

Innovating Antibodies, Improving Lives

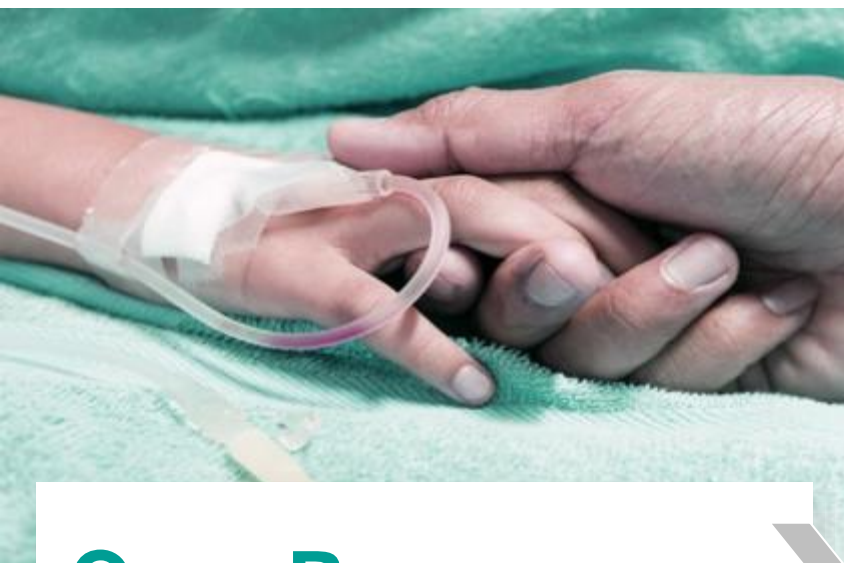
Investor Presentation
August 2020



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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

