

The background features several complex molecular structures, likely representing pharmaceutical compounds, rendered in a semi-transparent style. These structures are primarily in shades of blue and purple, with some appearing as glowing spheres connected by lines. They are scattered across the dark background, with a concentration on the left and right sides, framing the central text.

J.P. Morgan Healthcare Conference

January 12, 2026

REGENERON[®]

This non-promotional presentation contains investigational data as well as forward-looking statements; actual results may vary materially.

J.P. Morgan Healthcare Conference



Dr. Leonard Schleifer, MD, PhD

Board Co-Chair, Co-Founder,
President, & Chief Executive Officer



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Note regarding forward-looking statements and non-GAAP financial measures

This presentation includes forward-looking statements that involve risks and uncertainties relating to future events and the future performance of Regeneron Pharmaceuticals, Inc. ("Regeneron" or the "Company"), and actual events or results may differ materially from these forward-looking statements. Words such as "anticipate," "expect," "intend," "plan," "believe," "seek," "estimate," variations of such words, and similar expressions are intended to identify such forward-looking statements, although not all forward-looking statements contain these identifying words. These statements concern, and these risks and uncertainties include, among others, competing drugs and product candidates that may be superior to, or more cost effective than, products marketed or otherwise commercialized by Regeneron and/or its collaborators or licensees (collectively, "Regeneron's Products") and product candidates being developed by Regeneron and/or its collaborators or licensees (collectively, "Regeneron's Product Candidates") (including biosimilar versions of Regeneron's Products); uncertainty of the utilization, market acceptance, and commercial success of Regeneron's Products and Regeneron's Product Candidates and the impact of studies (whether conducted by Regeneron or others and whether mandated or voluntary) or recommendations and guidelines from governmental authorities and other third parties or other factors beyond Regeneron's control on the commercial success of Regeneron's Products and Regeneron's Product Candidates; the nature, timing, and possible success and therapeutic applications of Regeneron's Products and Regeneron's Product Candidates and research and clinical programs now underway or planned, including without limitation EYLEA HD® (afibercept) Injection 8 mg, EYLEA® (afibercept) Injection, Dupixent® (dupilumab), Libtayo® (cemiplimab), Praluent® (alirocumab), Kevzara® (sarilumab), Evkeeza® (evinacumab), Veopoz® (pozelimab), Ordspono™ (odronextamab), Lynozyfic™ (linvoseltamab), other clinical programs discussed in this presentation, Regeneron's and its collaborators' earlier-stage programs, and the use of human genetics in Regeneron's research programs; the likelihood and timing of achieving any of the anticipated milestones discussed or referenced in this presentation; safety issues resulting from the administration of Regeneron's Products and Regeneron's Product Candidates in patients, including serious complications or side effects in connection with the use of Regeneron's Products and Regeneron's Product Candidates in clinical trials; the likelihood, timing, and scope of possible regulatory approval and commercial launch of Regeneron's Product Candidates and new indications for Regeneron's Products, such as those listed above; the extent to which the results from the research and development programs conducted by Regeneron and/or its collaborators may be replicated in other studies and/or lead to advancement of product candidates to clinical trials, therapeutic applications, or regulatory approval; ongoing regulatory obligations and oversight impacting Regeneron's Products, research and clinical programs, and business, including those relating to patient privacy; 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changes to drug pricing regulations and requirements and Regeneron's drug pricing strategy; other changes in laws, regulations, and policies affecting the healthcare industry; unanticipated expenses; the costs of developing, producing, and selling products; Regeneron's ability to meet any of its financial projections or guidance and changes to the assumptions underlying those projections or guidance; Regeneron's estimates of market opportunities for Regeneron's Products and Regeneron's Product Candidates; the potential for any license or collaboration agreement, including Regeneron's agreements with Sanofi and Bayer (or their respective affiliated companies, as applicable), to be cancelled or terminated; the impact of public health outbreaks, epidemics, or pandemics on Regeneron's business; and risks associated with litigation and other proceedings and government investigations relating to the Company and/or its operations (including the pending civil proceedings initiated or joined by the U.S. Department of Justice and the U.S. Attorney's Office for the District of Massachusetts), risks associated with intellectual property of other parties and pending or future litigation relating thereto (including without limitation the patent litigation and other related proceedings relating to EYLEA), the ultimate outcome of any such proceedings and investigations, and the impact any of the foregoing may have on Regeneron's business, prospects, operating results, and financial condition. A more complete description of these and other material risks can be found in Regeneron's filings with the U.S. Securities and Exchange Commission. Any forward-looking statements are made based on management's current beliefs and judgment, and the reader is cautioned not to rely on any forward-looking statements made by Regeneron. Regeneron does not undertake any obligation to update (publicly or otherwise) any forward-looking statement, including without limitation any financial projection or guidance, whether as a result of new information, future events, or otherwise.

This presentation includes projected 2026 non-GAAP R&D expense, which is a financial measure that is not calculated in accordance with U.S. Generally Accepted Accounting Principles ("GAAP"). This and other non-GAAP financial measures are computed by excluding certain non-cash and/or other items from the related GAAP financial measure. The Company also includes a non-GAAP adjustment for the estimated income tax effect of reconciling items. The Company makes such adjustments for items the Company does not view as useful in evaluating its operating performance. Management uses this and other non-GAAP measures for planning, budgeting, forecasting, assessing historical performance, and making financial and operational decisions, and also provides forecasts to investors on this basis. Additionally, such non-GAAP measures provide investors with an enhanced understanding of the financial performance of the Company's core business operations. However, there are limitations in the use of such non-GAAP financial measures as they exclude certain expenses that are recurring in nature. Furthermore, the Company's non-GAAP financial measures may not be comparable with non-GAAP information provided by other companies. Any non-GAAP financial measure presented by Regeneron should be considered supplemental to, and not a substitute for, measures of financial performance prepared in accordance with GAAP. A reconciliation of the non-GAAP financial measure used in this presentation is provided herein.

