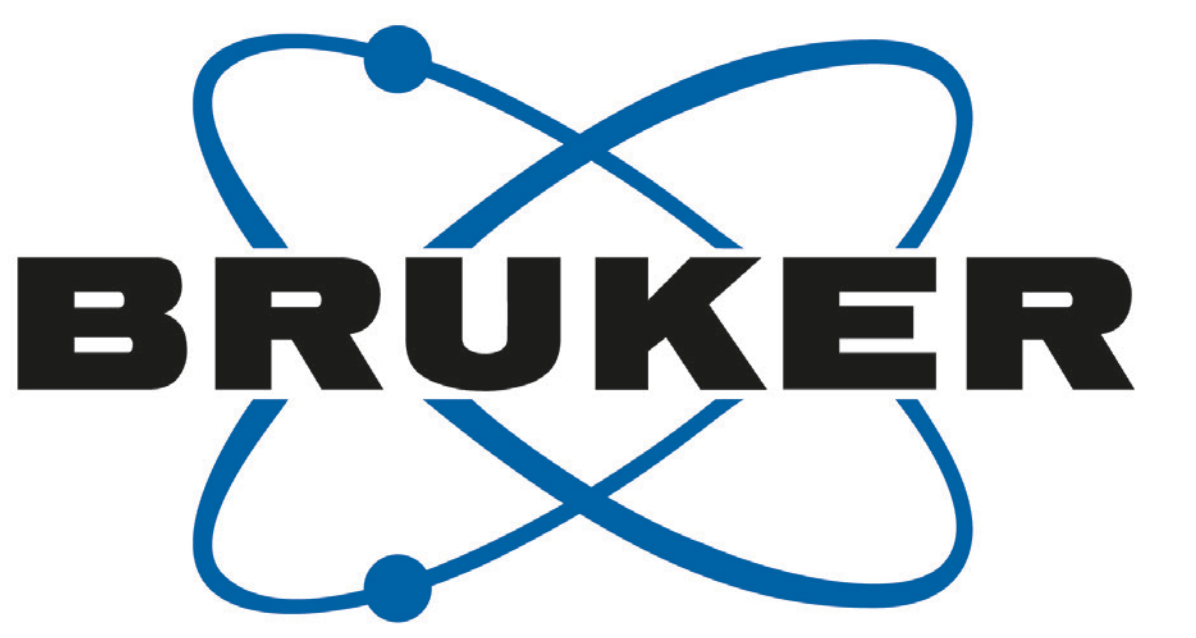
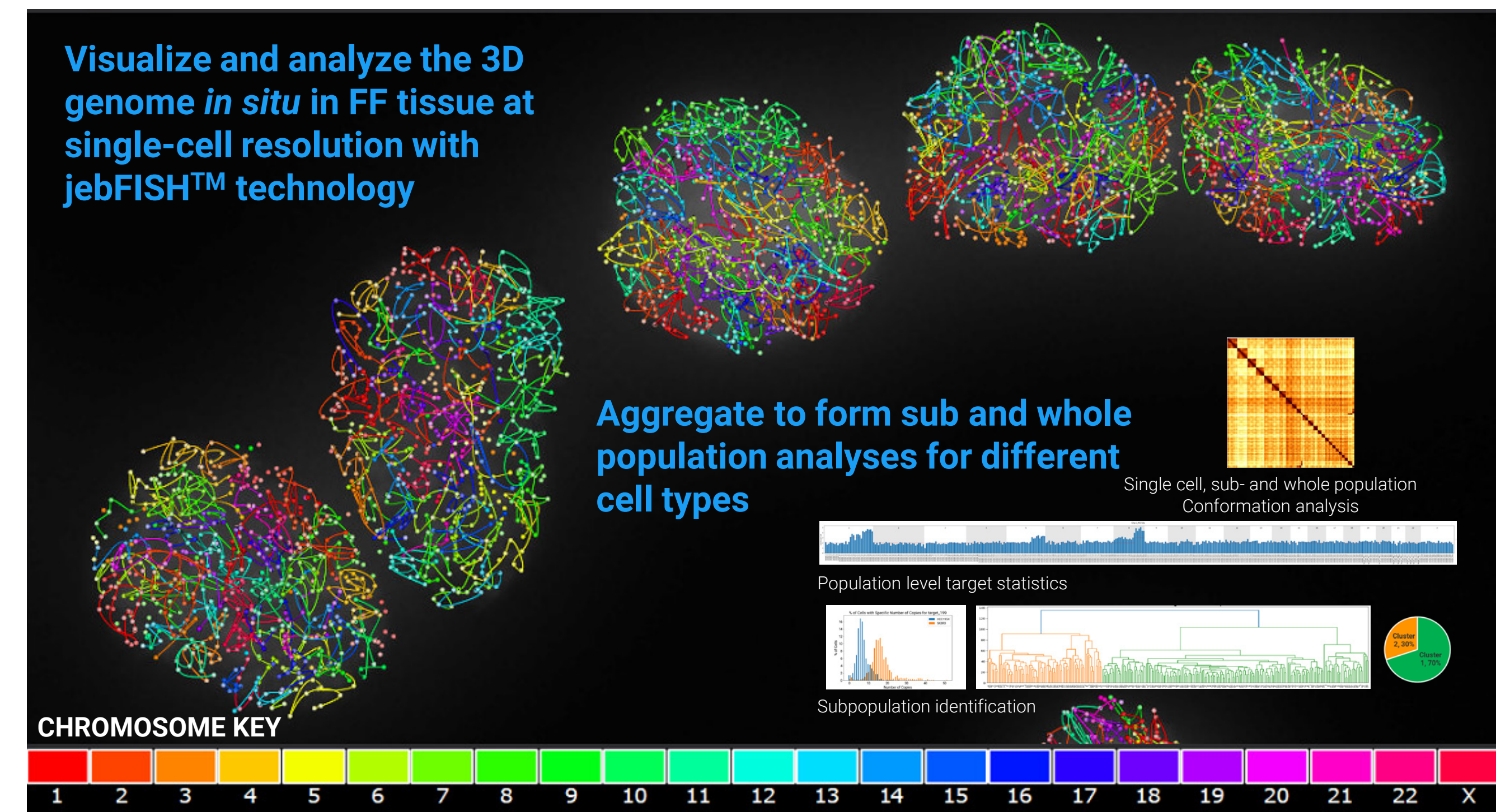


