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, the nature, timing, and possible success and therapeutic applications of Regeneron’s products, product candidates, and research and clinical programs now underway or planned, including without limitation EYLEA® (aflibercept) Injection, Dupixent® (dupilumab) Injection, Praluent® (alirocumab) Injection, Kevzara® (sarilumab) Injection, Libtayo® (cemiplimab) Injection, fasinumab, evinacumab, Regeneron’s immuno-oncology programs (including its costimulatory bispecific portfolio), Regeneron’s earlier-stage product candidates, and the use of human genetics in Regeneron’s research programs; the extent to which the results from Regeneron’s research programs or preclinical testing may lead to advancement of product candidates to clinical trials or therapeutic applications; unforeseen safety issues resulting from the administration of products and product candidates in patients, including serious complications or side effects in connection with the use of Regeneron’s products and product candidates in clinical trials; the likelihood and timing of possible regulatory approval and commercial launch of Regeneron’s late-stage product candidates and new indications for marketed products, including without limitation EYLEA, Dupixent, Praluent, Kevzara, Libtayo, fasinumab, and evinacumab; the likelihood and timing of achieving any of the anticipated milestones described in this presentation; the extent to which the results from the research and development programs conducted by Regeneron or its collaborators may be replicated in other studies and lead to therapeutic applications; ongoing regulatory obligations and oversight impacting Regeneron’s marketed products (such as EYLEA, Dupixent, Praluent, Kevzara, and Libtayo), research and clinical programs, and business, including those relating to patient privacy; determinations by regulatory and administrative governmental authorities which may delay or restrict Regeneron’s ability to continue to develop or commercialize Regeneron’s products and product candidates; competing drugs and product candidates that may be superior to Regeneron’s products and product candidates; uncertainty of market acceptance and commercial success of Regeneron’s products and product candidates and the impact of studies (whether conducted by Regeneron or others and whether mandated or voluntary) on the commercial success of Regeneron’s products and product candidates; the availability and extent of reimbursement of the Company’s products from third-party payers, including private payer healthcare and insurance programs, health maintenance organizations, pharmacy benefit management companies, and government programs such as Medicare and Medicaid; coverage and reimbursement determinations by such payers and new policies and procedures adopted by such payers; the ability of Regeneron to manufacture and manage supply chains for multiple products and product candidates; the ability of Regeneron’s collaborators, suppliers, or other third parties to perform filling, finishing, packaging, labeling, distribution, and other steps related to Regeneron’s products and product candidates; unanticipated expenses; the costs of developing, producing, and selling products; the ability of Regeneron to meet any of its sales or other financial projections or guidance and changes to the assumptions underlying those projections or guidance, including financial guidance, revenue, GAAP financial measures, GAAP unreimbursed R&D, non-GAAP SG&A, effective tax rate, and capital expenditures; risks associated with intellectual property of other parties and pending or future litigation relating thereto, including without limitation the patent litigation proceedings relating to EYLEA, Dupixent, and Praluent, the ultimate outcome of any such litigation proceeding, and the impact any of the foregoing may have on Regeneron’s business, prospects, operating results, and financial condition; and the potential for any license or collaboration agreement, including Regeneron’s agreements with Sanofi, Bayer, and Teva Pharmaceutical Industries Ltd. (or their respective affiliated companies, as applicable), to be cancelled or terminated without any further product success. 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, including its Form 10-K for the fiscal year ended December 31, 2018 in the section thereof captioned “Item 1A. Risk Factors.” 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 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 uses non-GAAP unreimbursed R&D and non-GAAP SG&A, which are financial measures that are not calculated in accordance with U.S. Generally Accepted Accounting Principles (“GAAP”). These non-GAAP financial measures are computed by excluding certain non-cash and other items from the related GAAP financial measure. Non-GAAP adjustments also include the income tax effect of reconcile items. The Company makes such adjustments for items the Company does not view as useful in evaluating its operating performance. For example, adjustments may be made for items that fluctuate from period to period based on factors that are not within the Company’s control, such as the Company’s stock price on the dates share-based grants are issued. Management uses these 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 these and other 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 Company’s full year 2019 non-GAAP to GAAP financial guidance is provided on slide 26.
**KEY 2018 MILESTONES AND ACHIEVEMENTS**