REGENERON

SCIENCE TO MEDICINE®

RGC
Regeneron Genetics Center

Integrating Genetics, Proteomics, and Big Data

World's largest DNA and proteomics-linked healthcare database, enabling advanced drug discovery, development, and healthcare analytics



Accelerating Innovation and R&D Productivity

Powerful toolkit of proprietary, turnkey technology platforms provides enduring competitive advantages

VELOCIMMUNE®
Leaders in human antibodies

VELOCI-BI®
Pioneers in bispecifics

Genetics Medicines

siRNA | gene editing | AAV gene therapy

Following the Science

~45 clinical programs across six core therapeutic areas provides a strong foundation for future growth



Delivering Breakthrough Medicines

14 internally-discovered therapies have been approved, poised to deliver many more...

DUPIXENT®
(dupilumab)

LIBTAYO®
(tisotumumab-axg)

LYNZOZYIC®
(lynestiprone-pyruvate)

EYLEA HD®
(aflibercept) injection 4mg

EYLEA®
(aflibercept) injection 2mg

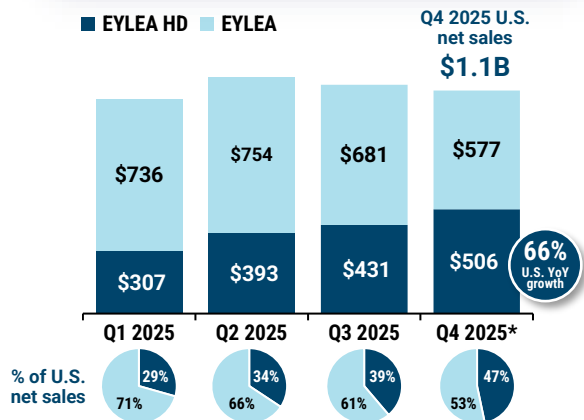
**Leveraging the power of science to bring transformative medicines to patients...
over and over again**

Portfolio of leading brands across diverse therapeutic areas

Delivering growth through leadership in key therapeutic categories



#1 in U.S. branded anti-VEGF category share



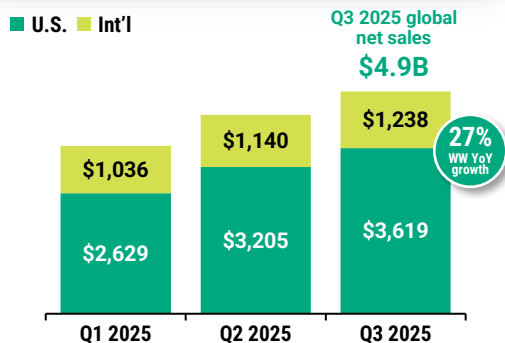
EYLEA HD physician demand grew **10%*** (Q4 vs. Q3)

FDA resubmission to include new EYLEA HD PFS filler completed; Q2 2026 decision anticipated

FDA approved addition of new EYLEA HD vial filler



#1 prescribed biologic for Type 2 inflammatory diseases



Sanofi records Dupixent global net sales

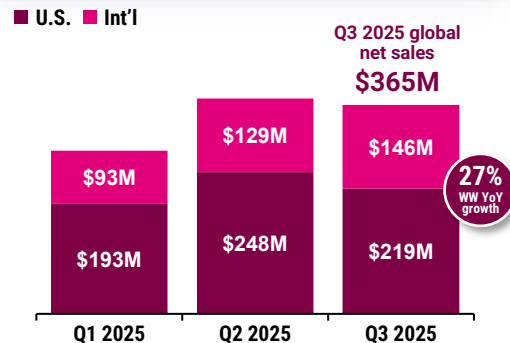
>1.3 million patients on therapy globally

#1 position in both **NBRx** and **TRx** in all established indications

Strong momentum from recent respiratory (**COPD**) and dermatology (**CSU, BP**) launches



#1 prescribed PD-1 antibody for non-melanoma skin cancers



Leading PD-1 antibody in **advanced CSCC & BCC**

Only PD-1 antibody approved in **adjuvant CSCC**

#2 most prescribed I/O treatment for metastatic NSCLC patients in U.S.

Deploying capital to maximize long-term value creation

Disciplined capital allocation approach laying the foundation for Regeneron's next wave of innovation

Internal Investment



Investing in world-class R&D capabilities and infrastructure to support sustainable growth

~\$6B Non-GAAP R&D* spend expected in 2026

\$7B+ committed to U.S. manufacturing and R&D infrastructure expansion over the coming years

Business Development



Leveraging external innovation to complement internal R&D

Expand through **complementary opportunities** across early and late development stages

- Collaboration with Alnylam, including in-licensing of **cemdisiran (C5 siRNA)**
- **GLP-1/GIP** in-licensed for obesity franchise expansion[†]
- Global collaborations for investigative **gene editing** therapies with Intellia, Mammoth and Tessera[‡]

Return Capital to Shareholders



Rewarding shareholders through opportunistic share repurchases and dividends

\$3.8B Capital returned to shareholders in 2025[§]

~\$3.4B share repurchases
~\$0.4B dividends

Quarterly dividend initiated in 2025 (\$0.88/share)

* Reflects estimated Non-GAAP R&D expense, which excludes SBC. GAAP R&D expense for 2026 is estimated to be ~\$6.5 billion. Formal financial guidance will be provided at Q4 2025 earnings.

