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Late clinical-stage biopharmaceutical company working to develop the first FDA-approved ophthalmic formulation of bevacizumab-vikg for use in retinal indications.



Investment Highlights

Advancing on Multiple Fronts Towards Potential FDA Approval



Potential FDA approval in wet AMD in 2022 with lead product candidate ONS-5010 / LYTENAVA[™] (bevacizumab-vikg)¹, an investigational ophthalmic formulation of bevacizumab-vikg, targeting \$13.1 billion global anti-VEGF market²

Phase 3 Clinical Program

- Demonstrated safety and efficacy in clinical experience trial
- Ongoing Phase 3 pivotal trial with topline data expected mid-2021

Manufacturing and Regulatory

- Partnered with Fujifilm and Ajinomoto as best-in-class cGMP global manufacturers
- Tentatively granted ATC code for ophthalmic bevacizumab by the World Health Organization

Commercial Planning Activities Underway

- Outreach to physicians, patients, KOLs and payors
- Market research indicates ONS-5010, if approved, will be a significant therapy in anti-VEGF market

Global Strategic Partnership

- Discussions with potential strategic partners progressing
- Signing of definitive agreement could be as soon as the end of 2020



Leadership Team: Global Ophthalmic Development and Commercial Launch Excellence



LAWRENCE KENYON President, CEO, CFO









JEFF EVANSON Chief Commercial Officer











TERRY DAGNON Chief Operating Officer









RANDY THURMAN Executive Chairman of the Board



MARK HUMAYUN, MD, PhD Medical Advisor





ONS-5010

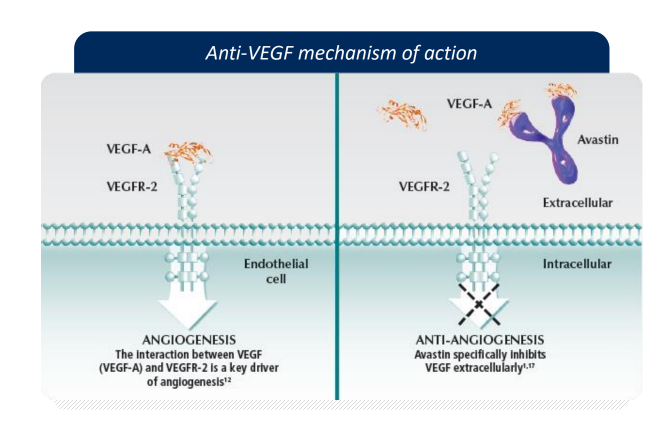
Addresses Significant Unmet Medical Need in a \$13.1 Billion Global Anti-VEGF Market



Standard of Care in Wet AMD

ONS-5010 / LYTENAVA™, if approved, will be the first on-label ophthalmic formulation of bevacizumab-vikg

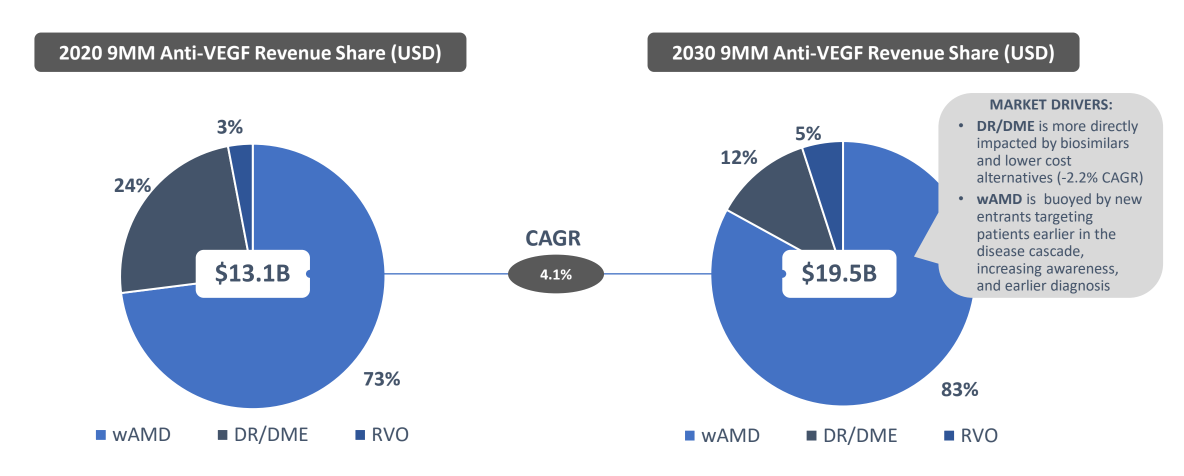
- Anti-VEGF drugs have been standard of care since 2006
 - Block growth of abnormal blood vessels and leakage of fluid from the vessels behind the retina
- Several new clinical-stage anti-VEGF drugs, including biosimilars, in development and/or recently approved
 - Require significant time and capital to achieve commercialization
 - New drugs expected to price at or near the high price points of current approved therapies





