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD84 CAA GCA GAA GAC GGC ATA CGA GAT TCC TGC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT CAA GCA GAA GAC GGC ATA CGA GAT CGC GGC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD85 CAA GCA GAA GAC GGC ATA CGA GAT ACC GCC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD86 ILL-RAD87 CAA GCA GAA GAC GGC ATA CGA GAT TAA TAC GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT CAA GCA GAA GAC GGC ATA CGA GAT CAC GTA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD88 ILL-RAD89 CAA GCA GAA GAC GGC ATA CGA GAT ATG TGA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD90 CAA GCA GAA GAC GGC ATA CGA GAT TAT AGA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT CAA GCA GAA GAC GGC ATA CGA GAT TTT GCA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD91 CAA GCA GAA GAC GGC ATA CGA GAT GTG CCA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT ILL-RAD92 ILL-RAD93 CAA GCA GAA GAC GGC ATA CGA GAT CTA ACA GTG ACT GGA GTT CAG ACG TGT GCT CTT CCG AT

qPCR quantification for mixing on the same HiSeq lane

Run Quant-IT picogreen DS DNA assay (Life Technologies P7589) to determine the final concentrations of the eluted product in order to mix libraries in equal proportions.

PicoGreen assay Protocol:

- 1) Place 100ul 1X TE into all first column_wells except B1.
- 2) Add 150ul of DNA standard (@ 2ug/ml, which is the same as 2ng/ul) into B1.
- 3) Serially dilute standards by taking 50ul of B1, mixing into C1, taking 50ul of C1, mixing into D1, and so on until taking 50ul from H1 and throwing it out.
- 4) To all sample wells, add 98ul of 1X TE.
- 5) Add 2ul sample DNA to sample wells.
- 6) Mix Pico Green Master mix: 99.5ul 1XTE + 0.5ul PicoGreen for one sample. Multiply accordingly (plus 8 wells for DNA standard).
- 7) Add 100ul of master mix to all standard and sample wells, bringing up final volumes in each well to 200.
- 8) Read the fluorescence (excitation 480nm, emission 520nm). We use SpectraMax M2 plate reader and Costar assay plates 96 well, no lid, flat bottom, non-treated black with black bottom (Corning 3650) or clear bottom (Corning 3631).
- 9) Save the data into txt file, assemble the results in Excel in two-column form well, reading save it as comma-delimited (.csv) file. The file must contain all A1-H1 wells (blank and calibrators) plus an arbitrary number of sample wells, in any order. See file picogreen.csv as an example.
- 10) Use picogreen.R script to calculate sample concentrations (ng/ul in the original sample).