**2018**

**RESEARCH & DEVELOPMENT**

**Key Regulatory Approvals***
- **LIBTAYO** Advanced CSCC
- **DUPIXENT** Moderate-to-severe Asthma
- **EYLEA** Q12 week dosing in wAMD after one year of effective therapy

**Key Regulatory Filings**
- **EYLEA** Diabetic Retinopathy
- **DUPIXENT** Atopic Dermatitis in adolescents
- **PRALUENT** Cardiovascular Risk Reduction

**Clinical Trial Readouts**
- **DUPIXENT** Ph3 Chronic Rhinosinusitis with Nasal Polyps
- **LIBTAYO** Ph1 Non Small Cell Lung Cancer
- **REGN1979 (CD20xCD3)** PoC in Follicular Lymphoma & Diffuse Large B-Cell Lymphoma
- **Fasinumab (NGF)** Ph3 Osteoarthritis
- **Pozelimab (C5)** Ph1 in Healthy Volunteers

**Ph2 and Ph3 Trial Initiations**
- **DUPIXENT** Ph2/3 Eosinophilic Esophagitis
- **Ph2** Grass Allergy
- **Ph2** Peanut Allergy
- **Ph2/3** AD in peds (6mo–5yr)
- **REGN3500 (IL-33)** Ph2 Chronic Obstructive Pulmonary Disease
- **Ph2** Asthma
- **Ph2** Atopic Dermatitis
- **KEVZARA** Ph3 Polymyalgia Rheumatica
- **Ph3** Giant Cell Arthritis

**INDs & Ph1 Trial Initiations**
- **REGN4018 (MUC16xCD3)** Ovarian Cancer
- **REGN5458 (BCMAxCD3)** Multiple Myeloma
- **REGN4659 (CTLA-4)** Cancer
- **REGN5069 (GFRα3)** Pain
- **REGN4461 (LEPR)** Metabolic Disease

**COMMERCIAL**

**Infectious Disease** Delivered REGN-EB3 to the Democratic Republic of the Congo for use in Ebola patients

**Genetics** Sequenced 500k human exomes to date

**New Partnerships/Collaborations** UK Biobank consortium, bluebird bio, Alnylam, Zoetis

**U.S. EYLEA** 2018 net sales of ~$4.08 Billion; 10% year-over-year growth

**DUPIXENT** Global 2018 net sales of $922 Million; Atopic Dermatitis launch continues to accelerate; Asthma launch progressing well, particularly among allergists

**LIBTAYO** U.S. 2018 net sales of $15 Million; physician interest and market uptake are encouraging

**PRALUENT** Working with payers to improve access and lower cost to patients

*Please see full Prescribing Information for all approved products.*

This slide includes pipeline drug candidates currently undergoing clinical testing in a variety of diseases. The safety and efficacy of these drug candidates have not been evaluated by any regulatory authorities for the disease categories described here.
EYLEA®: STRENGTHENING MARKET LEADERSHIP POSITION

Building on leadership position in wAMD and diabetic eye disease, both of which are increasing in prevalence

- We believe there are no near-term potential agents that can provide substantially different dosing flexibility, duration or visual gains than are already achievable with EYLEA

Label expansions and line extensions
Innovating next generation therapeutics

Our strategy is to maximize EYLEA growth opportunities and develop next generation therapeutics

Net Sales:

<table>
<thead>
<tr>
<th></th>
<th>4Q18</th>
<th>FY18</th>
</tr>
</thead>
<tbody>
<tr>
<td>U.S.</td>
<td>$1,078.9MM</td>
<td>$4,076.7MM</td>
</tr>
<tr>
<td>Global</td>
<td>$1,803.3MM</td>
<td>$6,745.6MM</td>
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</table>

U.S. Net Sales*, $Billion

<table>
<thead>
<tr>
<th>Year</th>
<th>2012</th>
<th>2013</th>
<th>2014</th>
<th>2015</th>
<th>2016</th>
<th>2017</th>
<th>2018</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>$0.8</td>
<td>$1.4</td>
<td>$1.7</td>
<td>$2.7</td>
<td>$3.3</td>
<td>$3.7</td>
<td>$4.1</td>
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</tbody>
</table>

* Outside the United States, EYLEA net product sales comprise sales by Bayer in countries other than Japan and sales by Santen Pharmaceutical Co., Ltd. in Japan under a co-promotion agreement with an affiliate of Bayer
Opportunities in Diabetic Eye Diseases

