

**BAUSCH+LOMB**

# ***FOCUS FORWARD***

**INVESTOR DAY**

**November 13<sup>th</sup>, 2025**

# Disclaimers

## Forward-Looking Statements

This presentation contains forward-looking information and statements, within the meaning of applicable securities laws (collectively, "forward-looking statements"), including, but not limited to, statements regarding future prospects and performance of Bausch + Lomb Corporation ("Bausch + Lomb", the "Company", "we", "us", or "B+L"), our 2025 full year guidance, our three year financial targets (including targets for Company and segment level cc revenue CAGRs, projected Adj. EBITDA margin (excl. Acq. IPR&D), annual adj. EPS (excl. Acq. IPR&D) growth rates, adjusted cash flow from operations to Adj. EBITDA (excl. Acq. IPR&D) conversion and net leverage), our strategic plan for meeting our financial targets and the steps thereof, our anticipated growth drivers and the expected timing and impact thereof, our investment grade rating target and framework for achieving same, our capital allocation priorities, our focus on pipeline innovation, the success of our pipeline products and R&D programs, the existence of potential game changers in our pipeline and our ability to be the best in class or first product to market of its kind, anticipated approval and launch dates for our pipeline products, our estimates for potential peak sales for our pipeline products, franchises and businesses, our ability to outperform the market, our ability to expand into new categories and new and existing markets and to address new indications and disease states, our ability to successfully launch next generation versions of our existing products, our ability to focus on and have success with disruptive innovation, the expected market acceptance and performance for certain of our pipeline products, the expected market size for certain of the markets in which we expect to have products, and the timing of commencement and completion of clinical studies and other development work. Forward-looking statements may generally be identified by the use of the words "anticipates," "expects," "predicts," "projects," "goals," "intends," "plans," "should," "could," "would," "may," "might" "will," "strive," "believes," "estimates," "potential," "target," "commit," "forecast," "outlook," "guidance," "tracking," or "continue" and positive and negative variations or similar expressions, and phrases or statements that certain actions, events or results may, could, should or will be achieved, received or taken or will occur or result, and similar such expressions also identify forward-looking information. These forward-looking statements, including the Company's 2025 full-year guidance and its three-year financial targets, are based upon the current expectations and beliefs of management and are provided for the purpose of providing additional information about such expectations and beliefs, and readers are cautioned that these statements may not be appropriate for other purposes. These forward-looking statements are subject to certain risks and uncertainties that could cause actual results to differ materially from those described in the forward-looking statements.

These risks and uncertainties include, but are not limited to, the risks and uncertainties discussed in Bausch + Lomb's filings with the U.S. Securities and Exchange Commission ("SEC") and the Canadian Securities Administrators (the "CSA") (including the Company's Annual Report on Form 10-K for the year ended December 31, 2024 (which was filed with the SEC and CSA on February 19, 2025) and its most recent quarterly filings), which factors are incorporated herein by reference. They also include risks relating to the development of our pipeline products, including the risk that our studies may not produce successful results or demonstrate safety and efficacy in humans, risks that our pre-clinical and clinical trials may be delayed (which, in turn, may delay the launch of these products) and risks that the regulatory approval of our products may be lengthy, costly and, ultimately, not successful. They also include risks relating to the launch and commercialization of our products, including risks relating to the costs, required resources and unpredictability of commercial launches, risks that our products may not achieve the anticipated levels of market acceptance, which can result from a number of factors, many of which are outside of our control, risks that our products may experience negative publicity or reputational harm, competitive risks, such as our competitors beating us to market or developing new or better technologies and risks that we may face supply interruptions with our finished products or components thereof, which, in turn, may impact our ability to successfully launch or commercialize our products. They also include risks and uncertainties respecting the proposed plan to separate the Company into an independent, publicly traded company, separate from the remainder of Bausch Health Companies Inc. ("BHC") (the "separation"), which include, but are not limited to, the expected benefits and costs of the separation, the expected timing of completion of the separation and its manner and terms (including that it may include the transfer of all or a portion of BHC's remaining direct or indirect equity interest in Bausch + Lomb to its shareholders (the "distribution")), the expectation that, if the separation is to be effected through a distribution, then it will be completed following the achievement of targeted debt leverage ratios, subject to market conditions and receipt of applicable shareholder and other necessary approvals and other factors (including those described in BHC's public statements), the ability to complete the distribution considering the various conditions to the completion of the distribution (some of which are outside the Company's and BHC's control, including conditions related to regulatory matters and receipt of applicable shareholder and other approvals), the impact of any potential sales or dispositions of the Company's common shares by BHC (including in connection with a foreclosure on the Bausch + Lomb common shares owned by BHC that are or maybe pledged as collateral for certain of BHC's debt), that market or other conditions are no longer favorable to completing the transaction, that applicable shareholder, stock exchange, regulatory or other approval is not obtained on the terms or timelines anticipated or at all, business disruption during the pendency of or following the separation, diversion of management time on separation-related issues, retention of existing management team members, the reaction of customers and other parties to the separation, the structure of the distribution, the qualification of the distribution as a tax-free transaction for Canadian and/or U.S. federal income tax purposes (including whether or not an advance ruling from the Canada Revenue Agency and/or the Internal Revenue Service will be sought or obtained), the ability of the Company and BHC to satisfy the conditions required to maintain the tax-free status of the distribution (some of which are beyond their control), other potential tax or other liabilities that may arise as a result of the distribution, the potential dis-synergy costs resulting from the separation, the impact of the separation on relationships with customers, suppliers, employees and other business counterparties, general economic conditions, conditions in the markets the Company is engaged in, behavior of customers, suppliers and competitors, technological developments and legal and regulatory rules affecting the Company's business. In particular, the Company can offer no assurance that the separation will occur at all, or that any such transaction will occur on the terms and timelines or in the manner anticipated by the Company and BHC. They also include risks and uncertainties relating to acquisitions and other business development transactions the Company has completed or may, in the future, pursue and complete, including risks that pending transactions may not close, risks that the Company may not realize the expected benefits of those transactions on a timely basis or at all and, where applicable, risks relating to increased levels of debt as a result of debt incurred to finance such transactions, including in regards to compliance with our debt covenants. They also include the expected impact of the tariffs imposed by the U.S. and counter-tariffs or other retaliatory measures imposed on the U.S. by other countries and disruptions to global supply chains and other potential results as a result of these developments and our ability to successfully manage the expected impact of such tariffs and counter-tariffs and other measures, including the success of our planned actions and levers to manage these matters. Finally, they also include, but are not limited to, risks and uncertainties caused by or relating to a potential recession and other adverse economic conditions (such as heightened inflation and interest rates, fluctuations in exchange rates,

imposition of and adverse changes to tariffs, duties and other trade protection measures and slower growth), which could adversely impact our revenues, expenses and resulting margins. In addition, certain material factors and assumptions have been applied in making these forward-looking statements, including the assumption that the risks and uncertainties outlined above will not cause actual results or events to differ materially from those described in these forward-looking statements. In addition, Management has also made certain assumptions regarding our 2025 full-year guidance with respect to expectations regarding base performance growth, business performance, currency impact, impacts of inflation, the company's ability to offset the impact of tariffs in 2025 (based on the current tariff policy and the actions the company is taking to manage these measures), adjusted gross margin (non-GAAP), adjusted SG&A expense (non-GAAP) and the Company's ability to continue to manage such expense in the manner anticipated, interest expense (which will vary based on, among other things, interest rates and our indebtedness), adjusted tax rate, and full year capex and the anticipated timing and extent of the Company's R&D expense. In addition, Management has also made certain assumptions regarding our three year financial targets, including those assumptions set out on Slide 96.

References in this presentation to future launch dates refer to the anticipated launch dates for the applicable product, based on management's estimates taking into account, among other things, the current stage of the development or regulatory pathway of such products, typical development, regulatory and launch timelines for similar products and assumptions regarding sufficient supply availability for launch purposes. In addition, unless otherwise indicated, references in this presentation to peak sales refer to the potential peak annual sales of the applicable product, franchise or business, based on management's estimates, taking into account, among other things, sales of similar products, franchises and business.

Readers are cautioned not to place undue reliance on any of these forward-looking statements. These forward-looking statements speak only as of the date hereof. Bausch + Lomb undertakes no obligation to update any of these forward-looking statements to reflect events or circumstances after the date of this presentation or to reflect actual outcomes, unless required by law.

The guidance and financial targets in this presentation are only effective as of the date given, November 13, 2025. Distribution or reference of this deck following November 13, 2025 does not constitute the Company updating or affirming such guidance or financial targets.

## Non-GAAP Information

Non-GAAP Information: To supplement the financial measures prepared in accordance with U.S. generally accepted accounting principles (GAAP), the Company uses certain non-GAAP financial measures and ratios, including (i) Adjusted EBITDA excluding Acquired IPR&D, (ii) Adjusted EBITDA Margin excluding Acquired IPR&D, (iii) EBITDA, (iv) Adjusted Cash Flow from Operations, (v) Adjusted Cash Flow from Operations to Adj. EBITDA (excl. Acq. IPR&D) Conversion, (vi) constant currency (cc) revenue growth, (vii) Net Leverage, (viii) Adjusted Earnings per Share (EPS) Attributable to Bausch + Lomb Corporation (excluding Acquired IPR&D) Growth, (ix) Adjusted Tax Rate and (x) Adjusted R&D expense. Management uses some of these non-GAAP measures and ratios as key metrics in the evaluation of Company performance and the consolidated financial results and, in part, in the determination of cash bonuses for its executive officers. The Company believes these non-GAAP measures and ratios are useful to investors in their assessment of our operating performance and the valuation of the Company. In addition, these non-GAAP measures and ratios, address questions the Company routinely receives from analysts and investors and, in order to assure that all investors have access to similar data, the Company has determined that it is appropriate to make this data available to all investors.

However, these measures and ratios are not prepared in accordance with GAAP nor do they have any standardized meaning under GAAP. In addition, other companies may use similarly titled non-GAAP financial measures and ratios that are calculated differently from the way we calculate such measures and ratios. Accordingly, our non-GAAP financial measures and ratios may not be comparable to such similarly titled non-GAAP measures and ratios of other companies. We caution investors not to place undue reliance on such non-GAAP measures and ratios, but instead to consider them with the most directly comparable GAAP measures and ratios. Non-GAAP financial measures and ratios have limitations as analytical tools and should not be considered in isolation. They should be considered as a supplement to, not a substitute for, or superior to, the corresponding measures calculated in accordance with GAAP.

The reconciliations of these historic non-GAAP financial measures and ratios to the most directly comparable financial measures and ratios calculated and presented in accordance with GAAP are shown in the appendix hereto. However, for outlook purposes, the Company does not provide reconciliations of projected Constant Currency Revenue Growth to projected GAAP Revenue Growth, projected Adjusted EBITDA excluding Acquired IPR&D (non-GAAP) to projected GAAP net income (loss), projected Adjusted EBITDA Margin excluding Acquired IPR&D (non-GAAP) to projected GAAP net income (loss) margin, projected Adjusted Cash Flow from Operations to projected Cash flow from operations/Cash used in operations (loss) attributable to Bausch + Lomb Corporation, or the components of projected net leverage to their GAAP equivalents (project net debt to debt and projected Adjusted EBITDA excluding Acquired IPR&D (non-GAAP) to projected GAAP net income (loss)), or projected Adjusted EPS Attributable to Bausch + Lomb (excl. Acq. IPR&D) to projected Diluted income per share attributable to Bausch + Lomb Corporation ("GAAP EPS") in each case, due to the inherent difficulty in forecasting and quantifying certain amounts that are necessary for such reconciliations. These amounts may be material and, therefore, could result in the GAAP measure Adjusted Earnings per Share (EPS) Attributable to Bausch + Lomb Corporation (excluding Acquired IPR&D) Growth and (iv) Adjusted Tax Rate. Management uses some of these or ratio being materially different from the projected non-GAAP measure or ratio.

For further information on non-GAAP financial measures and ratios, please see the Appendix.

# AGENDA

**Welcome & Introduction**

**Brent Saunders**, *Chairman and CEO*

**Financial Outlook**

**Sam Eldessouky**, *Executive Vice President and CFO*

**R&D Innovation**

**Yehia Hashad, MD**, *Executive Vice President, R&D and CMO*

**Consumer**

**John Ferris**, *President, Consumer*

**Mayssa Attar, Ph.D.**, *Senior Vice President, Pharmaceuticals and Consumer R&D*

**Pharmaceuticals**

**Andrew Stewart**, *President, Global Pharmaceuticals and International Consumer*

**Mayssa Attar, Ph.D.**, *Senior Vice President, Pharmaceuticals and Consumer R&D*

**Break & Product Exhibition**

**Contact Lens**

**Yang Yang**, *President, Vision Care*

**Bryan Reed**, *Vice President, Vision Care R&D*

**Surgical**

**Luc Bonnefoy**, *President, Surgical*

**Kelly Swaim, MD**, *Senior Vice President, Surgical R&D*

**Physician Panel**

**Moderator: Cathleen McCabe, MD**, *Strategic Medical Advisor, Bausch + Lomb*

**Q&A**

**Bausch + Lomb Leadership**

# WELCOME & INTRODUCTION

**BRENT SAUNDERS**

CHAIRMAN & CHIEF EXECUTIVE OFFICER



“

Mediocrity hides behind ‘good enough.’  
**Excellence** burns right through it.

”

# FINANCIAL OUTLOOK

## SAM ELDESSOUKY

EXECUTIVE VICE PRESIDENT &  
CHIEF FINANCIAL OFFICER



# Now *Entering Next Phase* of Our Strategy<sup>1</sup>

## Maintain Growth Momentum

---

Continue To  
Deliver Above  
Market Revenue  
Growth

## Expand Profitability

---

Drive Step  
Change Margin  
Expansion  
Profile

## Advance R&D Pipeline



*Significant  
Upside Potential  
Beyond 2028*

# Strategy to Deliver *Financial Excellence* Across All Key Metrics<sup>1,4</sup>

**Above-Market  
Revenue  
Growth**

**Meaningful  
Margin  
Expansion**

**Robust  
EPS  
Growth**

**Solid  
Cash Flow  
Generation**

**Strong  
Balance  
Sheet**

**5 – 7%**

2025-2028  
CC REVENUE  
CAGR<sup>2,3</sup>

**~23%**

ADJ. EBITDA MARGIN  
(EX. ACQ. IPR&D)<sup>3</sup>  
IN 2028

**Double  
Digit**

ADJ. EPS GROWTH<sup>3,5</sup>  
2026-2028

**~50%**

ADJ. CASH FLOW  
FROM OPERATIONS  
TO ADJ. EBITDA  
CONVERSION IN 2028<sup>3,5</sup>

**~3.5x**

NET LEVERAGE  
BY END OF 2028<sup>3</sup>

# Growth & *Meaningful* Margin Expansion

Reaffirming  
2025 Guidance<sup>1</sup>

**\$5.050 – \$5.150B**

FY25 Revenue

Preliminary  
2026 View<sup>1</sup>

**~5 – 7%**

FY25-28 CC Revenue CAGR<sup>2,3</sup>

2028  
Outlook<sup>1</sup>

**\$870 – \$910M**

FY25 Adj. EBITDA

(ex. Acq. IPR&D)<sup>2</sup>

~17% Margin

**~19%**

FY26 Adj. EBITDA Margin

(ex. Acq. IPR&D)<sup>2</sup>

+200bps vs FY25

**~23%**

FY28 Adj. EBITDA Margin

(ex. Acq. IPR&D)<sup>2</sup>

+600bps vs FY25

# *Broad-Based* Growth Across Business<sup>1</sup>

## Consumer

**5-7%**

FY25-28

CC REVENUE CAGR<sup>1,2</sup>

Expand Leadership with  
Launch of Lumify Luxe

Capture Expanded AMD  
Market with AREDS3

Continue Market Leading  
Growth in Dry Eye

## Contact Lens

**5-7%**

FY25-28

CC REVENUE CAGR<sup>1,2</sup>

Maintain Strong Growth  
Trajectory in DD SiHy

Drive Growth in Ultra FRP  
and Biotrue ONEday

Position to Launch Segment  
Creating Innovation

## Pharma

**5-7%**

FY25-28

CC REVENUE CAGR<sup>1,2</sup>

Continue Miebo  
Growth Momentum

Drive Growth in Xiidra  
TRx and Revenue

Deliver Durable Growth  
in International Pharma

## Surgical

**6-8%**

FY25-28

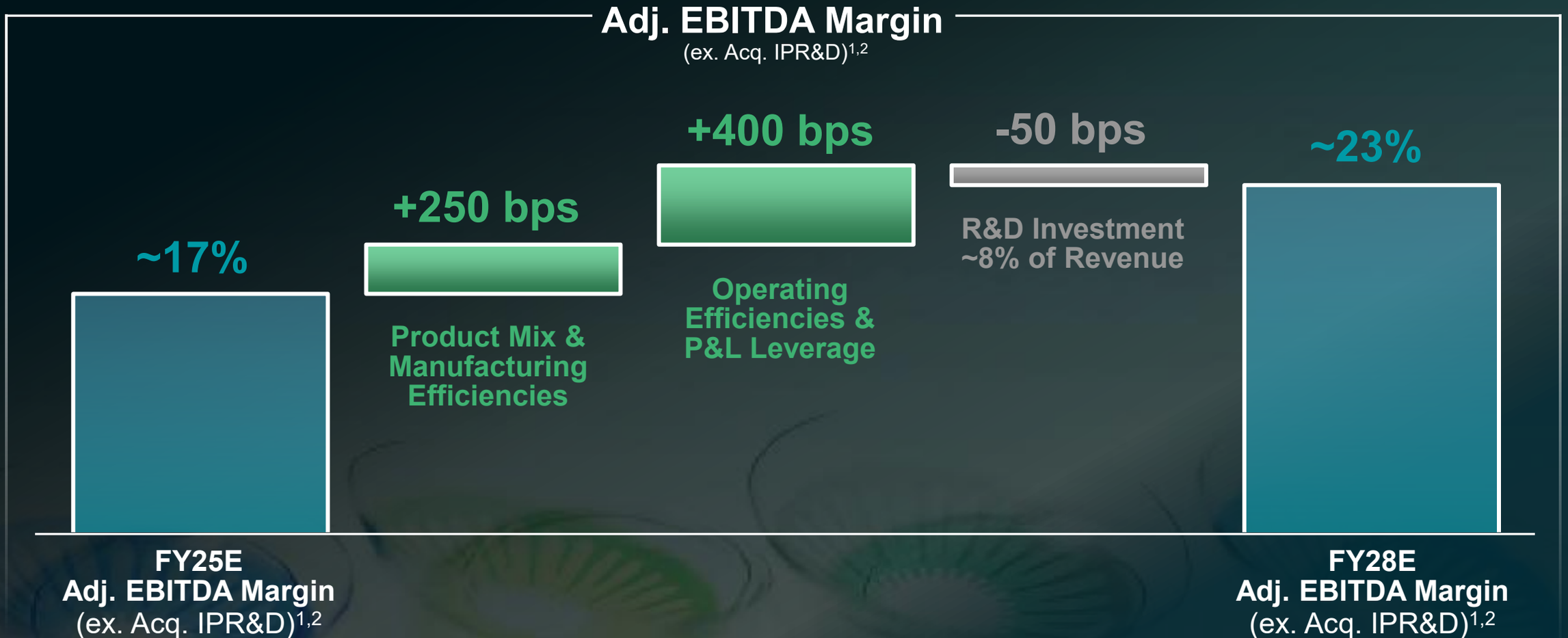
CC REVENUE CAGR<sup>1,2</sup>

Growth in Premium  
IOL Portfolio

Launch Into New  
Product Categories

Unlock Pull-Through  
Sales in Consumables

# Expanding Margins and Maintaining Focus on Advancing Pipeline



# *Strong* Cash Flow Generation

Adj. Cash Flow From Operations  
To Adj. EBITDA Conversion<sup>1,2,3</sup>



Strategic Focus to Optimize Working Capital Management

**BAUSCH+LOMB**

Net Leverage<sup>2</sup>

**~3.5x**

by end of 2028<sup>1</sup>

Targeting Investment  
Grade Rating  
in the Long-Term<sup>1</sup>

1. See Slide 2 for further information on forward-looking statements. The financial targets in this presentation are only effective as of the date given, November 13, 2025, and will not be updated or affirmed unless and until the Company publicly announces updated or affirmed financial targets. Distribution or reference of this deck following November 13, 2025, does not constitute the Company re-affirming financial targets.

