Manual

Introduction to the terminology and methods used in transgene & integration site TLA analyses & ddPCR





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TLA and sequencing

The samples submitted for TLA analysis are used and processed according to Solvias' Targeted Locus Amplification (TLA) protocol, developed by Cergentis (de Vree et al. Nature Biotechnology 2014). A summary of the TLA technology is shown in **Figure 1**.

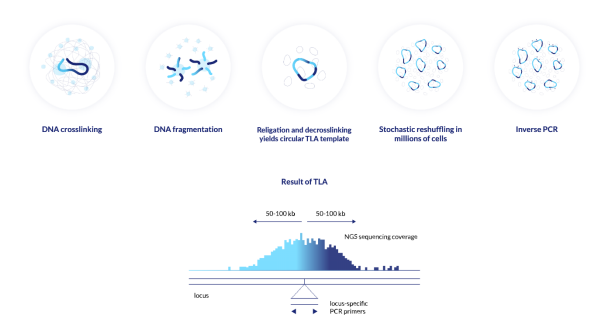


Figure 1: Schematic representation of TLA technology.

In brief, the TLA technology uses the physical proximity of sequences within a locus as the bases of selection. First, DNA is crosslinked, fragmented and re-ligated to generate the circular fragments of DNA. Then, using one TLA primer pair complementary to a sequence within a locus / vector containing a transgene, sequence information is generated across the entire vector and its integration site(s) (**Figure 2**). Solvias recommends the use of two vector-specific primer pairs complementary to two different sequences within a vector. The primer sets are used in individual TLA amplifications and subsequent analysis. This results in two independent data sets.

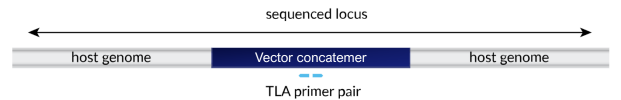


Figure 2: TLA-based transgene and integration site sequencing using transgene specific primer pairs.

PCR products are purified, library prepped using the Nextera® DNA Flex Library Prep protocol (Illumina), and pooled. The resulting pool contains unique barcodes (Nextera® DNA CD Indexes, Illumina) for each PCR product. NGS sequencing (paired-end, 2X149 bases) is performed on an Illumina System. For a standard analysis, ~1 million reads are generated per PCR product.



Alignment and processing of sequencing reads

The Illumina® System, together with the bcl2fastq Conversion Software (Illumina), performs base calling and demultiplexing (converting the base call information into the read information). Using the barcode information, paired-end FASTQ files are generated for each individual amplification of a TLA sample. Reads are mapped to the vector sequence and host genome using BWA-MEM, version 0.7.15-r1140, settings bwa mem -M -t 4 -B 7 -w 33 -O 5 -E 2 -T 33 -Y (Li H., 2013, arXiv:1303.3997).

Vector sequence coverage

Definition of vector and transgene coverage

Vector refers to the entire sequence that was used to modify the cells, so including a transgene/region of interest and backbone. Solvias recommends sending the full vector reference sequence in order to optimally assess its integration and the presence of sequence and structural variants.

Coverage is defined as the number of NGS reads that cover a base (in a vector sequence or genomic locus). The coverage determines the sensitivity with which sequence and/or structural variants can be detected.

Partial/complete vector integrations

A gap in vector coverage indicates partial integrations of the vector caused by (un)intended deletion(s) in the original vector or introduced during integration (**Figure 3**, left panel).

In case homology arms/ITRs/LTRs were used for (targeted) integrations, TLA data will, depending on how clean such integrations have been, show the presence or absence of coverage on the backbone sequences (**Figure 3**, right panel).

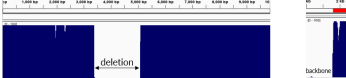




Figure 3: NGS sequencing coverage across the vector in the random (left) and targeted integration (right, homology arms are shown in red).

In samples with virus-mediated gene transfer, the obtained complete vector coverage might be a result of amplification of the episomal vector DNA remaining in the cells. In this case, the identification of the numerous integration sites in heterogeneous samples might be not possible due to loss of sequencing capacity on the non-integrated sequences. The experimental set-up can therefore be adjusted to eliminate episomal DNA before integration site enrichment.