**Diabetic Macular Edema (DME)**
- Targeted commercial strategy to increase anti-VEGF penetration

**Diabetic Retinopathy (DR) without DME – PDUFA date May 13, 2019**
- Phase 3 PANORAMA study shows potential to change clinical practice
  - High unmet need: ~40% of untreated patients developed VTCs or CI-DME through week 52
- EYLEA reduced vision deterioration by more than 75% in the overall patient population
  - 72-76% reduction in VTCs and CI-DME: 10-11% EYLEA vs. 41% sham
  - 65-80% of EYLEA-treated patients experienced ≥ two-step improvement from baseline on the Diabetic Retinopathy Severity Scale (DRSS) vs. 15% sham
- Of the 3.5M people in the U.S. with DR without DME, ~1M individuals have moderately severe to severe disease and are at greatest risk

Next Generation Strategy

**Our strategy is to make even better treatments than our market-leading anti-VEGF therapy, EYLEA**
- High Dose Formulation of EYLEA
- Other new molecular entities and gene therapies

VTCs – Vision-threatening complications; CI-DME – center-involved diabetic macular edema

This slide includes pipeline drug candidates currently undergoing clinical testing in a variety of diseases. The safety and efficacy of these drug candidates have not been evaluated by any regulatory authorities for the disease categories described here.
**DUPIXENT®: BUILDING LEADERSHIP IN ATOPIC DERMATITIS AND LAUNCHING IN ASTHMA**

**Atopic Dermatitis: Practice-Changing Advance in Management**

- In the U.S., less than 15% of adult AD patients with the greatest need have used DUPIXENT
- High persistence and compliance indicate patient and physician satisfaction
- Ex-U.S. launch in early stage and progressing well
- Encouraging prescription trends following commencement of DTC TV campaign in 3Q18

**Moderate-to-Severe Asthma: High Unmet Need**

- Only asthma biologic approved for:
  - Self administration
  - Moderate-to-severe asthma with an eosinophilic phenotype
  - Oral corticosteroid-dependent asthma regardless of phenotype
  - AD patients with comorbid asthma
- Clinically meaningful improvements in lung function, asthma attacks and oral steroid sparing
- Up to 900K patients (≥12 years) in the U.S. with moderate-to-severe asthma may be suitable for biologic therapy
- Encouraging initial prescription trends, particularly among allergists treating asthma

---
*A Source: IQVIA
Please see full Prescribing Information for all approved products*
DUPIXENT®: DELIVERING ON THE “PIPELINE IN A PRODUCT” PROMISE

### APPROVED INDICATIONS

<table>
<thead>
<tr>
<th>Disease</th>
<th>Approval Status</th>
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</thead>
<tbody>
<tr>
<td>Atopic Dermatitis</td>
<td>Approved in Adults</td>
</tr>
<tr>
<td>Moderate-to-Severe Asthma</td>
<td>Approved in Adults and Adolescents</td>
</tr>
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</table>

### NEAR-TERM OPPORTUNITIES

<table>
<thead>
<tr>
<th>Disease</th>
<th>Milestone</th>
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<tbody>
<tr>
<td>Atopic Dermatitis in Adolescents (12–17 years)</td>
<td>PDUFA date March 11, 2019</td>
</tr>
<tr>
<td>Atopic Dermatitis in Pediatrics (6–11 years)</td>
<td>Ph3 results expected in 2019</td>
</tr>
<tr>
<td>Chronic Rhinosinusitis with Nasal Polyps (CRSwNP)</td>
<td>Two Positive Ph3 studies reported 2H18 sBLA filing expected in 1Q19</td>
</tr>
<tr>
<td>Eosinophilic Esophagitis</td>
<td>Positive Ph2 results; pivotal Ph2/3 initiated 3Q18</td>
</tr>
<tr>
<td>Chronic Obstructive Pulmonary Disease (COPD)</td>
<td>Initiate Ph2/3 in 2019</td>
</tr>
</tbody>
</table>

