Innovating Antibodies, Improving Lives

Investor Presentation August 2020



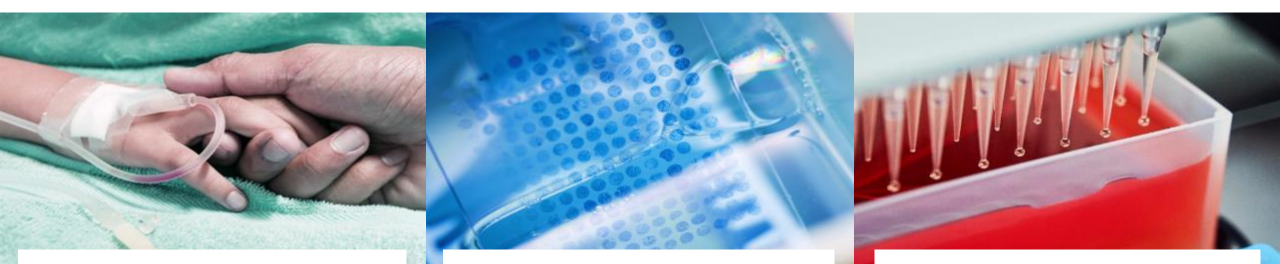


Forward Looking Statement

This presentation contains forward looking statements. The words "believe", "expect", "anticipate", "intend" and "plan" and similar expressions identify forward looking statements. All statements other than statements of historical facts included in this presentation, including, without limitation, those regarding our financial position, business strategy, plans and objectives of management for future operations (including development plans and objectives relating to our products), are forward looking statements. Such forward looking statements involve known and unknown risks, uncertainties and other factors which may cause our actual results, performance or achievements to be materially different from any future results, performance or achievements expressed or implied by such forward looking statements. Such forward looking statements are based on numerous assumptions regarding our present and future business strategies and the environment in which we will operate in the future. The important factors that could cause our actual results, performance or achievements to differ materially from those in the forward looking statements include, among others, risks associated with product discovery and development, uncertainties related to the outcome of clinical trials, slower than expected rates of patient recruitment, unforeseen safety issues resulting from the administration of our products in patients, uncertainties related to product manufacturing, the lack of market acceptance of our products, our inability to manage growth, the competitive environment in relation to our business area and markets, our inability to attract and retain suitably qualified personnel, the unenforceability or lack of protection of our patents and proprietary rights, our relationships with affiliated entities, changes and developments in technology which may render our products obsolete, and other factors. Further, certain forward looking statements are based upon assumptions of future events which may not prove to be accurate. The forward looking statements in this document speak only as at the date of this presentation. Genmab does not undertake any obligation to update or revise forward looking statements in this presentation nor to confirm such statements to reflect subsequent events or circumstances after the date made or in relation to actual results, unless required by law.



Our Core Purpose, Strategy & Vision Guide Our Work



Core Purpose

To improve the lives of patients by creating & developing innovative antibody products

Our Strategy

Turn science into medicine

Build a profitable & successful biotech

Focus on Core Competence

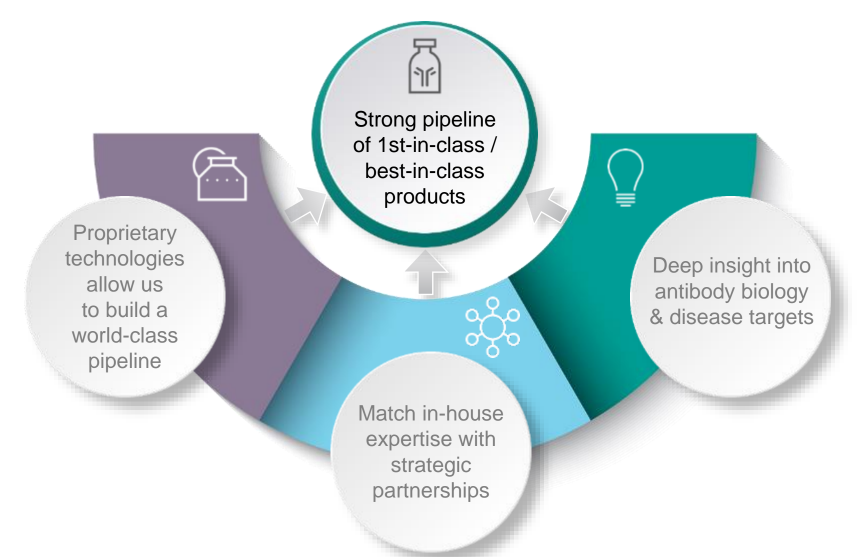
Vision

By 2025, our own product has transformed cancer treatment and we have a pipeline of knock-yoursocks off antibodies