2. This is a non-GAAP measure or ratio. See Slide 2 and Appendix for further information on non-GAAP measures and ratios.

3. Ex. Acq. IPR&D.

# Capital Allocation Priorities<sup>1</sup>

## Strengthen Balance Sheet

- Reduce net leverage<sup>2</sup> to ~3.5x by end of 2028
- Maintain framework for investment grade profile

## Invest in Organic Growth

- Drive commercial and operational excellence
- Advance pipeline to deliver sustainable growth
- Capacity expansion

## Strategic M&A / BD&L

- Disciplined M&A / BD&L opportunities
- Strategic partnerships to advance innovation

# Drive Value with *Execution*<sup>1,2</sup>

## 2025 - 2028

### 5-7%

2025-2028  
CC Revenue CAGR<sup>1,2,3</sup>

Above-Market  
Revenue Growth

### ~23%

2028 Adj. EBITDA  
Margin (ex. Acq. IPR&D)<sup>3</sup>

~600bps EBITDA  
Margin Expansion  
2025-2028

# Significant Pipeline *Upside*<sup>1</sup>

## 2028+

Pipeline to Drive  
Transformative Value with  
Potential Peak Sales

### ~\$7B<sup>4</sup>

# 3 Key Takeaways<sup>1</sup>

1

Durable  
Above-Market  
Growth

2

Meaningful  
Margin  
Expansion

3

Significant  
Upside Potential  
Beyond 2028

# R&D INNOVATION

## YEHIA HASHAD, MD

EXECUTIVE VICE PRESIDENT,  
R&D & CHIEF MEDICAL OFFICER



# Growth Engine with *Top* Industry Talent

FOCUSED ON DISRUPTIVE INNOVATION:  
NEW CAPABILITIES IN DISEASE BIOLOGY  
ADVANCED FORMULATIONS  
MATERIAL SCIENCE

~1K

R&D EMPLOYEES ACROSS  
12 SITES GLOBALLY

>20

RECORD NUMBER OF NEW  
PRODUCT LAUNCHES IN  
LAST TWO YEARS

>60

R&D PROJECTS IN PIPELINE WITH  
POTENTIAL GAME CHANGERS<sup>1</sup>

Driving  
**innovation**

across all  
business units<sup>1</sup>

Steady stream of  
**launches**

into the  
next decade<sup>1</sup>

**Game  
changers**

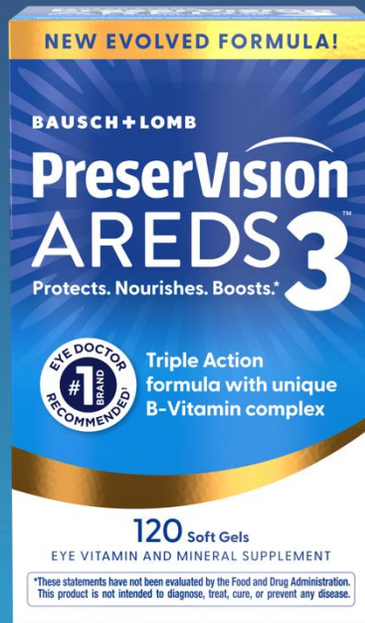
~\$7B potential  
peak sales<sup>1,2</sup>

# Pipeline Designed to *Raise* the Standard of Care

# Clinically-Differentiated Consumer Brands<sup>1</sup>

## Expand Market by Addressing all Stages of AMD

Launch Date: 1H26  
Franchise Peak Sales: ~\$600M



## Enhanced Comfort with Addition of Hyaluronic Acid

Launch Date: 1H27  
Franchise Peak Sales: ~\$450M



## Advanced Preservative-Free Lipid Based Formulation

Launch Date: 1H26  
Franchise Peak Sales: ~\$300M



# Potential *Game Changers* in Pharma<sup>1</sup>

**First**

dual-action therapeutic  
for evaporative and  
inflammatory DED

Launch Date: ~2029  
Peak Sales: ~\$0.7B

**First**

neurosensory agent  
to address  
ocular surface pain

Launch Date: ~2030  
Peak Sales: ~\$1.4B

**First**

glaucoma therapy  
to improve visual  
function through  
neuroprotection

Launch Date: ~2031  
Peak Sales: ~\$0.8B

**First**

therapeutic for  
intermediate AMD  
and best-in-class  
treatments for GA

Launch Date: 2030+  
Peak Sales: >\$1.0B

# Expanding in *Premium* Surgical Categories<sup>1</sup>



▶ **EDOF LENS TO COMPLETE PREMIUM IOL PORTFOLIO**

Launch Date: 2027  
enVista Platform  
Peak Sales: ~\$300M

elios



▶ **FIRST CLINICALLY VALIDATED IMPLANT FREE MIGS EXCIMER LASER**

Launch Date: 2H26 in U.S.  
Peak Sales: ~\$175M

see)NOVA



▶ **BEST-IN-CLASS CATARACT / RETINA COMBO SYSTEM**

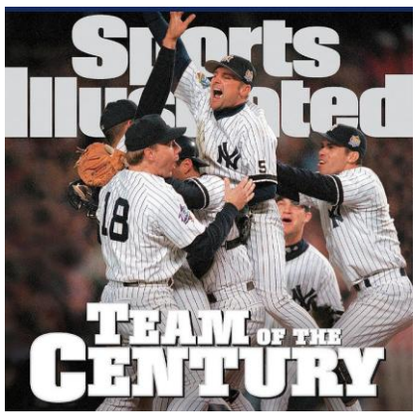
Launch Date: 2028  
seeNOVA and Stellaris  
Peak Sales: ~\$450M

see)LYRA



▶ **NEXT GENERATION FEMTOSECOND LASER**

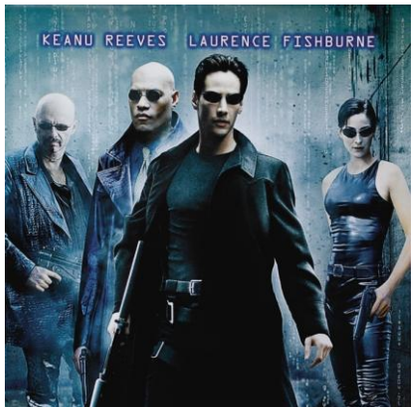
Launch Date: 2H26  
Peak Sales: ~\$50M



1999



No Breakthrough  
Innovation in  
Contact Lens  
Industry for  
*Over 25 Years*



# Contact Lens Pipeline Designed with *Purpose*<sup>1</sup>

DRIVE REVENUE GROWTH  
ACROSS ALL MARKET  
CATEGORIES

DRIVE MARGIN EXPANSION  
BY LEVERAGING EXISTING  
MANUFACTURING PLATFORMS



PROJECT  
HALO

FIRST-OF-ITS-KIND  
BIOACTIVE  
CONTACT LENS  
MATERIAL

Launch Date: 2028  
Peak Sales: ~\$500M



2ND DD SIHY LENS  
INNOVATION  
DESIGNED FOR  
AFFORDABILITY

Launch Date: 2029  
Peak Sales: ~\$250M



PREMIUM FRP  
SIHY PROVIDING  
UNSURPASSED  
LENS COMFORT

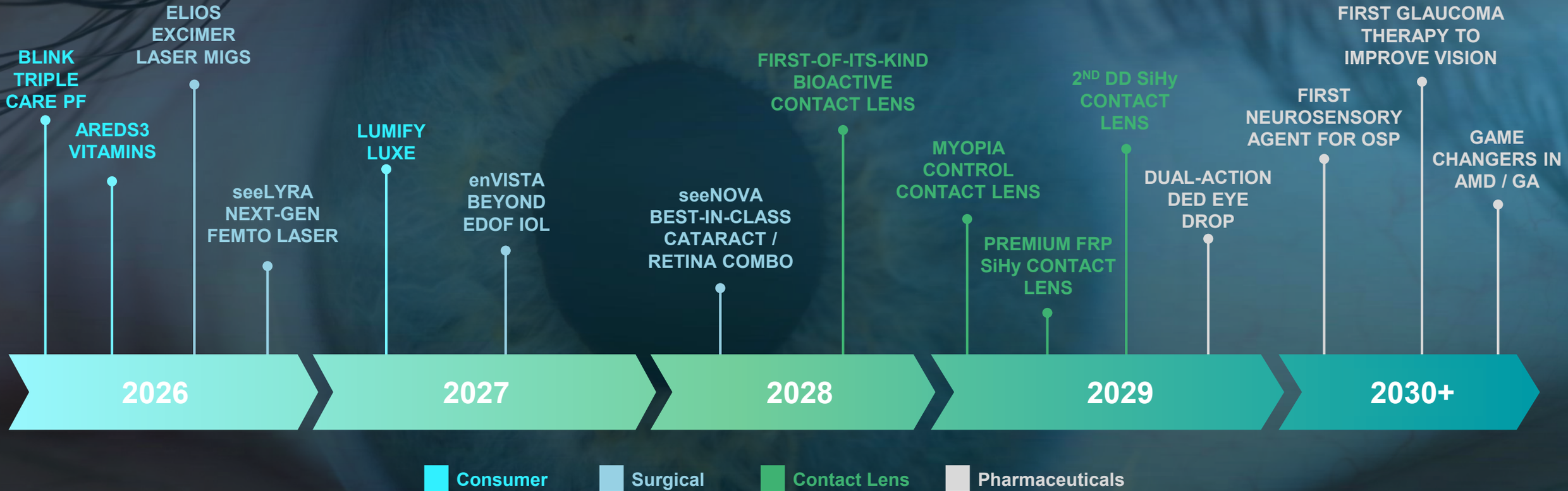
Launch Date: 2029  
Peak Sales: ~\$300M



CUTTING-EDGE SIHY  
LENS DESIGN TO  
SLOW PROGRESSION  
OF MYOPIA

Launch Date: 2029  
Peak Sales: ~\$200M

# *Steady Stream* of Launches into Next Decade<sup>1</sup>



# 3 Key Takeaways<sup>1</sup>

1

Expanding  
Talent &  
Capabilities

2

Driving  
Disruptive  
Innovation

3

Sustained  
Growth from  
New Launches



# CONSUMER

**JOHN FERRIS**

PRESIDENT, CONSUMER

**MAYSSA ATTAR, PH.D.**

SENIOR VICE PRESIDENT,  
PHARMACEUTICALS AND CONSUMER R&D





# Track Record

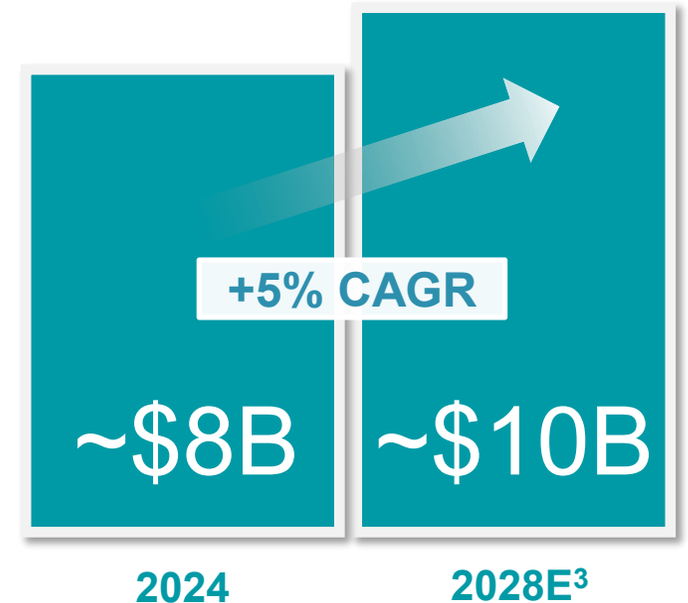
of Outperformance in a Growing Market

B+L Consumer

# #1

Global OTC  
Eye Health Company<sup>1</sup>

Global Consumer Market<sup>2</sup>



Top Factors Fueling  
Category Growth

Growing consumer  
self care mindset

Rising prevalence  
of dry eye

Digital lifestyle and  
environmental factors

Aging  
population

Contact  
lens wear

# Our *Winning Playbook*

SUPERIOR VISION SCIENCE

PROFESSIONAL RECOMMENDATION

PREMIUM INNOVATION

MODERN BRAND BUILDING  
POWERED WITH AI

DIGITAL COMMERCE EXCELLENCE



## Best-in-Class Portfolio of Hero Brands

# PreserVision:

20 Years of Eye Vitamin Leadership, Backed by Extensive Clinical Studies<sup>1,2,3</sup>

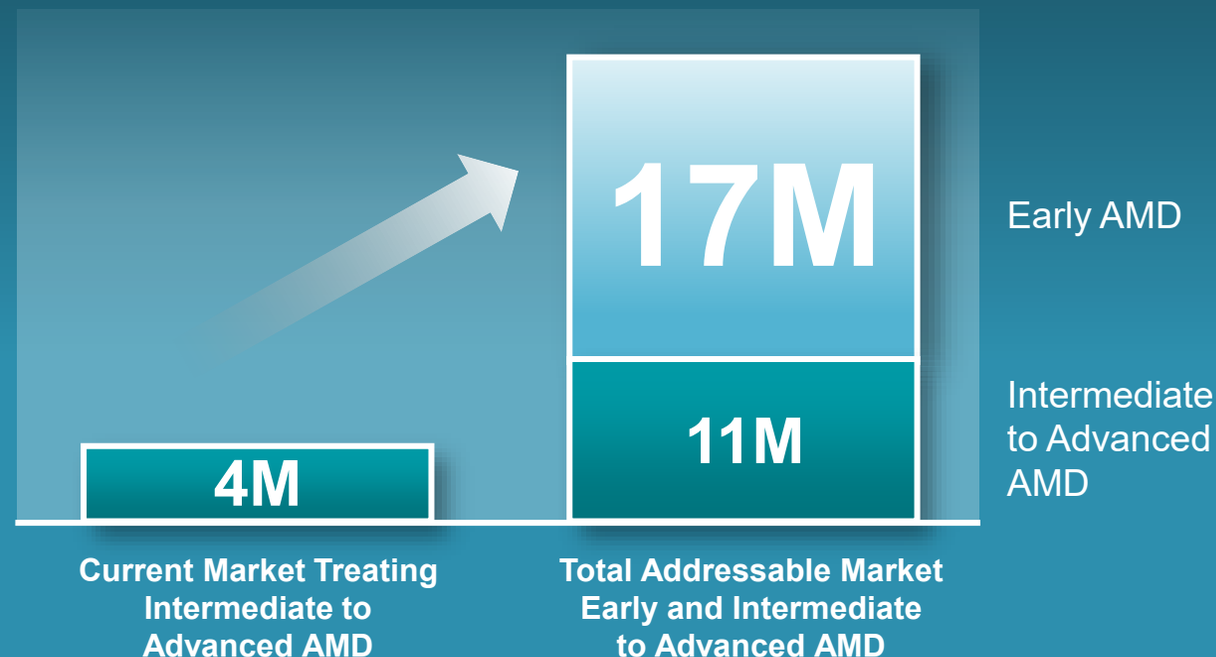
Collaboration  
with the National  
Eye Institute (NEI)

**+8%**  
5-Year CAGR  
(2019-2024)

**#1**  
Eye Vitamin Brand:  
Most Studied and  
Recommended  
Formulation<sup>1,2,3</sup>

**~93%**  
Market Share<sup>4</sup>

Potential to Expand Addressable Market  
by Treating Early AMD<sup>5,6</sup>



# The *Next* Generation of PreserVision: AREDS3

Advancing AMD Care with B-Vitamin Science

## How it Works

B-Vitamins

Boosts Cell  
Metabolism

Reduces  
Inflammation

AREDS2

Lowers  
Oxidative Stress

## B-Vitamin Clinical Data

A growing body of pre-clinical and clinical evidence highlights the critical role B-Vitamins play in supporting retinal health and reducing the risk of AMD