### LONGER-TERM OPPORTUNITIES

<table>
<thead>
<tr>
<th>Disease</th>
<th>Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pediatric Asthma (6-11 years)</td>
<td>Ph3 ongoing</td>
</tr>
<tr>
<td>Food Allergies</td>
<td>Ph2 in Peanut Allergy initiated; more planned</td>
</tr>
<tr>
<td>Airborne Allergies</td>
<td>Ph2 in Grass Allergy enrollment complete</td>
</tr>
<tr>
<td>Combinations with REGN3500 (IL-33)</td>
<td>Ph2 initiated in AD and Asthma; Asthma Ph2 results expected in 2019</td>
</tr>
</tbody>
</table>

This slide includes pipeline drug candidates currently undergoing clinical testing in a variety of diseases. The safety and efficacy of these drug candidates have not been evaluated by any regulatory authorities for the disease categories described here.
**DUPIXENT®: DELIVERING ON THE “PIPELINE IN A PRODUCT” PROMISE**

**ADOLESCENT AND PEDIATRIC ATOPIC DERMATITIS – HIGH DISEASE BURDEN WITH LIMITED TREATMENT OPTIONS**

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### Adolescent Atopic Dermatitis (Ages 12–17 years)

Positive Ph3 data reported; PDUFA date March 11, 2019

![Graph showing percent improvement in IGA and EASI-75](image)

- **IGA: 0-1***
  - Dupilumab q2w n=82: 24%
  - Placebo n=85: 2%

- **EASI-75***
  - Dupilumab q2w n=82: 41.5%
  - Placebo n=85: 8%

**Key Points**

- Overall rate of treatment-emergent adverse events was comparable between the dupilumab group (72%) and placebo (69%). The rate of overall infections and infestations was numerically lower in the dupilumab group (11%) vs. placebo (20%).

- No SAEs or events leading to discontinuation in the treatment group

**IGA: Investigator’s Global Assessment, EASI: Eczema Area and Severity Index**

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**Before DUPIXENT**

Prior treatments included cycles of prednisone, oral anti-Staph antibiotics, triamcinolone and chronic daily sedating antihistamines

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**After DUPIXENT**

Patient had significantly improved overall disease severity, skin clearing and reduced itching

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**Regeneron**

This slide includes pipeline drug candidates currently undergoing clinical testing in a variety of diseases. The safety and efficacy of these drug candidates have not been evaluated by any regulatory authorities for the disease categories described here.

For illustrative purposes only. Results are not representative of all patients; and individual results vary.
LIBTAYO®: NEW HOPE FOR PATIENTS WITH ADVANCED CSCC

Cutaneous squamous cell carcinoma (CSCC) is the second most common form of skin cancer (after Basal Cell Carcinoma) and is responsible for an estimated 7,000 deaths per year in the U.S.

LIBTAYO is the only FDA approved treatment option for advanced CSCC, a life-threatening condition

Regeneron reported 4Q18 net product sales of $15 Million

June 2018 NEJM publication details pivotal Phase 2 study results in 59 metastatic CSCC patients:

- Primary endpoint: 47.5% Overall Response Rate by independent review
- Durable Disease Control Rate of 61%
- Median duration of response and progression-free survival have not been reached
- LIBTAYO was associated with adverse events similar to other PD-1 inhibitors

An 83-year-old patient who had undergone multiple surgeries for CSCC, at baseline and after 8 weeks of treatment with LIBTAYO

Please see full Prescribing Information for all approved products
LIBTAYO®: THE FOUNDATION OF OUR IO STRATEGY

<table>
<thead>
<tr>
<th>DEVELOPMENT STRATEGY</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Maximize Skin Cancer Opportunity</strong></td>
</tr>
<tr>
<td>2L Basal Cell Carcinoma (BCC) – Ph2 (potentially pivotal) ongoing</td>
</tr>
<tr>
<td>CSCC – Ph3 adjuvant trial to start in 1H19; neo-adjuvant studies to follow</td>
</tr>
<tr>
<td>Melanoma – regulatory discussions anticipated in 1H19</td>
</tr>
<tr>
<td><strong>Non Small Cell Lung Cancer (NSCLC)</strong></td>
</tr>
<tr>
<td>1L NSCLC Monotherapy (≥50% PD-L1) (n=700) – Ph3 ongoing</td>
</tr>
<tr>
<td>1L NSCLC Combination therapy (non-squamous and squamous, stratified by PD-L1 status) – Ph3 amended</td>
</tr>
<tr>
<td>• LIBTAYO + Chemo vs. Chemo</td>
</tr>
<tr>
<td><strong>HPV Positive Cancers</strong></td>
</tr>
<tr>
<td>2L Cervical Cancer – Ph3 ongoing</td>
</tr>
<tr>
<td><strong>Additional Solid &amp; Liquid Tumor Indications</strong></td>
</tr>
<tr>
<td>Pediatric Glioblastoma (GBM) – Ph1/2 initiated</td>
</tr>
<tr>
<td>1L Classical Hodgkin Lymphoma – Ph1 anticipated in 2019</td>
</tr>
<tr>
<td><strong>Combinations</strong></td>
</tr>
<tr>
<td>Immune modulators, vaccines, cell therapies, kinase inhibitors, chemotherapy and bispecifics</td>
</tr>
</tbody>
</table>

