

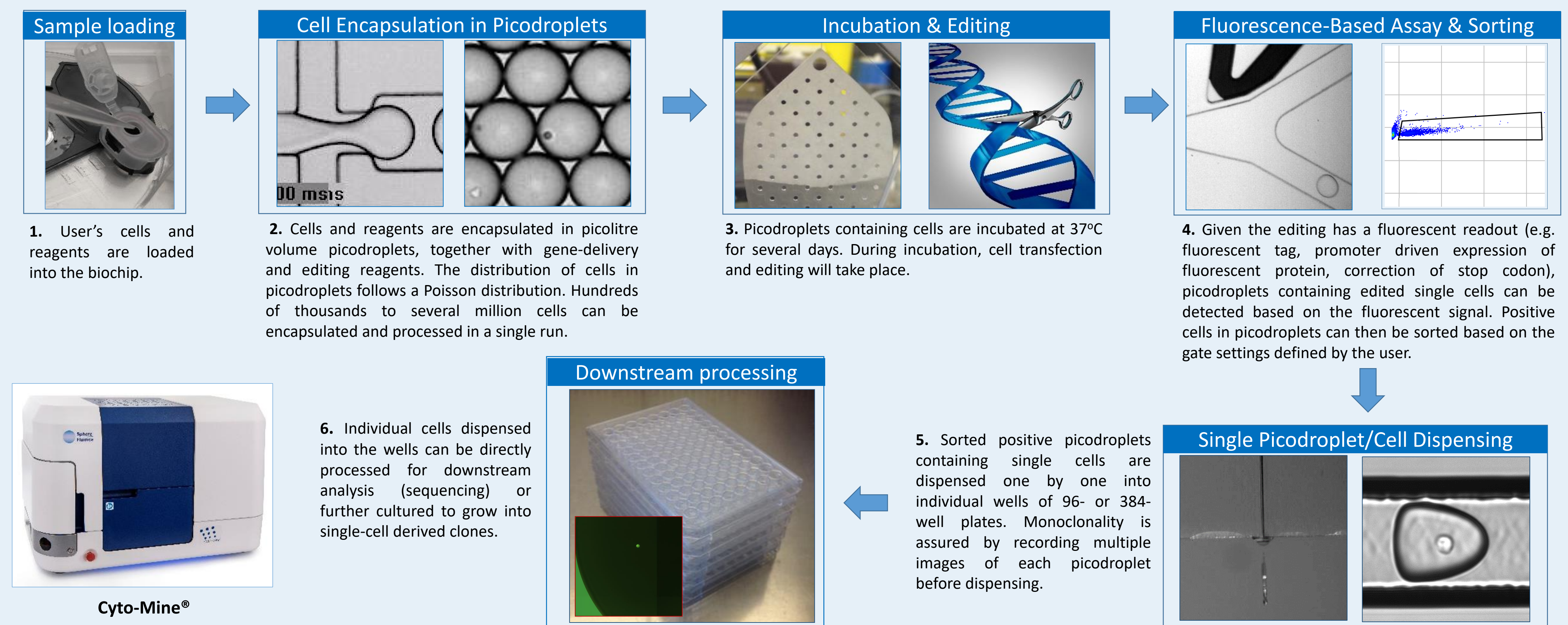
INTRODUCTION

Genome editing is rapidly becoming an essential tool across all areas of life sciences R&D such as gene therapy and regenerative medicine, synthetic biology or biomanufacturing.