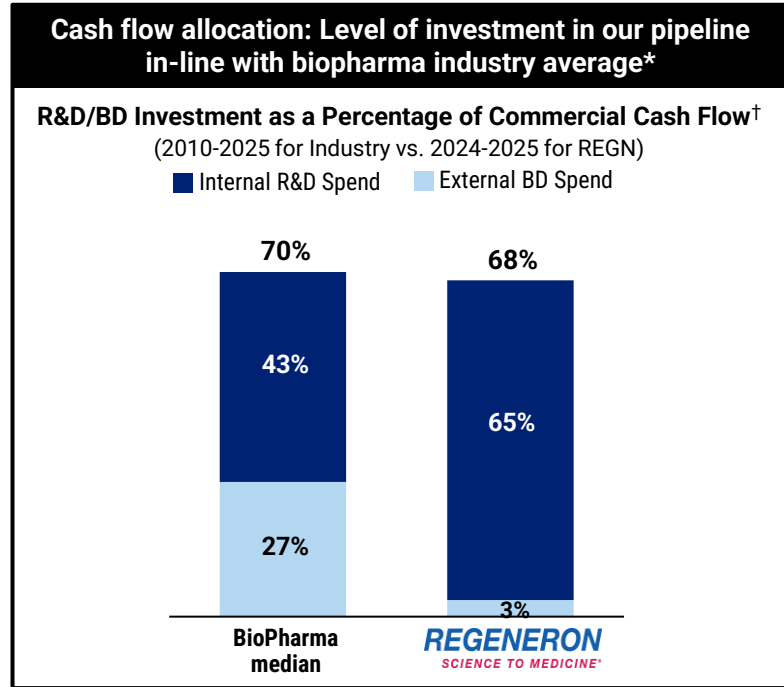
[†] License agreement with Hansoh Pharma.

[‡] Global collaboration with Tessera Therapeutics, Inc. is subject to customary closing conditions, including applicable regulatory agency clearances under the Hart-Scott-Rodino Antitrust Improvements Act of 1976 in the U.S.

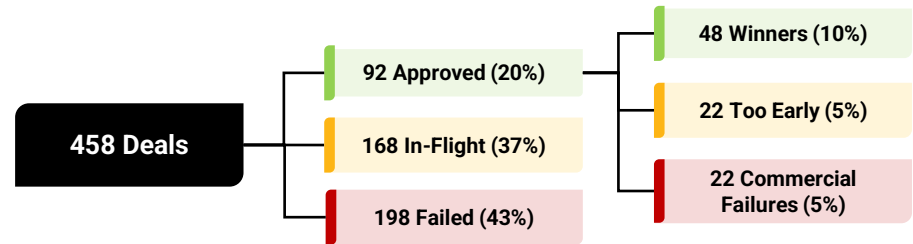
[§] Based on preliminary, unaudited results. As of December 31, 2025, ~\$1.5B was remaining under current share repurchase program.

Driving shareholder value with internal innovation while continuing our disciplined and opportunistic approach to business development

Over-reliance on business development (BD) to build pipelines poses challenges to delivering long-term shareholder value



Regeneron internal analysis suggests that a large majority of the 450+ Big Biopharma deals‡ since 2010 could end up being failures§



Looking at the return on the ~\$350B+ that was spent on the 290 deals where the outcome is now known

Internal Rate of Return (IRR) on 290 deals

~8% Overall | 4% M&A deals (n=125) | 18% Licensing deals (n=165)

* Biopharma group includes Abbvie, Amgen, AstraZeneca, Biogen, Bristol Myers Squibb, Celgene, GSK, Eli Lilly, Gilead, Merck, Pfizer, Sanofi, Novartis, Roche, J&J, and Novo Nordisk.

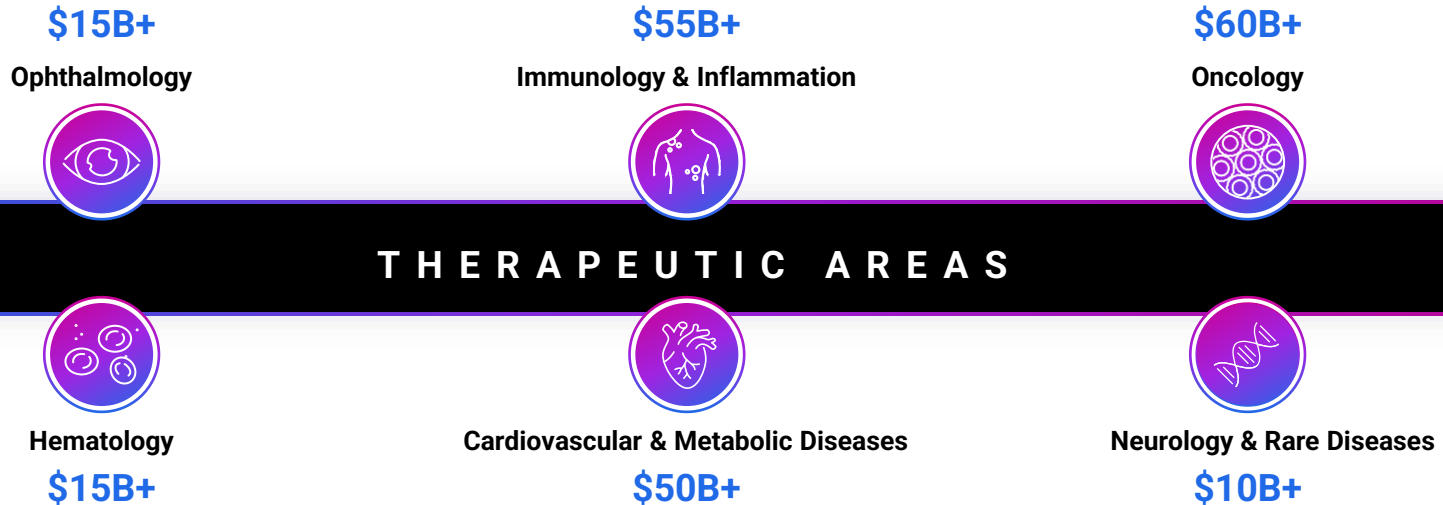
† Commercial cash flow is calculated as cash from operations before R&D spend; figures for Biopharma companies are based on 2010-2025 reported actuals; Internal R&D spend reflects reported GAAP figures; External BD spend includes upfront payments and all contingent milestone payments that we estimate will be incurred.