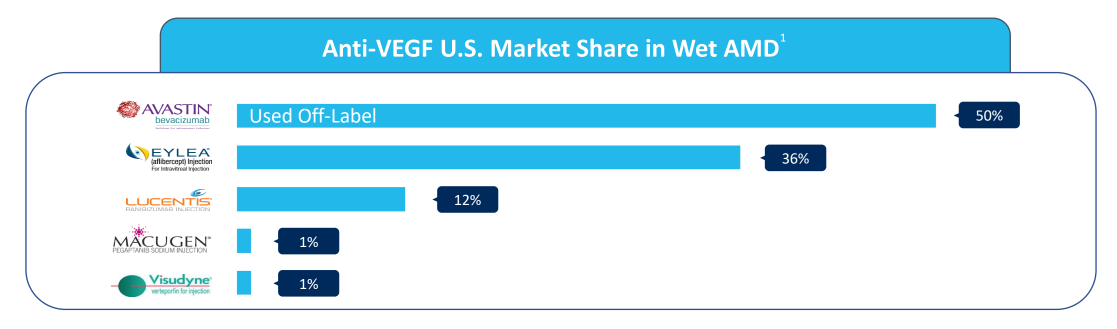
Targeting Large and Growing Ophthalmic Markets

ONS-5010, if approved, will be a significant therapy in the retinal anti-VEGF market, currently estimated to be in excess of \$13.1 billion worldwide





Unapproved Bevacizumab Represents 50% of U.S. Wet AMD Market



Expected Drivers to Compete Across All Anti-VEGF Therapeutics

- 1 Provide safe and cost-effective on-label bevacizumab
- Become first-line "step-edit" drug of choice

- 3 12 years market exclusivity under new BLA
- 4 Penetrate EU and developing markets



ONS-5010

Potential to be the first ophthalmic formulation of bevacizumab-vikg approved as an anti-VEGF therapy addressing vision loss from wet agerelated macular degeneration (wet AMD)



Unapproved Repackaged IV Bevacizumab Presents Safety Issues

Once approved, ONS-5010 will reduce the need for use of unapproved repackaged IV Avastin® from compounding pharmacists

Variability in Potency¹

- 81% of samples had lower protein concentrations than required
- Samples had statistically significant variations in protein concentration among samples

JAMA Ophthalmology

Safety and Sterility Adverse Events²

- Unvalidated hold times in syringes not designed to be primary packages
- Patients have lost eyesight due to infections
- Multiple unapproved repackaged IV bevacizumab recalls due to unsterile compounding practices