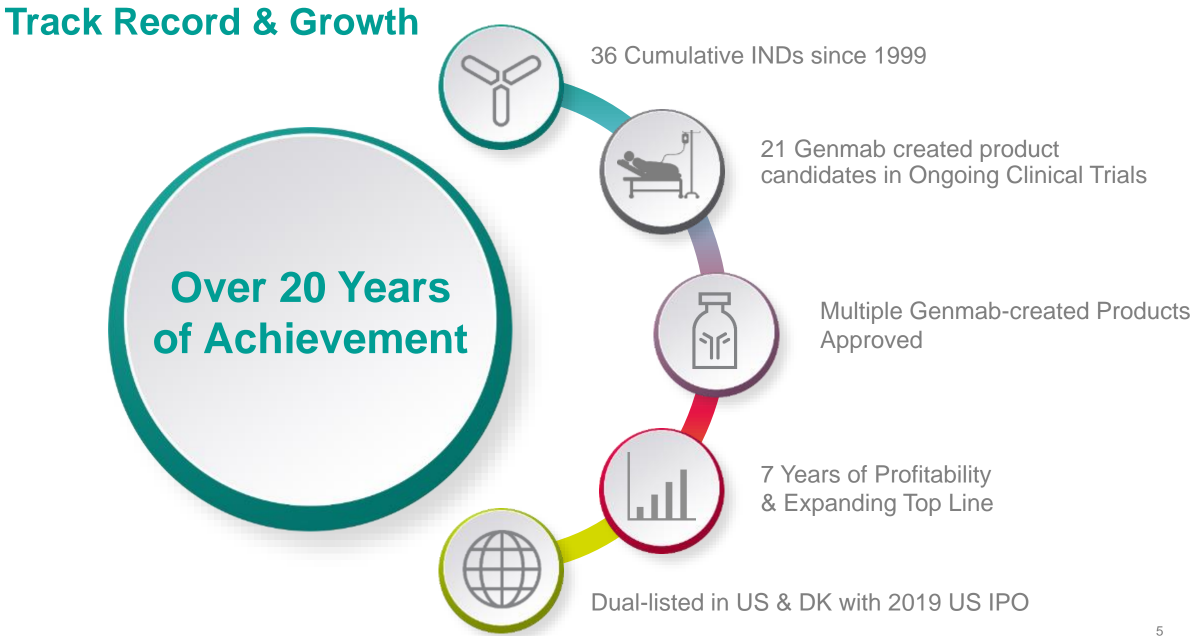


The Genmab Difference

Innovation Powerhouse Transforming Cancer Treatment & Creating Value









Solid Foundation Built on a Differentiated Pipeline

Potential 1st-in-Class/Best-in-Class

Our Own Clinical Pipeline

Tisotumab Vedotin⁴
Enapotamab Vedotin

HexaBody[®]-DR5/DR5

- DuoBody-CD40x4-1BB⁶
- DuoBody-PD-L1x4-1BB⁶
- DuoHexaBody[®]-CD37⁵
- Epcoritamab (DuoBody[®]-CD3xCD20)⁵
- DuoBody-CD3x5T4⁵

Solid Financial Base

Approved Partnered Products

- •DARZALEX[®] (daratumumab) / DARZALEX *FASPRO*[™] (daratumumab and hyaluronidase-fihj)¹
- •Kesimpta[®] (ofatumumab)²
- •TEPEZZA® (teprotumumab)³

Programs Built on Genmab's Innovation

Partner-owned Programs in the Clinic

- 11 product candidates in clinical development w/ partners
- Incl. 6 DuoBody products with Janssen, 1 with Novo Nordisk

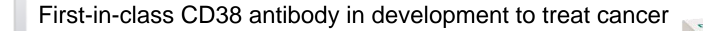
R&D Engine

Technologies & Pre-Clinical

- DuoBody technology
- HexaBody technology
- HexElect[®] technology
- DuoHexaBody[®] technology
- Rich Pre-Clinical Pipeline incl. HexaBody-CD387



DARZALEX[®] (daratumumab) & DARZALEX *FASPRO*[™] (daratumumab and hyaluronidase-fihj): Redefining Treatment of Multiple Myeloma



Collaboration with Janssen: Genmab entitled to tiered royalty of 12-20% of net sales

ADC 57894-502-20 RX ONY DARZALEX (daratumumab) Injection 400 mg/20 mL (20 mg/mL) For Intravenous Infusion Only



Approved in certain territories for various multiple myeloma (MM) indications¹

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DARZALEX FASPRO first and only SC CD38 mAb approved in U.S. for treatment of MM



RZALEX



2019 WW net sales by J&J: \$2,998M

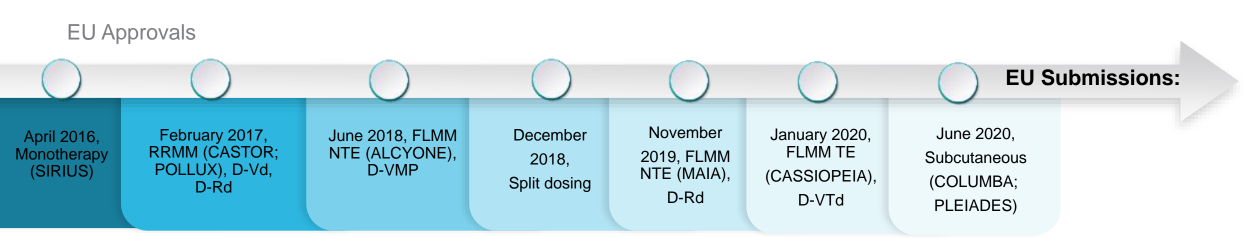


DARZALEX Approvals: US and EU

On Track for Approval Across All Lines of MM Treatment

US Approvals

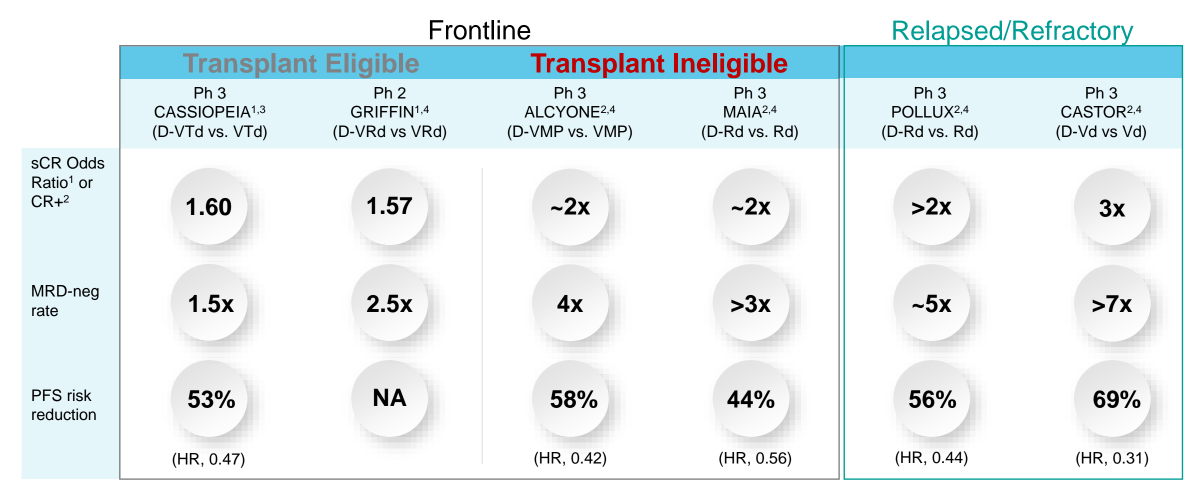
\bigcirc	\bigcirc	\bigcirc	\bigcirc	\bigcirc	\bigcirc	\bigcirc	U	S Submissions:
November 2015, Monotherapy (SIRIUS)	November 2016, RRMM (CASTOR; POLLUX), D-Vd, D-Rd	June 2017, RRMM (EQUULEUS), D-Pd	May 2018, FLMM NTE (ALCYONE), D-VMP	February 2019, Split dosing	June 2019, FLMM NTE (MAIA), D-Rd	September 2019, FLMM TE (CASSIOPEIA), D-VTd	May 2020, DARZALEX FASPRO (COLUMBA; PLEIADES) Subcutaneous	August 2020 RRMM (CANDOR), D-Kd