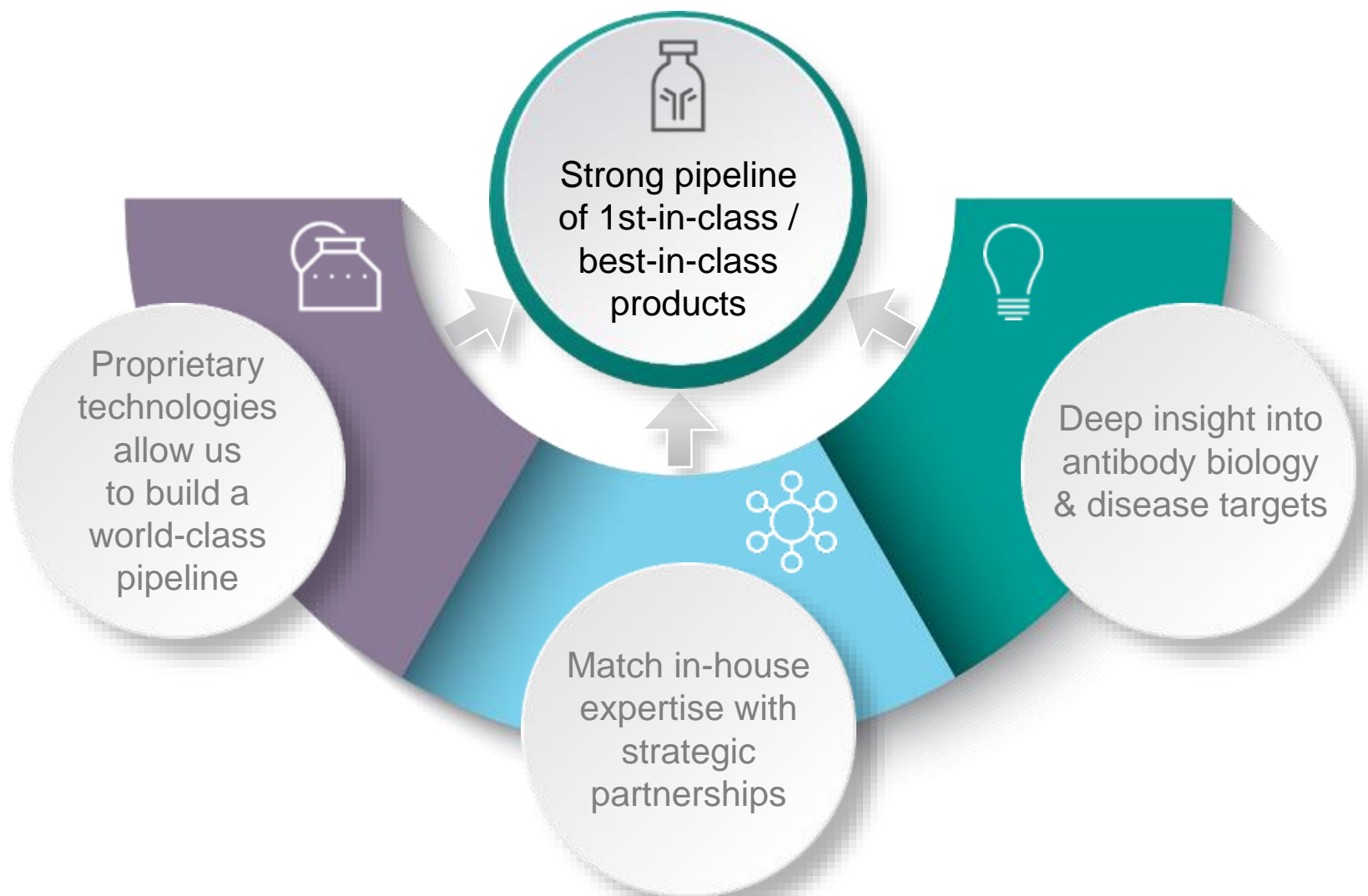


Vision

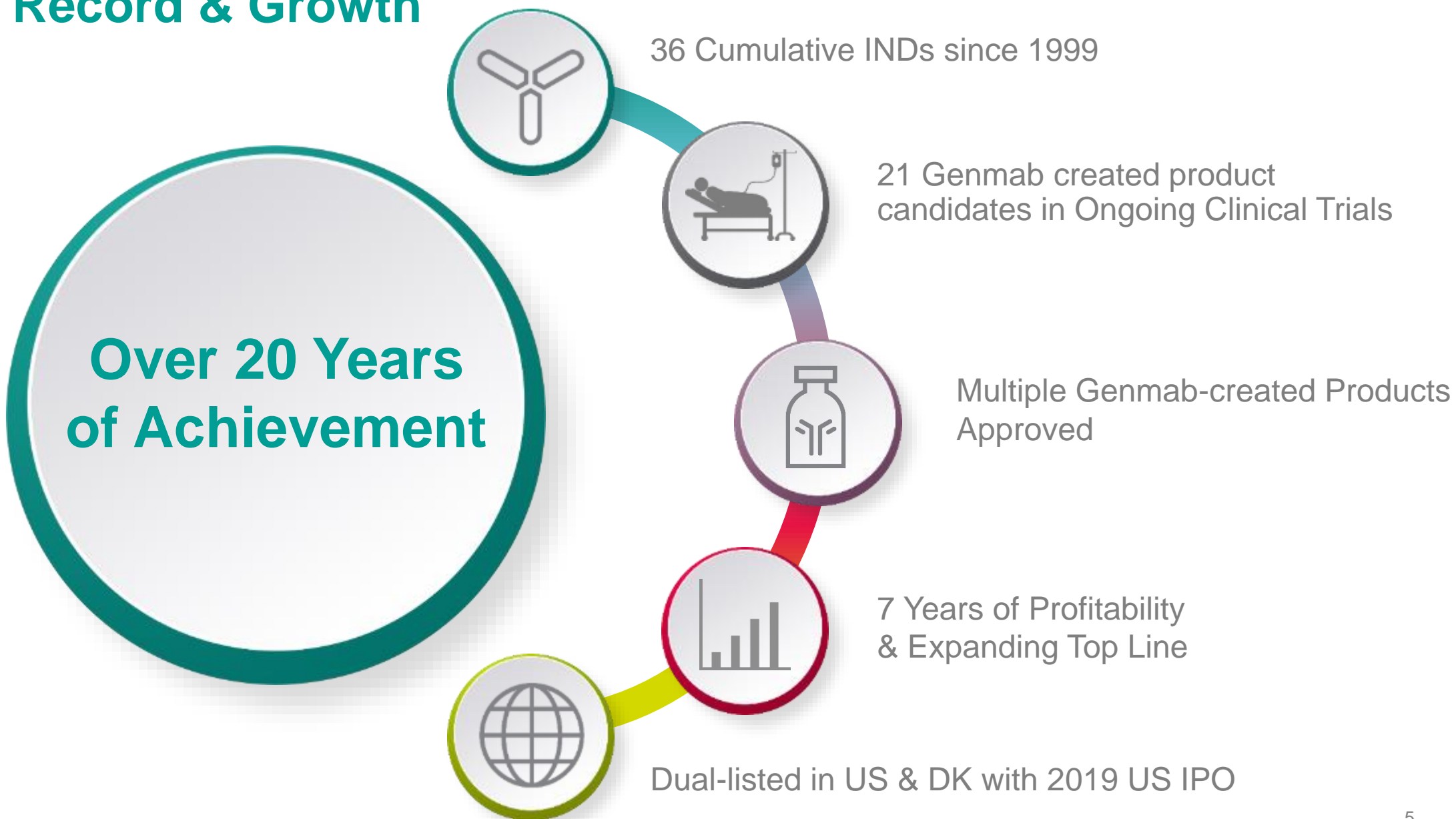
By 2025, our own product has transformed cancer treatment and we have a pipeline of knock-your-socks off antibodies

