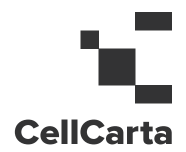
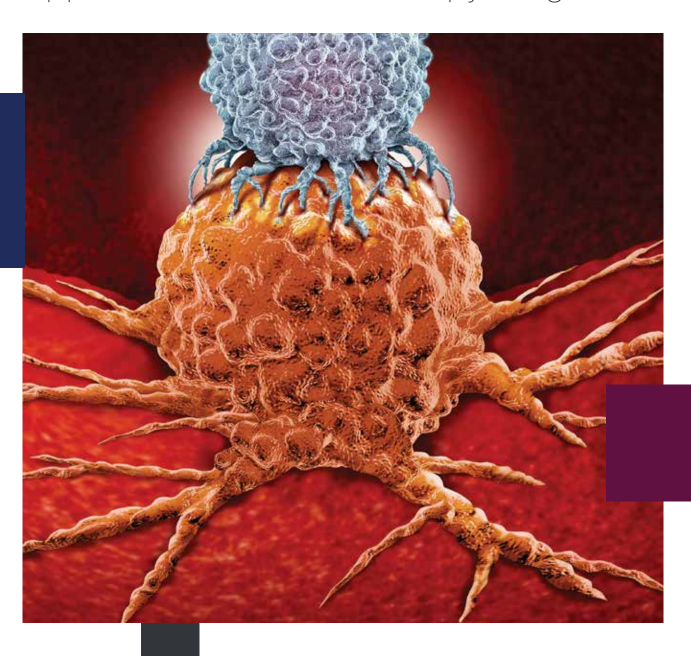
## Cell Therapy



Accelerate the Development and Approval of Your Cell Therapy Program



## Addressing the Challenges of Cell Therapy

Cell therapies have revolutionized treatment of some hematological malignancies, yet the newness of the technologies represents major challenges during their development and clinical testing. The diverse spectrum of cellular therapies includes CAR-T cell therapy and tumor-infiltrating lymphocytes (TIL) therapy, both FDA-approved, as well as engineered T cell receptor (TCR) therapy, and natural-killer (NK)-cell therapy. To navigate this complex and evolving field you need a scientific partner at the forefront of technologies.

At CellCarta, we collaborate closely with you and provide a critical edge to your studies, anticipating clinical testing challenges and providing a quick turnaround time essential to these studies.

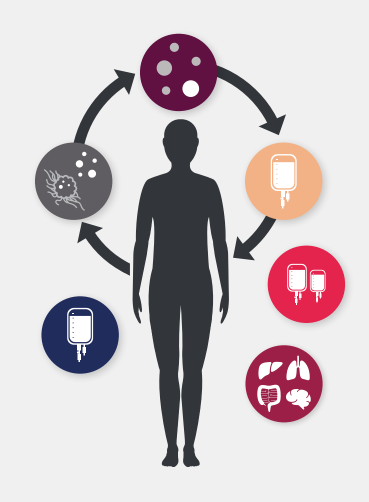
Our biomarker solutions and customized assays will give you with the insight you need to overcome cellular exhaustion and immunosuppressive tumor microenvironments (TME), enhance trafficking to solid tumors and much more.

QUALITY DATA TO MOVE YOUR CELL THERAPY PROGRAM FORWARD

# Providing a Strategic Advantage at Every Step with Our Cross-Platform Expertise

Your cell therapy is unique so it makes sense that your bioanalysis would be customized. CellCarta is a biomarker expert with +20 years of proficiency in developing customer specific bioanalytical solutions.

Benefit from our comprehensive range of platform technologies during all steps of clinical development, from **product** characterization, patient selection, to therapy and response monitoring, and toxicity assessment.



#### Achieving the Next Level in **Product Characterization**

Our genomics-based solutions, such as targeted RNA profiling, next-generation sequencing (NGS) and qPCR, can provide guidance on the numerous **genetic modifications (CAR or TCR constructs)** of your target cells.

At the cellular level, flow cytometry and CyTOF can be used to provide **surface profiling** of the engineered cells, and assess the expression levels of CAR constructs, immune checkpoints, as well as activation and exhaustion markers. Trucount<sup>™</sup> tubes are used to determine **absolute cell count**.

Our immune monitoring tools can provide deeper assessment of cellular functionality with antigen-specificity and cytokine expression monitored through intra-cellular cytokine staining (ICS). Multiplex cytokine response can also be measured using the Meso Scale Discovery® (MSD®) and the ELISpot/FluoroSpot platforms.

As the cellular expansion and function of the overall treatment can be influenced by the **clonal composition of the product**, we offer RNA sequencing (RNAseq) at the single-cell level to investigate clonal kinetics.

## Improving Patient Selection Using Complementary Platforms

Patient stratification is key in limiting toxicity and improving efficacy of treatment. CellCarta's **in-house team of board-certified pathologists** is a major asset to support patient stratification. The team provides their support to quantify new and established genetic biomarkers as well as key tumor targets in tissue samples.

Profiling the **general inflammatory state** of the patient as well as **lymphodepletion** status can support stratification and is achieved at the cellular level with inflammatory cytokine expression and TBNK enumeration.

Disease recurrence and development of cytokine release syndrome is correlated with tumor mutation burden (TMB) prior to treatment, which is why we offer **genetic biomarkers tracking** services, including mutational drivers.