Genmab

Daratumumab

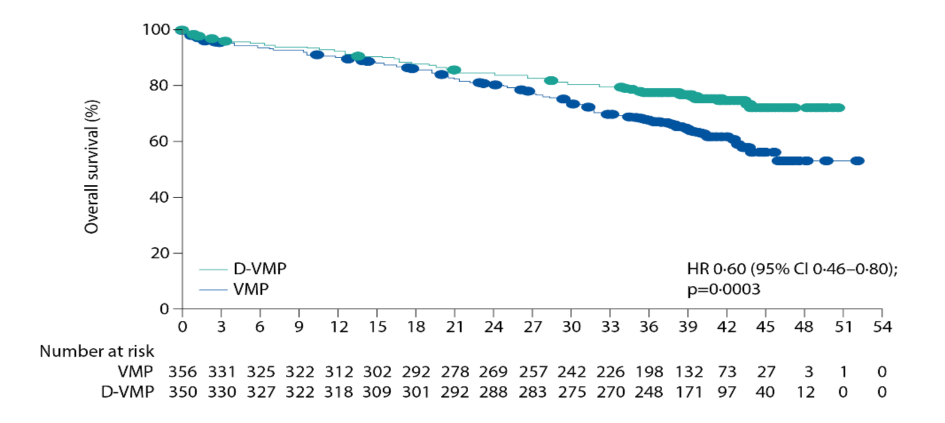
Proving to be the Critical Driver Across Different Combinations & Treatment Lines



Ongoing Phase 3: CEPHEUS (D-VRd, NDMM NTE), PERSEUS (D-VRd, NDMM TE)



Improved Survival for Patients with Multiple Myeloma Overall Survival Analysis from ALCYONE Trial



Kaplan-Meier estimates of overall survival in intention-to-treat population. Mateos, MV et al, 'Overall survival with daratumumab, bortezomib, melphalan, and prednisone in newly diagnosed multiple myeloma (ALCYONE): a randomized, open-label, phase 3 trial,' *The Lancet*, published online December 9, 2019



Kesimpta[®] (ofatumumab) Approved in Relapsing Multiple Sclerosis

Human CD20 Antibody – well validated target

Injection for SubQ use approved for RMS in the US

First B-cell therapy that can be self-administered by patients at home using Sensoready[®] autoinjector pen

Developed by Novartis: Regulatory submission also made in EU

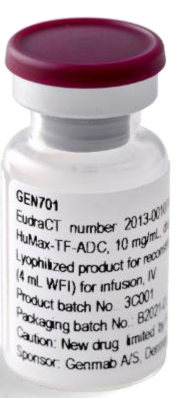
Genmab 10% royalty payment of net sales

Second Genmab-created product with blockbuster potential



Tisotumab Vedotin

Genmab's Most Advanced Asset with Potential in Solid Tumors





Fully human antibody-drug conjugate (ADC) targeting Tissue Factor (TF) in development to treat solid tumors



License and collaboration agreement with Seattle Genetics 50:50

Very favorable topline results, Phase 2 recurrent or metastatic cervical cancer



Ongoing trials in cervical, ovarian cancer, other solid tumors

Expanding development, additional studies planned



Tisotumab Vedotin in Cervical Cancer

Designed to Address a High Unmet Medical Need

Recurrent or metastatic cervical cancer

- Poor prognosis advanced / recurrent cervical cancer
 - RR standard therapies generally <15%
 - Median OS 6-8 months
- Data ORR & survival after progression on 1L bevacizumab + doublet chemotherapy are limited

Conclusions*

(previously treated recurrent or metastatic cervical cancer)

- Manageable AEs + encouraging antitumor activity
- ORR 35% (confirmed + unconfirmed, IRC)
- Confirmed ORR 22%
- Median DOR 6.0 months
- 6-month PFS of 40%

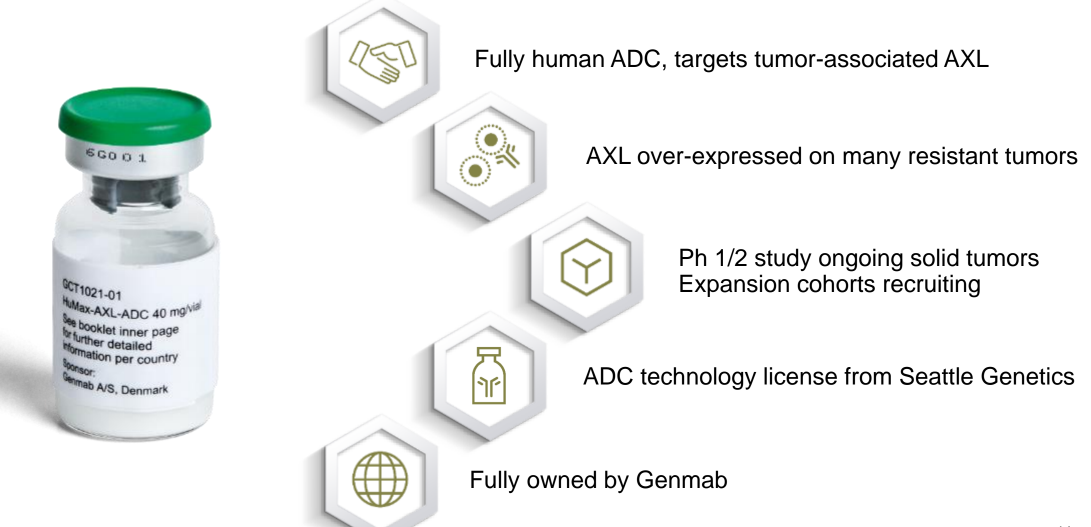
Encouraging	Antitumor	Activity	Observed
			0.000.000