The Genmab Difference

Innovation Powerhouse Transforming Cancer Treatment & Creating Value



Track Record & Growth



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

R&D Engine

Technologies & Pre-Clinical

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

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

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



First-in-class CD38 antibody in development to treat cancer



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



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



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



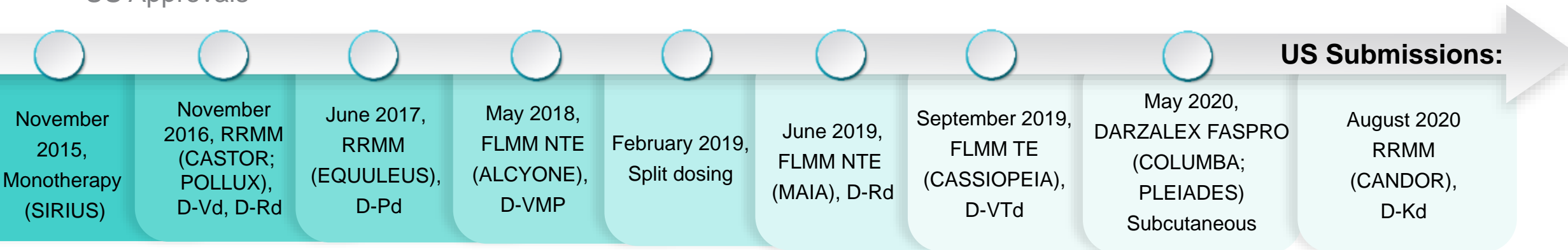
2019 WW net sales by J&J: \$2,998M



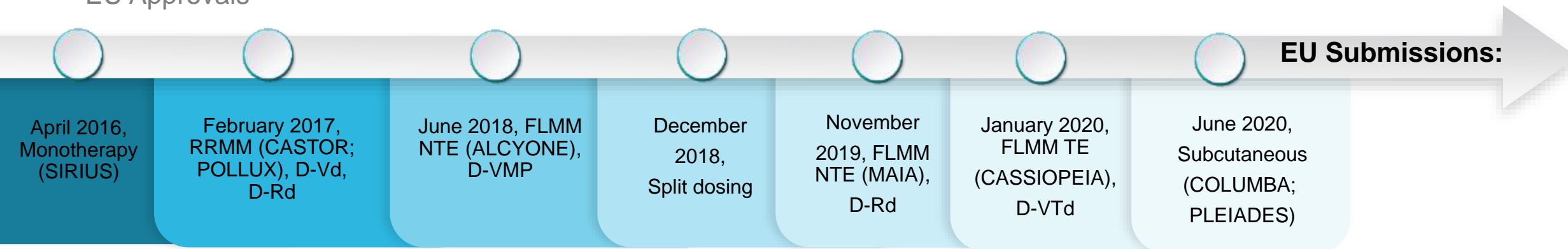
DARZALEX Approvals: US and EU

On Track for Approval Across All Lines of MM Treatment

US Approvals



EU Approvals



Daratumumab

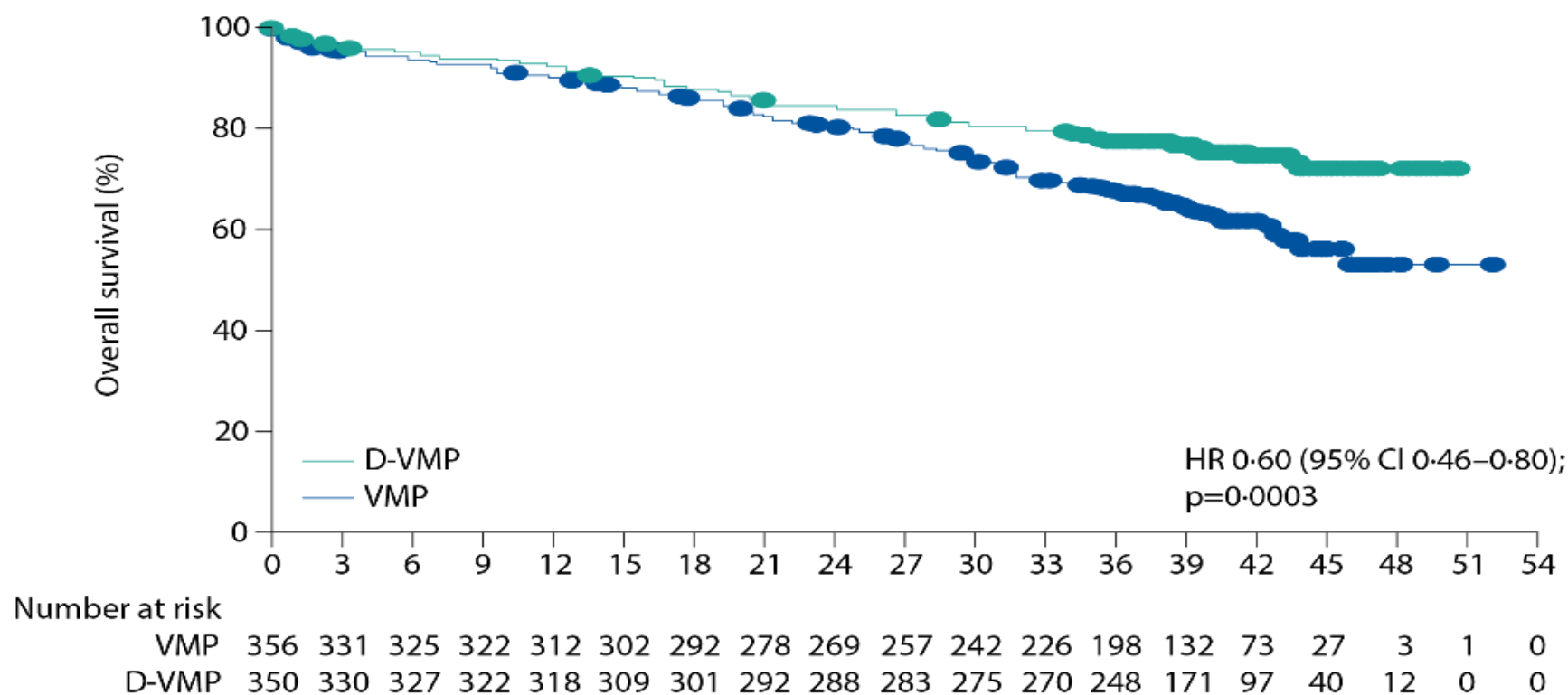
Proving to be the Critical Driver Across Different Combinations & Treatment Lines

	Frontline				Relapsed/Refractory	
	Transplant Eligible		Transplant Ineligible			
	Ph 3 CASSIOPEIA ^{1,3} (D-VTd vs. VTd)	Ph 2 GRIFFIN ^{1,4} (D-VRd vs VRd)	Ph 3 ALCYONE ^{2,4} (D-VMP vs. VMP)	Ph 3 MAIA ^{2,4} (D-Rd vs. Rd)	Ph 3 POLLUX ^{2,4} (D-Rd vs. Rd)	Ph 3 CASTOR ^{2,4} (D-Vd vs Vd)
sCR Odds Ratio ¹ or CR+ ²	1.60	1.57	~2x	~2x	>2x	3x
MRD-neg rate	1.5x	2.5x	4x	>3x	~5x	>7x
PFS risk reduction	53% (HR, 0.47)	NA	58% (HR, 0.42)	44% (HR, 0.56)	56% (HR, 0.44)	69% (HR, 0.31)

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



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*

	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



Fully human ADC, targets tumor-associated AXL



AXL over-expressed on many resistant tumors



Ph 1/2 study ongoing solid tumors
Expansion cohorts recruiting



ADC technology license from Seattle Genetics



Fully owned by Genmab

Epcoritamab (DuoBody-CD3xCD20)