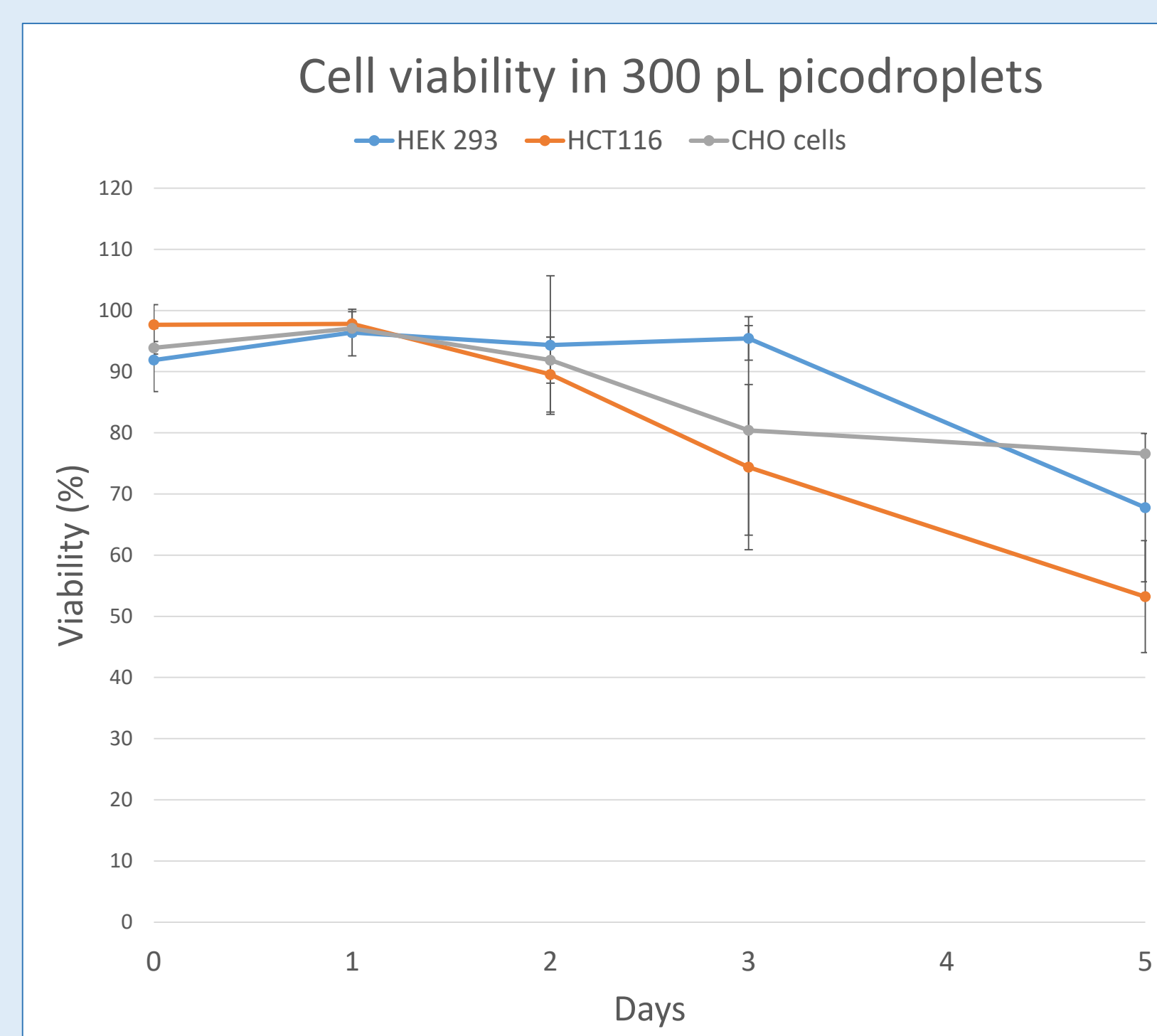
Current approaches, dependent upon manual labour and extensive screening, are highly inefficient and time-consuming, thus new methods that reduce costs and handling time for the generation and recovery of edited cells would be highly welcome.

Using picodroplet microfluidic technology, Sphere Fluidics is developing a novel system that will automate and miniaturise many of the steps of the genome editing pipeline, yielding high-quality, precision engineered cells in a high-throughput manner.