	N	=55
	IRC-Assessed	INV-Assessed
ORR confirmed + unconfirmed (95% CI), %	35 (22-49)	31 (19-45)
ORR confirmed (95% CI), %	22 (12-35)	24 (13-37)
CR, n (%)	1 (2)	0
PR, n (%)	11 (20)	13 (24)
SD, n (%)	19 (35	21 (38)
PD, n (%)	17 (31)	17 (31)
Not evaluable, ^b n (%)	5 (9)	4 (7)
DCR confirmed (95% CI), %	56 (42-70)	62 (48-75)
Median DOR (range), months	6.0 (+1.0 -9.7)	4.2 (+1.0 -9.7)
Median PFS (95% CI, months	4.1 (1.7-6.7)	4.2 (2.1-5.3)
6-month PFS rate (95% CI), %	40 (24-55)	29 (17-43)



Enapotamab Vedotin

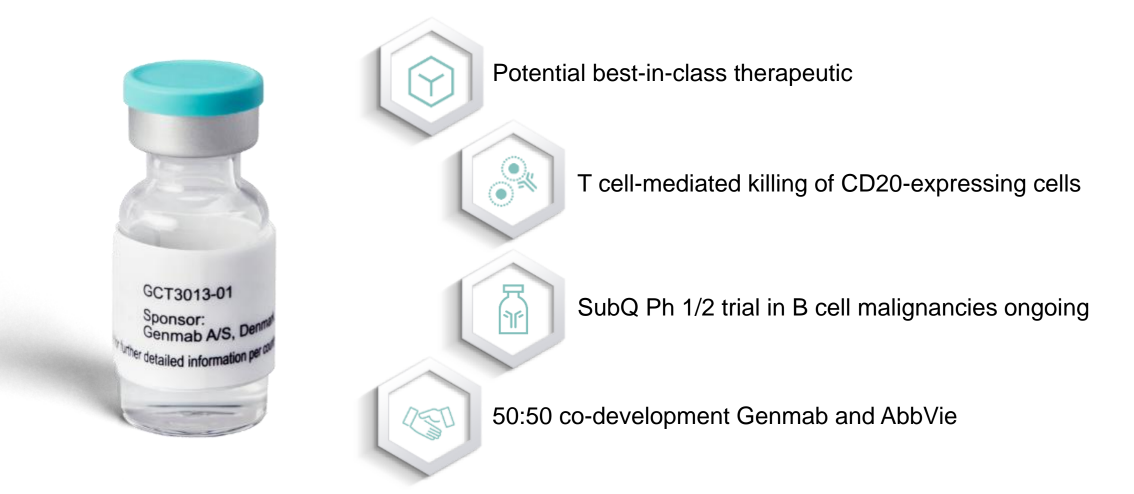
Potential in Solid Tumors





Epcoritamab (DuoBody-CD3xCD20)

Potential for Improved Efficacy & Safety in B Cell Malignancies



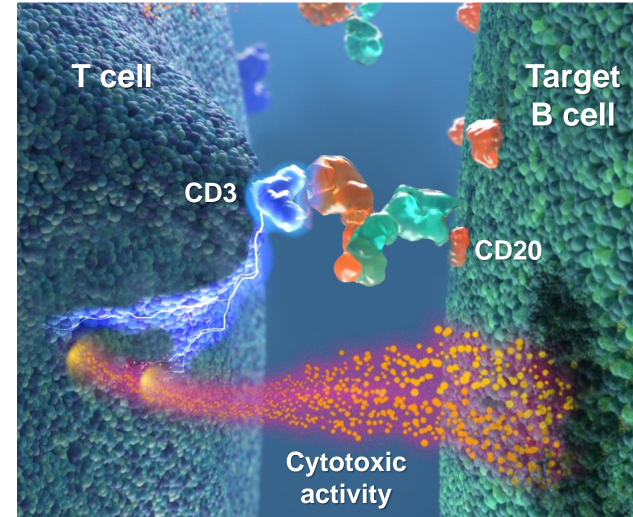
Epcoritamab: Dose Escalation Data Presented at EHA25 Virtual Congress 2020*

Anti-tumor activity

- 86% ORR in FL \geq 0.76mg
- 60% ORR, incl. 3 pts who failed prior CAR-T treatment, in DLBCL/HGBCL ≥12 mg
- Emerging prelim. data highly encouraging with substantial single-agent efficacy
- Induces rapid and deep responses in heavily pretreated pts with B-NHL across different subtypes

Safety

- No DLTs observed; MTD has not been reached
- · No treatment-related deaths
- No discontinuation due to AEs unrelated to disease progression
- No Grade ≥ 3 CRS events observed



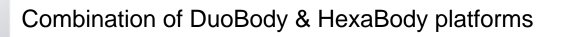
Dose-escalation data with subcutaneous epcoritamab indicate potential for best-in-class therapy



DuoHexaBody-CD37 (GEN3009)

Building Our Pipeline: First DuoHexaBody in the Clinic





Novel target for hematologic malignancies

Unique mechanism-of-action

Dose escalation ongoing

50:50 co-development Genmab and AbbVie



DuoBody-CD3x5T4 (GEN1044)

Latest in the Clinic



Based on proprietary DuoBody technology



CD3 bispecific, T cell mediated cytotoxicity of 5T4+ tumor cells

5T4 expressed on multiple solid tumors limited expression in healthy tissue

Potent anti-tumor activity in diversity pre-clinical models



50:50 co-development Genmab and AbbVie



DuoBody-PD-L1x4-1BB (GEN1046)

Bispecific Next Generation Checkpoint Immunotherapy





Potential as differentiated Genmab PD-L1 product

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Combining checkpoint blockade with T cell stimulation

Ph 1/2 study ongoing in solid tumors

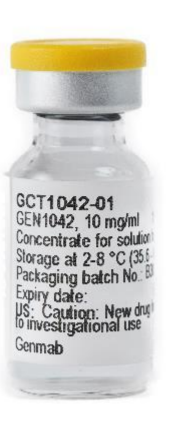
50:50 co-development Genmab and BioNTech



DuoBody-CD40x4-1BB (GEN1042)

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Bispecific Agonistic Antibody



Bispecific antibody targeting CD40 & 4-1BB (CD137)

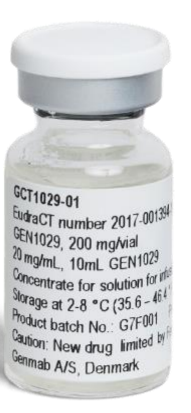
Conditionally activates T cells and APC in presence of CD40-expressing cells