Potential for Improved Efficacy & Safety in B Cell Malignancies



Potential best-in-class therapeutic



T cell-mediated killing of CD20-expressing cells



SubQ Ph 1/2 trial in B cell malignancies ongoing



50:50 co-development Genmab and AbbVie

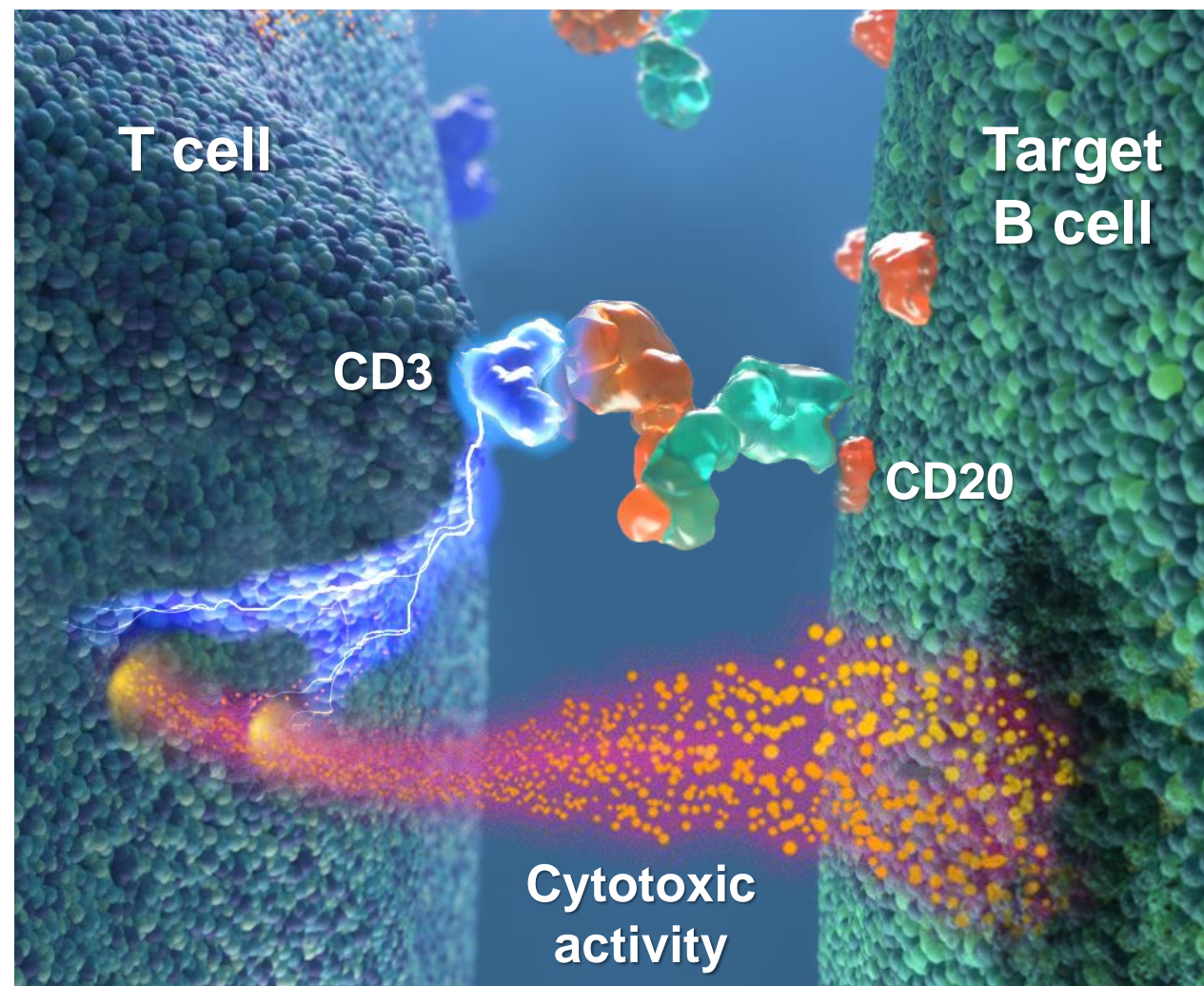
Epcoritamab: Dose Escalation Data Presented at EHA25 Virtual Congress 2020*

Anti-tumor activity

- 86% ORR in FL \geq 0.76mg
- 60% ORR, incl. 3 pts who failed prior CAR-T treatment, in DLBCL/HGBCL \geq 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 \geq 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



Combination of DuoBody & HexaBody platforms



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



Bispecific antibody targeting PD-L1 & 4-1BB (CD137)



Potential as differentiated Genmab PD-L1 product



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)

Bispecific Agonistic Antibody



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



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



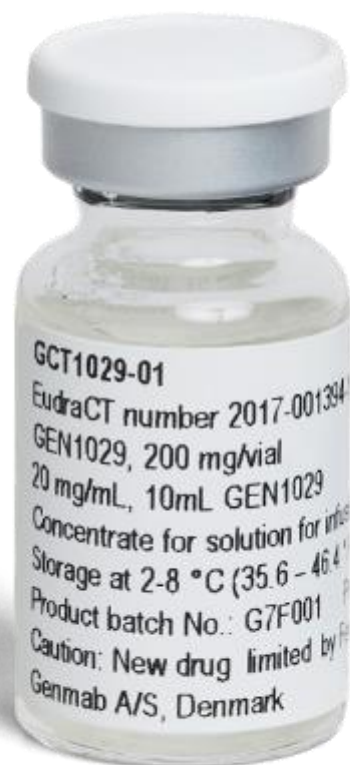
Phase 1/2 study ongoing in solid tumors



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



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 <ul style="list-style-type: none"> DARZALEX royalties of ~DKK 4.1bn to ~DKK 4.5bn to drive recurring revenue growth Nearly 90% of USD 750M upfront from AbbVie collab. recognized immediately Growth in operating expenses driven by expanding and accelerating our clinical pipeline DARZALEX Sales of USD 3.9bn – USD 4.2bn <ul style="list-style-type: none"> 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
Operating Expenses	(3,850) – (3,950)	(592) – (608)	
Operating Income	5,350 – 5,950	823 - 915	

