

Example Report

Prepared for:

Customer name:

Internal project number:
Quote number:

Version:
1
Date:

## Goal

In this study, 1 transgenic CHO sample with the vector xxxxx sequence was analyzed.
The aim of this analysis was to:

1. Study the vector integrity;
1) Determine the presence of sequence variants and their allele frequency.
2) Determine the presence of vector-vector breakpoints that represent concatemers of multiple copies of the vector and/or structural rearrangements in a single vector sequence.
2. Identify vector integration site(s) and breakpoint sequences between the vector and genome.
3. Assess the presence of structural variants surrounding the vector integration site(s).
4. Estimate the copy number of the vector.

An overview of the TLA technology and technical details of the performed analyses is provided in the manual "Introduction to the terminology and methods used in TLA analyses v2".

## Summary

| Sample | Vector Integrity | Integration site(s) | Copy number <br> estimation | Notes |
| :--- | :--- | :--- | :--- | :--- |
| Sample 1 | 3 sequence <br> variants, <br> 2 structural <br> variants | 28 | At least 30 (partial) <br> copies | Backbone <br> integration |

## Conclusion

In samples 1, 28 integration sites were observed. Backbone coverage was also observed. 3 sequence variants and 2 structural variants were seen in this sample.

## Methods

## TLA, sequencing and data mapping

Viable frozen CHO-K1 cells were used and processed according to Cergentis' TLA protocol (de Vree et al. Nat Biotechnol. Oct 2014). An overview of the TLA technology and technical details of the performed analyses is provided in the manual "Introduction to the terminology and methods used in TLA analyses_v2".

TLA was performed with 2 independent primer sets specific for the vector sequence (Table 1).
Table 1: Primers used in TLA analysis

| Primer set | Name/Viewpoint | Direction | Binding position | Sequence |
| :--- | :--- | :--- | :--- | :--- |
| $\mathbf{1}$ | GS | Rv | 725 | X |
|  |  | Fw | 650 | X |
| $\mathbf{2}$ | CMV | Rv | 2,745 | X |
|  |  | Fw | 3,186 | X |

The NGS reads were aligned to the vector sequence and host genome. The Chinese Hamster CriGriPICRH1.0 genome assembly GCF_003668045.3 was used as host reference genome sequence.

## Results Sample 1

## Vector integrity

Figure 1 depicts the NGS coverage across the vector sequence using primer set 1. Similar results were obtained with primer set 2.


Figure 1: NGS sequencing coverage across the vector with primer set 1. The black arrow indicates the primer location. Y -axis is limited to 1000 x . In an actual report the data of all primer sets will be presented.

High coverage is observed across the complete vector sequence Vector: 1-10,593. Coverage observed between 1-583 and 9,256-10,593 indicates the backbone has integrated in at least one location in this sample. Local dips in coverage are due to GC rich regions that are less efficiently sequenced.

Sequence variants and structural variants were called in the covered regions.

## Sequence variants

Detected sequence variants are presented in table 2. A total of 3 sequence variants were identified in the sample.

Table 2: Identified sequence variants

|  |  | Primer set 1 |  | Primer set 2 |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Region | Position | Ref | Mut | Coverage | $\%$ | Coverage | $\%$ |
| CMV | 141 | A | C | 21,254 | 20 | 788 | 25 |
| Not annotated | 1,013 | A | $+1 G$ | 751 | 18 | 2,221 | 15 |
| KAN | 10,037 | G | A | 2,145 | 20 | 854 | 20 |

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## Vector concatemerization and structural variants

The identified vector-vector breakpoint sites are shown in table 3. In the accompanying excel tables the sequences and frequencies of the breakpoints are presented. In total, 2 structural variants were identified. Note, the breakpoint 2 indicates that the vector has recombined with itself using ITRs and the TTAA sequence present at position 9,256.

Table 3: Vector-vector breakpoints

| Breakpoint | Vector |  |  | Vector | Orientation of the <br> breakpoint | Homology | Insert |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| $\mathbf{1}$ | $\boldsymbol{\rightarrow}$ | 1,457 | 7,657 | $\boldsymbol{\rightarrow}$ | tail to head | - | $\mathbf{1}$ |
| $\mathbf{2}$ | $\boldsymbol{Z}$ | 2,500 | 9,256 | $\leftarrow$ | tail to tail | 4 | - |

## Integration sites

Whole genome coverage plot


Figure 2: TLA sequence coverage across the Chinese Hamster genome using primer set 1. The chromosomes are indicated on the $y$-axis, the chromosomal position on the $x$-axis.

As shown in figure 2, the vector has integrated on multiple chromosomes. Similar results were obtained with primer set 2 . The identified integration sites are listed in Table 4. Most integration site breakpoints are identified at the expected ITR locations. A random integration event has occurred on chromosome 2 (breakpoint 6). Vector concatemerization most likely has occurred at this random integration site. In the accompanying excel tables the sequences and frequencies of the breakpoints are presented.