Multiple vectors and co-integrations

Depending on the nature of multiple integrated vector sequences, TLA primers can be used on sequences common to all integrated vectors, or primers can be designed that are specific for each individual vector. The specific TLA primer sets allow identification of vector-specific integration sites or demonstrate cointegration of individual vectors.

TLA analyses can be used to sequence and identify (un)expected co-integrations of (partial) vector sequences, unknown vector sequences and/or of other DNA sequences (e.g. *E. coli*). Detailed analyses of unexpected sequences can be performed by mapping generated NGS data on appropriate reference (genome) sequences.



Large vectors

In the analysis of large vectors (>50 kb), additional TLA primer pairs can be added to ensure sufficient sequencing coverage is generated on the vector and its integration site(s).

Variations in vector sequence

TLA allows identification of sequence variants (single nucleotide variants, insertions and deletions of one or several nucleotides), structural variants within the integrated vector, and its concatemerization. Heterogeneity of the sample, as well as copy number of the integrated vector and sequencing depth will affect the sensitivity of calling sequence and structural variants.

Sequence variants

The presence of sequence variants is determined using samtools mpileup (samtools version 1.11) (Li *et al.* Bioinformatics, Jun 2009 [PMID: 19505943], Li H. Bioinformatics, Nov 2011 [PMID: 21903627]). Only readbases with a minimal Q-score of 20 (Base call accuracy of >99%) are used for the detection.

Sequence variants are reported that meet the following criteria:

- Allele frequency (relative amount of reads with the variant, compared to total coverage on the variant position) of at least 5% for cell lines/cells in culture and 20% for primary cells;
- The variant is present in the data generated with at least 2 primer sets;
- For at least one of the primer-sets the coverage is ≥30x;
- The variant is identified in both forward and reverse aligning sequencing reads (variants present in >95% reads of the same orientation in the data of at least 1 primer set with ≥30x mutated bases are filtered out);
- The variant is not located in a repetitive sequence: variants surrounded by ≥4 bases of the mutated base are removed unless they show >50% frequency in at least 1 primer sets; InDels followed by ≥6 of the same bases/set of bases are removed unless they show >50% frequency in at least 1 primer sets;
- Low frequency variants (between 5-20% mutant allele frequency) are not found with similar frequencies in an independent control (Solvias recommends the use of an independent control for more reliable filtering).

Sequence variants are reported in a table which includes the following columns:

- Region annotated region in the reference sequence that was used for mapping;
- Position position within the reference sequence;
- Reference nucleotide present in the reference sequence at this position;
- Mutation observed mutation, "+" indicates an insertion, "-" indicates a deletion right downstream of the reference nucleotide:
- Coverage total number of reads passing Q20 that map to the indicated position;
- Percentage (%) percentage of reads containing the mutation (=mut/cov*100%).

Please note that using these filtering criteria, the reported frequencies for an individual sequence variant represent the fraction of all the occurrences of that variant among all vector copies integrated in all loci in the entire cell population.

In very heterogeneous samples (non-clonal cell population with viral/transposon-mediated integrations), detection of variants in individual integration sites is not possible. The identified variants are then categorized as follows:



- A. variants that occur in all samples with >80% mutant allele frequency represent general deviations
 that were present in the supplied reference sequence of the vector before its introduction in the
 cells.
- B. variants that are found in all samples with 5 80% mutant allele frequency can indicate
 heterogeneity in the sequence of the virus that was used. Variants in this category that have low
 allele frequencies (<20%) can also represent systematic sequencing errors. These errors can be
 filtered out by including an independent control for the analysis or by performing independent
 validation experiments.
- C. sample specific variants found in XX, but not in YY or ZZ with 5 100% mutant allele frequency represent specific mutations that occurred in this sample in 5 -100% of the integrated vector.

Vector concatemerization and structural variants

Structural variants within the integrated vector sequence are identified by detecting vector-vector breakpoint sequences.