Phase 1/2 study ongoing in solid tumors

50:50 co-development Genmab and BioNTech



HexaBody-DR5/DR5 (GEN1029) First HexaBody in Clinical Development





Targets 2 distinct DR5 epitopes

HexaBody platform - DR5 clustering & DR5 agonist activity

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First 100% Genmab-owned HexaBody product in clinic

Phase 1/2 study ongoing in multiple solid tumors



2020 Guidance*: Recurring Revenue Growth and Focused R&D Investments

Income Statement	DKKM	~USDM**	Key Observations
Revenue	9,250 – 9,850	1,423 – 1,515	 Summary P&L DARZALEX royalties of ~DKK 4.1bn to ~DKK 4.5bn to drive recurring revenue growth
Operating Expenses	(3,850) – (3,950)	(592) – (608)	 Nearly 90% of USD 750M upfront from AbbVie collab. recognized immediately
Operating Income	5,350 – 5,950	823 - 915	 Growth in operating expenses driven by expanding and accelerating our clinical pipeline
			 DARZALEX Sales of USD 3.9bn – USD 4.2bn Significant opportunity for growth in 1L MM market
			 SubQ DARZALEX approvals in H1 in U.S. & EU
			 Market share gain in the U.S. and RoW driven by uptake in all lines of treatment
			 8 approved indications in U.S., late stage to 1L MM
			 Growth expected to normalize in H2 2020



Key 2020 Priorities

Building a Strong Differentiated Product Pipeline

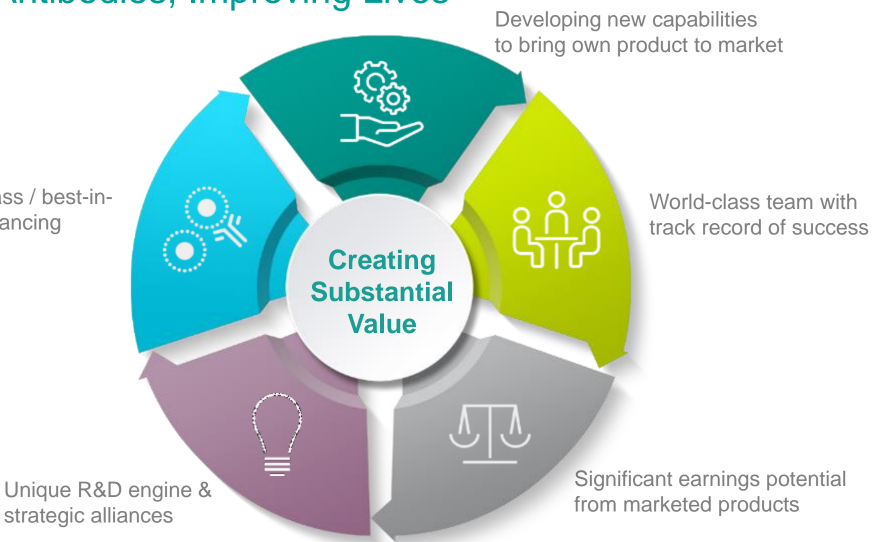
Priority	✓	Targeted Milestones
Genmab proprietary* products	✓	 initiate expansion cohorts » HexaBody-DR5/DR5 Phase 1/2 - advance dose escalation
Daratumumab ⁴	~	 » U.S. FDA and EMA decision on Phase 3 COLUMBA multiple myeloma SubQ submission » sBLA and MAA Submission Phase 3 ANDROMEDA amyloidosis » sBLA and MAA submission Phase 3 APOLLO multiple myeloma
Ofatumumab ⁵	\checkmark	» U.S. FDA decision on regulatory dossier submission in multiple sclerosis
Teprotumumab ⁶	\checkmark	» U.S. FDA decision on Phase 3 OPTIC active thyroid eye disease submission



Delivering on Genmab's Promise:



Pipeline of 1st-in-class / best-inclass therapies advancing through clinic



Innovating Antibodies, Improving Lives



Appendix



A Leading International Biotech With Large Free Float





Genmab & AbbVie: Collaboration Overview

A broad, long-term oncology collaboration with Genmab and AbbVie working together to jointly make all strategy, clinical development and commercialization decisions



50/50 partnership across three clinical nextgeneration bispecific antibody product candidates (epcoritamab, DuoHexaBody-CD37, DuoBody-CD3x5T4)



Genmab to book epcoritamab sales in the U.S. and Japan; AbbVie to commercialize epcoritamab RoW - Genmab to receive tiered royalties on RoW net sales



Worldwide co-commercialization and profit split of all other programs



Discovery Research Collaboration



Fourth* largest oncology partnership with total potential value ~USD 3.9bn (up-front cash + milestone payments) to Genmab

Genmab

Advancing Pipeline: Delivering on Our Promise & Creating Value Accelerating Development of Potential "Next Winners"

Delivering on Genmab's Promise to Patients

DuoBody-CD3xCD20 (epcoritamab)

- Potential best-in-class: SubQ administration
- Pre-clinical / preliminary clinical data shows encouraging safety & efficacy
- Expeditious and comprehensive clinical development plan
- RP2D decision & expansion cohorts initiation
- 50:50 AbbVie

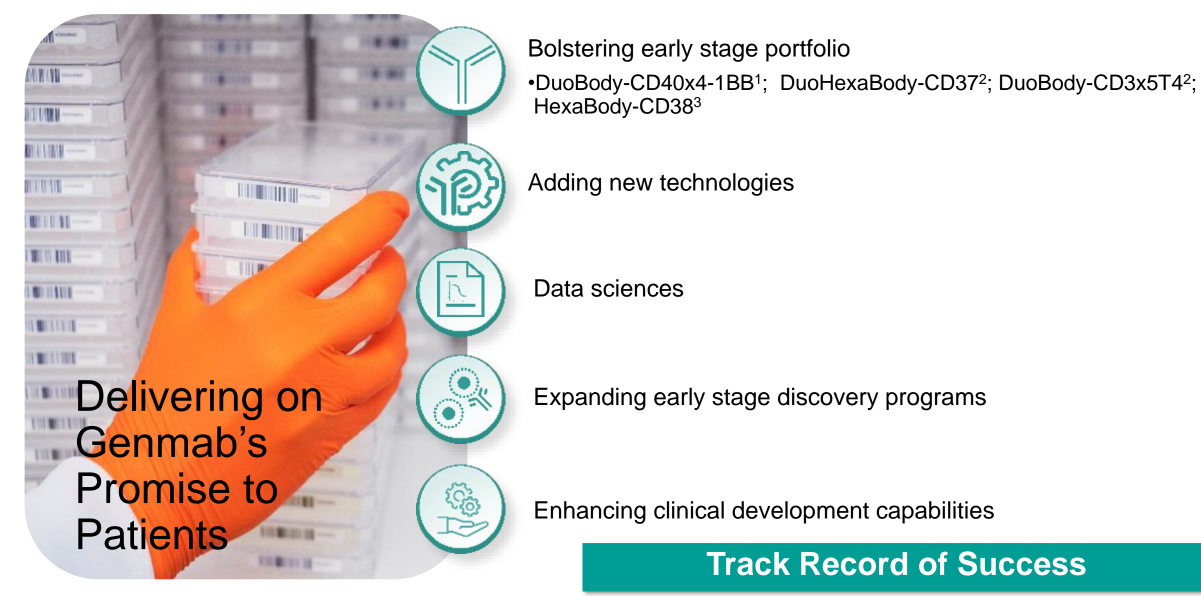
DuoBody-PD-L1x4-1BB (GEN1046)