41%

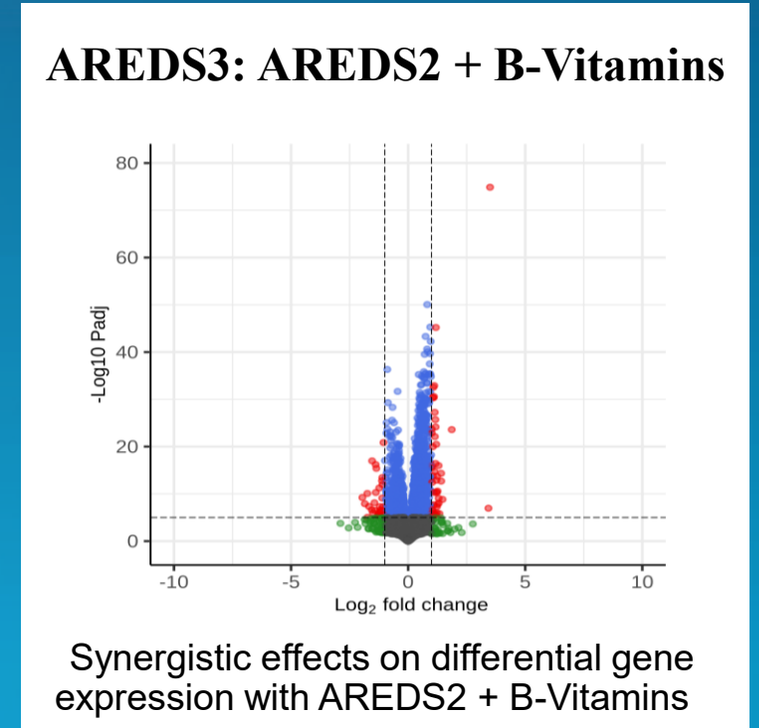
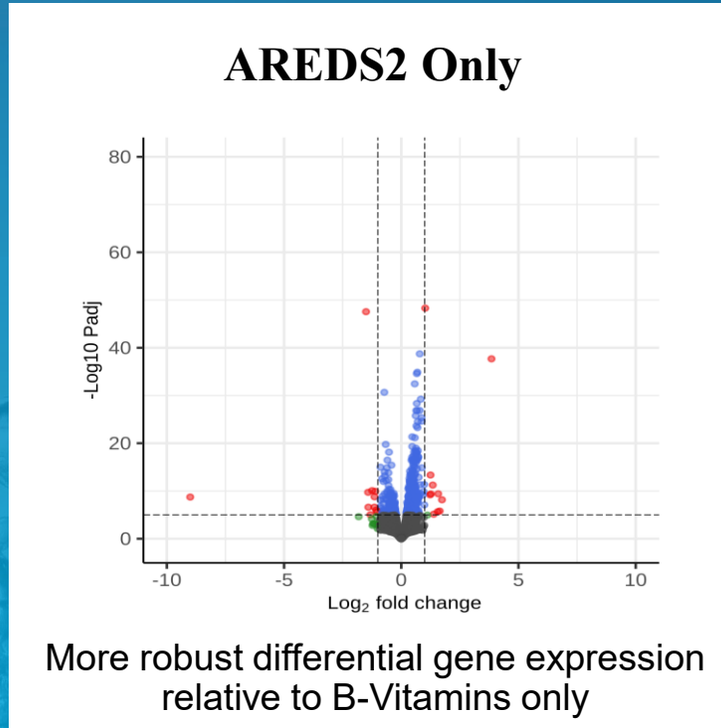
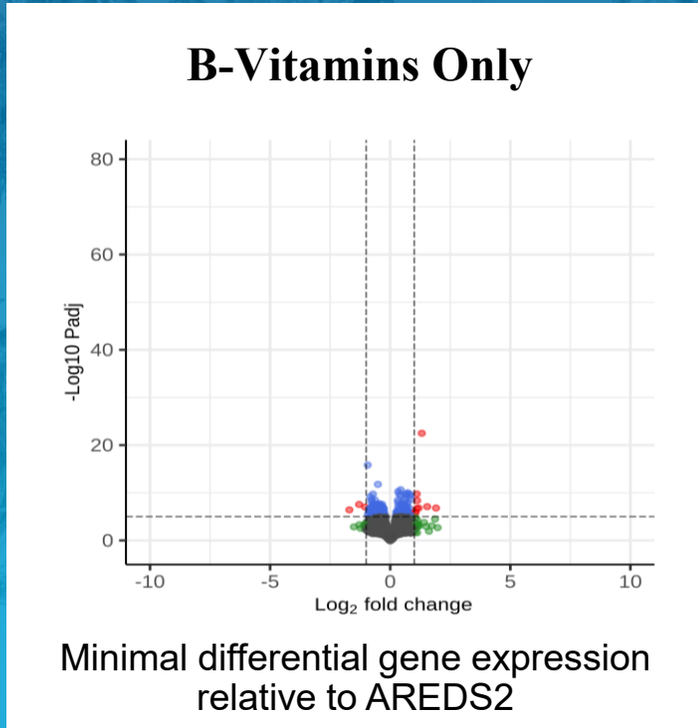
Decreased risk in developing visually significant AMD through daily supplementation with specific B-Vitamins<sup>1</sup>

51%

Reduced risk of developing AMD with normal serum folate levels ( $\geq 10$  nmol/L)<sup>2</sup>

# Human Genetics Support Inclusion of B-Vitamins in AREDS3<sup>1,2</sup>

- 1 Gene Expression Profile following administration of nutritional supplement to iPSC-RPE cells derived from AMD patients. Distinct Transcriptomic Signatures related to retinal health.



- 2 Human genetic analysis identified a genome-wide significant association pointing to Vitamin B metabolism as a potential protective factor in AMD progression



# Lumify is Now a Beauty Routine Essential

#1

eye doctor recommended<sup>1</sup>

#1

share in redness relief<sup>2</sup>

95%

consumer satisfaction<sup>4</sup>

Beauty Enthusiasts Represent Continued Growth Engine<sup>3</sup>

BEAUTY ENTHUSIASTS

100M

2.9M

LUMIFY USERS

Beauty Positioning Supports Premium Pricing

MEET  
THE

**LUMIFY**<sup>TM</sup>  
*Lovers*



# Introducing the Next Generation of Lumify<sup>1</sup>

Help Your Eyes Look Their Best  
& Feel Their Best



Expected  
Launch:

1H27



HA<sup>2</sup> is a moisturizer found naturally in the eye and a **well-established ingredient** in beauty and dry eye drops



Formula selected for a balance of **comfort, viscosity, and stability performance**

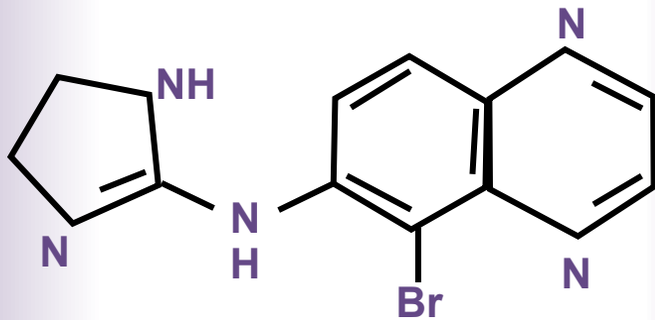


Viscosity optimized for an even more **luxurious feel**

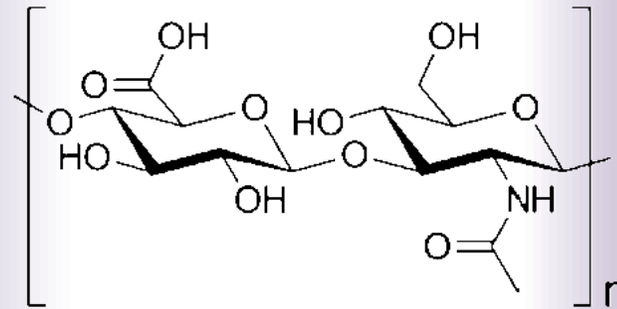
# Lumify Innovation

Brimonidine's Proven Redness Relief Combined with Well Established Safety and Tolerability<sup>1</sup>

## Brimonidine



## Hyaluronic Acid



B+L has filed for patent protection on scientific insights gained during the development and manufacturing of this novel innovative formulation

### Best-in-class molecule

*Highly selective alpha 2 adrenergic receptor agonist*

Potent redness reduction

Low risk of rebound redness or loss of efficacy

### Moisture Retaining

*Natural tear component*

Binds up to 1000X its weight in water to keep the eye moisturized

Provides viscosity and elasticity that protects the eye from mechanical stress

**Novel formulation constituents allow for a 60% reduction in preservatives<sup>2</sup>**

# Lumify + HA Phase 3 Results

## Successful Phase 3 Randomized Controlled Trial Complete



**N=578**

289 study participants treated with novel Lumify + HA



**4 weeks**

of treatment with 1 week follow up post-treatment



**11**

sites

**NDA  
Submission**

Planned for

**1Q26<sup>1</sup>**

Onset as early as



**30 seconds**

sustained redness relief, with improvement compared to baseline through 10 hours<sup>2</sup>



**84%**

of study participants chose **comfortable, cool or refreshing** as the first word they would use to describe the new product<sup>2</sup>

150M

adults in the U.S. experience symptoms of dry eye<sup>1</sup>

33%

of patients with dry eye symptoms treating with OTC products<sup>2</sup>

40%

of patients fail to administer eye drops correctly<sup>3</sup>

# Significant Opportunity to Expand in OTC Dry Eye



**Strong Global Position with Full Suite of Products  
Fastest Growing OTC Dry Eye Portfolio<sup>4</sup>  
+19% LTM Revenue Growth<sup>5</sup>**

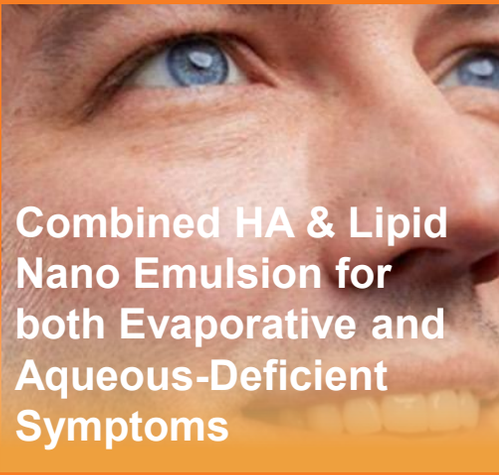
1. MultiSponsor Surveys, Inc. The 2022 Study of Dry Eye Sufferers. August 2022.  
2. 2022 Dry Eye Summary Report; Circana HH Panel % of HH's; Total US All Outlets data ending Sept. 2024.  
3. Mehuys E, C Delaey C, T Christiaens T, et al. Eye drop administration technique and problems reported by eye drop users. Eye. November 5, 2019.  
4. B+L Consumer Data Science: Circana, L52we 7-2025 + IQVIA Flexview MAT June 2025.  
5. Last twelve months as of Sept. 30, 2025.

**blink**<sup>®</sup>  
TRIPLE CARE PF

INNOVATION IN  
FORMULATION



INNOVATION IN  
DELIVERY TECHNOLOGY



Work.  
Play.  
Blink.  
Relief.

RELIEF WITH EVERY  
blink



## Beat the Blink

Novel delivery system designed to improve dosing precision and enhance patient experience by “beating the blink” reflex

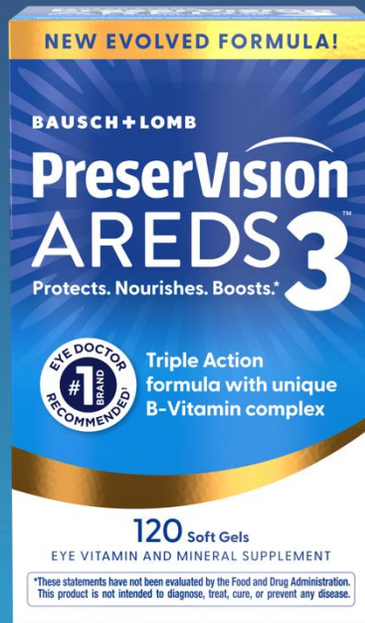
Potential to become platform across Consumer and Pharma products<sup>1</sup>

1. See Slide 2 for further information on forward-looking statements.

# Clinically-Differentiated Consumer Brands<sup>1</sup>

## Expand Market by Addressing all Stages of AMD

Launch Date: 1H26  
Franchise Peak Sales: ~\$600M



## Enhanced Comfort with Addition of Hyaluronic Acid

Launch Date: 1H27  
Franchise Peak Sales: ~\$450M



## Advanced Preservative-Free Lipid Based Formulation

Launch Date: 1H26  
Franchise Peak Sales: ~\$300M



# 3 Key Takeaways<sup>1</sup>

1

Proven Track  
Record of  
Above-Market  
Performance

2

Winning  
Brands with  
Clear Runway  
for Growth

3

Strong Pipeline  
of Clinically-  
Differentiated  
Products

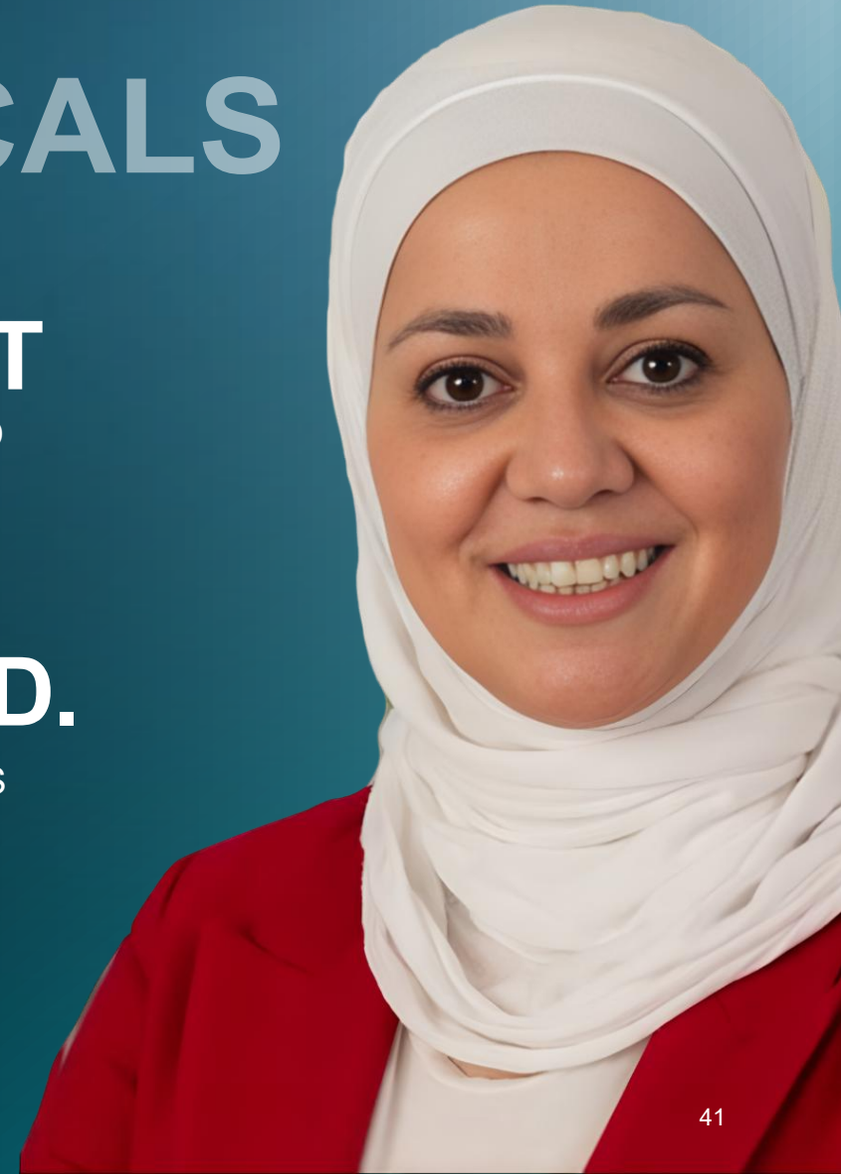
# PHARMACEUTICALS

**ANDREW STEWART**

PRESIDENT, GLOBAL PHARMACEUTICALS AND  
INTERNATIONAL CONSUMER

**MAYSSA ATTAR, PH.D.**

SENIOR VICE PRESIDENT, PHARMACEUTICALS  
AND CONSUMER R&D



# Building on the *Strength* of our Eye Care Platform<sup>1</sup>

Expanding leadership in ocular surface...

...advancing pipeline of novel therapeutics in retina

## Front of the Eye

### **DRY EYE**

First dual-action therapeutic to address both evaporative and inflammatory pathology

### **OCULAR SURFACE PAIN**

First-in-class neurosensory targeted therapeutic for ocular surface pain

## Back of the Eye

### **GLAUCOMA**

First therapy to improve visual function while lowering intraocular pressure

### **AMD / GA**

Multiple “shots-on-goal” to transform AMD treatment and disrupt GA landscape



# Finely-Tuned & Proven Commercialization Engine

**Miebo**  
(perfluorohexyloctane  
ophthalmic solution)

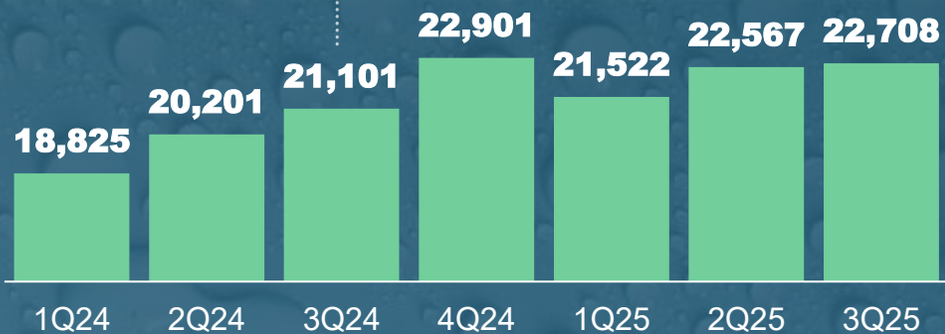
+110%



Avg. Weekly TRx & Growth (% Y/Y)<sup>1</sup>

**xiidra**  
(lifitegrast  
ophthalmic solution)<sup>5%</sup>

+8%



**Highest  
Rated**  
field force<sup>2</sup>

**Best**  
eye health launch  
2022-2024<sup>3</sup>

**Top 10**  
launches across  
all disease states<sup>4</sup>

1. IQVIA NPA Rapid Rx. Historical data was restated.  
2. ZoomRx PET.  
3. National Sales Perspective, National Prescription Audit, Patient Insights; Launch Center of Excellence, IQVIA.  
4. National Sales Perspective; Launch Center of Excellence, IQVIA.

# Dry Eye Disease:

## Rising Unmet Need, *Big Opportunity*

Comprehensive B+L Platform to Drive Market Penetration and Patient Adherence

**93%**

of estimated U.S. population with DED is not treated with an Rx product<sup>1</sup>

**7%**

of diagnosed patients receive Rx DED treatment<sup>1</sup>

**~150M**

U.S. adults experience symptoms of dry eye, with ~38M living with DED<sup>2,3</sup>

**~2x**

Global dry eye disease market expected to nearly double by 2030<sup>4</sup>

**~90%**

Of DED patients discontinue initial medication within one year<sup>5</sup>

# Therapeutic Strategy to Elevate Standard of Care

  
**Dry Eye is a Complex Disease**



Dry eye is a **multifactorial**, symptomatic disease characterized by a **loss of homeostasis of the tear film and/or ocular surface**, in which tear film instability and hyperosmolarity, **ocular surface inflammation and damage**, and **neurosensory abnormalities** are etiological factors

  
Updated TFOS DEWS III Definition

  
**Disease Drivers and Therapeutics**

Most dry eye is evaporative in nature



Indicated to treat signs and symptoms of dry eye. Unique MOA.

Inflammation can be both a cause and a consequence of dry eye



Only anti-inflammatory eye drop approved for chronic use on the basis of treating a sign and symptom of dry eye.