Terminology

A breakpoint sequence or fusion sequence is a sequence containing the breakpoint of and fusion between, two sequences originating from different origins compared to the reference sequence (**Figure 4**). This can be from two different genes/genomic regions, two vector regions where structural variation takes place or a vector sequence that fuses to a genomic sequence. In case of integration sites, it can also happen when deletions, insertions and inversions take place around the integration site. The reads that are indicative of a fusion sequence are called fusion reads, split reads, breakpoint reads or chimeric reads. In TLA procedure, DNA digestion and ligation results in the procedure-induced fusions of the sequences that are in proximity but are of different origin. These TLA induced fusions are not analyzed.

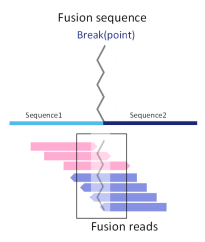


Figure 4: Schematic overview of a fusion or breakpoint sequence. Light blue and the dark blue sequence (on top) fuse together creating a fusion sequence. The point where the sequences fuse together is called the breakpoint. Based on the alignments of fusion reads (pink reads in forward orientation, pink reads in reverse orientation from the sequencer) you can reconstruct the fusion sequence and determine where sequence 1

There are three different breakpoint sequences derived from fusion reads that are reported. The first one (**Figure 5**, number 1) is when there is a perfect junction between sequence 1 (light blue) and sequence 2 (dark blue) and the exact breakpoint positions where the two sequences fuse together is known.



Perfect breakpoint

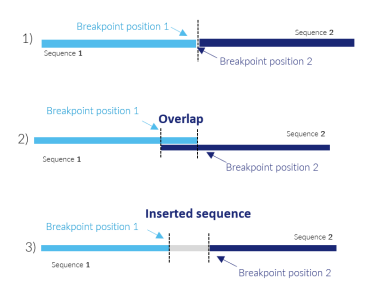


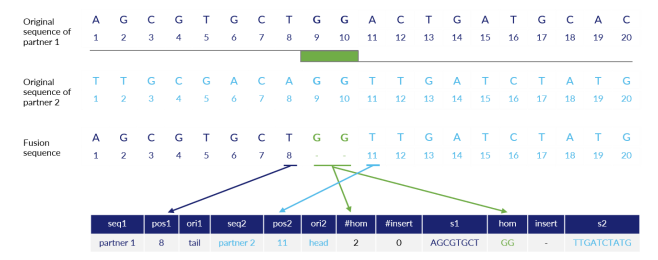
Figure 5: Schematic examples of different fusions. 1 - perfect breakpoint, 2 - fusion with homologous or overlapping bases, 3 - fusion with novel or inserted bases.

The second type of breakpoint sequences are the fusion sequences where there is homology and thus an overlap between sequence 1 and sequence 2 in the alignments of the fusion reads (**Figure 5**, number 2, **Figure 6A**). Bases from sequence 1 also align to sequence 2. With the current data set is not known if the overlapping bases belong to sequence 1 or sequence 2. Thus the overlapping based are removed from the reported breakpoint positions and reported as homology between sequence 1 and sequence 2.

The third type of breakpoint sequences are the fusion sequences that have an inserted sequence between sequence 1 and sequence 2 (**Figure 5**, number 3, **Figure 6B**). The length of the inserted sequence can be varying in size (just a few bases or even50 bases or more). The inserted sequence can be an unaligned sequence from unknown source / origin or align (partially) to the vector or other known sequence (genomic).



A. Example of a breakpoint/fusion sequence with homology



B. Example of a breakpoint/fusion sequence with an insert

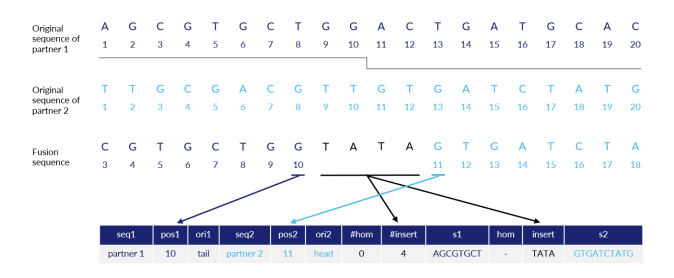


Figure 6: Explanation of the terms 'homology' (A) and 'insert' (B), used to describe details of the sequences found at vector-vector and vector-genome breakpoints.