This slide includes pipeline drug candidates currently undergoing clinical testing in a variety of diseases. The safety and efficacy of these drug candidates have not been evaluated by any regulatory authorities for the disease categories described here.
Regeneron’s IO strategy is built on a deep foundation of science and technology.

- 619 manuscripts published, 9,351 patent applications filed and 4,945 patents issued over the last 10 years.
- 500,000 exomes sequenced by Regeneron Genetics Center (RGC).

**Velocigene**
- Rapid, automated and high-scale manipulation of mouse DNA to identify and validate therapeutic targets.
- Genetically-humanized immune system in a mouse, producing a diverse range of fully human monoclonal antibodies.

**Velocimmune**
- Fully human T-cell receptors (TCR) against tumor and viral antigens.

**VelociBi**
- Proprietary method that uses Velocimmune with proprietary antibody manufacturing processes to generate full-length human bispecific antibodies.

**VelociMouse**
- Genetically altered mice derived from modified embryonic stem cells.

**Velocisuite**
- High-throughput screening of antibodies and rapid generation of production cell lines.
REGENERON’S IO STRATEGY CONNECTS MULTIPLE INDIVIDUAL PIECES…

- Kinase Inhibitors
- Costimulatory bispecifics
- CD3 bispecifics
- Tcell Rx
-Chemo
-Vaccines
-CTLA-4
-LAG-3
-GITR
-LIBTAYO
...LOGICALLY AND RATIONALLY INTO A COHESIVE WHOLE

...like pieces in a puzzle, bringing order to chaos

Regeneron’s IO puzzle is evolving and not yet complete; based on science and experimental data, the shape, components and configuration may change
REGN1979, OUR EXCLUSIVELY-OWNED CD20xCD3 BISPECIFIC ANTIBODY, DEMONSTRATES HIGH ORR/CR

In our dose escalation Ph1 study, treatment with ≥5 mg of REGN1979 demonstrated 100% ORR and 80% CR in 10 pts with R/R FL.

At higher doses in R/R DLBCL we are seeing response rates that make us optimistic about achieving activity comparable to CAR-Ts.

At doses tested, REGN1979 was well-tolerated in B-NHL: 75% patients had Grade 3/4/5 AEs, no DLTs, 3% discontinued due to AE, no discontinuations due to CRS or immune-related events, no clinically significant neurotoxicity (no seizures/encephalopathy), 1 death due to related AE*.

Safety and toxicity profile is encouraging and supports further dose escalation.

Data presented at the 2018 American Society of Hematology (ASH) Annual Meeting

<table>
<thead>
<tr>
<th>Relapsed/Refractory Follicular Lymphoma (R/R FL) Grade 1-3a</th>
<th>REGN1979 dose groups</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;5 mg (n=7)</td>
<td>≥5≤12 mg (n=5)</td>
</tr>
<tr>
<td>ORR</td>
<td>1/7 (14%)</td>
</tr>
<tr>
<td>CR</td>
<td>1/7 (14%)</td>
</tr>
<tr>
<td>PR</td>
<td>0/7 (0%)</td>
</tr>
<tr>
<td>Responding patients who did not progress during study treatment, n/N (% of responders)</td>
<td>1/1 (100%)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Relapsed/Refractory Diffuse Large B-Cell Lymphoma (R/R DLBCL)</th>
<th>REGN1979 dose groups</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;5 mg (n=15)</td>
<td>≥5≤12 mg (n=11)</td>
</tr>
<tr>
<td>ORR</td>
<td>3/15 (20%)</td>
</tr>
<tr>
<td>CR</td>
<td>0/15 (0%)</td>
</tr>
<tr>
<td>PR</td>
<td>3/15 (20%)</td>
</tr>
<tr>
<td>Responding patients who did not progress during study treatment, n/N (% of responders)</td>
<td>1/3 (33%)</td>
</tr>
</tbody>
</table>