# PaintScape™ enables *in situ*, single cell spatial multiomic visualization of 3D genome organization in fresh frozen colorectal carcinoma tissue in spatially resolved tissue microenvironments



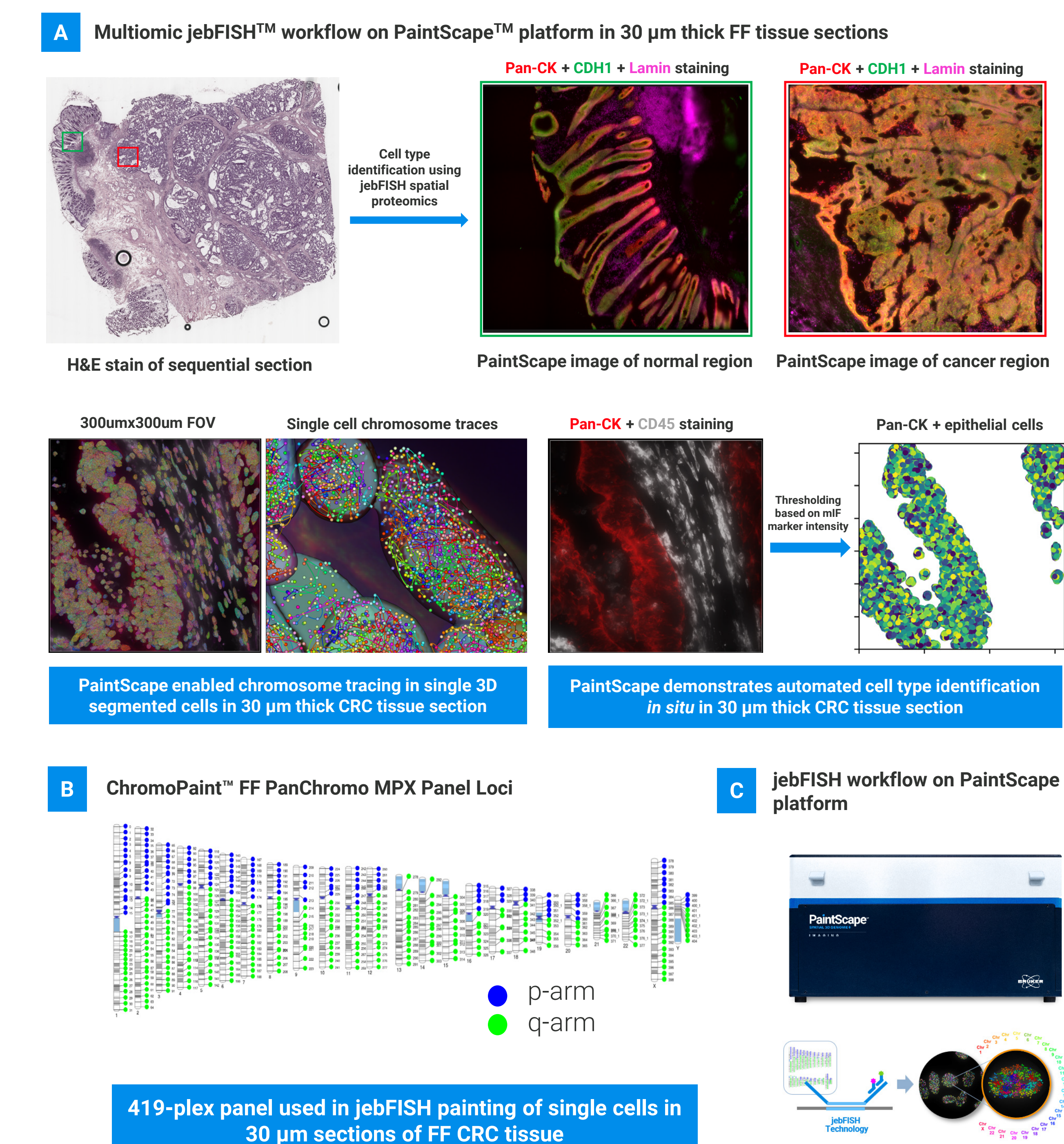
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## Introduction