Detection of a vector-vector breakpoint sequence

Breakpoint sequences consisting of two parts of the vector, are identified using a proprietary Solvias script. Breakpoint sequences resulting from the TLA procedure itself are recognized by the restriction enzymespecific sequence at the junction site and removed.

Vector-vector breakpoint sequences are reported that meet the following criteria:

- The breakpoint sequence is present in =>1% of the reads at one of the positions of the fusion;
- The breakpoint sequence is observed in data of at least two primer sets, unless the data provides a clear explanation why it is not found in one of the data sets;
- The breakpoint sequence is not present in independent control sample(s) (if included);
- Visual inspection of the breakpoint sequence in a NGS data browser is performed to remove those that are sequencing artifacts, e.g. breakpoints found at hairpin structures or low-complexity regions.

In heterogeneous samples, the detection of structural variants within individual integration sites is not possible. Only very abundant events like loss of the specific (backbone) sequences in the majority of integrations is identified.

Details regarding vector-vector breakpoint sequences

Vector-vector breakpoints and integration site breakpoints (see below) are described using the following terms:

- The number of homologous, common, or inserted, novel, bases at the junction site (for details see Figure 6).
- "head" and "tail" orientations of the vector/genome fragments at the junction site (for details see **Figure 7**).

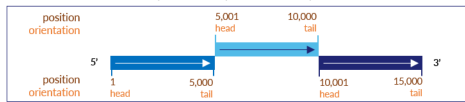
Please note that based on TLA data is it not possible to reconstruct the order and size of individual copies in the integrated concatemer sequence.



Complete sequence



Examples of breakpoints in the sequence



Examples of fusions

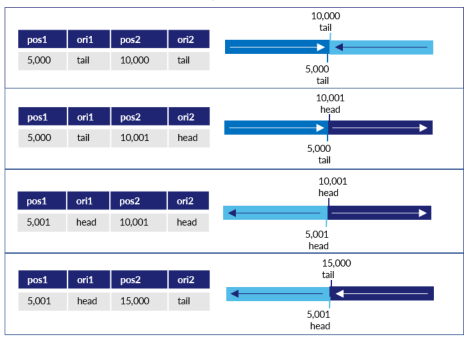


Figure 7: Schematic representation of the terms 'head' and 'tail', used to describe the orientation of sequences found at vector-vector and vector-genome breakpoints.



Integration site detection

Integration sites are detected based on coverage peak(s) in the genome and breakpoint sequences between a vector sequence and host genome.

Coverage peak(s) in the genome

TLA results in high coverage across the genomic positions of vector integration sites. Integration sites are therefore clearly visible in whole genome coverage plots (**Figure 8**).

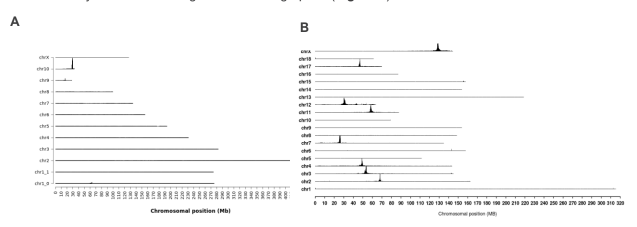


Figure 8: TLA sequence coverage across the Chinese Hamster Ovary (CHO) (A) and mouse (B) genomes. The chromosomes are indicated on the y-axis, the chromosomal position on the x-axis. High coverage peaks represent a single integration (A) and 8 integration sites (B) of a vector.

Partial vector integrations

Partial integrations can be identified if the integrated sequence contains a primer binding sequence.

Breakpoint sequences

A genome-vector breakpoint sequence will consist of a combination of a vector sequence and genomic sequence. Breakpoint sequences are reported in the same manner as vector-vector breakpoint sequences (**Figure 9**).