‡ Includes all M&A and licensing deals from 2010-2025 with >\$50M upfront for assets at IND stage or later; excludes discovery deals, deals for platform technologies, and deals for commercial stage assets.

§ Criteria for classifying the approved deals – Winner: >\$500M estimated peak WW sales, Commercial Failure: <\$500M estimated peak WW sales, Too Early: Launch 2023 or later (unless already at >\$500M WW sales, in which case deal is classified as a winner).

Source: Regeneron internal analysis

Regeneron pipeline targets large market opportunities across key therapeutic categories



THERAPEUTIC AREAS

Global market opportunity ~\$200B annually by 2030

Regeneron pipeline targets large market opportunities across key therapeutic categories

Ophthalmology

Cemdisiran (C5 siRNA) ± Pozelimab (C5 Ab)*	Geographic atrophy
REGN7041 (CD3)	Uveitis
Undisclosed Target	Glaucoma
Undisclosed Target	Thyroid Eye Disease, Graves



Immunology & Inflammation

Cemdisiran (C5 siRNA)*	gMG
IL-13	Type 2 Indications
IL-4	Type 2 Indications
IL-4xIL-13 bispecific§	Type 2 Indications
REGN1908-1909 (FelD1)	Cat Allergy
REGN5713-5715 (BetV1)	Birch Allergy
Multiple Agents§	Food Allergy
Itepekimab (IL-33)†	COPD, CRSwNP
Undisclosed Target	Lupus, Sjogren's, PBC, others



Oncology

Lyenzoic (BCMAxCD3)	Multiple myeloma
Fianlimab (LAG3) + Libtayo (PD-1)	1L metastatic melanoma, adjuvant melanoma
Ordspono (CD20xCD3)	Lymphoma
Ubamatomab (MUC16xCD3)	Ovarian Cancer



THERAPEUTIC AREAS



Hematology

Cemdisiran (C5 siRNA) ± Pozelimab (C5 Ab)*	Paroxysmal nocturnal hemoglobinuria
REGN7508 ^{CAT} (FXI)	Post-TKR VTE, Cancer VTE, PICC-associated thrombosis, SPAF, PAD
REGN9933 ^{A2} (FXI)	PICC-associated thrombosis, SPAF, PAD



Cardiovascular & Metabolic Diseases

Olatorepatide (GIP/GLP-1)	Obesity, T2D
Olatorepatide (GIP/GLP-1) + Praluent (PCSK9)	Obesity, T2D with dyslipidemia
GLP-1 + Trevogrumab (GDF8)	Muscle Sparing
Nex-z (TTR)†	ATTR
MASH siRNA* (CIDEB, PNPLA3, HSD17B13)	MASH



Neurology & Rare Diseases

DB-OTO (AAV-based gene therapy)	Hearing loss
Garetosmab (Activin A)	FOP
SNCA siRNA*	Parkinson's Disease
SOD1 siRNA*	ALS
MAPT (Tau) siRNA*	Alzheimer's Disease
HTT siRNA*	Huntington's Disease

Sustaining I&I leadership and unlocking new growth opportunities

Leveraging learnings from Dupixent and disease biology to advance next-gen approaches to treat inflammatory diseases

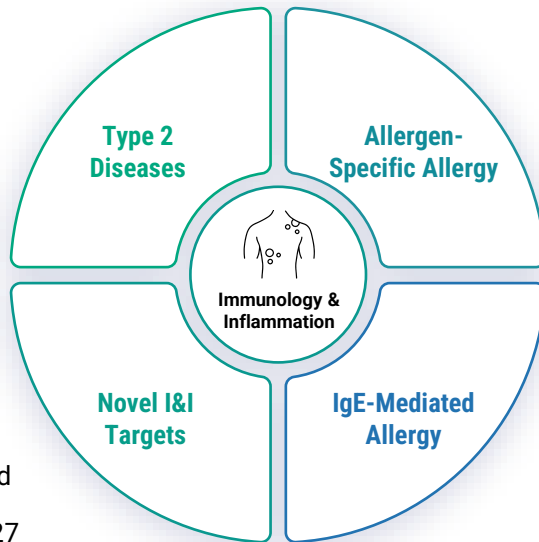
Pursuing multi-pronged approach to sustain I&I leadership into the next decade

'Lifecycle' opportunities

- **Longer Dupixent*** dosing intervals
- **Novel long-acting IL-4Ra⁺** antibody
- Long-acting, fully-human **IL-13 & IL-4** antibodies with optimized binding properties
 - Expedited AD development plan for IL-13; FIH expected in 1H 2026
- Long-acting **IL-4xIL-13 bispecific**