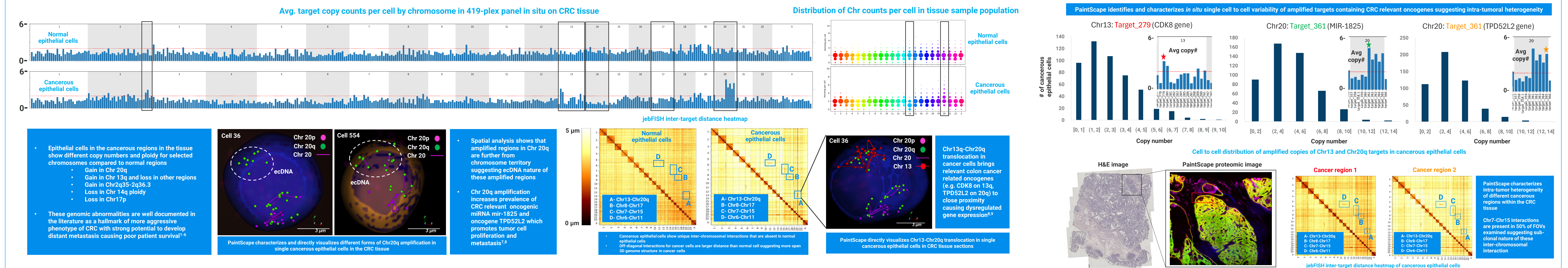


- Here, we present a novel multiomic jebFISH™ protocol on the PaintScape™ platform that can be used to characterize intratumoral heterogeneity of 3D chromatin architecture of fresh frozen human colon tissue from colorectal cancer (CRC) patient samples in different tissue microenvironments across different cell types and cell states at single cell, sub-population and population level.
- CRC is one of the most common cancers worldwide and a second leading cause of cancer mortality.<sup>1-6</sup>
- Tumor evolution and intra-tumor heterogeneity in CRC is fueled by chromosomal instability (CIN),<sup>1-5</sup> 3D genome rewiring and epigenetic dysregulation.<sup>6</sup>
- Many colon cancer cells show high CIN including gains in Chr8q, Chr 13q, Chr 20q and losses in Chr 8p, Chr17p, Chr14q.<sup>1-6</sup>
- Such selective copy gain/losses and rearrangement of those genomic regions cause dysregulation of oncogenic signaling pathways in a cell type dependent manner in the tumor microenvironment and increases cancer cell proliferation providing positive selection for cancer progression.

