# Merus

## **Closing In On Cancer**



## **Disclaimer**

This presentation (including any oral commentary that accompanies this presentation) contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995. All statements contained in this presentation that do not relate to matters of historical fact should be considered forward-looking statements, including without limitation statements regarding the impact our Biclonics® platform can have on cancer, our product candidates' potential to treat certain types of tumors, the timing of regulatory filings and the timing and anticipated data read outs or results from our clinical trials and our collaborations. These forward-looking statements are based on management's current expectations. These statements are neither promises nor guarantees, but involve known and unknown risks, uncertainties and other important factors that may cause our actual results, performance or achievements to be materially different from any future results, performance or achievements expressed or implied by the forward-looking statements, including, but not limited to, the following: we have incurred significant losses, are not currently profitable and may never become profitable; our need for additional funding, which may not be available and which may require us to restrict out operations or require us to relinquish rights to our technologies or bispecific antibody candidates; potential delays in regulatory approval, which would impact our ability to commercialize our product candidates and affect our ability to generate revenue; the unproven approach to therapeutic intervention of our Biclonics® technology; our limited operating history; economic, political, regulatory and other risks involved with international operations; the lengthy and expensive process of clinical drug development, which has an uncertain outcome; the unpredictable nature of our

early stage development efforts for marketable drugs; potential adverse public reaction to the use of cancer immunotherapies; potential delays in enrollment of patients, which could affect the receipt of necessary regulatory approvals; failure to obtain marketing approval internationally; failure to compete successfully against other drug companies; potential competition from other drug companies if we fail to obtain orphan drug designation or maintain orphan drug exclusivity for our products; our reliance on third parties to conduct our clinical trials and the potential for those third parties to not perform satisfactorily; our reliance on third parties to manufacture our product candidates, which may delay, prevent or impair our development and commercialization efforts; protection of our proprietary technology; our patents being found invalid or unenforceable; potential lawsuits for infringement of third-party intellectual property; our ability to attract and retain key personnel; managing our growth could result in difficulties; and we may lose our foreign private issuer status and incur significant expenses as a result.

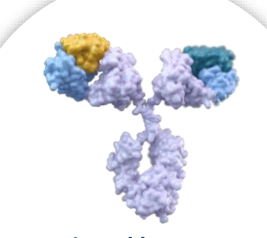
These and other important factors discussed under the caption "Risk Factors" in our Annual Report on Form 20-F filed with the Securities and Exchange Commission, or SEC, on April 3, 2019, and our other reports filed with the SEC, could cause actual results to differ materially from those indicated by the forward-looking statements made in this presentation. Any such forward-looking statements represent management's estimates as of the date of this presentation. While we may elect to update such forward-looking statements at some point in the future, we disclaim any obligation to do so, even if subsequent events cause our views to change.



## Merus: pioneering bispecific antibodies since 2006

4 clinical-stage bispecific antibodies in oncology

Multiple clinical milestones anticipated in the next 12 months



Sophisticated proprietary bispecific and trispecific technology platforms

Designed by nature. Selected by Merus.

Discovery of novel modes of action based on target combinations and functional screening

Fully integrated discovery-to-manufacturing bi-/tri-specific technology platforms



# Merus clinical pipeline, near term milestones and program status

PROGRAM	<b>TARGETS</b>	INDICATION / DRUG COMBINATION	PRE-IND	PHASE 1	PHASE 2	MILESTONE/STATUS
NACI A 139	LIEDZ VIJEDZ	NRG1 Solid tumors (monotherapy)				Update expected around end of 2020
MCLA-128	HER3 x HER2	Metastatic Breast (combination in 2 cohorts)				Phase 2 Results in 2020
MCLA-117	CLEC12A x CD3	Acute Myeloid Leukemia (AML)				Initial Data 1H 2020
MCLA-158	Lgr5 x EGFR	Solid tumors				Phase 1 Trial Ongoing
MCLA-145	CD137 x PD-L1	Solid tumors (ex- U.S.)				Phase 1 Trial Ongoing
MCLA-129	EGFR x c-MET	Solid tumors (China)				IND Enabling Studies Ongoing
ONO-4685	PD-1 x CD3	Autoimmune disease Ono				Phase 1 Trial Ongoing
	Undisclosed	Autoimmune disease Ono				