- Potential first-in-class: Next generation IO
- Unmet medical need
- FiH clinical study: escalation phase is ongoing
- 50:50 BioNTech

Track Record of Success

Genmab

Advancing Pipeline: Delivering on Our Promise & Creating Value



¹GEN1042, 50:50 w/ BioNTech; ²50:50 w/ AbbVie; ³Genmab is developing HexaBody-CD38 in an exclusive worldwide license and option agreement w/ Janssen Biotech, Inc



Genmab's Commitment to Society

Building a Socially Responsible & Sustainable Company

Anchored in our Core Purpose & Vision

- To improve the lives of patients by creating and developing innovative antibody products
- By 2025 our own product has transformed cancer treatment and we have a pipeline of knock-your-socksoff antibodies

CSR Committee comprised of representatives from variety of functions, chaired by CEO

- Ensures that Genmab carries out CSR activities effectively & communicates clearly and openly
- Focus on Environment, Society and Governance reporting

Focus on four main areas

- Employee well-being, including health, safety & development
- Ethics in relation to pre-clinical and clinical studies
- Environment, including waste management & recycling
- Business ethics & transparency



Innovation Powerhouse: Cutting Edge Proprietary Technologies

Technology		Principle	Applications
DuoBody		Bispecific antibodies	Dual targeting
HexaBody	20000 20000 20000	Target-mediated enhanced hexamerization	Enhanced potency
DuoHexaBody	2×14	Bispecific antibodies with target- mediated enhanced hexamerization	Dual targeting + enhanced potency
HexElect		Two co-dependent antibodies with target-mediated enhanced hexamerization	Dual targeting + enhanced potency & selectivity



Innovative Clinical and Pre-Clinical Pipeline

Genmab's Proprietary¹ Products

Product	Target	Developed By	Disease Indications	Most Advanc	ed Devel;	opment F	Phase			
				Pre-Clinical		1	1/2	2	3	Approved
Tisotumab vedotin	TF	50:50 Genmab / Seattle	Cervical cancer						 	
		Genetics	Ovarian cancer							
			Solid tumors							
Enapotamab vedotin	AXL	Genmab	Solid tumors							
Epcoritamab (DuoBody-CD3xCD20)	CD3, CD20	50:50 Genmab / AbbVie	Hematological malignancies	3				 	 	
DuoBody-PD-L1x4-1BB (GEN1046)	PD-L1, 4-1BB	50:50 Genmab / BioNTech	Solid tumors						 	
HexaBody-DR5/DR5 (GEN1029)	DR5	Genmab	Solid tumors					 		
DuoBody-CD40x4-1BB (GEN1042)	CD40, 4-1BB	50:50 Genmab / BioNTech	Solid tumors							
DuoHexaBody-CD37 (GEN3009)	CD37	50:50 Genmab / AbbVie	Hematologic malignancies							
DuoBody-CD3x5T4 (GEN1044)	CD3, 5T4	50:50 Genmab / AbbVie	Solid tumors						 	
IND/CTAs in 2020 HexaBody-CD38 (GEN3014) ²		Genmab						 	 	32

¹Certain product candidates in development with partners, as noted. ²Genmab is developing HexaBody-CD38 in an exclusive worldwide license and option agreement with Janssen Biotech, Inc



Products Created by Genmab*

Including Proposed Label Expansions for Marketed Products

Product	Target	Developed By	Disease Indications Most Advanced Development Phase						
				Pre-Clinical	1	1/2	2	3	Approved
DARZALEX (daratumumab) & DARZALEX FASPRO (daratumumab and hyaluronidase-fihj)	CD38	Janssen (Tiered royalties to Genmab on net global sales)	Multiple myeloma ¹						
Daratumumab			AL Amyloidosis						
			Non-MM blood cancers						
Kesimpta (ofatumumab)	CD20	Novartis (Royalties to Genmab on net global sales)	Relapsing Multiple Sclerosis ¹						
Arzerra (ofatumumab)	CD20	Novartis	Chronic lymphocytic leukemia ^{1,2}						
TEPEZZA (teprotumumab-trbw)	IGF-1R	Horizon Therapeutics (under sublicense from Roche, royalties to Genmab on net global sales)	Thyroid eye disease ¹						

*Out-licensed products marketed by partner ¹See local country prescribing information for precise indications, ²Not in active development