WORKFLOW:

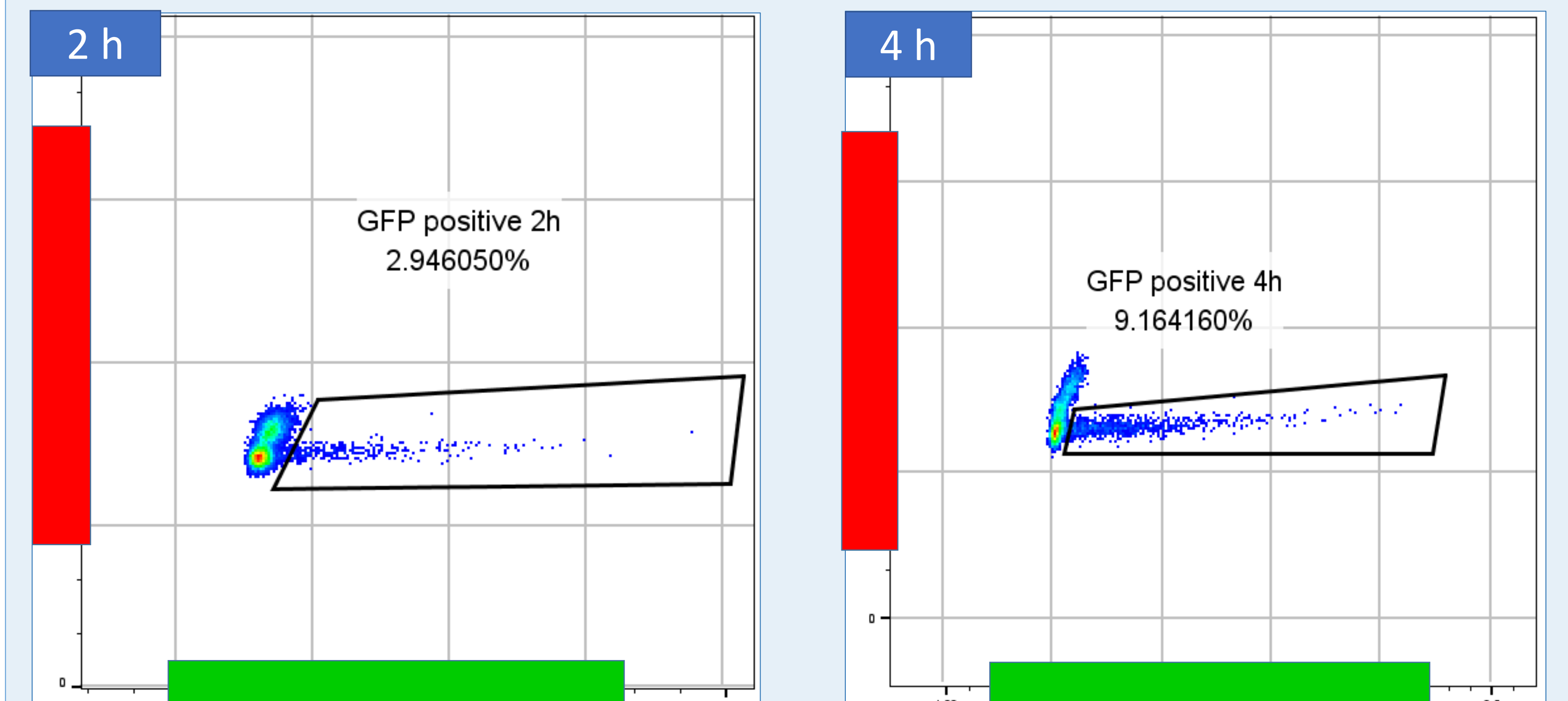


CELL VIABILITY IN 300 pL PICODROPLETS



- Cells were encapsulated in 300pL picodroplets using a PDMS biochip and pump driven flow.
- Cell concentration: 1×10^6 cells/ml.
- Incubation in a 1.5 mL microtube in static incubator, 37°C, 6% CO₂.
- Around 50 μ L of sample was tested per time point. To break the emulsion, 1H,1H,2H,2H-Perfluoro-1-octanol was added and the aqueous phase was collected after a brief spin.
- Viability was tested immediately after breaking emulsion using staining with Acridine Orange (all cells) and DAPI (dead cells).