During patient selection, the abundance, distribution, and localization of antigens of interest in tissues can be explored for **key tumor targets** such as MAGE4, NY-ESO-1, CD19, CD22, CD123, BCMA, using single or multiplex IHC. Some specific targets, such as **BCMA in multiple myeloma**, can be quantified in plasma samples as well as FFPE samples using mass spectrometry.

For patients **previously treated** prior to cellular therapy, CellCarta can monitor monoclonal antibody levels with ELISA.

## Seizing the Power of Biomarker Analysis to Monitor Your Product and Its Therapeutic Response

Precision biomarkers to evaluate therapeutic response can be monitored with our wide variety of genomics, proteomics, histopathological, and immune monitoring techniques.

Once your therapy is delivered, its progression can be tracked over time (cellular kinetics) with various genomic services such as single-cell RNA sequencing or with construct-specific antibodies by flow cytometry. Further immune profiling including complete surface profiling and absolute cell counts can be achieved in parallel with key biomarkers.

Our team can provide **critical biomarker information** on T-cell activation and exhaustion, antigen presentation, tumor immune infiltration, as well as characterization of the TME using multiplex or singleplex IHC.

CellCarta can **monitor the patient's response to treatment** by measuring remaining tumor
cells or tumor-specific targets. Some known
targets can be measured by mass
spectrometry (e.g. BCMA) or IHC with
quantification supported by our **in-house team of pathologists**. The reconstitution of patient's
immune system following treatment can be
monitored by simply using a TBNK panel by
flow cytometry.

## **Toxicity Assessment Simplified**

For multiplexed, high-sensitivity, quantitative measurement of key inflammatory cytokines and chemokines, CellCarta offers the MSD® and the ELISA platforms. Our multiplex MSD® panels are readily available and aid in the screening of cytokine release syndrome.

## **Evaluating Combination Therapy**

We monitor **critical biomarkers of efficacy** with high-throughput assays (MSD® and ELISA platforms) for sensitive and precise evaluation of combination therapies and their impact on overall treatment efficacy.

#### **Partner with CellCarta**

#### to Accelerate the Development and Approval of Your Cell Therapy Program

- An established track record, with over 150 completed and ongoing projects in cell therapy
- Unparalleled scientific expertise with cross-platform capabilities your onestop-shop for expertise in cell therapy
- Customized solutions. Every project is unique, and we have the flexibility and agility to adapt to your needs
- Highest standards in quality with CAP accreditations/CLIA certifications and validation processes that meet the requirements for primary/secondary endpoints
- Providing full data insights with Al-driven analysis and biological interpretation support
- Global presence with ten facilities located in Canada, USA, Belgium, Australia, and China



#### **Clinical Development Steps**

#### **Platforms**

Product Characterization	
Genomic profiling of target cells	<ul> <li>Targeted RNA profiling</li> <li>NGS –TCR sequencing</li> <li>qPCR</li> <li>Single-cell RNA sequencing</li> </ul>
Cellular profiling  ■ Absolute cell count (Trucount™ tubes)  ■ Off-the-shelf panels including key markers (activation/exhaustion, immune checkpoints)  ■ Customized construct-specific panel  ■ Functionality by ICS  ■ in vitro antigen-specific functionality by cytokine response	■ Flow cytometry ■ CyTOF ■ MSD® ■ ELISpot/FluoroSpot
Patient Selection	
Characterization of patients' inflammatory state Immune profiling – general panels Lymphodepletion – TBNK panel Cytokine profiling	■ Flow cytometry ■ MSD®
Tracking genetic biomarkers ■ Key patient mutations ■ Key tumor biomarkers ■ Microsatellite instability (MSI) ■ Tumor mutational burden (TMB)	<ul><li>■ qPCR</li><li>■ NGS (TSO500 panel by Illumina)</li><li>■ Targeted RNA profiling</li></ul>
Monitoring tumor targets ■ MAGE4, NY-ESO1, CD19, CD22, CD123, BCMA ■ soluble and FFPE BCMA	<ul><li>Immunohistochemistry (IHC)</li><li>Mass Spectrometry</li></ul>
Monitoring previous treatments ■ Detection of rituximab	■ ELISA
Monitoring Product and Therapeutic Response	
Tracking transgene or engineered cell	<ul> <li>Targeted RNA profiling</li> <li>NGS – TCR sequencing</li> <li>qPCR – preceded or not by cell sorting (FACS)</li> <li>Single-cell RNA sequencing</li> </ul>
Immune profiling / cellular kinetics  ■ Absolute cell counts (proliferation panels - TBNK)  ■ Complete surface profiling - including immune checkpoints (PD-1, TIM-3, LAG-3), activation/exhaustion markers  ■ ICS assays  ■ Tissue biomarkers for T-cell activation/exhaustion, antigen presentation, immune infiltration, TME	■ Flow cytometry ■ single-plex and multiplex IHC
Tracking remaining tumor cells or tumor-specific targets	<ul> <li>single-plex and multiplex IHC</li> <li>Mass spectrometry (BCMA)</li> <li>Flow cytometry</li> </ul>
Toxicity Assessment	
Measurement of cytokine release syndrome	■ MSD® ■ ELISA
Combination Therapy	
PK of other biologics	■ MSD® ■ ELISA

For more information on how CellCarta can partner with you, please contact us:

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