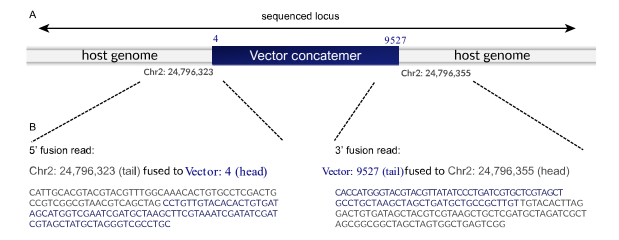


Figure 9: Vector-genome breakpoint sequences. A. Schematic depiction of a vector carrying a transgene and integrated into the host genome at the indicated positions on chromosome 2. B. Breakpoint sequences as depicted in a TLA report. For each breakpoint, the exact sequence as well as the relative orientation of the partner is provided.



Targeted sequencing of individual integration sites

TLA analysis with primers complementary to a wild-type sequence next to an individual integration site will provide sequence information across both transgenic and wild-type alleles (**Figure 10**). This enables determination of zygosity and characterization of the vector integrity (i.e. sequence variants and vector-vector fusions) in that locus.

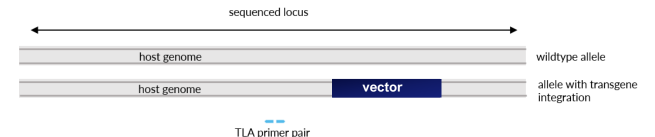


Figure 10: TLA-based analyses of individual integration sites. A TLA analysis with a primer pair in close proximity to the integration site provides sequence information across the entire locus on all alleles.

Integration sites in heterogeneous samples

Detection of the integration sites

In heterogeneous samples, the coverage per integration site that occurred in an individual cell or a small subset of the analysed cells is limited and no integration peak is seen on a genome-wide scale. Therefore, only the breakpoint reads containing vector-genome breakpoint sequences are used for the analyses.

When transduction is performed using a viral/transposon vector, the breakpoints are expected to occur at the boundaries of the LTR/ITR sequences. The breakpoints at these locations are selected and further filtered according to the following criteria:

- The reads in which the part that aligns to the genome is <20 bp are removed to ensure specificity.
- The reads with the wrong orientation at the expected location are filtered out
- The breakpoint sequences containing any inserted, novel, bases between the vector and the genome are filtered out (if the integration mechanism is not designed to generate/allow insertions at the integration sites).
- The duplicates due to homology of LTRs are removed.
- The breakpoint sequences present in the independent control (if included) are filtered out.
- For some breakpoint sequences, two or more hits are found on the genome and it cannot be determined which is the real integration site. In these cases, all breakpoints are reported, so that the total number of identified integrations is slightly over-estimated (typically 1-5%).

Interpretation of the data

After 5-6 million cells are crosslinked, fragmented and circularized, TLA amplification is performed on a subset of the total input, ~100,000 cells. Therefore, if a sample contains rare integration sites, only a subset of the integration sites is present in the amplification reaction. In addition, only a subset of the amplified TLA circles includes vector-genome fusion sequences and only a subset of amplified DNA sequences is sequenced. Therefore, TLA analyses of heterogeneous samples with unique integration sites that occur infrequently, results in identification of numerous integration sites, but each of them is found in the data of only one primer set and with the low number of sequencing reads. Additional amplifications and/or additional sequencing can be considered for a more complete list of the identified integrations. However, increasing the amount of input material per amplification is not required.



Identification of structural variants at integration sites

The presence of structural variants (genomic rearrangements) at the integration site is assessed by analysing the coverage profile and positions of vector – genome breakpoint sequences (**Figure 11**). If the genome sequence is intact, TLA will generate a bell-shaped coverage profile around the integration site. Deviations in this profile indicate genomic rearrangements with respect to the reference sequence.

If complex structural rearrangements are identified, paired-end sequencing analyses can be used to determine which breakpoints occur in physical proximity to each other and constitute the same integration site.