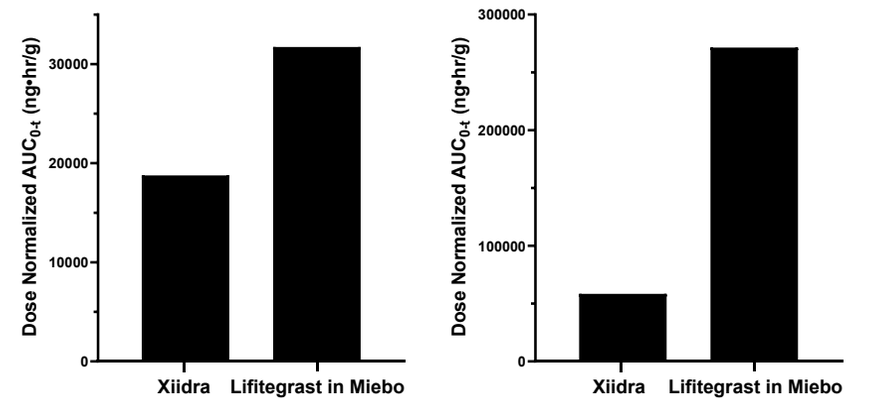
  
**Rationale for Dual-Action Eye Drop**

-  Lifitegrast (active in Xiidra) and perfluorohexyloctane (active in Miebo) act with distinct mechanisms
-  Today, only single-action products
-  Faster relief and larger treatment effect may be achieved with first dual-action eye drop to treat dry eye
-  Anticipate superior improvement in signs and symptoms with improved tolerability

# Miebo-Based Formulation Delivers Lifitegrast More Efficiently

## Ocular Surface Tissue Concentrations

Cornea<sup>1</sup>      Conjunctiva<sup>1</sup>



### More Efficient Drug Delivery

More tissue penetration when lifitegrast (active in Xiidra) is formulated with the active in Miebo



#### Unique physiochemical properties of new formulation allows:

- Reduced total lifitegrast dose compared to Xiidra
- Smaller drop size that delivers efficacious drug levels



#### These attributes are designed to achieve:

- Dual-action efficacy superior to Xiidra or Miebo
- Improved tolerability



#### Clinical study ongoing with data anticipated 2H26<sup>2</sup>

- Designed to test dual-action of novel lifitegrast in Miebo formulation and superiority to Xiidra and Miebo in treating dry eye



Improved formulation designed to achieve superior dual-action efficacy with improved tolerability

1. Data on file. Single topical dose rabbit pharmacokinetics comparing Xiidra to lifitegrast in Miebo.  
2. See Slide 2 for further information on forward-looking statements.

# #1 Reason for ECP Visit is Ocular Discomfort & Pain<sup>1</sup>

## Acute OSP

**Steroids and NSAIDs not sufficient treatment for many patients developing AOSP**

**10-15%**

Post-cataract and refractive surgery patients (4.5-5M patients annually)<sup>2</sup>

Other relevant conditions include

- Corneal abrasions
- Ocular infections
- Dry Eye flares

## Chronic OSP

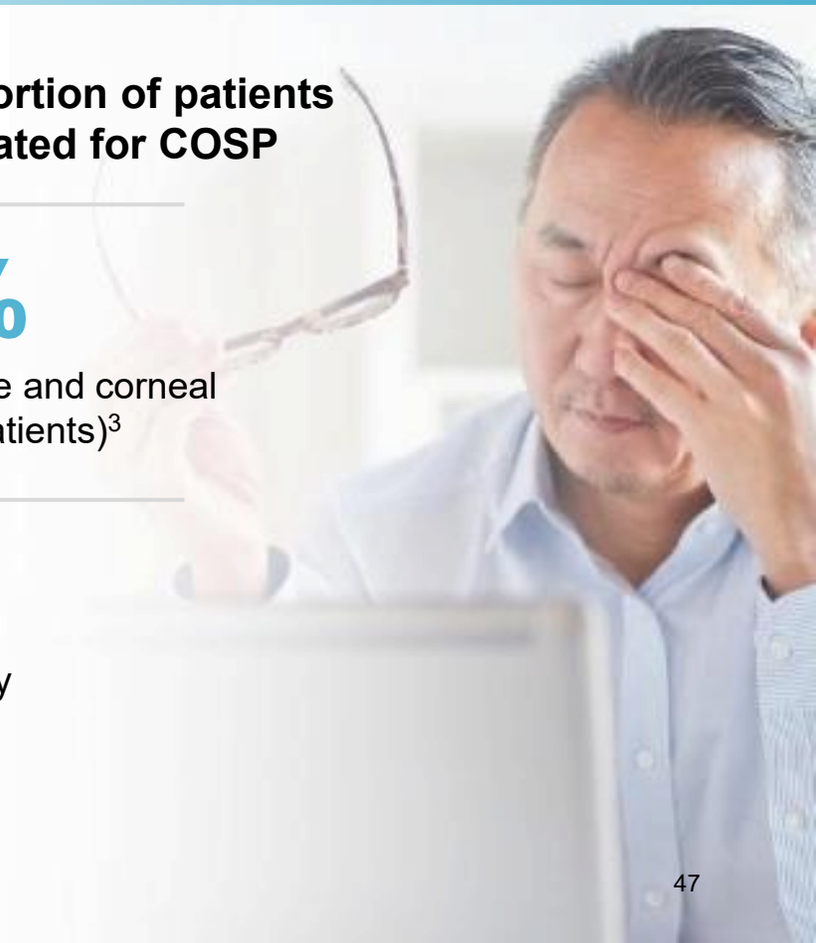
**Significant proportion of patients inadequately treated for COSP**

**20-30%**

Patients with Dry Eye and corneal diseases (10-15M patients)<sup>3</sup>

**5-15%**

Patients post-surgery (0.4M patients)<sup>3</sup>



# No Glaucoma Therapeutic Exists to Treat the Vision Threatening Neurodegenerative Pathology

**4.2M**

individuals in U.S. have glaucoma<sup>1</sup>

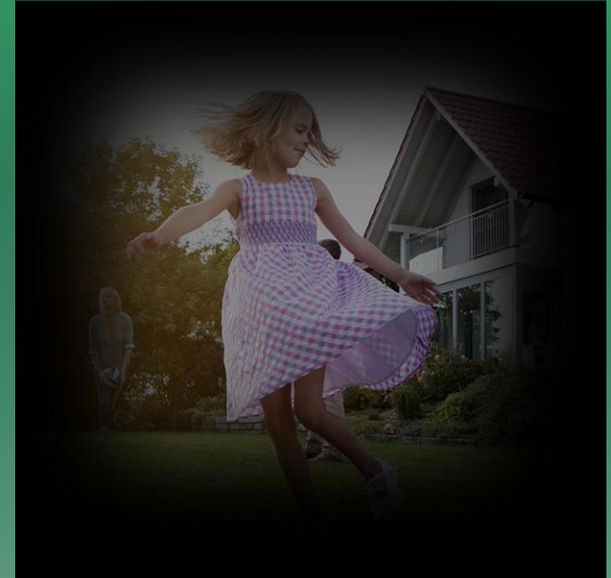
**35%**

of patients with glaucoma have vision-affecting glaucoma<sup>1</sup>

GLAUCOMA CONTINUES TO BE THE “SILENT THIEF OF SIGHT”



Normal Vision

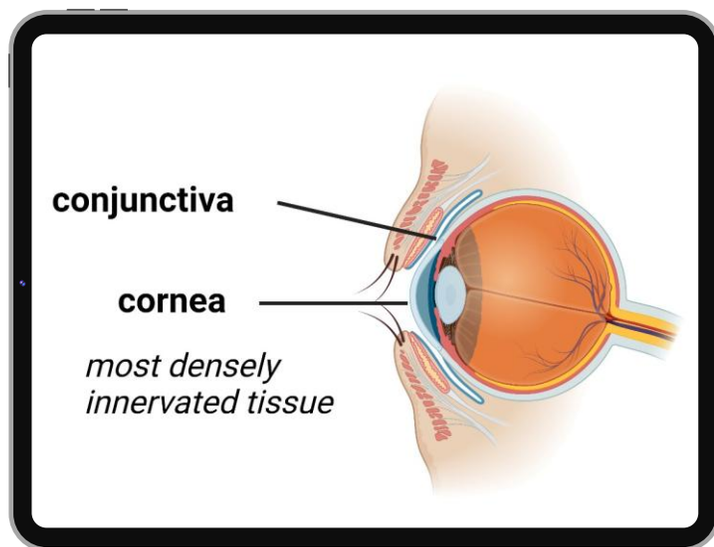


Glaucoma

1. <https://jamanetwork.com/journals/jamaophthalmology/fullarticle/2824476>.

# TRPV1 Antagonism to Treat Ocular Surface Pain

## Cornea has Highest Density of Sensory Nerve Endings



TRPV1 ion channels represent a primary sensor of pain

TRPV1 antagonists block the cascade leading to pain

## Target Rationale



**Target** TRPV1; transient receptor potential vanilloid subtype 1; nociceptor



**Function** Cell surface receptor ion channel critical for sensing of nociceptive and thermal inflammatory pain<sup>1</sup>



**Ocular Tissue Distribution** Cornea, Conjunctiva, peripheral and central terminals of sensory neurons<sup>2</sup>



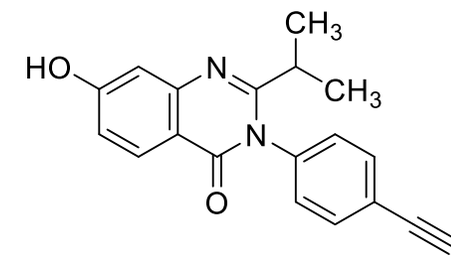
**MOA** Non-competitive antagonist of TRPV1

## Molecules



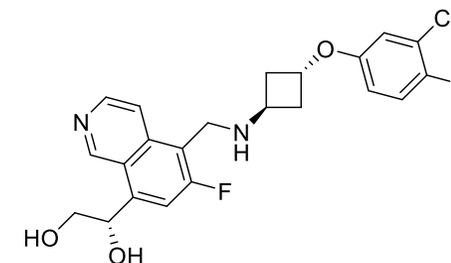
**BL1312**

Achieved positive clinical POC



**BL1332**

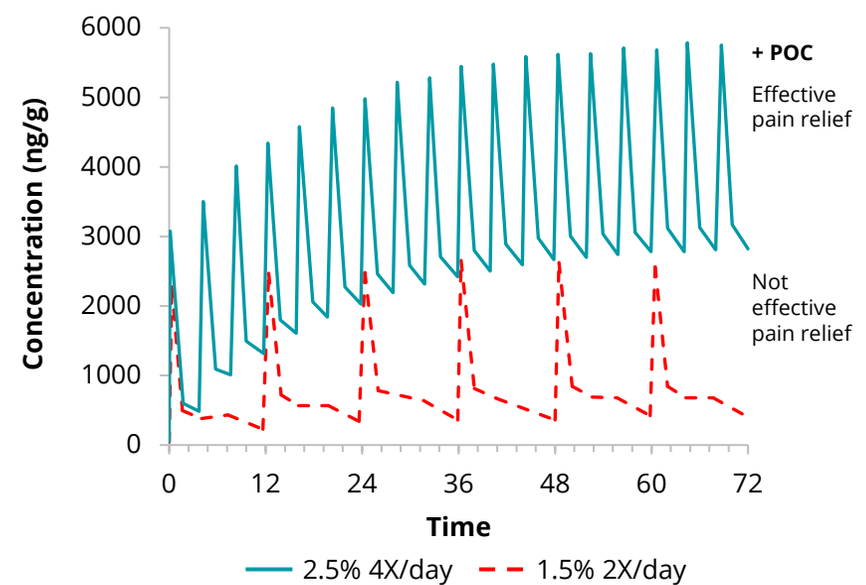
More potent water-soluble molecule



# Data-Informed Development of TRPV1 Antagonist

## BL1312 Clinical Data Guide Strategy

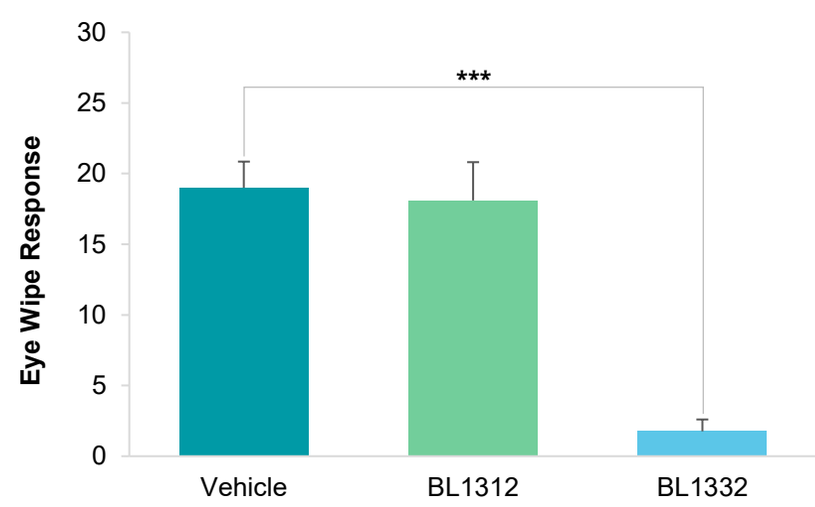
Simulated Human Cornea BL1312 Concentrations<sup>1</sup>



Maximize efficacy by increasing drug exposure

## Developing More Potent BL1332

Capsaicin-Induced Nonclinical Model for Evaluating Analgesic Efficacy<sup>1</sup>



Increased potency translates to increased efficacy

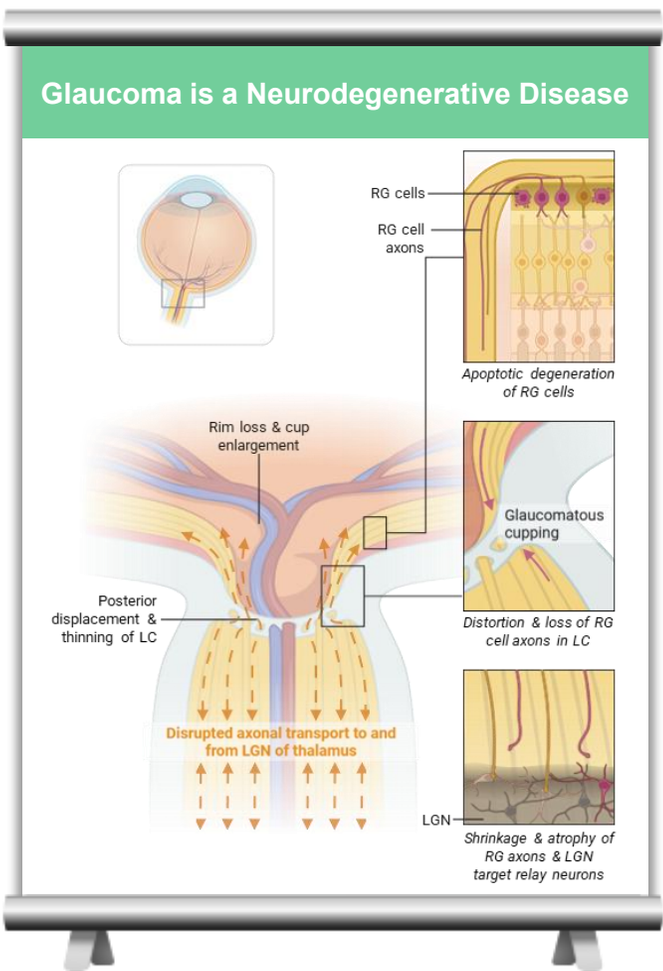
1 BL1332 Phase 1 complete and highest dose was safe and well tolerated

2 BL1332 Phase 2 POC planned to initiate 1H26<sup>2</sup>

More potent BL1332 molecule formulated to maximize tissue exposure to block ocular pain signaling 

1. Data on file pharmacokinetic human cornea simulation and nonclinical capsaicin model.  
2. See Slide 2 for further information on forward-looking statements.

# Addressing Neurodegenerative Vision Loss in Glaucoma




Glaucoma is a **neurodegenerative disease** with loss of retinal ganglion cells and optic nerve injury



Studies show most patients with mild to moderate glaucoma have **macular damage** and **vision complaints under low luminance**<sup>1,2</sup>



**No approved therapeutics to treat vision loss** associated with glaucoma



**Brimonidine-treated glaucoma patients were less likely to develop visual field loss** compared to timolol even when IOP was decreased to the same extent.<sup>3</sup> Brimonidine is an **alpha2 adrenergic receptor (α2AR) agonist**



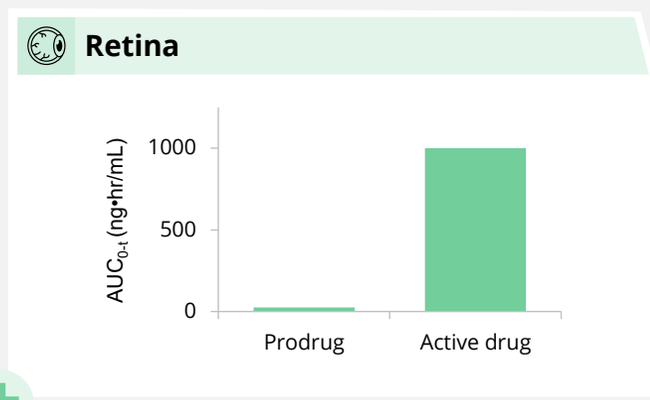
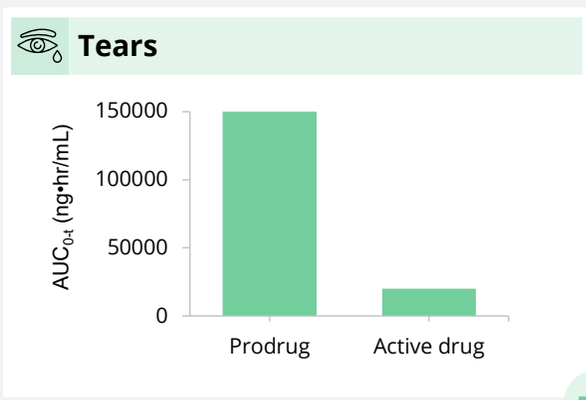
**Mechanisms α2AR agonists improve vision:**

1. Neurofunctional enhancement (improve vision in days to weeks)
2. Neuroprotection (protect from vision-loss over months)

1. DOI: [10.1016/j.ajo.2019.08.024](https://doi.org/10.1016/j.ajo.2019.08.024).  
 2. [doi.org/10.1111/aos.13695](https://doi.org/10.1111/aos.13695).  
 3. DOI: [10.1016/j.ajo.2010.09.026](https://doi.org/10.1016/j.ajo.2010.09.026); glaucoma neurodegeneration figure created in BioRender.