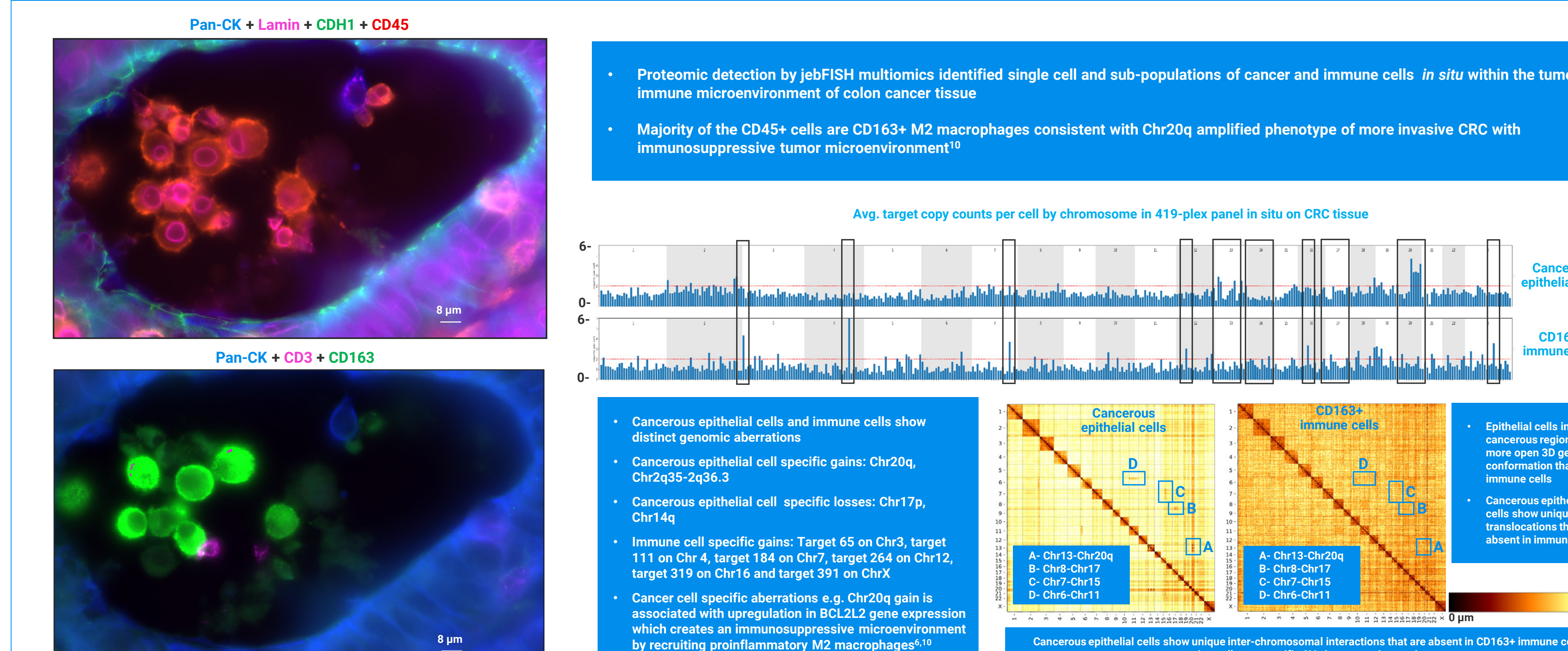
## Technology and Methods



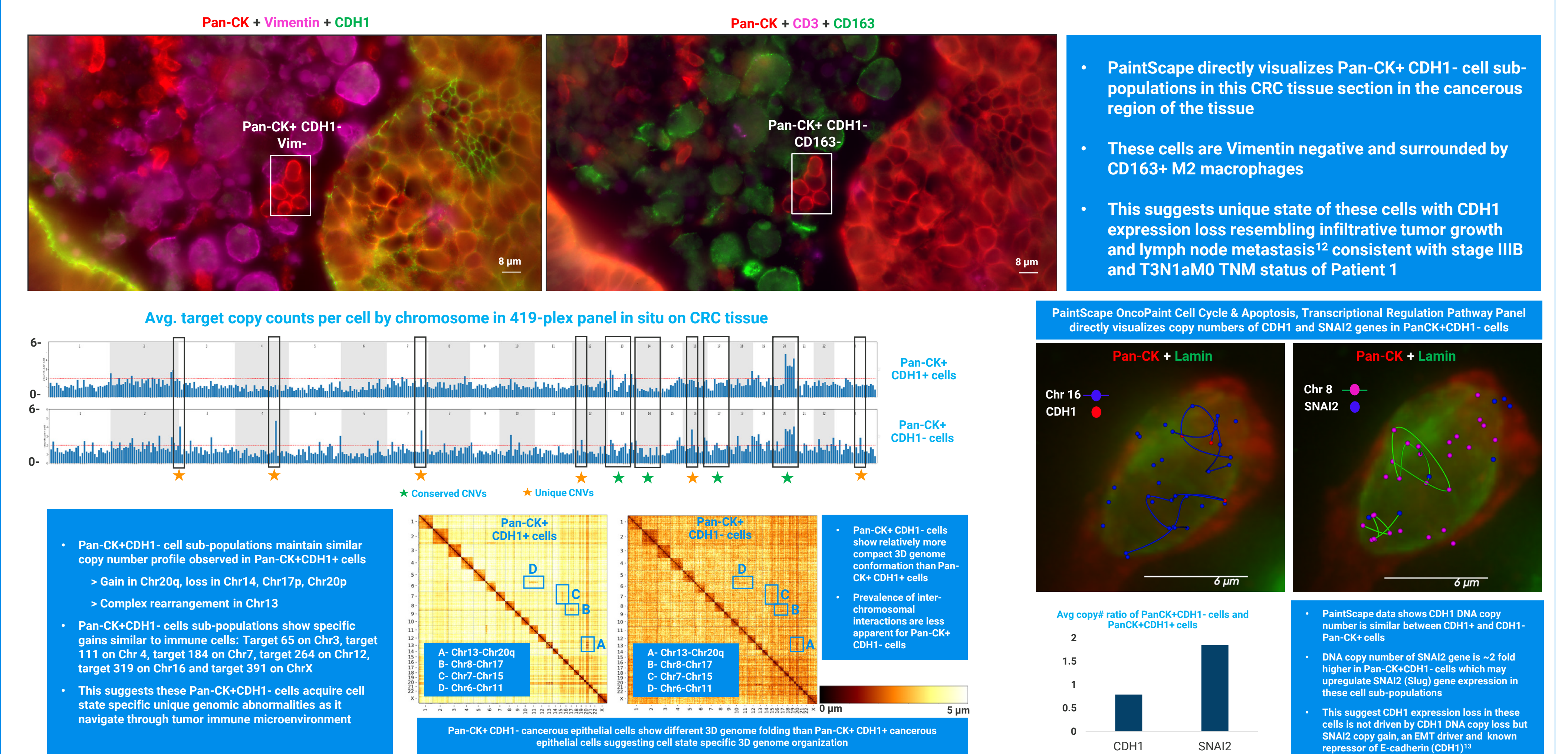
## PaintScape™ directly visualizes and characterizes *in situ*, single cell 3D genome structural genotype and conformational changes of normal and cancerous epithelial cells in 30 μm thick FF CRC tissue section