Investigating novel I&I targets

- **Itepekimab* (IL-33)**: Advancing in respiratory indications with strong genetic associations
 - Phase 3 CRSwNP data anticipated in 2027
- Additional **genetic-defined targets** discovered by RGC, each with pipeline-in-a-product potential, expected to enter clinic in 2026-2027



Advancing broader allergy pipeline into large commercial opportunities

Allergen-specific antibody approaches

- **Cat (FelD1)** and **birch (BetV1)** allergy programs each demonstrated positive Phase 3 results in 2025
- Registration-enabling studies initiating in 2026 for both programs; data anticipated in 2027

Severe IgE-mediated food allergy

- **Lynozifyf (BCMAxCD3) + Dupixent*** achieved proof-of-principle; demonstrated sustained >90% reductions in IgE in 4 of 4 evaluable patients
- Advancing **novel therapeutic candidates** to develop more-targeted and/or specific approaches to potentially **eliminate IgE-mediated allergies**; FIH expected by 2027

* In collaboration with Sanofi

† Covered by the Sanofi collaboration

Key programs positioned to deliver over the next few years

Late-stage opportunities spanning multiple therapeutic areas

FIANLIMAB + LIBTAYO

LAG-3 + PD-1

Combining two potentially best-in-class checkpoint inhibitors

- Potential for **differentiated efficacy** vs. current standards-of-care in **melanoma** without exacerbating safety

Program Status

Pivotal data from **1L metastatic melanoma** trial anticipated in **1H 2026**



BCMAxCD3

Transform the **multiple myeloma** treatment paradigm

- **Monotherapy** & simplified combinations in **early-line** myeloma settings
- Goal to **prevent** myeloma by treating precursor conditions

Program Status

4 registrational studies underway, 4 more expected to initiate in 2026

Pivotal data anticipated starting in 2027

CEMDISIRAN ± POZELIMAB

C5 siRNA ± C5 antibody

PNH: combination approach for complete C5 blockade and potentially best-in-class efficacy

gMG: siRNA monotherapy delivers potentially best-in-class efficacy and convenience

GA: monotherapy and combination approaches being explored

Program Status

gMG: on track for FDA submission in **Q1 2026**

PNH: pivotal data expected in **Q4 2026/Q1 2027**

GA: initial results from lead-in cohort anticipated in **2H 2026**

REGN7508 & REGN9933

Two Factor XI antibodies allow for customized approach

REGN7508^{Ca1}: **optimizes anticoagulation activity** with reduced bleeding risk vs. SOC

REGN9933^{A2}: effective anticoagulation with further **reduced bleeding risk**

Program Status

2 registrational studies underway, 6 more expected to initiate in 2026

Pivotal data anticipated starting in 2027

OLATOREPATIDE (OLA) ± VARIOUS AGENTS

GIP/GLP-1, combinations

Multi-faceted approach including GIP/GLP-1

Prioritizing combo with Praluent (PCSK9): potential to achieve >50% LDL lowering along with weight loss, dosed via similarly-convenient weekly injection as leading GLP-1s

Program Status

Phase 3 results for Ola in obesity in China* expected in 1H 2026

Comprehensive global clinical development plan initiating in 2026

*Hansoh Pharmaceuticals retains development and commercialization rights to olatorepatide in China.

This slide contains investigational drug candidates that have not been approved by any regulatory authority.

Combining two potentially best-in-class checkpoint inhibitors: Fianlimab (anti-LAG-3) & LIBTAYO (anti-PD-1)

Potentially differentiated 1L metastatic melanoma treatment option; additional data readouts across other settings expected in 1H 2026

1H 2026
Anticipated Milestones:

Phase 3 1L metastatic melanoma data

Phase 3 adjuvant melanoma data (1st interim)


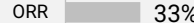
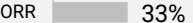
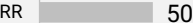
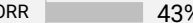
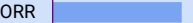


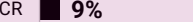


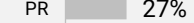

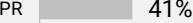
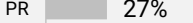
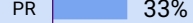






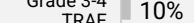
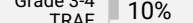