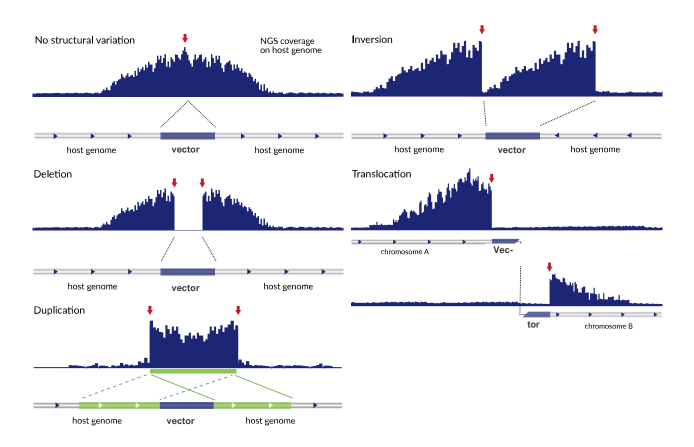


Figure 11: Schematic representation of the changes in the coverage profile and vector-genome breakpoint sequences (red arrows) that are observed in the most common types of rearrangements.



Paired-end information and paired-end analyses

TLA samples are sequenced in paired-end: generated TLA amplicons are sheared to ~600 bp fragments and then both the first 149 bp and the last 149 bp of these fragments are sequenced. The two sequences resulting from the same DNA fragment are called "mates". Both sequences are used in the analyses, and the mating information is kept. The two mates within a pair will originate from the same physical DNA locus. TLA paired-end sequence information can thus be used to assess which vector and/or integration site sequences originate from the same allele (**Figure 12**).

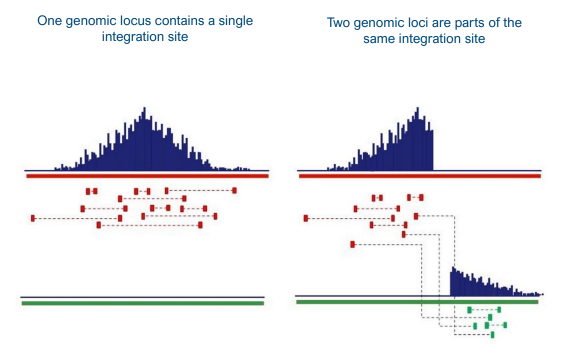


Figure 12: Schematic representation of the use of paired-end information in integration site analyses.



Copy number estimation

TLA is a targeted sequencing method. Routine vector and integration site sequencing analyses using a vector specific primer pair therefore only provide information about the present vector copies and do not provide information about the abundance of cells without the vector sequence.

Based on TLA data itself, copy numbers can only be estimated. For more accurate copy number determination ddPCR is recommended (see below).

Copy number estimations are based on three variables: the number of integration sites, the number of vector-vector junctions in a concatemer, and the ratio of the coverage on the vector-side and genome-side of the integration site (**Figure 13**).

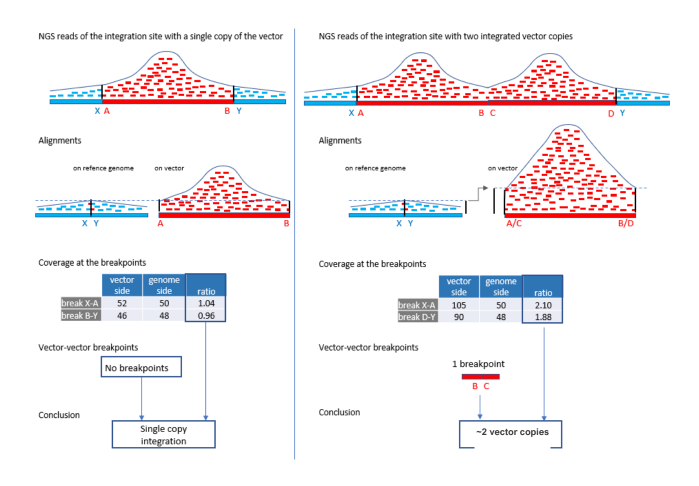


Figure 13: Schematic representation of vector copy number estimation based on TLA data.

If multiple integration sites are identified, standard TLA analyses cannot determine whether these occur in the same or different cells in the analysed sample. Standard TLA analyses can also not identify duplications of complete genomic loci/chromosomes in which vectors have integrated.

Therefore, the provided estimation gives information about the copy number present at an individual locus, which is not necessarily the same as the copy number per cell (**Figure 14-I**). In some cases, a TLA amplification from the genomic side of the integration site(s) can provide more detailed information on the vector copy number per cell (**Figure 14**).