*Initiating potentially pivotal studies in 2019

This slide includes pipeline drug candidates currently undergoing clinical testing in a variety of diseases. The safety and efficacy of these drug candidates have not been evaluated by any regulatory authorities for the disease categories described here.
### Anti-PD-1 Responsive Tumors
TCR binds tumor MHC/peptide

- **Anti-PD-1 mAb monotherpay or combination**

### Anti-PD-1 Unresponsive Tumors
TCR does not recognize tumor MHC/peptide

- **CD3 bispecific alone, or in combination with PD-1 and/or costimulatory bispecifics**

### Additional Strategic Opportunities

- **CAR-T therapies alone or in combination**
  - Major collaboration with bluebird bio to empower and extend CAR-T therapies with novel tumor targeting moieties such as TCRs or reagents that bind peptide/MHC complexes
  - Can complement with soluble reagents such as anti-PD-1 and CD3 or costimulatory bispecifics

---

**REGENERON’S IO STRATEGY IS BASED ON RATIONAL COMBINATIONS**

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REGENERON’S CD3 & COSTIMULATORY BISPECIFICS ARE OFF-THE-SHELF DRUGS WITH POTENTIAL TO TURN PATIENTS’ T CELLS INTO CAR-T-LIKE CANCER KILLERS

CAR-T Mechanism

The combination of CD3 and costimulatory bispecifics has the potential to activate T cells into highly effective, targeted cancer killers.

Bispecific/Costimulatory Mechanism

T cell activation can be inhibited by PD-1 signaling.
REGENERON’S CD3 & COSTIMULATORY BISPÉCIFICS ARE OFF-THE-SHELF DRUGS WITH POTENTIAL TO TURN PATIENTS’ T CELLS INTO CAR-T-LIKE CANCER KILLERS

CAR-T Mechanism

**LIBTAYO** blocks the stop signal

**CAR-T**

**Tumor Target**

**CD3ζ**

**scFv**

**PD-L1/2**

**PD-1**

**Signal 1**

**Signal 2**

Bispecific/Costimulatory Mechanism

**LIBTAYO** blocks the stop signal

**LIBTAYO**

**Tumor Target**

**Tumor Target**

**CD3 bispecific**

**Costimulatory bispecific**

**PD-L1/2**

**PD-1**

**Signal 1**

**Signal 2**

**CD3ζ**

**CD3ε**

**TCRα**

**TCRβ**

**TCRγ**

**TCRδ**

**α**

**β**

**Using LIBTAYO to block PD-1 signaling can further enhance the efficacy of CD3 and costimulatory bispecifics**
**ADDING COSTIMULATORY BISPECIFICS TO CD3 BISPECIFICS OR TO ANTI-PD-1 SHOWS SYNERGY IN PRECLINICAL TUMOR MODELS**

**TSA1xCD3 + TSA1xCD28**  
*in vivo* xenogeneic humanized TSA1 mouse model

**anti-PD-1 + TSA2xCD28**  
*in vivo* syngeneic humanized TSA2 mouse model

- Unlike superagonist CD28 mAbs, our CD28 bispecifics have no toxicity, and little or no activity on their own, but when clustered on cells expressing their target, activate signal 2 and synergize with signal 1 (via CD3 bispecific) and/or anti-PD-1.
- In 2019, Regeneron plans to advance two distinct CD28 bispecific antibodies into clinical development.

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**TSA = Tumor Specific Antigen**

---

This slide includes pipeline drug candidates currently undergoing clinical testing in a variety of diseases. The safety and efficacy of these drug candidates have not been evaluated by any regulatory authorities for the disease categories described here.
### BROADENING OUR IMMUNO-ONCOLOGY PIPELINE