# BL1107 is Designed to Target Vision-Threatening Disease

Prodrug *drives* tissue penetration



## Next-Generation α2B AR Agonist

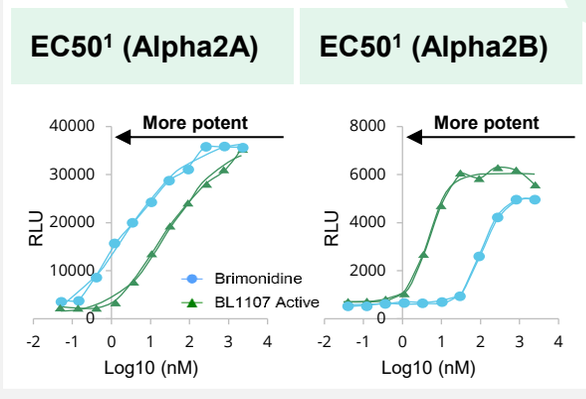


Molecular attributes to target vision-threatening disease

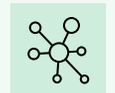
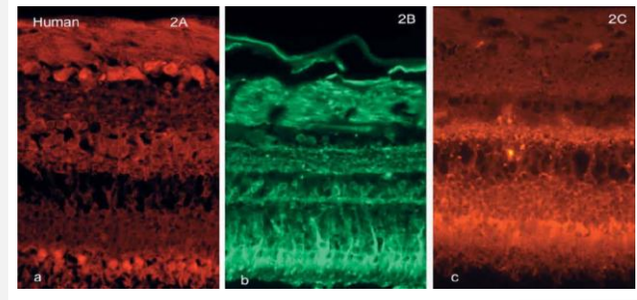
Better penetration to retina

More potent and selective agonism of relevant receptor

Selective and potent α2B agonism



α2B is widely expressed in retina & mediates beneficial effects on neuronal function<sup>2</sup>



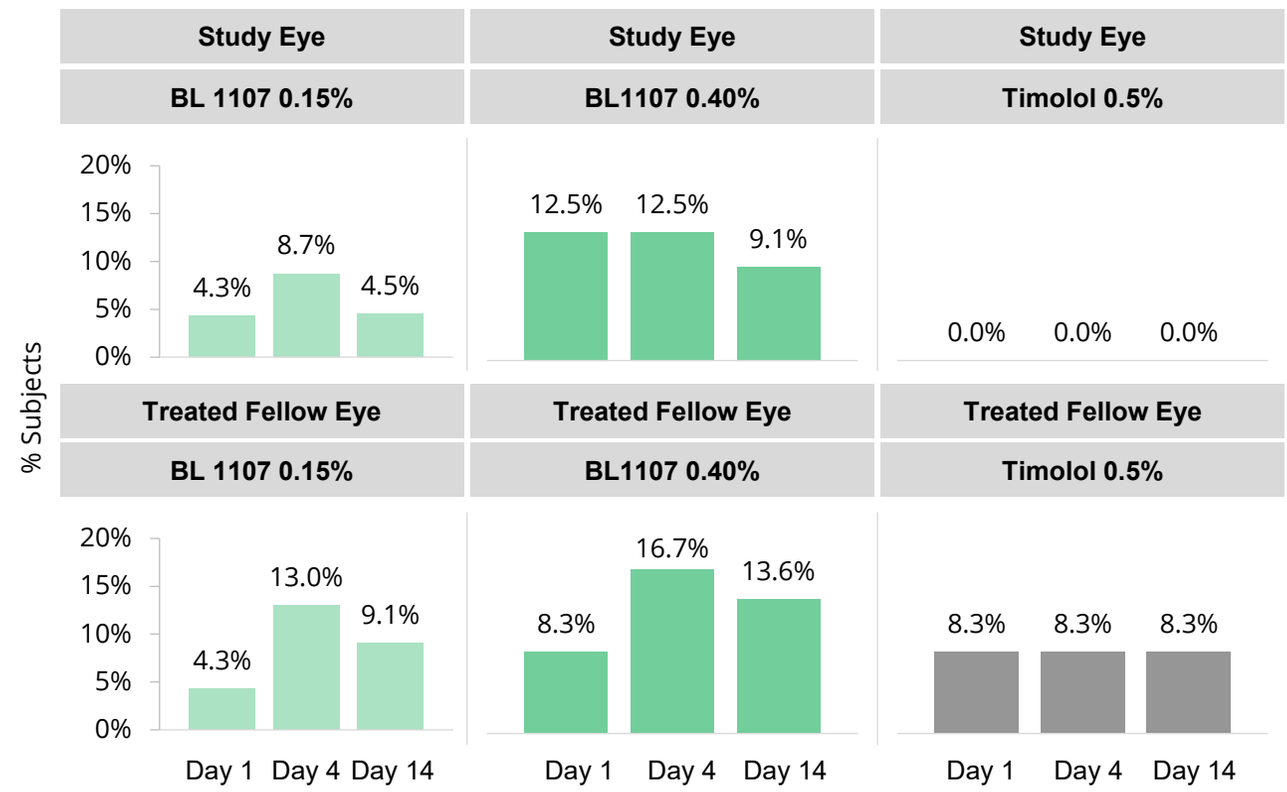
Rapidly improve vision via neurofunctional enhancement and long-term preserve vision via neuroprotection

Enhanced retinal penetration combined with selective and potent α2B agonism should maximize vision benefits

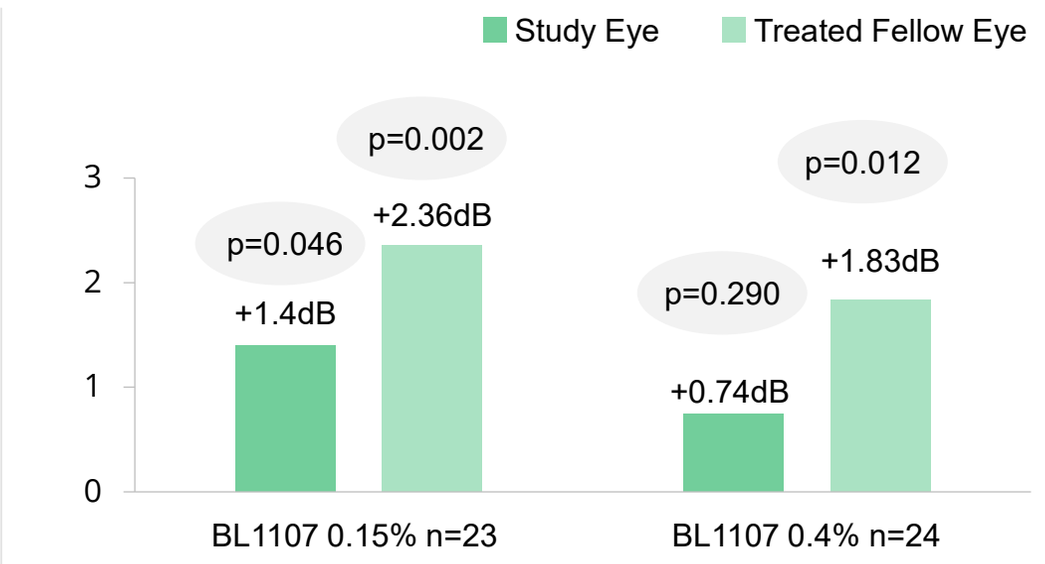
1. Data on file.  
2. DOI: 10.1017/S0952523807070605.

# BL1107 Achieved Clinical POC: Improves Vision in Glaucoma

## LL-BCVA ≥15-Letter Gain<sup>1</sup>



## Treatment Difference in Mean CFB at Day 14 in VF MD<sup>1</sup>



**Comparing patients treated with BL1107 to timolol:**

- Greater proportion experienced ≥15-letter gain in LL-BCVA
- Statistically significant improvement in VF MD

 Ongoing larger Phase 2 study powered to demonstrate visual neuroenhancement effects is anticipated to readout 2H26<sup>2</sup>. These data will guide Phase 3 study design.

<sup>1</sup> Data on file. LL-BCVA: low luminance best corrected visual acuity; CFB: change from baseline; VF MD: visual field mean deviation.  
<sup>2</sup> See Slide 2 for further information on forward-looking statements.

# Addressing Massive Unmet Needs for ~20M Americans with AMD/GA <sup>1</sup>

Intermediate  
Dry AMD



Geographic  
Atrophy



Currently approved therapies only slow lesion growth. B+L is targeting new treatments that could<sup>2</sup>:



Preserve vision by slowing disease progression



Show greater reduction in rate of lesion growth and/or visual function



Reduce CNV risk and lower rate of intraocular inflammation



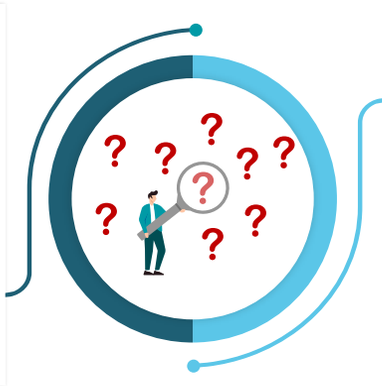
Reduce injection burden

1. Rein DB, Wittenborn JS, Burke-Conte Z, Gulia R, Robalik T, Ehrlich JR, Lundeen EA, Flaxman AD. Prevalence of Age-Related Macular Degeneration in the United States in 2019. JAMA Ophthalmology. 2022.  
2. See Slide 2 for further information on forward-looking statements.

# Strategy for Developing Complex Disease Treatments

*Drugging a complex disease with many implicated mechanisms, no animal models recapitulating disease etiology, and slow disease progression*

Difficult to select pharmacological targets that impact clinical disease endpoints in humans



Very slow and expensive to find out if preclinical science translates to humans

## Strategy

1 Leverage human data to identify targets and patient selection criteria

2 Design optimal therapeutic modality and molecule

3 Best-in-class drug delivery approaches

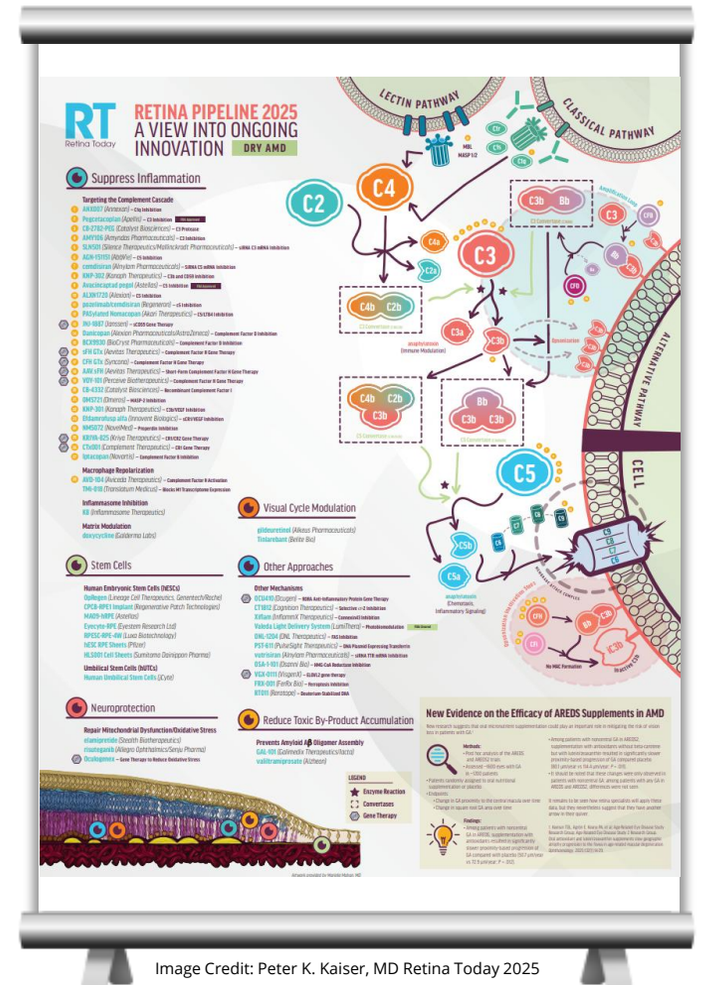


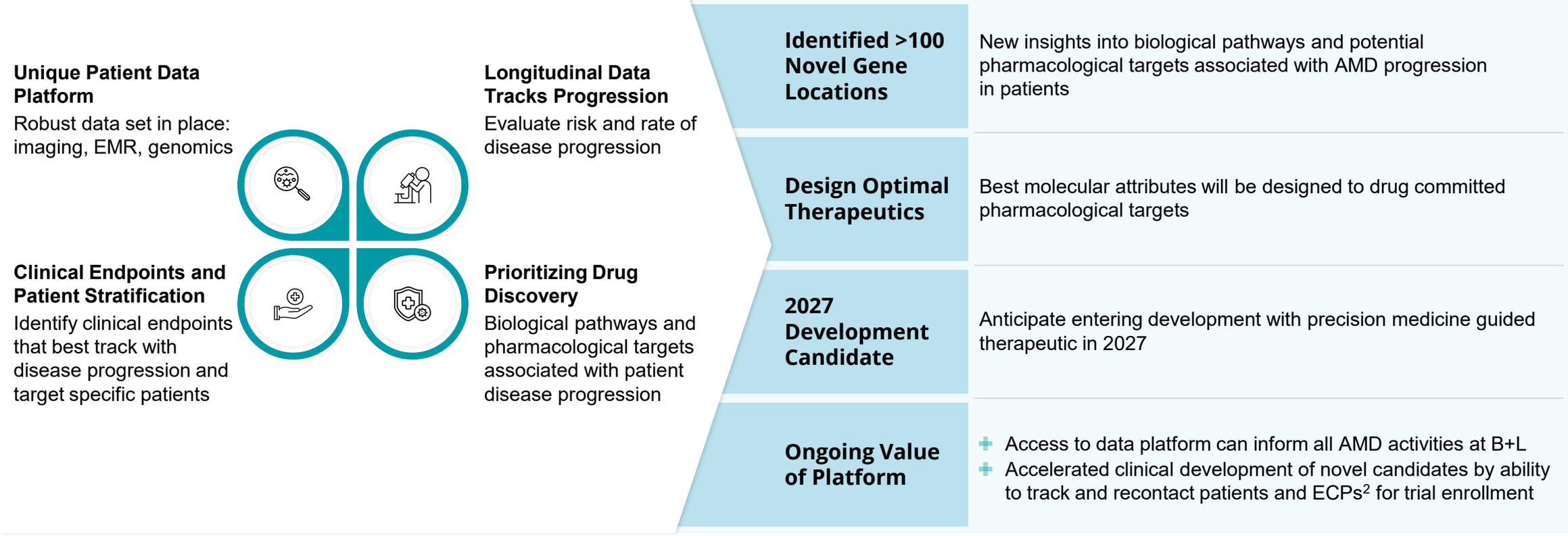
Image Credit: Peter K. Kaiser, MD Retina Today 2025

# Precision Medicine Will Transform AMD Treatment<sup>1</sup>

Partnership grants exclusive access to integrated patient data platform and AI-powered analytical engine to drive novel intermediate AMD<sup>2</sup> and GA<sup>2</sup> drug discovery and development



## Strategic Advantage      Progress



Insights enable us to select the right patients for each target, accelerating development and improving the probability of success

1. See Slide 2 for further information on forward-looking statements.  
2. AMD: age related macular degeneration; GA: geographic atrophy; ECP: eye care professional.

# Gene Silencing for Unprecedented Potency and Durability

Advantages of genetic medicine approach without the challenges of viral gene delivery

**Local delivery into a small closed physiological space enhances target engagement and minimizes systemic safety risks**

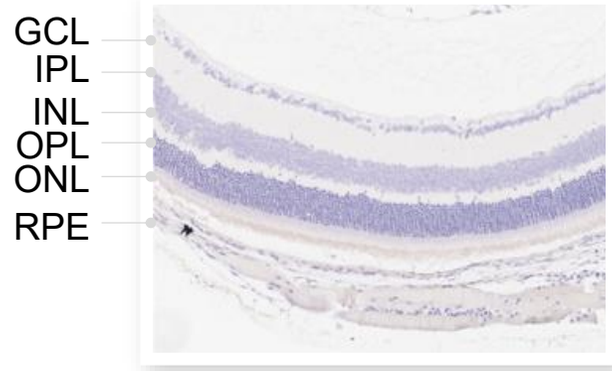
**Optimize drugging of clinically validated targets to de-risk overall development**

**Better molecules: more potent, efficient and specific**

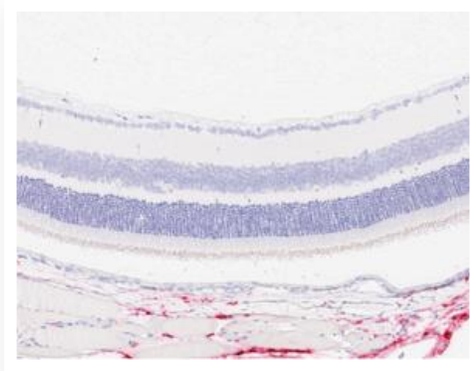
**Better delivery: novel targeting ligands to enable robust delivery**



## Broad retina distribution and robust target knockdown following intravitreal injection into mouse eye<sup>1</sup>



Saline



Novel siRNA molecule



Novel siRNA molecule



Targeting ligand

**Anticipate development candidate in 2H26**

1. Data on file for mouse IVT study.

# BL1107 Sustained-Release Implant is a Superior GA Treatment



## α2AR Agonist Slows GA Progression

- In 2 RCT, brimonidine slowed GA lesion growth but treatment effect was relatively small<sup>1,2</sup>
- Sustained release of an α2AR agonist feasible
- No increased risk of neovascular AMD or intraocular inflammation



## BL1107 a Perfect Candidate for GA

- Specific for α2B in the retina
- Better retina penetration
- Compatible with sustained-release
- Potential for both improving vision and decreasing GA lesion growth



## Development Path

Development enabled through partnership with



Anticipate development candidate in 2026 and approval in 2030+ for first small molecule sustained release implant for GA<sup>3</sup>

Criteria		Target Product Profile
 <b>Product Attributes</b>	<b>Indication</b>	Treatment of GA secondary to AMD
	<b>Treatment Frequency</b>	Single IVT administration every 3-6 months
	<b>Efficacy Outcomes</b>	GA Lesion Growth and Vision
	<b>Safety</b>	Reduced rate of conversion to neovascular AMD Absence of severe complications such as retinal vasculitis and/or retinal vein occlusion

# Opportunity for Pipeline to Deliver ~\$3.9B in Peak Sales<sup>1,2</sup>

Indication	Program	Thesis	Phase 3 Data	Launch Date	Peak Sales
Dry Eye	Dual-Action Eye Drop	First dual-action therapeutic for evaporative and inflammatory DED	~2028	~2029	~\$0.7B
OSP	BL1332	First neurosensory agent to address ocular surface pain	~2029	~2030	~\$1.4B
Glaucoma	BL1107	First glaucoma therapy to improve visual function through neuroprotection	~2030	~2031	~\$0.8B
AMD/GA	Retina Pipeline	First therapeutic for intermediate AMD and best-in-class treatments for GA	2030+	2030+	>\$1.0B

# 3 Key Takeaways<sup>1</sup>

1

Strengthening  
Leadership in  
Dry Eye

2

Expanding  
Front of the  
Eye Platform

3

Building Pipeline  
of Therapeutics  
in Retina

# CONTACT LENS

**YANG YANG**

PRESIDENT, VISION CARE

**BRYAN REED**

VICE PRESIDENT, VISION CARE R&D

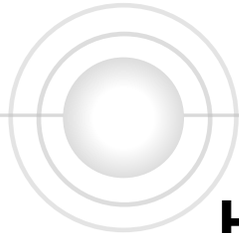
NO BREAKTHROUGH  
INNOVATION IN  
CONTACT LENS INDUSTRY  
FOR OVER 25 YEARS

# Introducing the *First* *Bioactive* Contact Lens Material



# Creating a *New Category* of Contact Lenses<sup>1</sup>

1971



**Hydrogel**



B+L launches SofLens

The first mass-produced soft contact lens in the market

1999



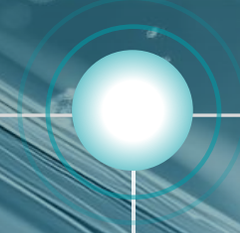
**Silicone Hydrogel**



B+L launches PureVision

The first silicone hydrogel contact lens

2028



**Bioactive Hyaluronic Acid Hydrogel**



**The first bioactive contact lens**

1. See Slide 2 for further information on forward-looking statements.

# *A First-of-its-Kind* Technology

Re-engineered from the inside out.