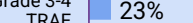
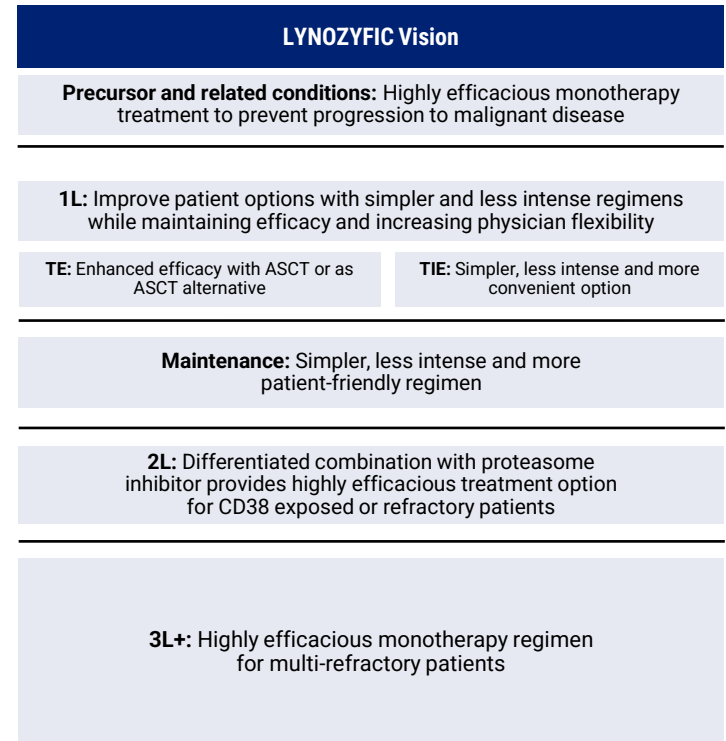
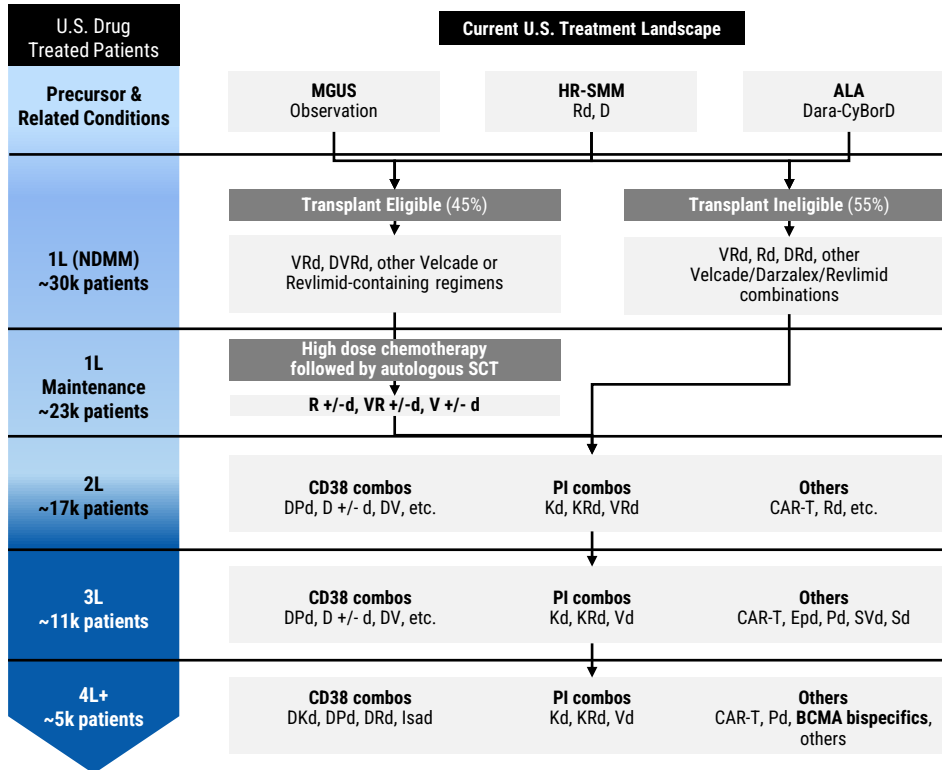
	Pembrolizumab (anti-PD-1) KEYNOTE-006 n=277 (Q3W)	Nivolumab (anti-PD-1) RELATIVITY-047 n=359	Ipilimumab (anti-CTLA4) + nivolumab CHECKMATE-067 n=314	Relatlimab (anti-LAG-3) + nivolumab RELATIVITY-047 n=355	Fianlimab + cemiplimab Pooled POC Cohorts n=98
 Efficacy	ORR  33%	ORR  33%	ORR  50%	ORR  43%	ORR  57%
	CR  6%	CR  14%	CR  9%	CR  16%	CR  25%
	PR  27%	PR  18%	PR  41%	PR  27%	PR  33%
mPFS (months)	4.1	4.6	11.7	10.1	24 (KM estimate)
mOS (months)	Not Reached	34.1	Not Reached	Not Reached	Not Reached
 Safety	All TRAE  73%	All TRAE  70%	All TRAE  96%	All TRAE  81%	All TRAE  81%
	Grade 3-4 TRAE  10%	Grade 3-4 TRAE  10%	Grade 3-4 TRAE  59%	Grade 3-4 TRAE  19%	Grade 3-4 TRAE  23%
Follow up	OS: final analysis with an additional FU of 9 mo	At the time of the final OS analysis	Minimum FU: 9 mo for ORR, 28 mo for PFS, 48 mo for OS	At the time of the final OS analysis	Median FU: 23 mo
Source	KEYTRUDA U.S. FDA PI; Robert et al., 2015 NEJM	OPDUALAG U.S. FDA PI; Tawbi et al., 2022 NEJM	YERVOY & OPDIVO U.S. FDA PI; Wolchok et al., 2017 NEJM	OPDUALAG U.S. FDA PI; Tawbi et al., 2022 NEJM	ESMO 2024 Data

Table depicts randomized Phase 3 data for four FDA-approved treatments in 1L metastatic melanoma as well as pooled, post-hoc data from three independent cohorts from initial trial of fianlimab + cemiplimab; there are no randomized, head-to-head clinical trials between these products. Study data being provided for descriptive purposes only. Caution is advised when drawing conclusions based on cross-trial comparisons.

*This slide contains investigational data for the combination of fianlimab + cemiplimab; this combination has not been approved by any regulatory authority. All other products listed are FDA-approved therapies.