MRNA TRANSFECTION IN PICODROPLETS



- HCT116 cells (1×10^6 cells/ml) were co-encapsulated in 300pL picodroplets with Cas9/GFP mRNA (100nM) and DharmaFect transfection reagent using Cyto-Mine[®].
- Picodroplets were incubated for 2h and 4h in Cyto-Mine[®], at 37°C.
- Picodroplets containing GFP-expressing cells were analysed at different time points.
- Following a 4h incubation, picodroplets were sorted based on fluorescent signal using Cyto-Mine[®].
- Positive picodroplets were then dispensed into five 96-well plates and successful cells outgrowth was monitored for 2 weeks post-dispensing.

Expected picodroplet occupancy (Poisson distribution)

	λ	0.12
% of empty picodroplets		88.69%
% of picodroplets contain 1 cell		10.64%
% of picodroplets contain 2 cells		0.64%
% of picodroplets contain 2+ cells		0.03%

2h incubation = ~30% of cells transfected
4h incubation = ~90% of cells transfected

GENE EDITING IN PICODROPLETS: TRANSFECTION

	λ	0.30
% of empty picodroplets		74.08%
% of picodroplets contain 1 cell		22.22%
% of picodroplets contain 2 cells		3.33%
% of picodroplets contain 2+ cells		0.36%

4.35% GFP positive picodroplets = 20% edited cells

48 h
GFP positive 48h: 4.351297%

plate#	recovery (%)
1	20.48
2	13.54
3	15.22
4	11.46
5	18.75
6	14.58
average	15.67
stdev	3.36

- HCT116 FIRE-line (kindly provided by Horizon Discovery): inactive GFP under control of CMV promoter integrated into HPRT locus.
- HCT116 cells, RNPs, DNA repair template and DharmaFect transfection reagent were co-encapsulated in 400 pL picodroplets using a Y-junction biochip and pump driven flow.
- Cells were incubated in picodroplets in 1.5 mL microtubes at 37°C, 6% CO₂ for 48 hours and analysed using Cyto-Mine[®].
- Picodroplets, containing GFP-expressing cells were dispensed into 96-well plates and cell outgrowth was analysed 3 weeks later.

GENE EDITING IN PICODROPLETS: VIRAL TRANSDUCTION

- 1 - Generation**: Cells are grown in a well.
- 2 - Transduction**: Cells are co-encapsulated with MBM-VSVG-Cas9 eGFP virus (MOI=0.25) via a Y-junction PDMS biochip and pump driven flow.
- 3 - Re-injection**: After 24h incubation at 37°C, picodroplets are re-injected into Cyto-Mine[®].
- 4 - Sorting**: GFP-positive picodroplets are sorted and dispensed using Cyto-Mine[®] into 96-well plates.
- 5 - Dispensing and outgrowth**: Cell outgrowth was inspected 4 days post-dispensing. RFP (mCherry) signal as readout of genome-editing (insertion of RFP into ACTB gene safe harbour) was detected using fluorescence microscope.

4 - Sorting
GFP positive: 10.36%

- HCT116 cells were co-encapsulated in 400 pL picodroplets with MBM-VSVG-Cas9 eGFP virus (MOI=0.25) via a Y-junction PDMS biochip and pump driven flow.
- After 24h incubation at 37°C, picodroplets were re-injected into Cyto-Mine[®].
- GFP-positive picodroplets were sorted and dispensed using Cyto-Mine[®] into 96-well plates.
- Cell outgrowth was inspected 4 days post-dispensing.
- RFP (mCherry) signal as readout of genome-editing (insertion of RFP into ACTB gene safe harbour) was detected using fluorescence microscope.

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CONCLUSIONS:

We successfully performed CRISPR-mediated gene-editing of adherent mammalian cells in water-in-oil picodroplets, either by lipofection or through viral gene delivery. These editing events were detected via a fluorescent readout; single, positive cells were selectively sorted, dispensed into MTP plates and subsequently grown into single-cell derived clones. The system enables monoclonality assurance, providing a comprehensive solution for accelerated cell line development. Further development work is ongoing, aiming to develop a multiplexing workflow that will allow simultaneous editing of libraries of genes and/or cell lines.