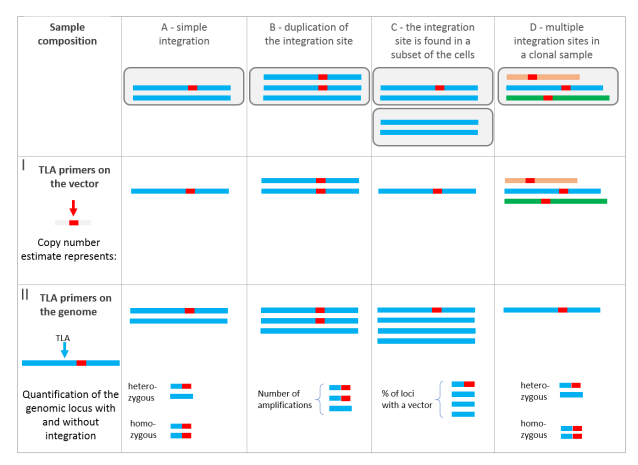
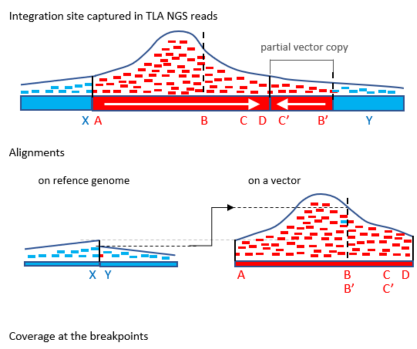


Figure 14: Copy number estimations in different sample types. I) Using vector specific primer sets, only the allele carrying the vector is interrogated. Based on this data alone, one cannot distinguish situations A, B and C. II) TLA amplifications performed from the genomic side of the integration site provides information on the relative frequency of alleles carrying a vector versus wild-type alleles. The combination of the information obtained with TLA amplifications targeting the vector and the genome can resolve the different sample types. Note that in a more complex situations (D), copy number estimations are not always possible.



Rearranged copies of the integrated vector can influence the measurements of the coverage ratio, as illustrated in **Figure 15**. Also, if many vector fragments are concatemerized, it can be difficult to estimate how many vector copies they jointly represent. Therefore, copy number estimations of high copy number vector integrations with complex vector-vector junctions are less accurate.



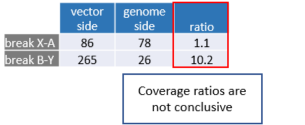


Figure 15: The effect of vector rearrangements on copy number estimations. If rearranged copies of the vector are present, the number of NGS reads produced on the breakpoint position in the vector can differ between the copies of the vector. In those cases, the total NGS coverage obtained on the breakpoint position is not a good representation of the number of vector copies. In a simple case represented here, the copy number can still be deducted, but in more complex cases this becomes increasingly difficult.

In heterogeneous samples with numerous integration sites, a copy number per integration site cannot be estimated based on the TLA data.



Accurate copy number assessment

Digital droplet PCR (ddPCR) allows accurate copy number quantifications and is performed to determine e.g.:

- vector copy number in clonal cell banks
- vector copy number stability over time
- average vector copy number in heterogeneous cell populations (like CAR engineered T-cells)
- presence of residual plasmid
- endogenous gene copy number

ddPCR workflow

ddPCR can be performed either in conjunction with TLA-based service or as a stand-alone experiment. ddPCR assays are designed on one or multiple selected regions of interest (i.e. vector element or endogenous gene) as well as two reference genes. Due to uncertainty about ploidy in modified cell lines two independent reference genes are used. For further confirmation of the ploidy of the cell line and improved vector copy number quantifications, an additional assay can be designed around a vector-genome breakpoint sequence in the modified cell line (previously determined by TLA analysis) as vector breakpoint sequences in random integration events are prone to occur on one allele.