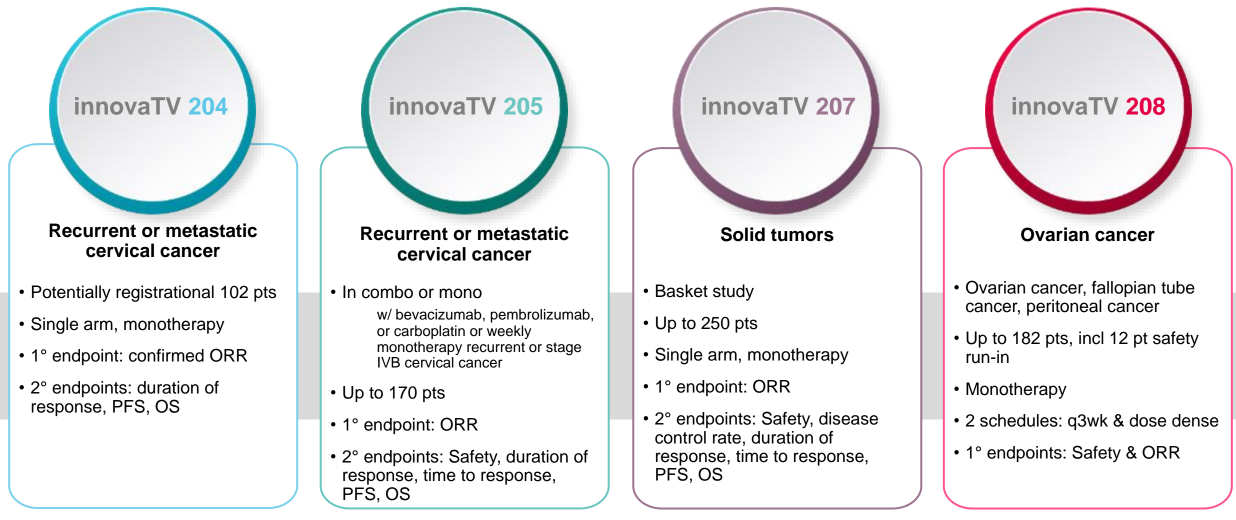
Partner-owned Products Incorporating Genmab's Innovation*

Product	Target	Developed By	Disease Indications	Most Advan	ced De	evelop	ment Phase			
				Pre-Clinical	1	l	1/2	2	3	Approved
Camidanlumab tesirine (ADCT-301)	CD25	ADC Therapeutics	Relapsed /Refractory Hodgkin Lymphoma							
			Solid tumors							
Mim8	FIX(a), FX	Novo Nordisk	Healthy volunteers & hemophilia A							
Amivantamab (JNJ-61186372)	EGFR, cMet	Janssen	Non-small-cell lung cancer (NSCLC)						
JNJ-63709178	CD123, CD3	Janssen	Acute Myeloid Leukemia (AML)							
Teclistamab (JNJ-64007957)	BCMA, CD3	Janssen	Relapsed or refractory MM							
Talquetamab (JNJ-64407564)	GPRC5D, CD3	Janssen	Relapsed or refractory MM							
JNJ-67571244	CD33, CD3	Janssen	Relapsed or refractory AML or MDS	;						
JNJ-63898081	PSMA, CD3	Janssen	Solid tumors							
HuMax-IL8	IL8	BMS	Advanced cancers							
Lu AF82422	alpha-Synuclein	Lundbeck	Parkinson's disease							
~20 active pre-clinical programs			Partnered & proprietary programs: HuMab, DuoBody, DuoHexaBody and HexaBody							



Solid Foundation Built on a Differentiated Pipeline

Tisotumab Vedotin Clinical Program





Tisotumab Vedotin

Cervical Cancer Market Size

United States ³	Japan ⁶	Europe ²		
New Diagnoses Deaths 12,578 4,115	New Diagnoses Deaths 9,390 3,654	New Diagnoses Deaths 58,373 24,404		
3rd most common gynecologic cancer in US ⁴	2nd most common gynecologic cancer in Japan ⁶	3rd most common gynecologic cancer in Europe ^{2*}		

In developed countries, incidence rates are low (<7.9 per 100,000 women) compared with *developing countries* in sub-Saharan Africa and Central and South America, where incidence is especially high (>30 per 100,000 women)⁵

*Europe is defined as the 40 countries in the four United Nations-defined areas of Europe and the European Union (EU-27). **References: 1**. American Cancer Society 2. EUCAN (2012) 3. Centers for Disease Control and Prevention. Cervical Cancer Statistics (2017) 4. UpToDate.
5. Ginsburg O et al. Lancet 2017 6. HPV Information Centre Japan (2017)



HexaBody-CD38 (GEN3014)

Expanding the Potential of CD38 Antibodies

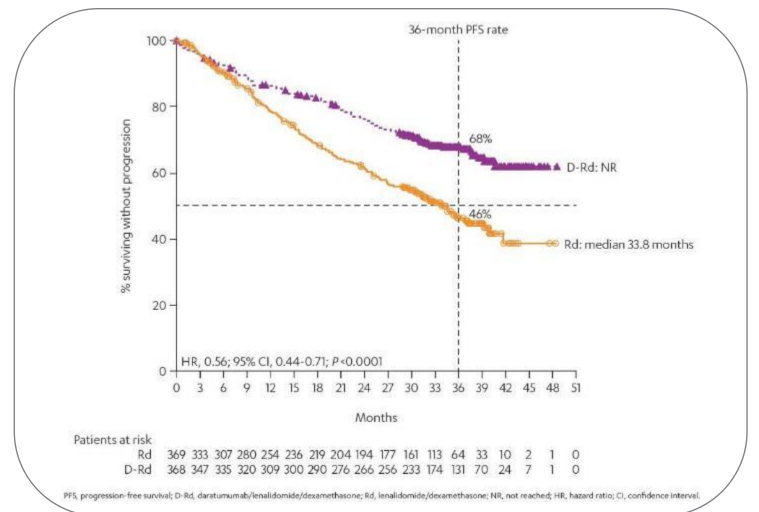
Highly promising Could potentially **IND/CTA** planned Incorporates add to and broaden in H2 2020 data pre-clinical proprietary models for MM, HexaBody DARZALEX technology lymphoma & AML franchise



Covering All Stages of MM and Beyond: Key Ongoing* Industry Sponsored Trials

Disease	Therapy	Development Phase							
		Pre-Clinical	1	1/2	2	3			
High Risk Smoldering MM	Subcutaneous	\checkmark	AQUILA						
	Monotherapy	\checkmark	CENTAURUS						
Front line MM (transplant & non-	Dara + VRd		CEPHEUS						
transplant)	Dara + VMP (Asia Pacific)	\checkmark	OCTANS						
	Dara + VRd	\checkmark	PERSEUS						
	Dara + R (maintenance)		AURIGA						
Relapsed or Refractory MM	Dara + combinations								
	Dara + I.O. (PD1 & PDL1)	Opdivo [®] (Ph	I/II), Tecentriq®	(Ph I)					
ALL	Dara + SoC chemo		DELPHINUS						

Daratumumab Efficacy in Newly Diagnosed Multiple Myeloma Updated Phase 3 MAIA Trial (D+Rd, NTE): ASH Dec 2019