Aiming to transform the multiple myeloma treatment landscape

4 registrational studies underway to potentially transform the treatment paradigm with convenient, simplified and less intense treatment regimens
 At Linozyfic 200 mg monotherapy, 100% of evaluable patients (n=21) achieved MRD-negativity in HRSMM and 1L multiple myeloma



Tailored C5 therapeutic approach: siRNA ± antibody provides flexibility to address multiple complement-mediated diseases

siRNA (cemdisiran) lowers C5 target burden while antibody (pezelimab) blocks circulating C5, enabling near-complete C5 inhibition

Paroxysmal Nocturnal Hemoglobinuria

2025 U.S. Prevalence (patients): ~6k
Worldwide market sales* (2025e): ~\$2.0B
Estimated market sales CAGR* (2025-2030): ~12%



- Cemdisiran + pezelimab combination demonstrated potential best-in-class profile (Phase 3 lead-in cohort presented at ASH 2024)
- Combination maximizes C5 inhibition and minimizes hemolysis
- Convenient monthly subcutaneous administration
- Ongoing Phase 3 study, data expected by Q4 2026/Q1 2027

Myasthenia Gravis

2025 U.S. Prevalence (patients): ~85k
Worldwide market sales* (2025e): ~\$5.0B
Estimated market sales CAGR* (2025-2030): ~17%



- Cemdisiran monotherapy reported best MG-ADL improvement among C5 inhibitors[†] (Phase 3 data)
- Convenient Q3M subcutaneous administration
- FDA submission for cemdisiran planned for Q1 2026 with decision expected by Q4 2026/Q1 2027

Geographic Atrophy

2025 U.S. Prevalence (patients): ~1.1M
Worldwide market sales* (2025e): ~\$1.1B
Estimated market sales CAGR* (2025-2030): ~34%



- Both cemdisiran monotherapy and cemdisiran + pezelimab combination being evaluated in ongoing Phase 3 pivotal program (initiated in 2H 2024)
- Interim data from Phase 3 lead-in cohort anticipated in 2H 2026

Program Status

Differentiated siRNA ± antibody approach has pipeline-in-a-product potential to deliver tailored, effective, and convenient treatments across multiple complement-mediated diseases

Addressing the bleeding risk in anticoagulation treatment: Regeneron's broad Factor XI clinical program

\$20B anticoagulation market remains underpenetrated due to bleeding risk; <50% of eligible patients receive therapy because of safety concerns

Regeneron's two antibodies allow customized approach: **REGN7508^{Cat}** optimizes anticoagulation activity with reduced bleeding risk vs. SOC, **REGN9933^{A2}** further reduces bleeding risk with comparable anticoagulation vs. SOC



Post-TKR
VTE

R7508

Two trials enrolling,
data expected in 2027



Cancer
VTE

R7508

Trials initiating in 1H 2026,
data expected in 2029+



PICC-associated
Thrombosis

R7508

R9933

Trial to initiate in 2026,
data expected in 2028+



Stroke
Prevention
in AF

R7508

R9933

Phase 2 enrolling, data expected in 2027
Phase 3 trials to initiate in 2026



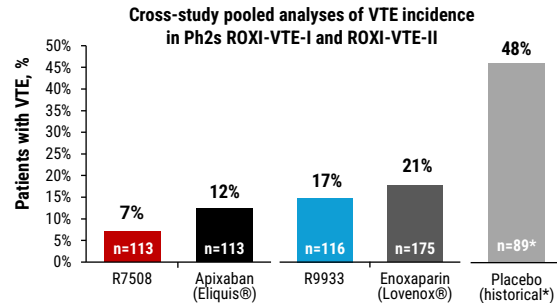
Peripheral
Artery Disease

R7508

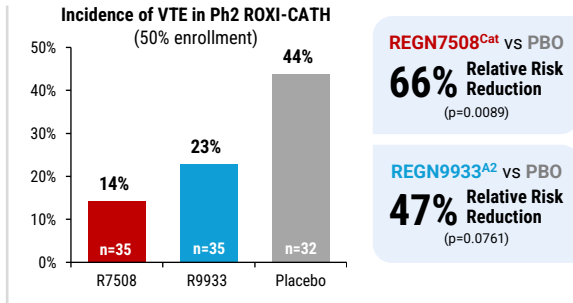
R9933

Trial initiating in 1H 2026,
initial data expected in 2029+

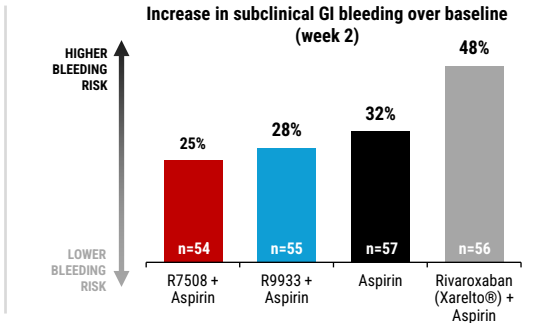
Phase 2 results in VTE prevention post-knee replacement surgery support broad Phase 3 development



Interim Phase 2 results in catheter-associated thrombosis support development in contact-mediated settings



Phase 1 GI Bleed Study results support favorable bleeding profile in a healthy volunteer provoked bleeding model



To date, no major bleeding events observed in Phase 1 or Phase 2 studies due to REGN7508 or REGN9933

Transforming patient care for obesity and related conditions

Three major opportunities for Regeneron in the rapidly growing obesity therapeutic area

1



GIP/GLP-1 Receptor Agonist monotherapy

In-licensing of olatorepatide (dual GIP/GLP-1 receptor agonist) enables initial monotherapy development

- Phase 3 program in obesity with and without T2D to initiate in 2026

Monotherapy

2



Address obesity comorbidities with novel combinations

Initiating olatorepatide combo with Praluent (PCSK9) in 2026:

- Approved GLP-1s lower LDL-C by less than 10%
- Combination to potentially achieve >50% LDL lowering along with weight loss
- To be administered via similarly-convenient weekly injection as leading GLP-1s