Key 2020 Priorities

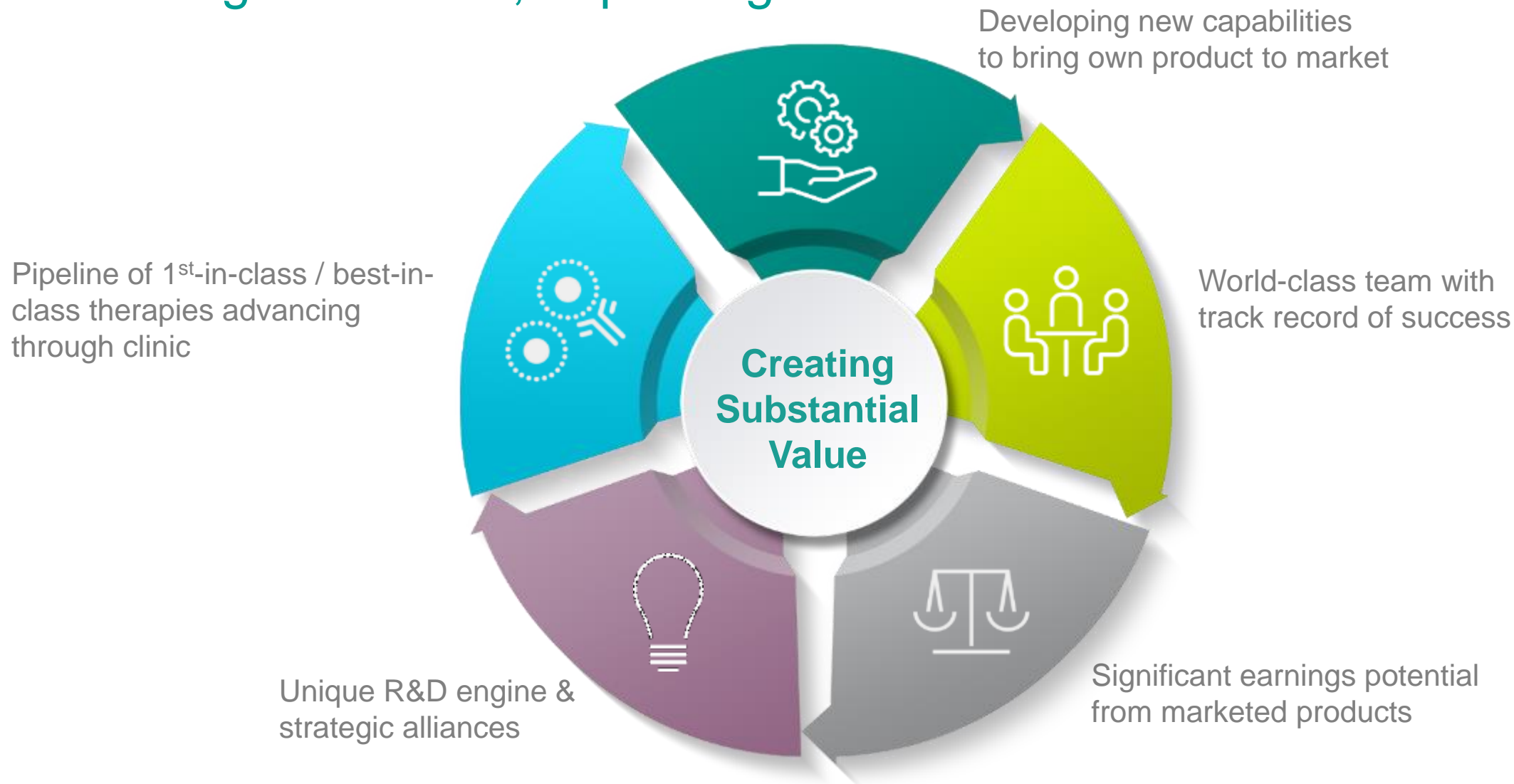
Building a Strong Differentiated Product Pipeline