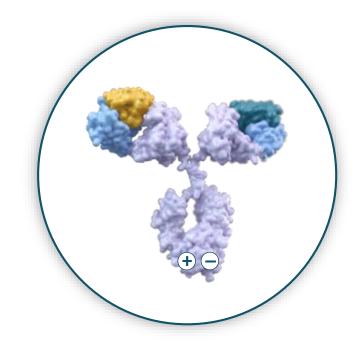
# Biclonics® — designed to look and perform like natural human antibodies

BICLONICS® - leveraging the attractive characteristics of natural antibodies Merus' Bispecific Antibodies are produced by a single cell

Common Light Chain

for 'unforced', natural pairing with 2 different heavy chains

to efficiently drive formation of Biclonics®



**IgG** Format

for efficient manufacturing <u>and</u> predictable *in vivo* behavior

Fc Modifications
for Improved functionality
(ADCC or silencing)



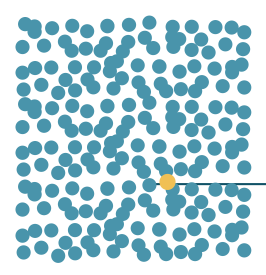
# Biclonics® — Selected in functional assays for differentiated activity

## HUMAN ANTIBODY GENERATION



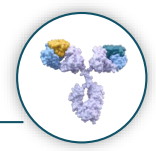
We use a proprietary transgenic mouse to generate panels of high-quality human antibodies

#### **PANEL GENERATION**



We can create up to 1,000 Biclonics® against any target pair of choice

#### **FUNCTIONAL SCREENING**





We use large-scale functional screening in cell-based assays to identify Biclonics® with novel modes of action



# Clinical Programs MCLA-128

**HER2 x HER3 Biclonics®** 



# Blocks primary tumor cell growth and escape to HER2/EGFR targeted therapy

## Unique DOCK & BLOCK® mechanism of action potently inhibits neuregulin (NRG)-driven tumor growth

## **Block HER3**

Blocks HER3 signaling even in high neuregulin tumor environments

## Enhanced ADCC

Mediates tumor elimination by immune killer cells

# KILL TUMOR CELLS Tumor

## **Dock HER2**

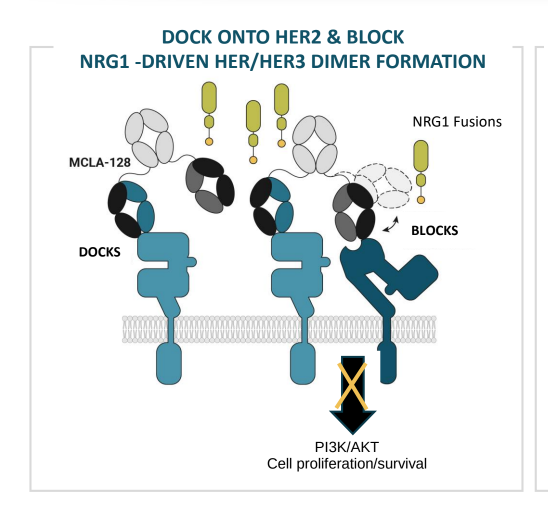
Docks to HER2 expressed on tumor cells to efficiently block HER2:HER3 dimer formation

## Combination

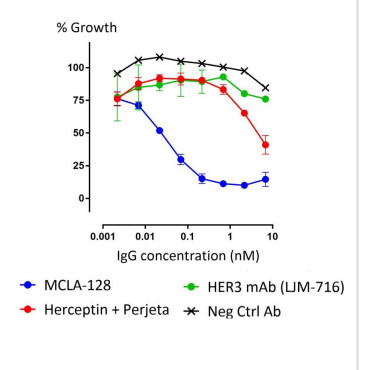
Combinations with HER2 targeted therapies possible