Novel combinations

3



Enhancing the quality of GLP-1-based weight loss

- Harness beneficial effects of muscle preservation in obesity
- POC data on anti-myostatin ± anti-activin A warrant potential future development
- Unimolecular solutions in preclinical development

Improving quality of weight loss

Q&A



**Dr. Leonard
Schleifer, MD, PhD**

Board Co-Chair,
Co-Founder, President,
and Chief Executive Officer



**Dr. George
Yancopoulos, MD, PhD**

Board Co-Chair,
Co-Founder, President,
and Chief Scientific Officer

2026 key milestones

Ophthalmology

- **EYLEA HD**: pre-filled syringe (PFS) FDA decision (2Q26)
- **Cemdisiran ± pozelimab**: interim results from lead in cohort of Phase 3 trial in GA (2H26)

Immunology & Inflammation

- **Cemdisiran**: NDA submission for gMG (1Q26); FDA decision (4Q26 / 1Q27)
- **Dupixent**: EC decision for BP (1H 2026), FDA decision for AFRS (Q1 2026)
- **IL-13**: Initiate clinical program in atopic dermatitis
- **R5713-5715**: Initiate second Phase 3 trial for birch allergy (1H26)
- **R1908-1909**: Initiate second Phase 3 trial for cat allergy (1H26)

Cardiovascular & Metabolic Diseases

- **Muscle preservation**: Report additional data from proof-of-concept data of combination of semaglutide and trevogrumab with and without garetosmab in obesity (2026)
- **Olatorepatide (monotherapy)**: Initiate Phase 3 program in obesity with and without T2D (2026)
- **Olatorepatide + Praluent**: Initiate clinical program (2026)

Hematology

- **R7508/R9933**: Initiate additional Phase 3 trials in anticoagulation (1H26)
- **Cemdisiran ± Pozelimab**: report results from Phase 3 trial in PNH (4Q26 / 1Q27)

Oncology

Solid Oncology

- **Fianlimab + cemiplimab**: Report results in 1L metastatic melanoma from Ph3 trial (1H26)
- **Fianlimab + cemiplimab**: Report initial Phase 2 data in 1L advanced NSCLC (1H26)

Heme-onc

- **Lynozytic**: Initiate additional Ph3 studies in multiple myeloma and precursor conditions (2026)

Neurology & Rare Diseases

- **DB-OTO**: FDA decision for genetic hearing loss (1H26)
- **Garetosmab**: FDA and EC decisions in FOP (2H26)

Abbreviations and definitions

Abbreviation	Definition
1L	First line
AAV	Adeno-associated virus
ALA	Amyloid light-chain amyloidosis
ALS	Amyotrophic lateral sclerosis
AI	Artificial Intelligence
AD	Atopic Dermatitis
AATR	Transthyretin amyloidosis
BCC	Basal cell carcinoma
BCMA	B-cell maturation antigen
BP	Bullous pemphigoid
CAR-T	Chimeric antigen receptor T-cell
CAGR	Compounded annual growth rate
CI	Confidence Interval
COPD	Chronic obstructive pulmonary disease
CR	Complete response
CRSwNP	Chronic sinusitis with nasal polyposis
CSCC	Cutaneous squamous cell carcinoma
CSU	Chronic spontaneous urticaria
DOAC	Direct oral anticoagulants
ESMO	European Society for Medical Oncology
EC	European Commission
FDA	U.S. Food And Drug Administration
FIH	First in human
FU	Follow-up
FOP	Fibrodysplasia Ossificans Progressiva

Abbreviation	Definition
GA	Geographic atrophy
GIP	Gastric inhibitory polypeptide
GLP-1	Glucagon-like peptide 1
gMG	Generalized myasthenia gravis
HR-SMM	High-Risk Smoldering Multiple Myeloma
HR	Hazard Ratio
HTT	Huntington
I/O	Immuno-Oncology
I&I	Immunology and Inflammation
IgE	Immunoglobulin-E
IND	Initial new drug application
KM	Kaplan-Meier curve
LAG-3	Lymphocyte-activation gene 3
LOF/GOF	Loss of function/ Gain of function
MAPT	Microtubule-associated protein tau
MASH	Metabolic Dysfunction-Associated Steatohepatitis
MGUS	Monoclonal gammopathy of unknown significance
M&A	Merger and Acquisitions
MM	Multiple myeloma
mOS	Median overall survival
mPFS	Median progression-free survival
MUC16	Mucin 16
NBRx	New to Brand Prescriptions
NDA	New Drug Application
NR	Not Reached
NSCLC	Non-small cell lung cancer

Abbreviation	Definition
ORR	Overall Response Rate
OS	Overall Survival
PAD	Peripheral Arterial Disease
PBC	Primary Biliary Cholangitis
PBO	Placebo
PD-1/PD-(L)1	Programmed cell death protein/(ligand) 1
PFS	Pre-filled Syringe
PFS	Progression Free Survival
PI	Prescribing Information
PICC	Peripherally Inserted Central Catheter
PNH	Paroxysmal nocturnal hemoglobinuria
POC	Proof-of-concept
PR	Partial response
R/R	Relapsed/Refractory
RGC	Regeneron Genetics Center
SBC	Stock-based compensation
SC	Subcutaneous
siRNA	Small interfering RNA
SOC	Standard of care
SPAF	Stroke Prevention in Atrial Fibrillation
T2D	Type 2 diabetes mellitus
TI	Transplant eligible
TIE	Transplant Ineligible
TKR	Total Knee Replacement
TRAE	Treatment-related adverse events
VEGF	Vascular endothelial growth factor
VTE	Venous thromboembolism