Double dose of bio-engineered HA.

HA polymerized into lens and in solution.

Hyaluronic acid (HA)  
 High-molecular-weight  
 polysaccharide that can  
 hold up to  
**1000x**  
 its weight in water<sup>1</sup>

Hydrating

Acts as "nature's sponge" to provide maximum hydration

Lubricating

Reduces friction and mechanical stresses

Natural

Found throughout the body in eyes, joints, skin

Cornea

**Hyaluronan Modulates the Biomechanical Properties of the Cornea**

Xiao Lin,<sup>1</sup> Taye Mekonnen,<sup>2</sup> Sudhir Verma,<sup>1,3</sup> Christian Zevallos-Delgado,<sup>2</sup> Manmohan Singh,<sup>2</sup> Salavat R. Aglyamov,<sup>4</sup> Tarsis F. Gesteira,<sup>1</sup> Kirill V. Larin,<sup>2</sup> and Vivien J. Coulson-Thomas<sup>1</sup>

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<sup>2</sup>Department of Biomedical Engineering, University of Houston, Houston, Texas, United States  
<sup>3</sup>Department of Zoology, Deen Dayal Upadhyaya College, University of Delhi, Delhi, India  
<sup>4</sup>Department of Mechanical Engineering, University of Houston, Houston, Texas, United States

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**Purpose.** Hyaluronan (HA) is a major constituent of the extracellular matrix (ECM) that has high viscosity and is essential for maintaining tissue hydration. In the cornea, HA is enriched in the limbal region and is a key component of the limbal epithelial stem cell niche. HA is upregulated after injury participating in the formation of the provisional matrix, and has a key role in regulating the wound healing process. This study investigated whether changes in the distribution of HA before and after injury affects the biomechanical properties of the cornea in vivo.

# Bioactive

*Interacts with the Natural Biology of the Eye to Release HA*



**Unmet Need:** Increased contact lens dryness symptoms as day progresses

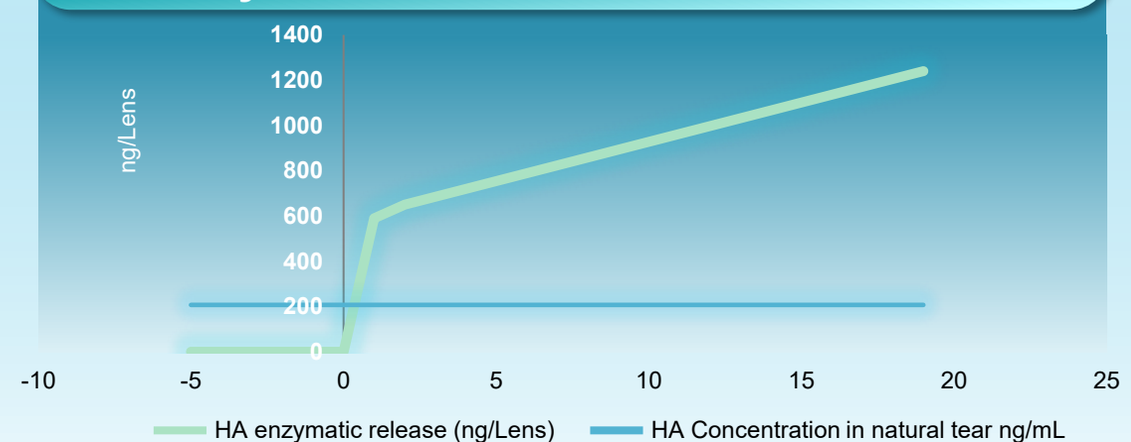
When exposed to hyaluronidase, a natural enzyme in tears, **HA is released** from the lens surface

Steady, consistent release profile through 19 hours of testing for **improved end of day hydration and comfort**

## % of Contact Lens Wearers Who Report Dryness Symptoms<sup>1</sup>



## Enzymatic Release of HA from Lens<sup>2</sup>



# Optimal Hydration and Eye Health

## *Cell and Evaporation Protection + Oxygen Permeability*

HA protects corneal epithelial cells' metabolic activity after exposure to dry environment

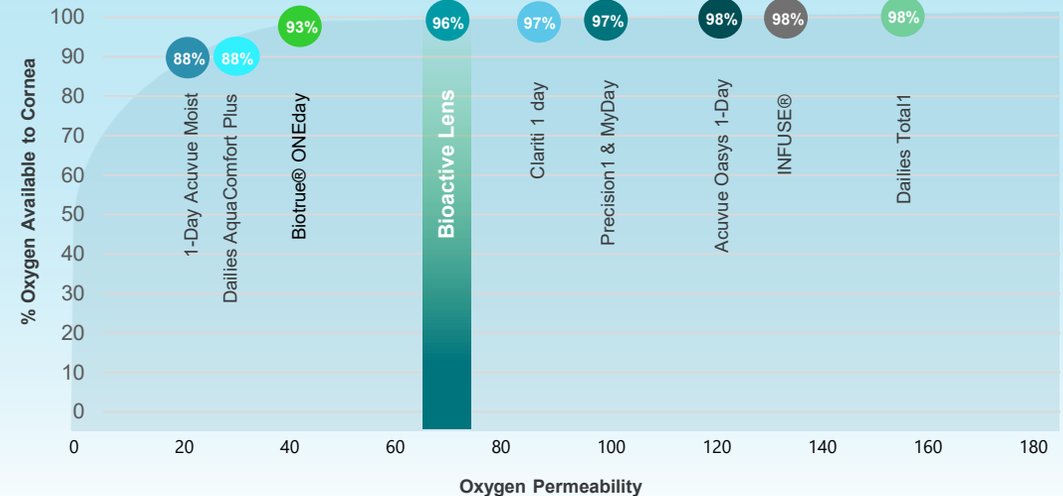
HA supports improved tear film stability by reducing evaporation

Healthy oxygen permeability achieved without requiring silicone (Dk 65% higher than ISO prediction)

### Desiccation Protection<sup>1</sup>



### % Oxygen Available to Cornea<sup>1</sup>



# Unparalleled Comfort

*Lowest Coefficient of Friction Minimizes Interaction Between Eyelid & Lens*

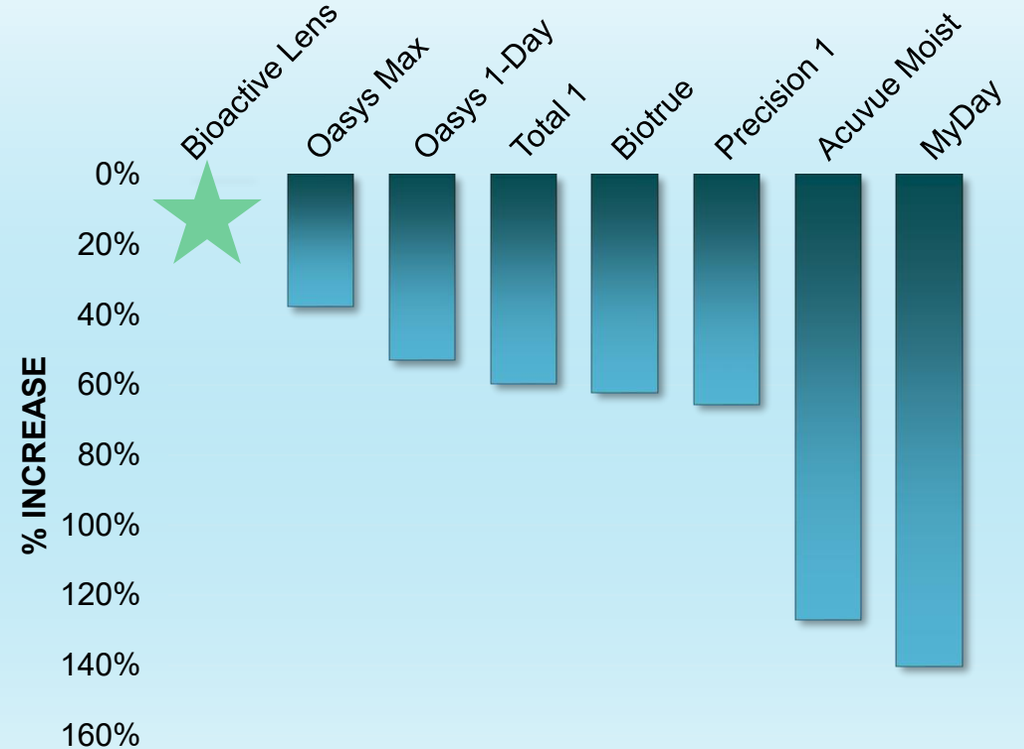
**14** blinks per minute

**840** blinks per hour

**13,440** blinks per 16-hour day

Initiated external performance clinical in 4Q25

## Coefficient of Friction<sup>1</sup>



# Contact Lens Pipeline Designed with *Purpose*<sup>1</sup>

DRIVE REVENUE GROWTH  
ACROSS ALL MARKET  
CATEGORIES

DRIVE MARGIN EXPANSION  
BY LEVERAGING EXISTING  
MANUFACTURING PLATFORMS



**PROJECT HALO**

**FIRST-OF-ITS-KIND  
BIOACTIVE  
CONTACT LENS  
MATERIAL**

Launch Date: 2028  
Peak Sales: ~\$500M



**2ND DD SIHY LENS  
INNOVATION  
DESIGNED FOR  
AFFORDABILITY**

Launch Date: 2029  
Peak Sales: ~\$250M



**PREMIUM FRP  
SIHY PROVIDING  
UNSURPASSED  
LENS COMFORT**

Launch Date: 2029  
Peak Sales: ~\$300M



**CUTTING-EDGE SIHY  
LENS DESIGN TO  
SLOW PROGRESSION  
OF MYOPIA**

Launch Date: 2029  
Peak Sales: ~\$200M

# Next-Generation DD SiHy

## *Accessible Innovation in Fastest Growing Lens Category*

Global Daily Silicone Hydrogel Market<sup>1</sup>



**Material Innovation**

Novel chemistry designed for high surface wettability

**Maintain Homeostasis**

Integrated with moisturizers, electrolytes and osmoprotectants

**Efficient Manufacturing**

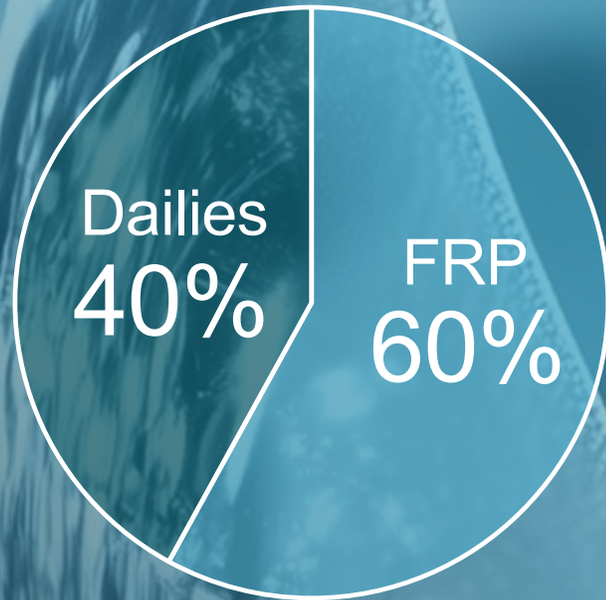
Leverages existing equipment, minimal capital required

Initiating external performance clinical study in 2026<sup>2</sup>

# Premium FRP SiHy

*Designed to be the Premium FRP Lens in the Market*

FRP User Base Remains Large,  
with Limited Innovation<sup>1</sup>



Contact Lens Wearers by Modality

## Unsurpassed Comfort

Comfort of a daily and affordability of a monthly – optimized to balance hydration, breathability and resist deposits

## Crisp Clear Stable Vision

Leverages proven optical designs across SVS, MF, Toric, MFT family

## B+L Lens Care Synergy

Engineered to integrate seamlessly with B+L lens care portfolio

Initiating external performance clinical study in 2026<sup>2</sup>

# DD SiHy Myopia Control Lens

## *Novel Design to Control the Progression of Myopia*

Premium DD SiHy Material

*Paired with*

Cutting-Edge Lens Design

Differentiated peripheral defocus design with

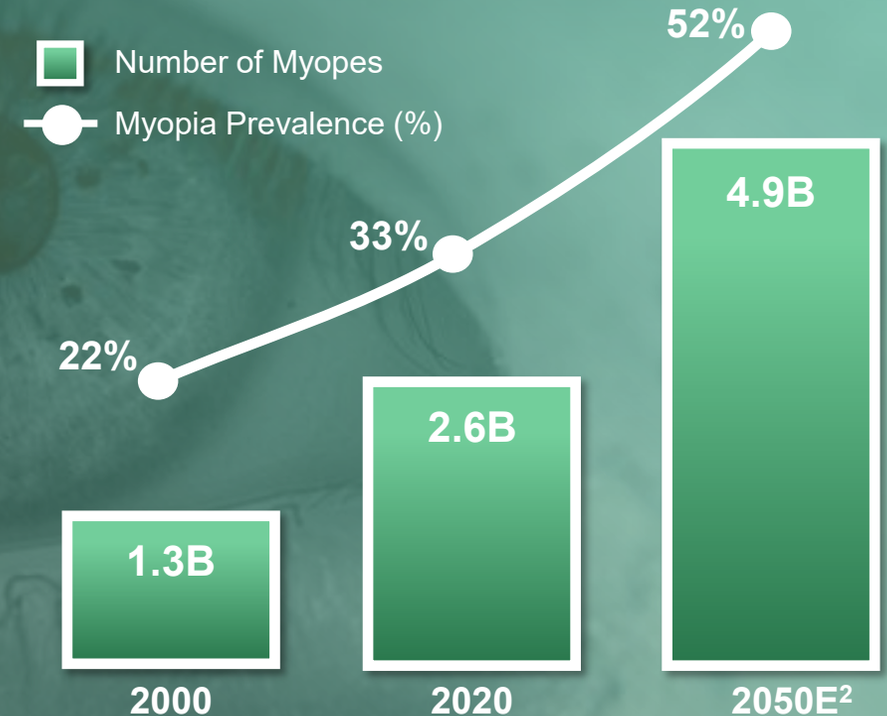
- 1) Multiple Zone Diameters
- 2) Dose Adjustment Across SKU Range

“Made for Kids”

Innovative customized approach to balance controlling myopia progression with daily vision needs

Initiating registration clinical study in U.S. in 1H26<sup>2</sup>

Myopia – Now and in 2050<sup>1,2</sup>



# Continuing to *Outperform* Market And Built Best-in-Class Pipeline to Sustain Leading Growth<sup>3</sup>



3Q25 YTD CC<sup>1</sup> Revenue Growth<sup>2</sup>

Capturing Share in **Fastest Growing DD SiHy** Market

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Driving **Segment-Creating** Material Innovation

---

Entering **New Categories** with High Unmet Need

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Addressing **All Wearer Needs** with Comprehensive Portfolio