## Preclinical efficacy demonstrated, well-tolerated in patients



## SUPERIOR ACTIVITY SHOWN IN PRECLINICAL MODELS



## SAFETY AND TOLERABILITY IN PHASE 1/2 TRIAL

#### 117 PATIENTS EVALUATED\*

MCLA-128 Dosing: 750 mg ranging from q1w-q3w

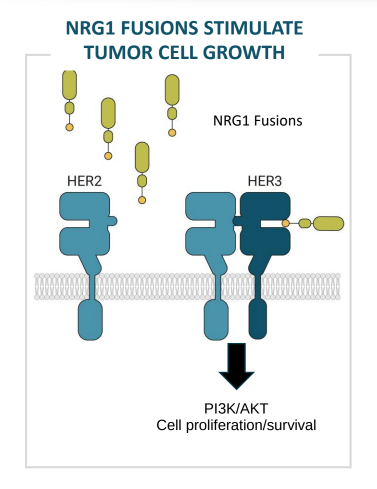
- Single agent well tolerated
- Low risk for immunogenicity

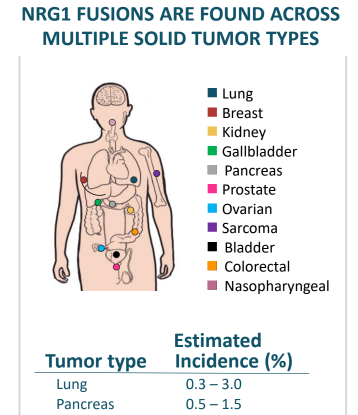
Please Refer to <a href="https://merus.nl/publications/">https://merus.nl/publications/</a> for full data presented. Refer to ASCO poster 2018 and AACR-NCI-EORTC International Conference on Molecular Targets and Cancer Therapeutics 2019

\* As of Jan. 2019

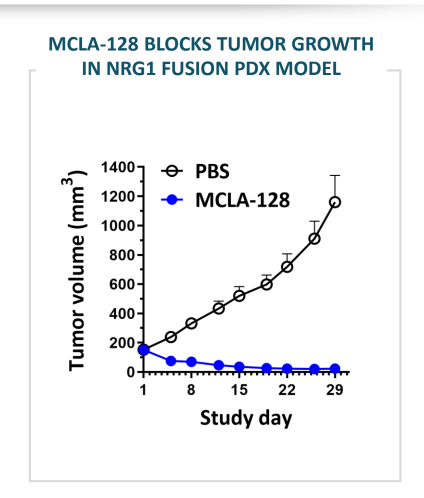


## Novel therapeutic paradigm for NRG1 Fusion cancers





Other



Projections of NRG1 Fusions occurrence (incidence) are based on limited published information

< 1.0

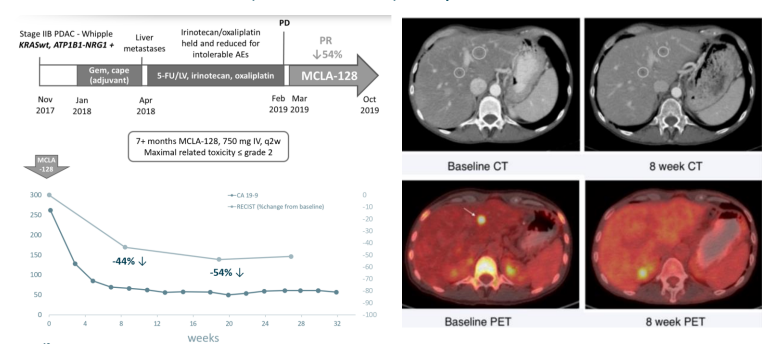


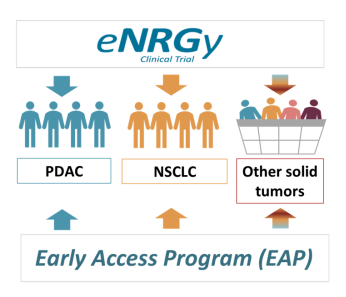
## Active in NRG1 fusion<sup>POS</sup> cancers