 Median PFS not reached in D-Rd arm

Genmab

- MRD-negativity significantly higher with D-Rd vs. Rd (29% vs 9%; P<0.0001)
- No new safety concerns
- Results continue to support use of D-Rd in 1st line treatment of TIE pts with NDMM



Ongoing Daratumumab Clinical Trials

Janssen Sponsored Phase 3 & 4

Daratumumab Trials Sponsored by Pharma / Biotech

Ct.gov Identifier	Phase	Sponsor	Indication	Therapy
NCT03768960	4	J&J Private Ltd	Relapsed or Refractory MM	Daratumumab (MMY4008)
NCT02252172	3	Janssen	Untreated MM	Daratumumab + Rd (MAIA)
NCT02195479	3	Janssen	Untreated MM	Daratumumab + VMP (ALCYONE)
NCT02541383	3	Janssen	Untreated MM	Daratumumab + VTd (CASSIOPEIA)
NCT02076009	3	Janssen	Relapsed or Refractory MM	Daratumumab + Rd (POLLUX)
NCT02136134	3	Janssen	Relapsed or Refractory MM	Daratumumab + Vd (CASTOR)
NCT03180736	3	Janssen	Relapsed or Refractory MM	Daratumumab + Pom-d (APOLLO)
NCT03201965	3	Janssen	Amyloidosis	Daratumumab + CyBorD (ANDROMEDA)
NCT03217812	3	Janssen	Untreated MM	Daratumumab + VMP (Asia Pacific) (OCTANS)
NCT03234972	3	Janssen	Relapsed or Refractory MM	Daratumumab + Vd vs Vd (LEPUS)
NCT03277105	3	Janssen	Relapsed or Refractory MM	Daratumumab SubQ vs IV (COLUMBA)
NCT03301220	3	Janssen	Smoldering MM	Daratumumab SC (AQUILA)
NCT03652064	3	Janssen	Untreated MM	Daratumumab + VRd (CEPHEUS)
NCT03710603	3	Janssen/EMN	Untreated MM	Daratumumab + VRd (PERSEUS)
NCT03901963	3	Janssen	Untreated MM / Maintenance	Daratumumab + R (AURIGA)



Ongoing Daratumumab Clinical Trials

Janssen Sponsored Phase 1 & 2

Daratumumab Trials Sponsored by Pharma / Biotech

Ct.gov Identifier	Phase	Sponsor	Indication	Therapy
NCT03384654	2	Janssen	Relapsed / Refractory ALL / LL	Dara + Vincristine + Prednisone + Doxorubicin (DELPHINUS)
NCT02951819	2	Janssen	Untreated and Relapsed MM	Daratumumab + CyBorD (LYRA)
NCT02874742	2	Janssen	Untreated MM	Daratumumab + VRd (GRIFFIN)
NCT02316106	2	Janssen	Smoldering MM	Monotherapy (CENTAURUS)
NCT02927925	2	Janssen	NKTCL, Nasal Type	Monotherapy (VOLANS)
NCT03412565	2	Janssen	Newly diag. & relapsed / refractory MM	Daratumumab SubQ + Rd, VMP & VRd (PLEIADES)
NCT03871829	2	Janssen	Dara retreatment	Daratumumab SubQ+ Kd vs Kd (LYNX)
NCT03011034	2	Janssen	MDS	Daratumumab (or talacotuzumab) (MDS2002)
NCT01615029	1/2	Janssen	Relapsed and Refractory MM	Daratumumab + Rd (GEN503)
NCT02852837	1	Janssen	Relapsed or Refractory MM	Monotherapy (in China) (MMY1003)
NCT02519452	1	Janssen	Relapsed or Refractory MM	Monotherapy, subcutaneous (PAVO)
NCT02918331	1	Janssen	Untreated MM	Daratumumab + Rd (Japan) (MMY1006)
NCT03242889	1	Janssen	Relapsed or Refractory MM	Daratumumab subq (Japan) (MMY1008)
NCT01998971	1	Janssen	Various MM	Daratumumab + backbone regimens (Vd, VMP, VTd, Pom-d, Kd, KRd) (EQUULEUS)
NCT04108195	1	Janssen	Multiple Myeloma	Daratumuamb + either talquetamab or teclistamab (MMY1002)
NCT04121260	1	Janssen	Multiple Myeloma	Subcutaneous monotherapy (in China) (MMY1010)



Ongoing Daratumumab Clinical Trials

Other Industry Sponsored Trials

Daratumumab Trials Sponsored by Pharma / Biotech

Ct.gov Identifier	Phase	Sponsor	Indication	Therapy
NCT03158688	3	Amgen	Relapsed or Refractory MM	Daratumumab + Kd (CANDOR)
NCT01946477	2	Celgene	Relapsed or Refractory MM	Daratumumab + Pom-d
NCT02807454	2	Celgene	Relapsed and Refractory MM	Daratumumab + Imfinzi (FUSION)
NCT03439293	2	Takeda	Relapsed or Refractory MM	Daratumumab + NINLARO (ixazomib) + Dex
NCT03314181	2	AbbVie	Relapsed or Refractory MM	Daratumumab + Venetoclax + Dex (w/ or w/out bortezomib)
NCT02807558	2	Syros Pharma	AML or MDS	Daratumumab + SY-1425
NCT02773030	1/2	Celgene	Relapsed or Refractory MM	Daratumumab + CC-220 + Dex
NCT02343042	1/2	Karyopharm	Relapsed or Refractory MM	Daratumumab + Selinexor + Dex (STOMP)
NCT03481556	1/2	Oncopeptides AB	Relapsed or Refractory MM	Daratumumab + Melflufen + Dex (ANCHOR)
NCT01592370	1/2	BMS	Relapsed or Refractory MM	Daratumumab + nivolumab
NCT03837509	1/2	Incyte	Relapsed or Refractory MM	Daratumumab + INCB001158
NCT03989414	1/2	Celgene	Various MM	Daratumumab + CC-92480
NCT02431208	1	Roche	Resistant or Refractory MM	Daratumumab + Tecentriq (atezolizumab)
NCT03068351	1	Roche	Resistant or Refractory MM	Daratumumab + RO6870810
NCT04045028	1	Genentech	Relapsed or Refractory MM	Daratumumab + tiragolumab
NCT04136756	1	Nektar Thera.	Salvage for MM	Daratumumab + NKTR-255

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