<table>
<thead>
<tr>
<th>Pre-IND</th>
<th>Clinical Development</th>
<th>LIBTAYO Potential Indications</th>
<th>Approved</th>
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<tbody>
<tr>
<td>TSAxCD28 Solid Tumor</td>
<td>REGN1979 (CD20xCD3) B-Cell NHL</td>
<td>LIBTAYO NSCLC, Cervical, BCC, Pediatric</td>
<td>LIBTAYO CSCC</td>
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<tr>
<td>TSAxCD28 B cell malignancy</td>
<td>REGN5458 (BCMAxCD3) Multiple Myeloma</td>
<td>LIBTAYO + REGN1979 (CD20xCD3) B-Cell NHL</td>
<td></td>
</tr>
<tr>
<td>GITR Solid tumors</td>
<td>REGN4018 (MUC16xCD3) Ovarian Cancer</td>
<td></td>
<td></td>
</tr>
<tr>
<td>And More To Come HLA/peptide (tumor and viral), etc.</td>
<td>REGN4659 (CTLA-4) NSCLC</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>REGN3767 (LAG-3) Solid/hematologic malignancies</td>
<td></td>
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</tr>
</tbody>
</table>

TSA = Tumor Specific Antigen

This slide includes pipeline drug candidates currently undergoing clinical testing in a variety of diseases. The safety and efficacy of these drug candidates have not been evaluated by any regulatory authorities for the disease categories described here.
MANY COMPANIES CAN DO ONE THING...

CD3 bispecifics

PD-1/L1

Costimulatory bispecifics

CTLA-4, LAG-3, GITR…
...FEW CAN DO MANY THINGS

PD-1/L1

Costimulatory bispecifics

CD3 bispecifics

CTLA-4, LAG-3, GITR...

REGENERON is one of the few
OSTEARTHRITIS IS A COMMON CONDITION ASSOCIATED WITH WEAR AND TEAR ON THE JOINTS, AND IS THE MOST COMMON INDICATION FOR KNEE AND HIP REPLACEMENT

Pain is a protective mechanism

NGF blockade treats pain, but not osteoarthritis itself

In clinical trials we observed a dose-dependent increase in rapidly progressive osteoarthritis (RPOA) and total joint replacement (TJR); we therefore limited development to lower dose regimens

We announced in August 2018 positive topline results showing a clinically meaningful reduction in pain and increased function in patients with chronic pain from osteoarthritis of the knee or hip

Based on our analysis of the data, we believe we have identified a minimally effective dose and are encouraged that under close and careful scrutiny, the independent data monitoring committee overseeing patient safety has supported continued development

Fasinumab* is a human monoclonal antibody that treats osteoarthritis pain by blocking nerve growth factor (NGF)

* Partnered with Teva and Mitsubishi Tanabe Pharma (MTPC) outside of the U.S.
This slide includes pipeline drug candidates currently undergoing clinical testing in a variety of diseases. The safety and efficacy of these drug candidates have not been evaluated by any regulatory authorities for the disease categories described here.
## 2019 GOALS AND MILESTONES

### KEY REGULATORY APPROVALS & SUBMISSIONS

<table>
<thead>
<tr>
<th>Drug</th>
<th>Key Event</th>
</tr>
</thead>
<tbody>
<tr>
<td>EYLEA</td>
<td>FDA decision on sBLA for the treatment of Diabetic Retinopathy (PDUFA date May 13, 2019); re-submission of Prior-Approval Supplement (PAS) for pre-filled syringe</td>
</tr>
<tr>
<td>DUPIXENT</td>
<td>FDA decision on sBLA for expanded Atopic Dermatitis indication in adolescent patients 12–17 years of age (PDUFA date March 11, 2019); EMA decision on regulatory application for Asthma; file sBLA for Chronic Rhinosinusitis with Nasal Polyps (CRSwNP)</td>
</tr>
<tr>
<td>LIBTAYO</td>
<td>EMA decision for advanced cutaneous squamous cell carcinoma (CSCC)</td>
</tr>
<tr>
<td>PRALUENT</td>
<td>FDA (PDUFA date April 28, 2019) and EMA decisions on applications for Cardiovascular Risk Reduction; FDA decision on sBLA for first-line treatment of Hyperlipidemia (PDUFA date April 29, 2019)</td>
</tr>
</tbody>
</table>