## **Clinical Proof of Concept Established**

## **Clinical Trial Ongoing**

PDAC (ATP1B1-NRG1): 52-year-old male\*







<sup>\*</sup> Data was presented on October 27, 2019, at the AACR-NCI-EORTC International Conference on Molecular Targets and Cancer Therapeutics by investigators from the Memorial Sloan Kettering Cancer Center in a presentation titled "Clinical proof-of-concept for MCLA-128, a bispecific HER2/3 antibody therapy, in NRG1 fusion-positive cancers"

**MCLA-128** 

# Phase 1/2 in NRG1 solid tumors, Phase 2 in metastatic breast cancer (combo)

WHOLLY-OWNED	TARGETS	INDICATION / DRUG COMBINATION	PRE-IND	PHASE 1	PHASE 2
NACI A 120	LIEDZ VIJEDZ	NRG1 Solid tumors (monotherapy)			
MCLA-128	HER3 x HER2	Metastatic Breast (2 cohorts)			
MRUS Fully Owned					

	DESIGN	ENDPOINTS	STATUS	
NRG1 Solid Tumors (Monotherapy)	Phase 1/2 Study Cohort 1: NRG1+ Pancreatic cancer Cohort 2: NRG1+ NSCLC Cohort 3: NRG1+ Other solid tumors Dose: 750mg every 2 weeks	<ul><li>Safety</li><li>Anti-tumor activity</li></ul>	Enrollment ongoing	
Metastatic Breast Cancer (MBC)	Phase 2 Study in combination with 2 cohorts in MBC Cohort 1: HER2+ (MCLA-128 + Herceptin + Chemo) Cohort 2: ER+/HER2 <sup>low</sup> (MCLA-128 + Hormone Therapy) Dose: 750mg every 3 weeks	<ul> <li>Clinical benefit at 24 weeks</li> </ul>	<ul> <li>To be presented at a medical conference in 2020</li> </ul>	



# Clinical Programs MCLA-117

CD3 x CLEC12A Biclonics®



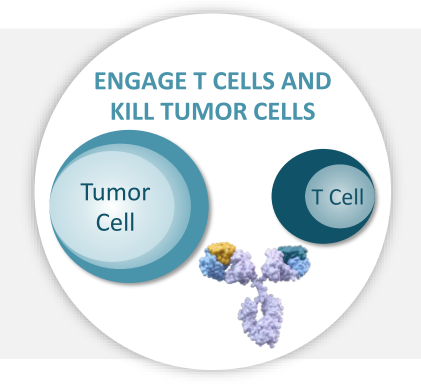
# T cell engager for acute AML addressing a potential first-in-class target

## MCLA-117 efficiently activates and redirects T cells to kill CLEC12A-expressing acute myeloid leukemia (stem) cells

#### CLEC12A

Expressed by tumor (stem) cells in ~ 90-95% of AML patients

Expression restricted to hematopoietic system = potential less off-tumor toxicity



CD3

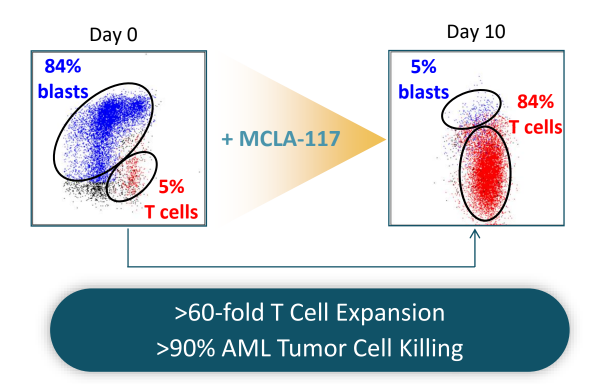
Low affinity CD3 arm and silenced Fc for controlled T cell activation to reduce toxicity and improve biodistribution



## Potent activation of T cells in preclinical studies

- MCLA-117 efficiently activates and redirects T cells to kill CLEC12A-expressing AML(stem) cells
- Designed to spare the formation of platelets and red blood cells during therapy
- Active across AML patient subpopulations