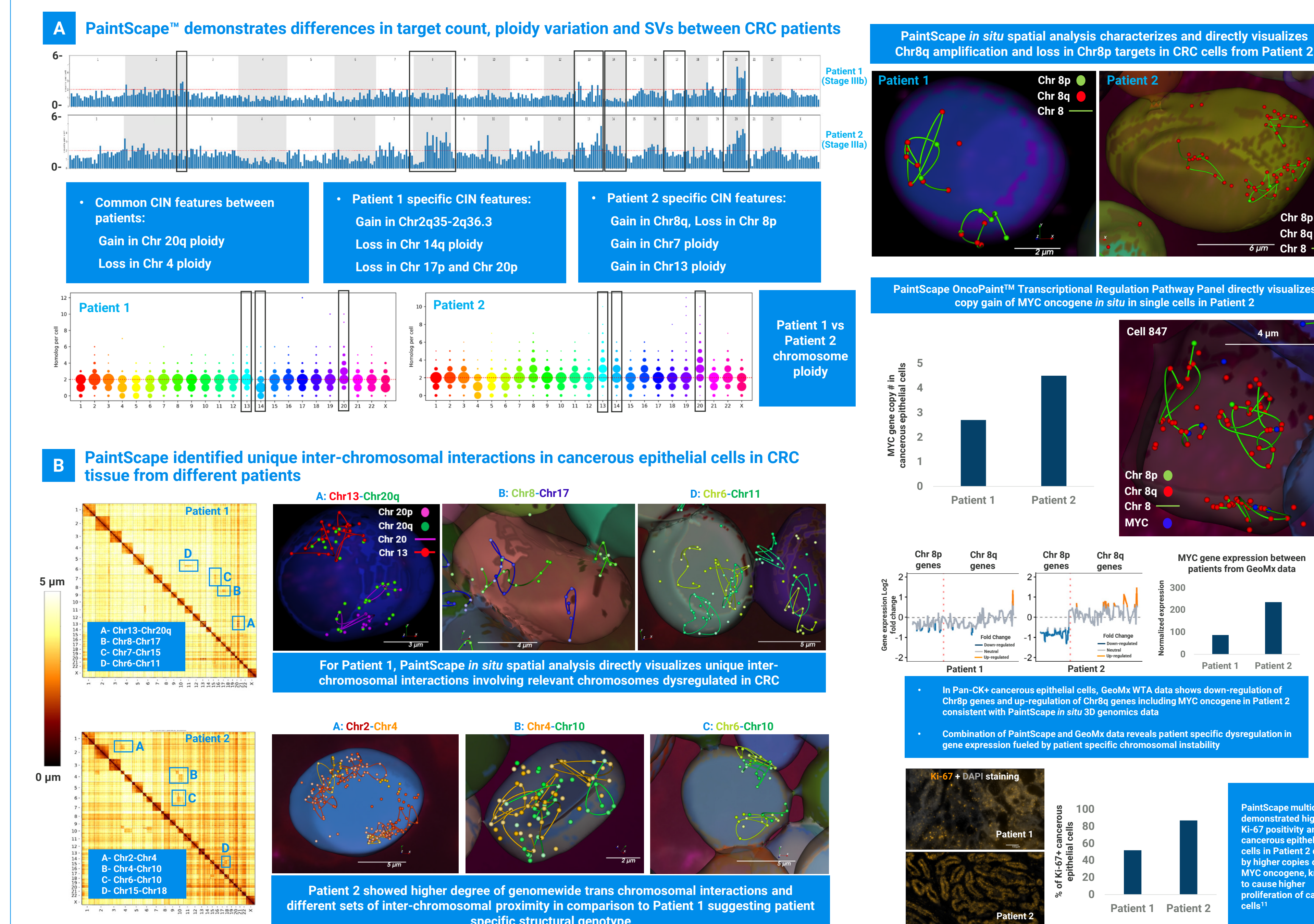
## PaintScape characterizes distinct 3D genome profiles across different cell types in tumor immune microenvironment (TIME)