Table 4: Identified integration sites

| Breakpoint | Vector |  | Chromosome |  |  | Orientation of the breakpoint | Hom | Insert | Structural variants at the integration site | Gene at integration Site |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | $\rightarrow$ | 9,256 | chr1_0 | 65,102,039 | $\rightarrow$ | tail to head | 4 | - |  | Pbxip1 |
|  | $\leftarrow$ | 583 | chr1_0 | 65,102,042 | $\leftarrow$ | head to tail | 4 |  |  |  |
| 2 | $\rightarrow$ | 9,256 | chr1_0 | 98,996,200 | $\leftarrow$ | tail to tail | 5 | - |  | - |
|  | $\leftarrow$ | 583 | chr1_0 | 98,996,207 | $\rightarrow$ | head to head | 5 | - |  |  |
| 3 | $\rightarrow$ | 9,256 | chr1_1 | 150,700,514 | $\rightarrow$ | tail to head | 5 | - |  | - |
|  | $\leftarrow$ | 583 | chr1_1 | 150,700,508 | $\leftarrow$ | head to tail | 4 | - |  |  |
| 4 | $\rightarrow$ | 9,256 | chr1_1 | 250,408,074 | $\rightarrow$ | tail to head | - | - |  | - |
|  | $\leftarrow$ | 579 | chr1_1 | 250,408,073 | $\leftarrow$ | head to tail | - | - |  |  |
| 5 | $\leftarrow$ | 583 | chr2 | 44,323,798 | $\leftarrow$ | head to tail | 4 | - |  | - |
|  | $\rightarrow$ | 9,256 | chr2 | 44,323,803 | $\rightarrow$ | tail to head | 4 | - |  |  |
| $6^{\#}$ | $\leftarrow$ | 1,250 | chr2 | 179,287,078 | $\leftarrow$ | head to tail | 4 | - |  | - |
|  | $\rightarrow$ | 9,256 | chr2 | 179,287,083 | $\rightarrow$ | tail to head | 4 | - |  |  |
| 7 | $\rightarrow$ | 9,256 | chr2 | 255,303,426 | $\leftarrow$ | tail to tail | 4 | - |  | Fam184a |