## MCLA-117 MEDIATED ACTIVATION OF T CELLS AND KILLING OF TUMOR CELLS





## **Phase 1 Trial**



DESIGN	ENDPOINTS	STATUS		
Single-arm, open-label, dose escalation w/ safety dose expansion	<ul> <li>Primary Endpoints: safety, tolerability</li> </ul>	<ul><li>Ongoing in Europe and the U.S.</li><li>Preliminary anti-tumor activity has</li></ul>		
<ul> <li>Up to 50 patients with relapsed / refractory AML</li> </ul>	<ul> <li>Secondary Endpoints: PK/PD, anti- tumor response, clinical benefit</li> </ul>	<ul><li>been observed</li><li>Initial data expected at medical</li></ul>		
<ul> <li>Starting dose determined using MABEL dose escalation requirements</li> </ul>		conference 1H 2020		
<ul> <li>Protocol amended July 2019 to allow for the exploration of higher doses</li> </ul>				



## Clinical Programs MCLA-158

Lgr5 x EGFR Biclonics®



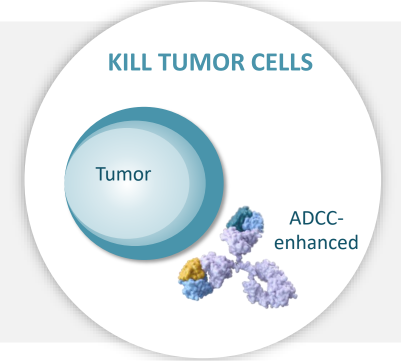
# Potently blocks EGFR signaling in Wnt dysregulated solid tumors

## Potential to be first colorectal cancer treatment to block growth of tumors with RAS mutations (~50% of patients), a high unmet need

## Lgr5

Expressed by intestinal cancer initiating cells

Identified through Merus functional screening and organoid discovery methods



#### **EGFR**

Blocks growth in Wnt dysregulated tumors including RAS<sup>mut</sup>

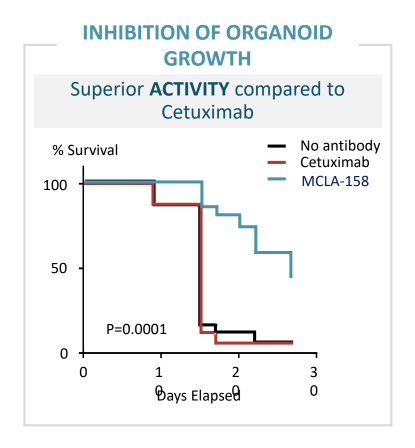
Preclinical data shows higher potency than Cetuximab

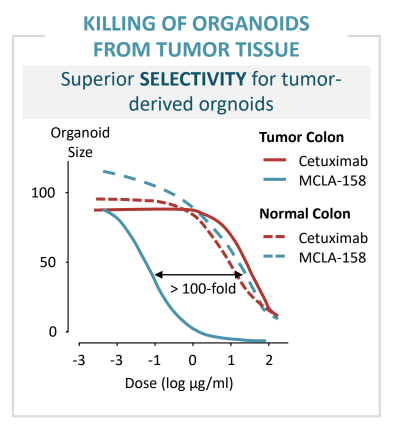


MCLA-158

# Superior growth inhibition and selectivity of tumor versus healthy tissue

- MCLA-158 is active in xenograft models that are resistant to treatment with Cetuximab
- Unlike Cetuximab, MCLA-158 discriminates between organoids derived from tumor and healthy tissue







## **Phase 1 Trial**

	TARGETS	INDICATION / DRUG COMBINATION	PRE-IND	PHASE 1	PHASE 2
MCLA-158	Lgr5 x EGFR	Solid tumors			
MRUS Fully Owned					

DESIGN	ENDPOINTS	STATUS
<ul> <li>Global open-label, multicenter dose escalation w/ safety dose expansion phase</li> <li>Patients with solid tumors</li> <li>Initial focus on metastatic colorectal cancer</li> </ul>	<ul> <li>Primary endpoint: safety and tolerability of defined dose</li> <li>Secondary endpoint: single-agent preliminary anti-tumor activity</li> </ul>	<ul> <li>Dose escalation is ongoing.</li> <li>Amended protocol to allow for the exploration of higher dose cohorts.</li> <li>Acceptable safety profile with no observed dose limiting toxicities to date.</li> </ul>