Priority	✓	Targeted Milestones
Genmab proprietary* products		<ul style="list-style-type: none"> » Tisotumab vedotin¹ - Phase 2 innovaTV 204 safety & efficacy analysis in recurrent/metastatic cervical cancer and engage U.S. FDA for BLA submission subject to trial results » Tisotumab vedotin - data on other solid tumor types » Enapotamab vedotin – data to support late stage development ✓ » Epcoritamab (DuoBody-CD3xCD20)² Phase 1/2 – decision on recommended Phase 2 dose & initiate expansion cohorts » HexaBody-DR5/DR5 Phase 1/2 - advance dose escalation ✓ » DuoBody-PD-L1x4-1BB³ Phase 1/2 – initiate expansion cohorts » DuoBody-PD-L1x4-1BB initial data in H2 2020 » File INDs and/or CTAs for 2 new products
Daratumumab ⁴	✓	<ul style="list-style-type: none"> » 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	✓	<ul style="list-style-type: none"> » U.S. FDA decision on regulatory dossier submission in multiple sclerosis
Teprotumumab ⁶	✓	<ul style="list-style-type: none"> » U.S. FDA decision on Phase 3 OPTIC active thyroid eye disease submission

*Certain product candidates in development with partners, as noted.

1. 50:50 dev. w/ Seattle Genetics; 2. 50:50 dev w/ AbbVie; 3. 50:50 dev. w/ BioNTech; 4. In dev. by Janssen; 5. In dev. by Novartis; 6. In dev. by Horizon Therapeutics

Delivering on Genmab's Promise: Innovating Antibodies, Improving Lives



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 next-generation 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

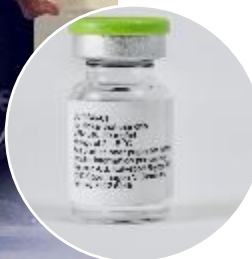
Advancing Pipeline: Delivering on Our Promise & Creating Value

Accelerating Development of Potential “Next Winners”



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

Advancing Pipeline: Delivering on Our Promise & Creating Value



**Delivering on
Genmab's
Promise to
Patients**



Bolstering early stage portfolio

• DuoBody-CD40x4-1BB¹; DuoHexaBody-CD37²; DuoBody-CD3x5T4²; HexaBody-CD38³



Adding new technologies



Data sciences



Expanding early stage discovery programs



Enhancing clinical development capabilities

Track Record of Success

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-socks-off 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		Target-mediated enhanced hexamerization	Enhanced potency
DuoHexaBody		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 Advanced Development Phase					
				Pre-Clinical	1	1/2	2	3	Approved
Tisotumab vedotin	TF	50:50 Genmab / Seattle Genetics	Cervical cancer						
			Ovarian cancer						
			Solid tumors						
Enapotamab vedotin	AXL	Genmab	Solid tumors						
Epcoritamab (DuoBody-CD3xCD20)	CD3, CD20	50:50 Genmab / AbbVie	Hematological malignancies						
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							

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) Daratumumab	CD38	Janssen (Tiered royalties to Genmab on net global sales)	Multiple myeloma ¹						
			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 Advanced Development Phase					
				Pre-Clinical	1	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

innovaTV 204

Recurrent or metastatic cervical cancer

- Potentially registrational 102 pts
- Single arm, monotherapy
- 1° endpoint: confirmed ORR
- 2° endpoints: duration of response, PFS, OS

innovaTV 205

Recurrent or metastatic cervical cancer

- In combo or mono
w/ bevacizumab, pembrolizumab, or carboplatin or weekly monotherapy recurrent or stage IVB cervical cancer
- Up to 170 pts
- 1° endpoint: ORR
- 2° endpoints: Safety, duration of response, time to response, PFS, OS

innovaTV 207

Solid tumors

- Basket study
- Up to 250 pts
- Single arm, monotherapy
- 1° endpoint: ORR
- 2° endpoints: Safety, disease control rate, duration of response, time to response, PFS, OS

innovaTV 208

Ovarian cancer

- Ovarian cancer, fallopian tube cancer, peritoneal cancer
- Up to 182 pts, incl 12 pt safety run-in
- Monotherapy
- 2 schedules: q3wk & dose dense
- 1° endpoints: Safety & ORR