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|  | $\leftarrow$ | 583 | chr2 | 255,303,431 | $\rightarrow$ | head to head | 4 | - |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 8 | $\leftarrow$ | 583 | chr2 | 315,900,623 | $\leftarrow$ | head to tail | 4 | - |  | $\begin{gathered} \text { XM_02740 } \\ 1667.2 \end{gathered}$ |
|  | $\rightarrow$ | 9,256 | chr2 | 315,900,629 | $\rightarrow$ | tail to head | 5 | - |  |  |
| 9 | $\rightarrow$ | 9,256 | chr2 | 444,590,878 | $\rightarrow$ | tail to head | 4 | - |  | - |
|  | $\leftarrow$ | 583 | chr2 | 444,590,871 | $\leqslant$ | head to tail | 6 | - |  |  |
| 10 | $\rightarrow$ | 9,256 | chr3 | 20,921,238 | $\rightarrow$ | tail to head | 4 | - |  |  |
|  | $\leftarrow$ | 583 | chr3 | 20,921,233 | $\leftarrow$ | head to tail | 4 | - |  |  |
| 11 | $\leftarrow$ | 583 | chr3 | 65,816,955 | $\leftarrow$ | head to tail | 4 | - |  | $\begin{gathered} \text { XM_02740 } \\ 9106.2 \end{gathered}$ |
|  | $\rightarrow$ | 9,256 | chr3 | 65,816,960 | $\rightarrow$ | tail to head | 4 | - |  |  |
| 12 | $\leftarrow$ | 583 | chr3 | 75,428,314 | $\leqslant$ | head to tail | 4 | - |  | - |
|  | $\rightarrow$ | 9,256 | chr3 | 75,428,319 | $\rightarrow$ | tail to head | 4 | - |  |  |
| 13 | $\leftarrow$ | 583 | chr3 | 85,891,055 | $\rightarrow$ | head to head | - | - |  | $\begin{gathered} \text { XM_02741 } \\ 0136.2 \end{gathered}$ |
|  | $\rightarrow$ | 9,256 | chr3 | 85,891,055 | $\leqslant$ | tail to tail | 3 | - |  |  |
| 14 | $\rightarrow$ | 9,256 | chr3 | 170,208,201 | $\leftarrow$ | tail to tail | 4 | - |  | Zswim4 |
|  | $\leftarrow$ | 585 | chr3 | 170,208,206 | $\rightarrow$ | head to head | 6 | - |  |  |
| 15 | $\leftarrow$ | 583 | chr3 | 213,761,150 | $\leftarrow$ | head to tail | 4 | - |  | - |
|  | $\rightarrow$ | 9,256 | chr3 | 213,761,155 | $\rightarrow$ | tail to head | 4 | - |  |  |
| 16 | $\rightarrow$ | 9,256 | chr3 | 255,204,534 | $\leftarrow$ | tail to tail | 4 | - |  |  |
|  | $\leftarrow$ | 583 | chr3 | 255,204,540 | $\rightarrow$ | head to head | 5 | - |  |  |
| 17 | $\leftarrow$ | 9,263 | chr3 | 255,641,056 | $\rightarrow$ | head to head | 5 | - |  | Rpp40 |
|  | $\leftarrow$ | 583 | chr3 | 255,641,056 | $\leqslant$ | head to tail | - | - |  |  |
| 18 | $\rightarrow$ | 9,256 | chr3 | 275,012,875 | $\rightarrow$ | tail to head | 1 | - | $\begin{gathered} \text { 4.8kb } \\ \text { deletion } \end{gathered}$ | - |
|  | $\leftarrow$ | 583 | chr3 | 275,012,875 | $\leftarrow$ | head to tail | 2 | - |  |  |
| 19 | $\rightarrow$ | 9,256 | chr4 | 125,860,215 | $\leqslant$ | tail to tail | 6 | - |  |  |
|  | $\leftarrow$ | 583 | chr4 | 125,860,222 | $\rightarrow$ | head to head | 4 | - |  |  |
| 20 | $\leftarrow$ | 584 | chr4 | 205,159,441 | $\leftarrow$ | head to tail | 5 | - |  | Rasa2 |
|  | $\rightarrow$ | 9,256 | chr4 | 205,159,446 | $\rightarrow$ | tail to head | 4 | - |  |  |
| 21 | $\leftarrow$ | 579 | chr5 | 120,097,581 | $\leftarrow$ | head to tail | 1 | - |  | Rbm25 |
|  | $\rightarrow$ | 9,256 | chr5 | 120,097,583 | $\rightarrow$ | tail to head | - | - |  |  |
| 22 | $\rightarrow$ | 9,256 | chr6 | 136,164,486 | $\leftarrow$ | tail to tail | 4 | - |  | Ehmt1 |
|  | $\leftarrow$ | 583 | chr6 | 136,164,491 | $\rightarrow$ | head to head | 4 | - |  |  |
| 23 | $\rightarrow$ | 9,256 | chr6 | 155,241,475 | $\leftarrow$ | tail to tail | 4 | - |  | Otog |
|  | $\leftarrow$ | 584 | chr6 | 155,241,480 | $\rightarrow$ | head to head | 5 | - |  |  |
| 24 | $\rightarrow$ | 9,256 | chr7 | 2,230,999 | $\leqslant$ | tail to tail | 4 | - |  | Ephx4 |
|  | $\leftarrow$ | 583 | chr7 | 2,230,004 | $\rightarrow$ | head to head | 4 | - |  |  |
| 25 | $\rightarrow$ | 9,256 | chr7 | 25,809,621 | $\leftarrow$ | tail to tail | 4 | - |  |  |
|  | $\leftarrow$ | 579 | chr7 | 25,812,758 | $\leqslant$ | head to tail | - | - |  |  |
| 26 | $\rightarrow$ | 9,256 | chr8 | 93,163,509 | $\rightarrow$ | tail to head | - | - |  | Ube3c |
|  | $\leftarrow$ | 583 | chr8 | 93,163,509 | $\leftarrow$ | head to tail | 3 | - |  |  |
| 27 | $\leftarrow$ | 583 | chr9 | 15,622,618 | $\leftarrow$ | head to tail | 4 | - |  | - |
|  | $\rightarrow$ | 9,256 | chr9 | 15,622,623 | $\rightarrow$ | tail to head | 4 | - |  |  |
| 28 | $\rightarrow$ | 9,256 | chr9 | 14,787,373 | $\leftarrow$ | tail to tail | 3 | - |  | - |
|  | $\leftarrow$ | 583 | chr9 | 14,622,618 | $\leqslant$ | head to tail | 4 |  |  |  |

## Copy number estimation

In this sample, the copy number is estimated based on the number of integration sites and the number of structural variants identified for the specific vector. 28 integration sites and 2 vector-vector junctions as well as backbone integration are found in this sample. Backbone integration indicates that multiple (partial) vector copies are found at some of the identified integration sites. The copy number is estimated to be at least 30 (partial) vector copies. The numbers provided here are the minimum expected copy numbers.

## QC information

Sample and Study details
Sample receipt date
Condition of sample at receipt
Start date in the lab
Sequencing run
Deviations from the protocol
TLApp version

## Study Personnel

Lab technician
Lab technician qPCR
Data Analyst
QC Analysis and Report

## Quality control

The results are independently verified and reviewed and are an accurate and complete representation of the study. TLA processing of cells, NGS
sequencing, and data analysis (except for copy number) are ISO/IEC 17025:2017 accredited by the Dutch Accreditation Council RvA, Registration number L671.

Scientific approval
Date
Signature


[^0]:    '+' indicates an insertion;