## PaintScape multiomics identifies and characterizes distinct 3D genome profiles of unique cell states in tumor microenvironment of Patient 1 with stage IIIB CRC



## PaintScape characterizes chromosomal instability (CIN) of cancerous epithelial cells in CRC tissue from different patients with different tumor stage



## Conclusions

- PaintScape™ enables identification, characterization and direct *in situ* visualization of single cell 3D genome features of different cell types in FF Colorectal Cancer tissue across different patients. Distinct structural genotypes are revealed:
- Copy number variation including amplification and deletions *in situ* in single cells and sub-populations
    - Identified cancer vs normal cell sub-populations with
      - gain in Chr 8q, Chr 13q and Chr 20q in cancer cells
      - loss in Chr 8p, Chr 14q, Chr 17p and Chr 20p in cancer cells
    - These CIN patterns are suggestive of more aggressive phenotype of CRC with metastatic potential
  - In situ* single cell ploidy variation of whole chromosomes and sub-chromosomal regions
    - Higher variation in sub-population of cells with variable chromosome ploidy in cancer cells compared to normal cells
    - Chr8, Chr13q and Chr20 show much higher chromosome ploidy in cancer cell sub-populations
    - LOH of Chr14q and Chr4 in cancer cell sub-populations
  - Inter-chromosomal interactions, potential translocations and simultaneous proximity of multi-loci interactions *in situ* in single cells
    - For patient 1: Chr13q-Chr20q, Chr8-Chr17, Chr7-Chr15 and Chr6-Chr1 interactions
    - For patient 2: Chr13q-Chr20q translocation in cancer cells brings relevant CRC related oncogenes (e.g. CDK8 on 13q, TP53L2 on 20q) to close proximity, potentially causing their dysregulated gene expression
  - Cell type specific differences in 3D genome structure *in situ* in single cells in tissue immune microenvironment
    - Identified and visualized unique 3D genome aberration of individual chromosomes in Pan-CK+ epithelial cells and CD45+CD163+ immune cells
    - Immune cells show relatively more compact 3D genome structure than cancerous epithelial cells
    - Cancer cells in stage IIIB patient show sub-populations of unique cell state with epithelial nature but loss in CDH1 expression suggesting metastatic potential of these cells
  - Multimic correlations of cancer cell states between patients
    - Patient 2 with Chr8q gain shows much higher Ki-67 positivity rate for cancerous epithelial cell sub-populations compared to Patient 1 lacking Chr8q gain
    - Chr8q gain increases prevalence of key oncogene MYC which upregulates Ki-67 leading to higher proliferation and greater genome wide instability

## PaintScape™ enables *in situ* direct visualization of 3D genome in FF Colorectal Cancer tissue in spatially resolved tumor microenvironments

## References

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