gDNA is either extracted in house from viable frozen or crosslinked cells or gDNA received already isolated. The ddPCR reactions are set up as a duplex or multiplex PCR where a primer/probe pair targeting the region of interest is combined with another primer/probe pair targeting the reference gene tagged with different fluorophores, FAM and HEX are the most used fluorophores. DNA digestion is performed directly in the ddPCR reaction mixture (buffer, dNTPs, primers, and probes) using a restriction enzyme, according to the protocol supplied by Bio-Rad. The mixture is randomly distributed in 20,000 droplets in the QX200 Droplet Generator®. Thereafter, PCR amplification is performed in all droplets simultaneously. The number of droplets being negative and positive for the fluorophore signal is measured using a QX200 Droplet Reader and analyzed with the Bio-Rad QuantaSoft software (QX manager regulatory edition). The Bio-Rad QuantaSoft software fits the fraction of positive droplets to a Poisson algorithm providing the total copies of the region of interest and the reference genes.

Copy number quantifications

The copy number is determined as the ratio of the total copies for the region of interest to the total copies for the reference gene, multiplied by the number of copies for the reference (**Figure 16**). The latter is known, assumed or determined as follows: similar total copies are expected for the reference genes if the cell line sample has a stable diploid or polyploid for their complete genome. The ploidy of the cell line sample may not be previously determined and therefore unknown. In such cases, the cell line sample is considered diploid presenting with 2 copies of the reference genes unless any literature stating otherwise is available. Alternatively, the copy number is inferred based on the ddPCR assay for the breakpoint sequence of the vector integration site (**Figure 16**). For the latter, the copy number of the reference gene is determined as the ratio of the total copies for the reference gene to the total copies for the breakpoint sequence, assuming there is 1 copy of the breakpoint sequence per cell.

```
= ( total copies region of interest total copies reference gene ) X 2 (assuming a diploid genome)

copy number region of interest (i.e. vector element)

= ( total copies region of interest total copies breakpoint sequence) X 1 (assuming integration site on 1 allele)
```

Figure 16: The formulas used to calculate the copy number of the region of interest using genomic ploidy or integration site breakpoint as reference.



Example of ddPCR results

In the following example, the vector copy number was determined for two related samples (MCB and EOPC). The copy number of the vector elements X and Y was determined using ddPCR. Endogenous gene 1 and gene 2 were used as reference genes. In addition, a ddPCR assay for the breakpoint found during TLA analysis was included. The total copies of reference genes 1 and 2 were 3 times higher than the total copies for the breakpoint sequence of each sample (**Figure 17A**), resulting in approximately 3 copies per cell for these reference genes as compared to the breakpoint sequence. Hence, the results suggest that these cell line samples are triploid.

The total copies for the vector elements X and Y were 3 and 4 times higher than the total copies of the reference genes and 9 and 12 times higher than the breakpoint sequence (**Figure 17A**). This indicates the presence of 9 copies per cell for vector element X and 12 copies per cell for vector element Y in each cell line sample (**Figure 17B**). The difference in copy number between the elements indicates partial copies are also present in this sample.

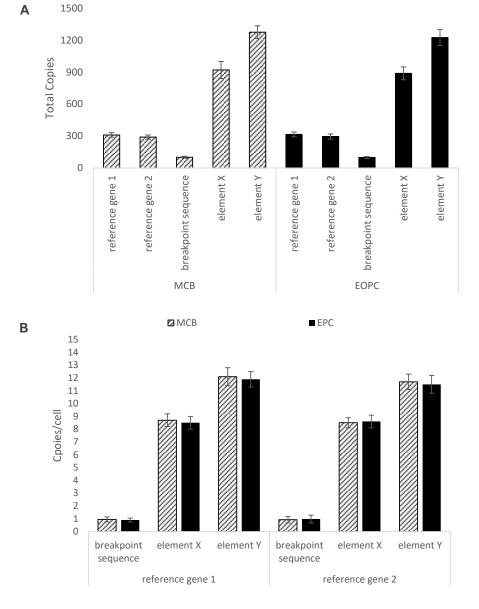


Figure 17: A. The total copies of the breakpoint sequence, genomic reference genes and element X and Y for MCB, and EOPC. B. The copies per cell of the elements X and Y as compared to the genomic reference genes 1 and 2 for MCB, and EOPC. The data represents mean ± SD of three experiments performed in duplicate.