## Clinical Programs MCLA-145

CD137 x PD-L1 Biclonics®

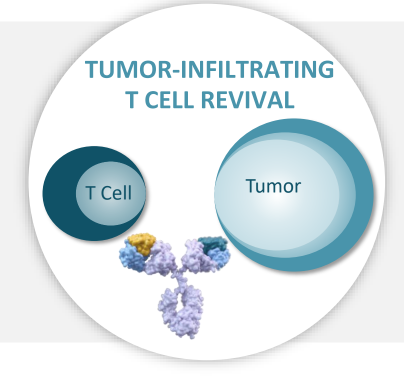


# Recruit, activate and prevent exhaustion of tumor-infiltrating T cells

## Triple action for potent and durable T cell activation in the tumor micro-environment

#### CD137

Potent activation of tumor infiltrating T cells dependent on PD-L1 expressing tumor cells and macrophages



#### PD-L1

Targeting to PD-L1 positive cells in the tumor

Blocking of T cell inhibitory PD-1/ PD-L1 interactions in tumor

Attracting T cells into the tumor

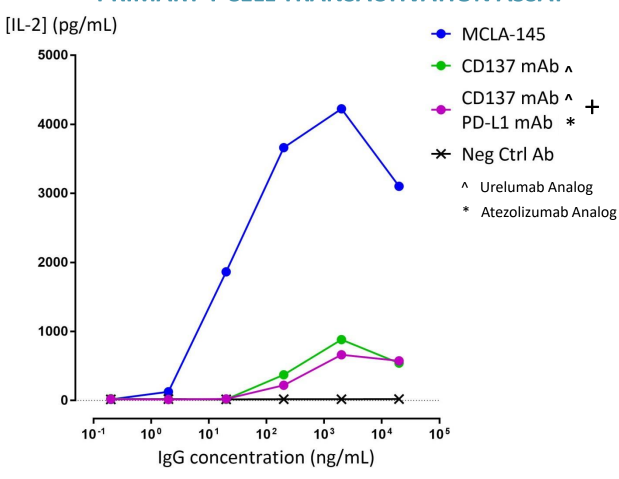


## **Demonstrated Potent T Cell Activation**

- Binds to PD-L1 and CD137
- Preclinical work demonstrates
  - recruitment of T cells into the tumor
  - blocking of inhibitory PD-1/PD-L1 axis
  - potent T cell activation
- Potential to overcome the known side effects of CD137 agonists in development

MCLA-145 preclinical data presented at AACR 2019

#### PRIMARY T CELL TRANSACTIVATION ASSAY





## Phase 1 trial initiated in May 2019

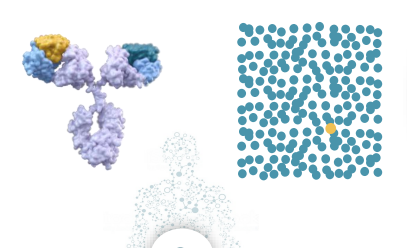
	TARGETS	INDICATION / DRUG COMBINATION	PRE-IND	PHASE 1	PHASE 2
MCLA-145	CD137 x PD-L1	Solid tumors			
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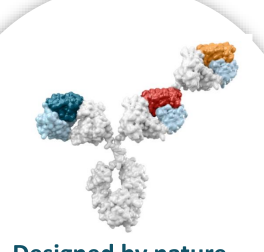
DESIGN	ENDPOINTS	STATUS
Global open-label, multicenter dose escalation with dose expansion phase  • Patients with advanced solid tumors	<ul> <li>Primary endpoint: dose finding, safety and tolerability</li> <li>Secondary endpoint: single-agent preliminary activity</li> </ul>	<ul> <li>IND cleared January 2019</li> <li>First patient dosed May 9, 2019</li> </ul>



# The new Triclonics™ platform for additional differentiated modes of action

## The BICLONICS® Beginning Our existing foundation...





Designed by nature. Improved by Merus.

## Next Generation TRICLONICS™ Platform for 2 or 3 different targets



Common light chain for unforced pairing with 3 (different) V<sub>H</sub> regions



Linker diversity for added functionality

1:1:1 or 2:1 format for new biology/modes of action



# Merus

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