### CLINICAL PROGRESS

<table>
<thead>
<tr>
<th>Drug</th>
<th>Key Event</th>
</tr>
</thead>
<tbody>
<tr>
<td>EYLEA</td>
<td>Initiate a study of higher dose formulations of aflibercept</td>
</tr>
<tr>
<td>DUPIXENT</td>
<td>Continue enrollment in pivotal eosinophilic esophagitis (EoE) study; initiate Ph2/3 program in Chronic Obstructive Pulmonary Disease (COPD)</td>
</tr>
<tr>
<td>LIBTAYO</td>
<td>Continue enrollment in NSCLC and various other studies</td>
</tr>
<tr>
<td>REGN1979 (CD20xCD3)</td>
<td>Initiate potentially pivotal Ph2 study in Follicular Lymphoma (FL) and potentially pivotal Ph2 study in Diffuse Large B-Cell Lymphoma (DLBCL)</td>
</tr>
<tr>
<td>Fasinumab (NGF)</td>
<td>Continue patient enrollment in Ph3 long-term safety study and Ph3 efficacy studies in Osteoarthritis</td>
</tr>
<tr>
<td>Pozelimab (C5)</td>
<td>Initiate Ph2 in Paroxysmal Nocturnal Hemoglobinuria (PNH)</td>
</tr>
</tbody>
</table>

### KEY DATA READOUTS

<table>
<thead>
<tr>
<th>Drug</th>
<th>Key Event</th>
</tr>
</thead>
<tbody>
<tr>
<td>DUPIXENT</td>
<td>Report results from Ph3 study for Atopic Dermatitis in pediatric patients 6–11 years of age</td>
</tr>
<tr>
<td>REGN3500 (IL-33)</td>
<td>Report results from Ph2 Asthma study</td>
</tr>
<tr>
<td>Trevogrumab (GDF8) + Garetosmab (Activin-A)</td>
<td>Report results from multi-dose portion of Ph1 study</td>
</tr>
</tbody>
</table>

### NEW INDs

Expect to advance 4-6 new molecules into clinical development (including more CD3 & CD28 bispecifics)
### 2019 FINANCIAL GUIDANCE*

* As of February 6, 2019. The guidance does not assume the completion of any significant business development transaction that had not been completed as of the date of the guidance. Regeneron does not undertake any obligation to update publicly any financial projection or guidance, whether as a result of new information, future events, or otherwise.

<table>
<thead>
<tr>
<th>Category</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>GAAP Sanofi Collaboration Revenue:</td>
<td>$510 – 560MM</td>
</tr>
<tr>
<td>Reimbursement of Regeneron Commercialization-Related Expenses</td>
<td></td>
</tr>
<tr>
<td>GAAP unreimbursed R&amp;D</td>
<td>$1.855 – $2.000B</td>
</tr>
<tr>
<td>Non-GAAP unreimbursed R&amp;D†</td>
<td>$1.590 – $1.710B</td>
</tr>
<tr>
<td>GAAP SG&amp;A</td>
<td>$1.700 – $1.830B</td>
</tr>
<tr>
<td>Non-GAAP SG&amp;A†</td>
<td>$1.500 – $1.600B</td>
</tr>
<tr>
<td>GAAP Effective Tax Rate</td>
<td>14 – 16%</td>
</tr>
<tr>
<td>GAAP Capital Expenditures</td>
<td>$410 – 490MM</td>
</tr>
</tbody>
</table>

† Please refer to slide 2 for important information regarding non-GAAP financial measures and to slide 26 for a reconciliation of these measures to GAAP financial measures.
## RECONCILIATION OF FULL YEAR 2019 NON-GAAP TO GAAP FINANCIAL GUIDANCE

<table>
<thead>
<tr>
<th>(in millions)</th>
<th>Projected Range</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Low</td>
</tr>
<tr>
<td>GAAP unreimbursed R&amp;D*</td>
<td>$1,855</td>
</tr>
<tr>
<td>R&amp;D: Non-cash share-based compensation expense</td>
<td>(265)</td>
</tr>
<tr>
<td>Non-GAAP unreimbursed R&amp;D</td>
<td>$1,590</td>
</tr>
<tr>
<td>GAAP SG&amp;A</td>
<td>$1,700</td>
</tr>
<tr>
<td>SG&amp;A: Non-cash share-based compensation expense</td>
<td>(200)</td>
</tr>
<tr>
<td>Non-GAAP SG&amp;A</td>
<td>$1,500</td>
</tr>
</tbody>
</table>

* Unreimbursed R&D represents R&D expenses reduced by R&D expense reimbursements from the Company’s collaborators and/or customers.