# 3 Key Takeaways<sup>1</sup>

1

Creating New  
Category of  
Contact  
Lenses

2

Advancing  
Pipeline to  
Sustain Market  
Leading Growth

3

Leveraging  
Existing  
Platforms to  
Drive Margin



# SURGICAL

**LUC BONNEFOY**

PRESIDENT, SURGICAL

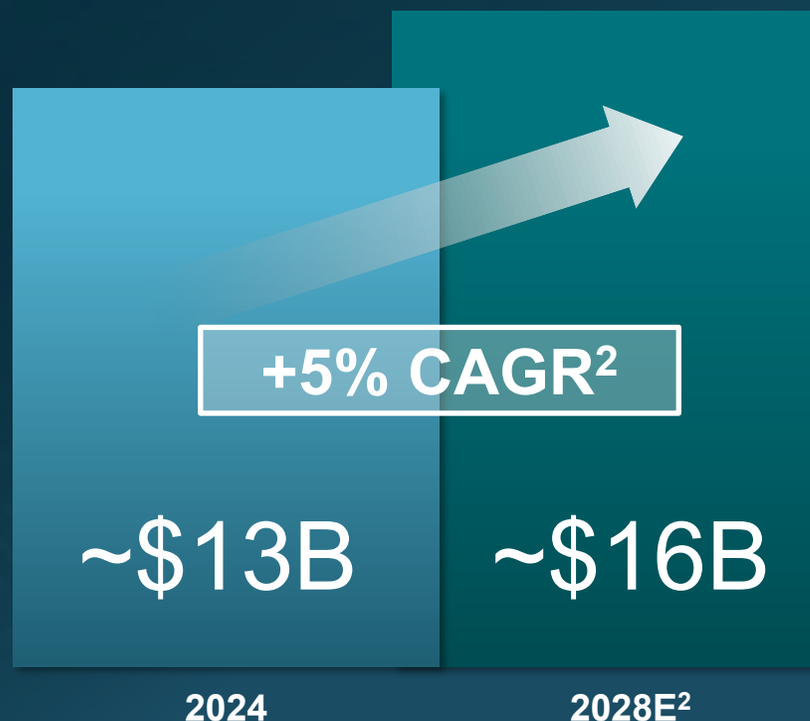
**KELLY SWAIM, MD**

SENIOR VICE PRESIDENT, SURGICAL R&D



# Strong Performance in a Durable & Growing Market

Global Surgical Market<sup>1</sup>



## Strategy to Drive *Growth* and *Margin Expansion*<sup>2</sup>

OWN THE OPERATING ROOM  
WITH “ONE-STOP SHOP”  
APPROACH

TRANSFORM PORTFOLIO WITH  
STEADY STREAM OF LAUNCHES  
IN PREMIUM CATEGORIES

RESHAPE MANUFACTURING AND  
SUPPLY NETWORK TO DRIVE **MARGIN  
EXPANSION** AND BEST-IN-CLASS SERVICE

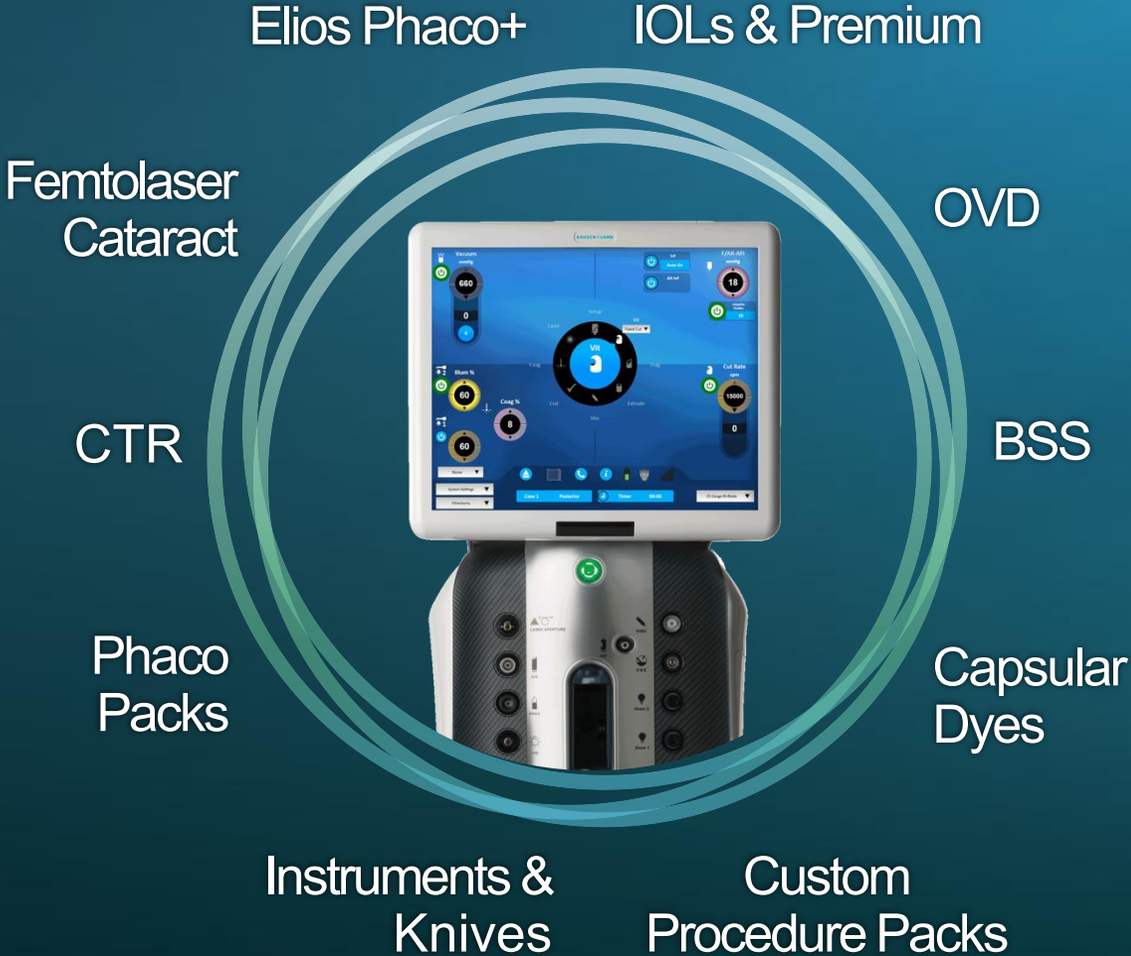
# Focused on Procedure Selling to Meet *All* Surgeon Needs

Own the Operating Room:  
*“One-Stop Shop”* Cataract Strategy

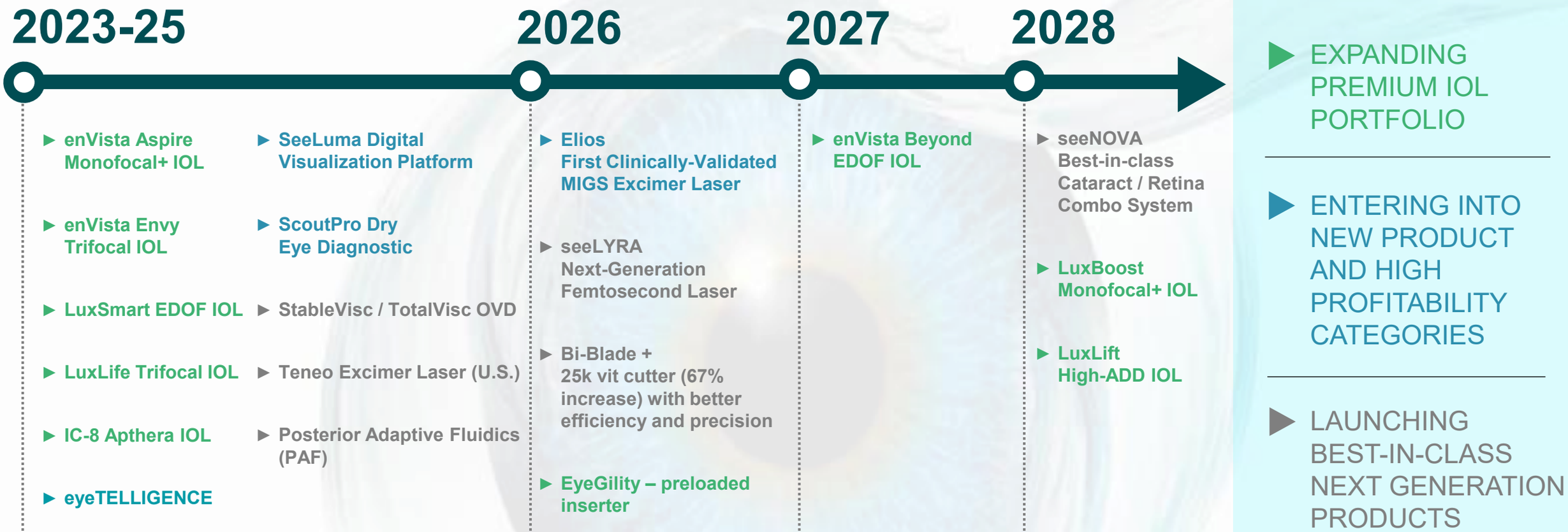
Cataract Surgery Devices<sup>1</sup>

Leading Cataract Equipment Companies	IOLs	OVD	Instruments	Knives	Procedure Packs	BSS	CTR	Fluid Mgmt.	Capsular Dyes
Bausch + Lomb	■	■	■	■	■	■	■	■	■
Alcon	■	■	■	■	■	■	■	■	■
J&J Vision	■	■		■	■	■	■		
Carl Zeiss	■	■	■				■		

1. Market Scope.



# Steady Stream of Launches in *Premium* Categories<sup>1</sup>



# *Expanding* Manufacturing Capabilities & *Optimizing* Surgical Supply Network<sup>1</sup>

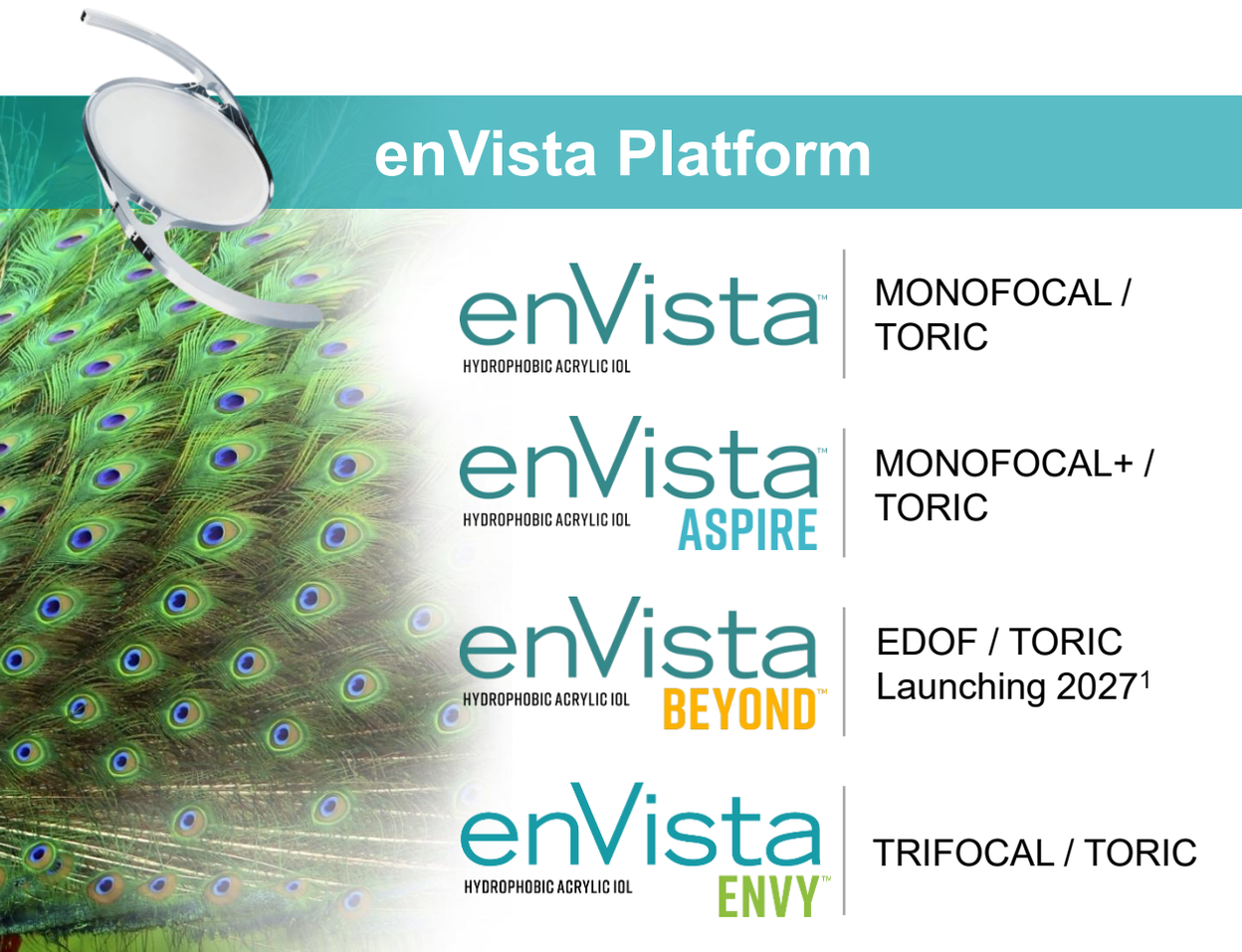
## Comprehensive Strategy to Drive Margin Expansion and Unlock Revenue Growth:

- ▶ Transition machine pack and custom pack assembly to low-cost supply base
- ▶ Vertical integration of custom pack production to reduce CMO costs
- ▶ Streamline production capacity across existing global supply network
- ▶ Optimizing partnerships of select products (viscoelastics, inserters, instruments)



# Building IOL Portfolio Across *All* Segments

Expanding in Premium IOLs – Market Projected to Grow +10% CAGR 2025-30<sup>1,2</sup>



## enVista Platform

<b>enVista™</b> HYDROPHOBIC ACRYLIC IOL	MONOFOCAL / TORIC
<b>enVista™</b> HYDROPHOBIC ACRYLIC IOL <b>ASPIRE</b>	MONOFOCAL+ / TORIC
<b>enVista™</b> HYDROPHOBIC ACRYLIC IOL <b>BEYOND</b>	EDOF / TORIC Launching 2027 <sup>1</sup>
<b>enVista™</b> HYDROPHOBIC ACRYLIC IOL <b>ENVY</b>	TRIFOCAL / TORIC



## Lux Platform

<b>LUXGOOD™</b>	MONOFOCAL / TORIC
<b>LUXBOOST™™</b>	MONOFOCAL+ / TORIC Launching 2028 <sup>1</sup>
<b>LUXSMART™</b>	EDOF / TORIC
<b>LUXlife™</b>	TRIFOCAL / TORIC
<b>LUXLIFT™</b>	EDOF High-Add / TORIC Launching 2028 <sup>1</sup>

# enVista Beyond

Completes Comprehensive IOL Portfolio

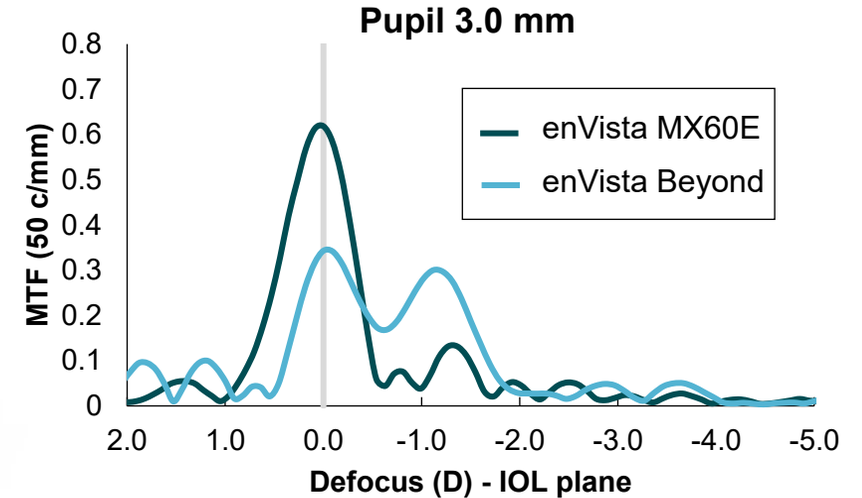
Extended Depth of Focus IOL

Pure Refractive Technology

Expected Launch in 2027<sup>1</sup>



## Designed to Build on the enVista Platform<sup>2</sup>



### Optical Bench Testing:

Visual Acuity

Greater DoF relative to enVista monofocal

Tilt & Decentration

Greater performance relative to Vivity, Symphony

Halos

Greater performance relative to Vivity, Symphony

# Unlocking Growth Opportunities in *MIGS*

~80M

People worldwide living with glaucoma<sup>1</sup>

~20%

of patients undergoing cataract surgery have POAG / OHT<sup>2</sup>

<50%

Cataract surgeons are performing MIGS today<sup>3</sup>

elios

The first clinically-validated excimer laser MIGS

Ten 210- $\mu$ m “microchannels” are created in the TM using a precision excimer laser

**Implant Free:** Laser-based technology does not require a surgical implant

**Lasting Results:** Sustained IOP reduction and medication burden

**Ease of Use:** Rapid learning curve with 2-4 cases to surgeon proficiency

**Safety Advantage:** Non-thermal laser ablation to protect surrounding tissue

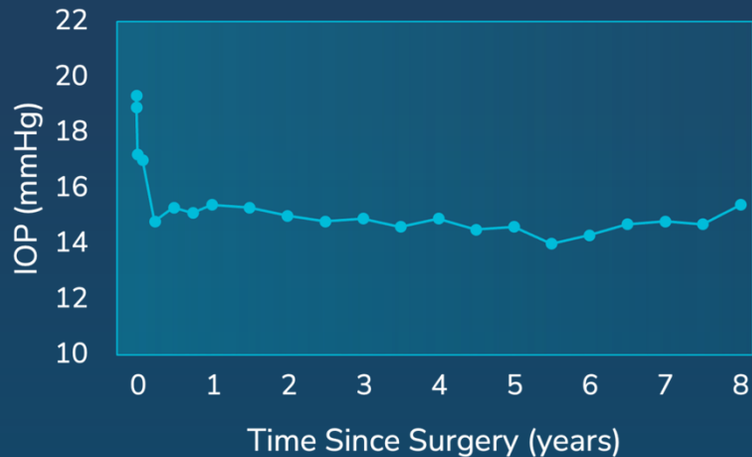


# Backed by *Extensive* Clinical Data

Elios is commercialized in Europe and clinically validated with 8 years of follow-up data

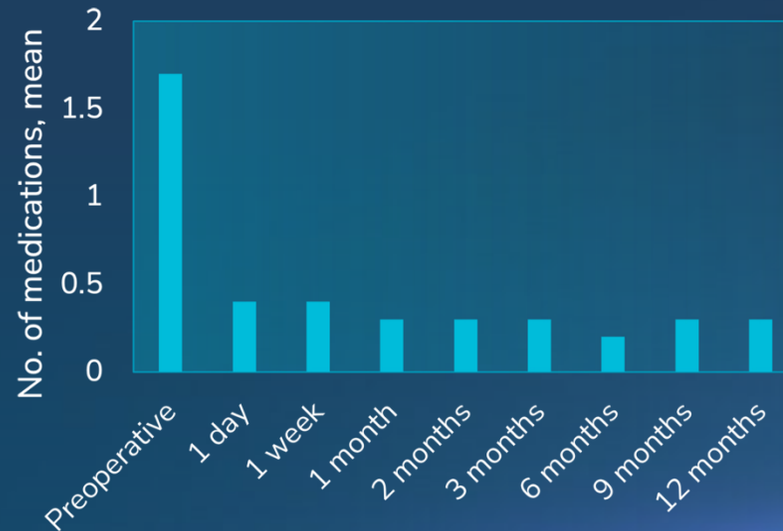
## Lowers IOP

IOP (>20%) for up to 8 years<sup>1</sup>



## Reduces Medication Burden

Up to 80% of patients are medication free one year after surgery<sup>2</sup>



## Sustained Results

Long-term micro channel patency<sup>3</sup>



31 Months Post-Op

A microscopic image showing a textured, brownish, cylindrical structure against a blue background. A white play button icon is centered over the structure. A dark blue semi-transparent box is on the right side of the image, containing white text.