Tisotumab Vedotin

Cervical Cancer Market Size

United States³

New Diagnoses	Deaths
12,578	4,115

3rd most common gynecologic cancer in US⁴

Japan⁶

New Diagnoses	Deaths
9,390	3,654

2nd most common gynecologic cancer in Japan⁶

Europe²

New Diagnoses	Deaths
58,373	24,404

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



Incorporates
proprietary
HexaBody
technology



Highly promising
data pre-clinical
models for MM,
lymphoma & AML



Could potentially
add to and broaden
DARZALEX
franchise



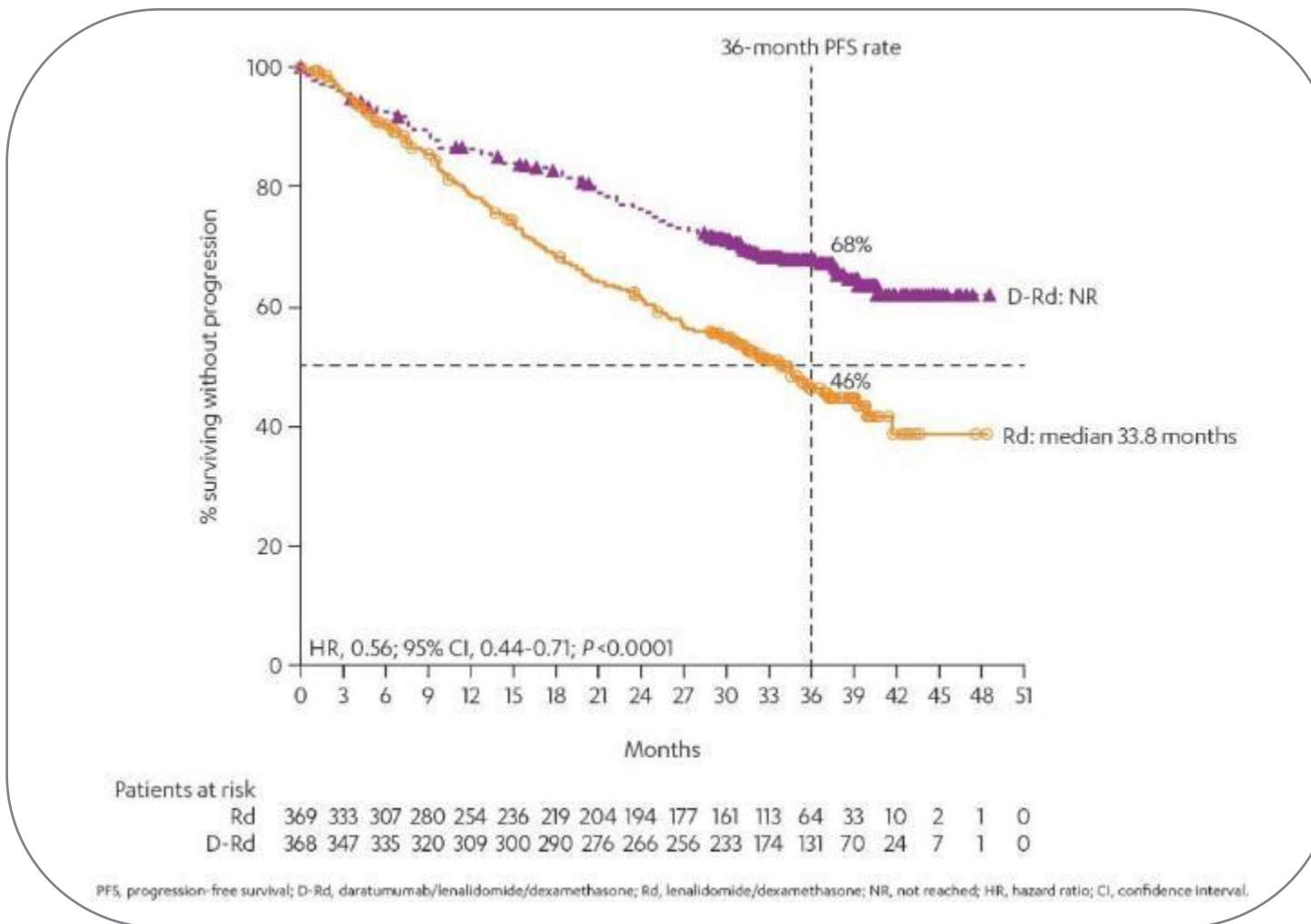
IND/CTA planned
in H2 2020

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	✓ AQUILA				
	Monotherapy	✓ CENTAURUS				
Front line MM (transplant & non-transplant)	Dara + VRd	✓ CEPHEUS				
	Dara + VMP (Asia Pacific)	✓ OCTANS				
	Dara + VRd	✓ PERSEUS				
	Dara + R (maintenance)	AURIGA				
		NINLARO® (Ph II), Venclexta® (Ph II), Selinexor (Ph I/II)				
Relapsed or Refractory MM	Dara + combinations	Opdivo® (Ph I/II), Tecentriq® (Ph I)				
	Dara + I.O. (PD1 & PDL1)					
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
- 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 T1E 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