# Elios Is the Next Generation of Glaucoma Technology

# Best-in-Class Cataract / Retina Combo System

**SEE)NOVA™**

**2028** Launch<sup>1</sup>



## DUAL-MODE ASPIRATION

Uncompromised post-occlusion chamber stability with customizable flow or vacuum control on demand

## CHALLENGE THE STATUS QUO

Embracing disruptive technologies and rethink phacoemulsification surgical paradigms with alternative lens removal technologies

## LASER-BASED ILLUMINATION

One million colors available, including four thousand shades of white, for optimal surgeon control and flexibility over contrast enhancement

# Delivering Improved Outcomes Over Current Microsurgical Systems



SEE NOVA™

# Next-Generation Femtosecond Laser



**SEE) LYRA™**

**2H26** Launch<sup>1</sup>

## **BUILDING ON 15 YEARS EXPERIENCE WITH VICTUS**

### **OPTIMIZED OUTCOMES**

Improved cutting for flaps under a curved PI, designed for future integration of phaco system → complete FLACS workstation

### **GREATER EFFICIENCY**

Designed for high volume cataract and refractive clinics: easy-to-use GUI, independency from a fixed patient bed, small footprint and mobility provides a simplified workflow in any OR environment

### **ENHANCED SAFETY**

Live optical coherence tomography (OCT) and camera

1. See Slide 2 for further information on forward-looking statements.

# Expanding in *Premium* Surgical Categories<sup>1</sup>



▶ **EDOF LENS TO COMPLETE PREMIUM IOL PORTFOLIO**

Launch Date: 2027  
enVista Platform  
Peak Sales: ~\$300M

elios



▶ **FIRST CLINICALLY VALIDATED IMPLANT FREE MIGS EXCIMER LASER**

Launch Date: 2H26 in U.S.  
Peak Sales: ~\$175M

see)NOVA



▶ **BEST-IN-CLASS CATARACT / RETINA COMBO SYSTEM**

Launch Date: 2028  
seeNOVA and Stellaris  
Peak Sales: ~\$450M

see)LYRA



▶ **NEXT GENERATION FEMTOSECOND LASER**

Launch Date: 2H26  
Peak Sales: ~\$50M

# 3 Key Takeaways<sup>1</sup>

1

Transforming  
Portfolio to a  
“One-Stop  
Shop”

2

Steady Stream  
of Launches  
in Premium  
Categories

3

Comprehensive  
Strategy to  
Drive Margin  
Expansion

# Physician Panel



**Moderator:**  
**Cathleen McCabe, MD**  
Strategic Medical Advisor,  
Bausch + Lomb

**Mark Schaeffer, OD, FAO**  
Optometrist, Ocular Disease  
& Contact Lenses,  
MyEyeDr

**Eric Donnenfeld, MD**  
Cornea, Laser Cataract  
& Refractive Surgeon,  
OCLI

**Lejla Vajzovic, MD, FASRS**  
Professor of Ophthalmology,  
Vitreoretinal Diseases & Surgery,  
Duke University

**William Trattler, MD**  
Cataract, Refractive, and Corneal  
Surgeon, Center for Excellence in  
Eye Care

**Q&A**

**FOCUS  
FORWARD**

# Executing Strategy to Drive Step-Change Performance

# Above-Market Revenue Growth & Meaningful Margin Expansion

# Elevating Standard of Care with Disruptive Innovation

**5-7%**

2025-2028 CC Revenue CAGR<sup>1,2,3</sup>

**ABOVE-MARKET REVENUE GROWTH**

**~23%**

2028 Adj. EBITDA Margin (ex. Acq. IPR&D)<sup>1,3</sup>

**~600 BPS EBITDA MARGIN EXPANSION 2025 to 2028**

**~\$7B**

Pipeline Potential Peak Sales<sup>1,4</sup>

**PIPELINE TO DRIVE TRANSFORMATIVE VALUE**

1. See Slide 2 for further information on forward-looking statements. The financial targets in this presentation are only effective as of the date given, November 13, 2025, and will not be updated or affirmed unless and until the Company publicly announces updated or affirmed financial targets, as the case may be. Distribution or reference of this deck following November 13, 2025, does not constitute the Company re-affirming financial targets.  
2. Constant currency. Compounded Annual Growth Rate.  
3. This is a non-GAAP measure or ratio. See Slide 2 and Appendix for further information on non-GAAP measures and ratios.  
4. Represents total projected peak sales of pipeline products, with anticipated peaks staggered based on launch dates.

# Appendix

# Key Modeling Assumptions (2025-2028)<sup>1</sup>

<b>Bausch + Lomb Revenue</b>	<ul style="list-style-type: none"> <li>• 5-7% FY25-28 CC CAGR<sup>2</sup></li> </ul>
Consumer Revenue	<ul style="list-style-type: none"> <li>• 5-7% FY25-28 CC CAGR<sup>2</sup></li> </ul>
Contact Lens Revenue	<ul style="list-style-type: none"> <li>• 5-7% FY25-28 CC CAGR<sup>2</sup></li> </ul>
Pharmaceuticals Revenue	<ul style="list-style-type: none"> <li>• 5-7% FY25-28 CC CAGR<sup>2</sup></li> </ul>
Surgical Revenue	<ul style="list-style-type: none"> <li>• 6-8% FY25-28 CC CAGR<sup>2</sup></li> </ul>
<b>Adj. R&amp;D (% of Revenue)<sup>2</sup></b>	<ul style="list-style-type: none"> <li>• Modest Increase to ~8% in FY28</li> </ul>
<b>Adj. EBITDA Margin (ex. Acq. IPR&amp;D)<sup>2</sup></b>	<ul style="list-style-type: none"> <li>• ~19% in FY26 Expanding to ~23% in FY28</li> </ul>
<b>Adj. Tax Rate<sup>2</sup></b>	<ul style="list-style-type: none"> <li>• ~19-20% FY26-28</li> </ul>
<b>Avg. Fully Diluted Share Count</b>	<ul style="list-style-type: none"> <li>• ~370M in FY28</li> </ul>
<b>Adj. EPS Attributable to B+L (ex. Acq. IPR&amp;D)<sup>2</sup></b>	<ul style="list-style-type: none"> <li>• Double Digit Growth FY26-28</li> </ul>
<b>Adj. Cash Flow From Operations to Adj. EBITDA Conversion<sup>2</sup></b>	<ul style="list-style-type: none"> <li>• ~50% Conversion in FY28</li> </ul>
<b>CapEx (% of Revenue)</b>	<ul style="list-style-type: none"> <li>• Mid-single digits FY26-28</li> </ul>
<b>Net Leverage<sup>2</sup></b>	<ul style="list-style-type: none"> <li>• ~3.5x Net Leverage by End of FY28</li> </ul>

1. See Slide 2 for further information on forward-looking statements.

2. This is a non-GAAP measure or ratio. See Slide 2 and Appendix for further information on non-GAAP measures and ratios.

# Reconciliation of Reported Revenue to Constant Currency Revenue<sup>1</sup> and Constant Currency Revenue Growth<sup>1</sup> (\$M)

	Calculation of Constant Currency Revenue for the Nine Months Ended			Change in		Change in		
	September 30, 2025		September 30, 2024	Reported Revenue		Constant Currency Revenue <sup>1</sup>		
	Revenue as Reported	Changes in Exchange Rates <sup>2</sup>	Constant Currency Revenue (Non-GAAP) <sup>1</sup>	Revenue as Reported	Amount	Pct.	Amount	Pct.
Contact Lens	766	(3)	763	720	46	6%	43	6%

# Non-GAAP Appendix

## Description of Non-GAAP Financial Measures

To supplement the financial measures prepared in accordance with U.S. generally accepted accounting principles (GAAP), the Company uses certain non-GAAP financial measures and ratios. These measures and ratios do not have any standardized meaning under GAAP and other companies may use similarly titled non-GAAP financial measures and ratios that are calculated differently from the way we calculate such measures and ratios. Accordingly, our non-GAAP financial measures and ratios may not be comparable to similar non-GAAP measures and ratios of other companies. We caution investors not to place undue reliance on such non-GAAP measures and ratios, but instead to consider them with the most directly comparable GAAP measures and ratios. Non-GAAP financial measures and ratios have limitations as analytical tools and should not be considered in isolation. They should be considered as a supplement to, not a substitute for, or superior to, the corresponding measures calculated in accordance with GAAP.

## EBITDA/Adjusted EBITDA/Adjusted EBITDA Margin/Adjusted EBITDA excluding Acquired IPR&D/Adjusted EBITDA Margin excluding Acquired IPR&D

EBITDA (non-GAAP) is Net income (loss) attributable to Bausch + Lomb Corporation (its most directly comparable U.S. GAAP financial measure) adjusted for interest, income taxes, depreciation and amortization. Adjusted EBITDA (non-GAAP) is EBITDA (non-GAAP) further adjusted for the items described below. Management believes that Adjusted EBITDA (non-GAAP), along with the GAAP measures used by management, most appropriately reflect how the Company measures the business internally and sets operational goals and incentives. In particular, the Company believes that Adjusted EBITDA (non-GAAP) focuses management on the Company's underlying operational results and business performance. As a result, the Company uses Adjusted EBITDA (non-GAAP) both to assess the actual financial performance of the Company and to forecast future results as part of its guidance. Management believes Adjusted EBITDA (non-GAAP) is a useful measure to evaluate current performance. Adjusted EBITDA (non-GAAP) is intended to show our unleveraged, pre-tax operating results and therefore reflects our financial performance based on operational factors. In addition, cash bonuses for the Company's executive officers and other key employees are based, in part, on the achievement of certain Adjusted EBITDA (non-GAAP) targets.

Adjusted EBITDA margin (non-GAAP) is Adjusted EBITDA (non-GAAP) divided by Revenues.

## Adjusted EBITDA (non-GAAP) Adjustments

**Adjusted EBITDA (non-GAAP) is net income (loss) attributable to the Company (its most directly comparable GAAP financial measure) adjusted for interest expense, net, (benefit from) provision for income taxes, depreciation and amortization and the following items:**

**Asset impairments:** The Company has excluded the impact of impairments of finite-lived and indefinite-lived intangible assets as such amounts are inconsistent in amount and frequency and are significantly impacted by the timing and/or size of acquisitions and divestitures. The Company believes that the adjustments of these items correlate with the sustainability of the Company's operating performance. Although the Company excludes impairments of intangible assets from measuring the performance of the Company and its business, the Company believes that it is important for investors to understand that intangible assets contribute to revenue generation.

**Restructuring, integration and transformation costs:** The Company has incurred restructuring costs as it implemented certain strategies, which involved, among other things, improvements to its infrastructure and operations, internal reorganizations and impacts from the divestiture of assets and businesses. With regard to infrastructure and operational improvements which the Company has taken to improve efficiencies in the businesses and facilities, these tend to be costs intended to right size the business or organization that fluctuate significantly between periods in amount, size and timing, depending on the improvement project, reorganization or transaction. Additionally, with the completion of the B+L IPO, as the Company prepares for post-Separation operations, the Company is launching certain transformation initiatives that will result in certain changes to and investment in its organizational structure and operations. These transformation initiatives arise outside of the ordinary course of continuing operations and, as is the case with the

Company's restructuring efforts, costs associated with these transformation initiatives are expected to fluctuate between periods in amount, size and timing. These out-of-the-ordinary-course charges include third party advisory costs, as well as certain compensation-related costs. Investors should understand that the outcome of these transformation initiatives may result in future restructuring actions and certain of these charges could recur. The Company believes that the adjustments of these items provide supplemental information with regard to the sustainability of the Company's operating performance, allow for a comparison of the financial results to historical operations and forward-looking guidance and, as a result, provide useful supplemental information to investors.

**Acquisition-related costs and adjustments excluding amortization of intangible assets:** The Company has excluded the impact of acquisition-related costs and fair value inventory step-up resulting from acquisitions as the amounts and frequency of such costs and adjustments are not consistent and are significantly impacted by the timing and size of its acquisitions. In addition, the Company excludes the impact of acquisition-related contingent consideration non-cash adjustments due to the inherent uncertainty and volatility associated with such amounts based on changes in assumptions with respect to fair value estimates, and the amount and frequency of such adjustments are not consistent and are significantly impacted by the timing and size of the Company's acquisitions, as well as the nature of the agreed-upon consideration.

**Share-based compensation:** The Company excludes costs relating to share-based compensation. The Company believes that the exclusion of share-based compensation expense assists investors in the comparisons of operating results to peer companies. Share-based compensation expense can vary significantly based on the timing, size and nature of awards granted.

# Non-GAAP Appendix

## Adjusted EBITDA (non-GAAP) Adjustments (continued)

**Separation costs and separation-related costs:** The Company has excluded certain costs incurred in connection with activities taken to: (i) separate the Bausch + Lomb business from the remainder of BHC and (ii) register the Bausch + Lomb business as an independent publicly traded entity. Separation costs are incremental costs directly related to effectuating the separation of the Bausch + Lomb business from the remainder of BHC and include, but are not limited to, legal, audit and advisory fees, talent acquisition costs and costs associated with establishing a new board of directors and audit committee. Separation-related costs are incremental costs indirectly related to the separation of the Bausch + Lomb business from the remainder of BHC and include, but are not limited to, IT infrastructure and software licensing costs, rebranding costs and costs associated with facility relocation and/or modification. As these costs arise from events outside of the ordinary course of continuing operations, the Company believes that the adjustments of these items provide supplemental information with regard to the sustainability of the Company's operating performance, allow for a comparison of the financial results to historical operations and forward-looking guidance and, as a result, provide useful supplemental information to investors.

**Gain (Loss) on extinguishment of debt:** The company has excluded gain (loss) on extinguishment of debt as this represents a loss from refinancing our existing debt and is not a reflection of our operations for the period. Further, the amount and frequency of such amounts are not consistent and are significantly impacted by the timing and size of debt financing transactions and other factors in the debt market that are not in management's control. Bausch + Lomb did not have any material losses on extinguishment of debt prior to the second quarter of 2025.

**Other Non-GAAP adjustments:** The Company also excludes certain other amounts, including IT infrastructure investment, litigation and other matters, gain/(loss) on sales of assets and certain other amounts that are the result of other, non-comparable events to measure operating performance if and when present in the periods presented. These events arise outside of the ordinary course of continuing operations. Given the unique nature of the matters relating to these costs, the Company believes these items are not routine operating expenses. For example, legal settlements and judgments vary significantly, in their nature, size and frequency, and, due to this volatility, the Company believes the costs associated with legal settlements and judgments are not routine operating expenses. The Company excluded these costs as this event is outside of the ordinary course of continuing operations and is infrequent in nature. The Company believes that the exclusion of such out-of-the-ordinary-course amounts provides supplemental information to assist in the comparison of the financial results of the Company from period to period and, therefore, provides useful supplemental information to investors. However, investors should understand that many of these costs could recur and that companies in our industry often face litigation.

Adjusted EBITDA excluding Acquired In-Process Research and Development (IPR&D) is Adjusted EBITDA (non-GAAP) further adjusted to exclude Acquired IPR&D. Adjusted EBITDA Margin excluding Acquired In-Process Research and Development (IPR&D) is Adjusted EBITDA (non-GAAP) further adjusted to exclude Acquired IPR&D divided by Revenues. The IPR&D expenditures represent costs directly resulting from business development transactions and not through the normal course of business. The Company believes that the exclusion of such out-of-the-ordinary-course amounts provides supplemental information to assist in the comparison of the financial results of the Company from period to period and, therefore, provides useful supplemental information to investors in assessing our performance. However, investors should understand that the Company may enter into additional business development transactions in the future and, as a result, such acquired IPR&D may recur in the future.

## Constant Currency

Constant currency change or constant currency growth is calculated by adjusting or further adjusting a measure or ratio by changes in or impact of foreign currency exchange rates. Constant currency impact is determined by comparing 2025 amounts adjusted to exclude currency impact, calculated using 2024 monthly average exchange rates, to the actual 2024 reported amounts. Constant currency revenue is GAAP revenue (its most directly comparable GAAP financial measure) adjusted for changes in foreign currency exchange rates. The Company uses Constant Currency Revenues (non-GAAP) and Constant Currency Revenue Growth (non-GAAP) to assess performance of its reportable segments, and the

Company in total, without the impact of foreign currency exchange fluctuations. The Company believes that such measures are useful to investors as they provide a supplemental period-to-period comparison. Although changes in foreign currency exchange rates are part of our business, they are not within management's control. Changes in foreign currency exchange rates, however, can mask positive or negative trends in the underlying business performance.

## Adjusted Cash Flow from Operations/Adjusted Cash Flow from Operations to Adj. EBITDA (excl. Acq. IPR&D) conversion

Adjusted cash flow from operations (non-GAAP) is Cash flow from operations/Cash used in operations (loss) attributable to Bausch + Lomb Corporation (its most directly comparable GAAP financial measure) adjusted for: (i) payments of legacy legal settlements, net of insurance proceeds, if any (ii) payments for separation costs, IPO costs, separation-related costs, and IPO-related costs (iii) payments for business transformation costs and (iv) payments for financing fees related to the modification of debt, if any. Management believes that Adjusted cash flow from operations (non-GAAP), along with the GAAP and non-GAAP measures used by management, most appropriately reflect how the Company measures the business internally. The Company uses Adjusted cash flow from operations (non-GAAP) both to assess the actual financial performance of the Company and to forecast future results as part of its guidance. Management believes Adjusted cash flow from operations (non-GAAP) is a useful measure to evaluate current performance amounts. As these payments arise from events outside of the ordinary course of continuing operations as discussed above, the Company believes that the adjustments of these items provide supplemental information with regard to the sustainability of the Company's cash from operations, allow for a comparison of the financial results to historical operations and forward-looking guidance and, as a result, provide useful supplemental information to investors. Adjusted cash flow from operations to Adj. EBITDA (excl. Acq. IPR&D) conversion is Adjusted cash flow from operations divided by Adjusted EBITDA (excluding Acquired IPR&D).

## Net Leverage

Net Leverage is the ratio of net debt (which is calculated as total debt (its GAAP equivalent) less cash and cash equivalents) over Adjusted EBITDA (excluding Acquired IPR&D). Management believes that net leverage is an important measure of our overall liquidity position and an indicator of our ability to meet financial obligations.

## Adjusted Tax Rate

Adjusted Tax Rate (the most directly comparable financial measure for which is our GAAP tax rate) includes the tax impact of the various non-GAAP adjustments used in calculating our non-GAAP measures. However, due to the differences in the tax treatment of items excluded from non-GAAP earnings, our adjusted tax rate will differ from our GAAP tax rate and from our actual tax liabilities.

## Adjusted Earnings Per Share (EPS)/Adjusted EPS excluding Acquired IPR&D

Adjusted earnings per share or Adjusted EPS (non-GAAP) is calculated as Diluted income per share attributable to Bausch + Lomb Corporation ("GAAP EPS") (its most directly comparable GAAP financial measure), adjusted for the per diluted share impact of each adjustment made to reconcile Net income (Loss) attributed to Bausch + Lomb Corporation to Adjusted net income (non-GAAP) as discussed above. Adjusted EPS excluding Acquired IPR&D (non-GAAP) is Adjusted EPS (non-GAAP) further adjusted for the per diluted share impact of Acquired IPR&D. Like Adjusted net income (non-GAAP), Adjusted EPS (non-GAAP) and Adjusted EPS excluding Acquired IPR&D excludes the impact of certain items that may obscure trends in the Company's underlying performance on a per share basis. By disclosing these non-GAAP measures, it is management's intention to provide investors with a meaningful, supplemental comparison of the Company's results and trends for the periods presented on a diluted share basis. Accordingly, the Company believes that Adjusted EPS (non-GAAP) and Adjusted EPS excluding Acquired IPR&D (non-GAAP) are useful to investors in their assessment of the Company's operating performance, the valuation of the Company and an investor's return on investment. It is also noted that, for the periods presented, our GAAP EPS was significantly lower than our Adjusted EPS (non-GAAP) and Adjusted EPS less Acquired IPR&D (non-GAAP).

## Adjusted R&D Expense

Adjusted R&D expenses (non-GAAP) represents research and development expenses ("R&D expenses") (its most directly comparable GAAP financial measure), adjusted to exclude certain separation-related costs. See the discussion under "Other NonGAAP adjustments" above. Management uses Adjusted R&D (non-GAAP), along with GAAP measures, as a supplemental measure for period-to-period comparison to understand and evaluate each segment's ability to control costs. The Company believes that Adjusted R&D (non-GAAP) is useful to investors as it provides consistency and comparability with our past financial performance and facilitates period-to-period comparisons of our R&D expenses, as this measure eliminates the effects of separation-related costs, which given their nature and frequency, are outside the ordinary course and relate to unique circumstances.