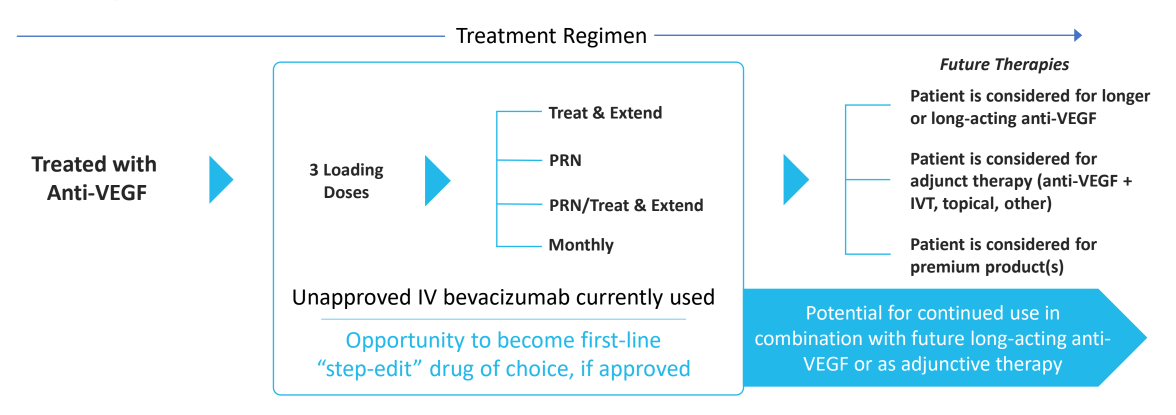
Syringe Malfunctioning³

- Variability in repackaging can lower quality of syringe products, resulting in adverse events
- Silicone oil droplets may be released by the syringe into the eye





ONS-5010: If Approved, Potential First Access in Treatment Paradigm with Step-Edit Therapy



Step-Edit is a Payor Cost Saving Measure

 Less expensive therapies are covered first

 Patient must "fail" medication before advancing to more costly treatments



Clinical Progress Drives ONS-5010 Towards U.S. and EU Filings in 2021

Recently completed clinical experience trial provides a high level of confidence in the outcome of the ongoing fully-enrolled pivotal trial



Ongoing U.S.-based Phase 3 pivotal trial

- Completed enrollment of 227 patients
- Pivotal data are expected mid-2021



Demonstrated safety and efficacy

 Recently reported data from clinical experience trial

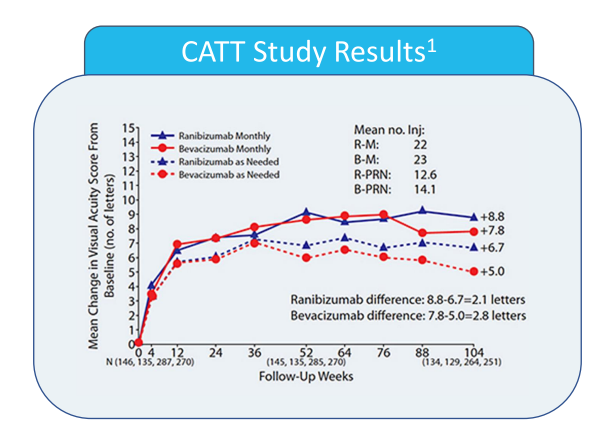


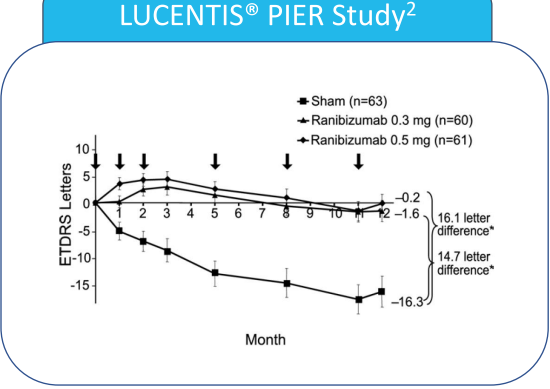
Regulatory strategy aligned with FDA

Pursuing new Biologics
 License Application (BLA)
 submission in wet AMD



Bevacizumab Demonstrated to be Equivalent to LUCENTIS® in CATT Trial







Completed Clinical Experience Trial

1st Registration Trial







Provides high level of confidence in the outcome of the ongoing fully-enrolled pivotal trial

Demonstrated anticipated safety and efficacy signals consistent with previously published results for ophthalmic bevacizumab

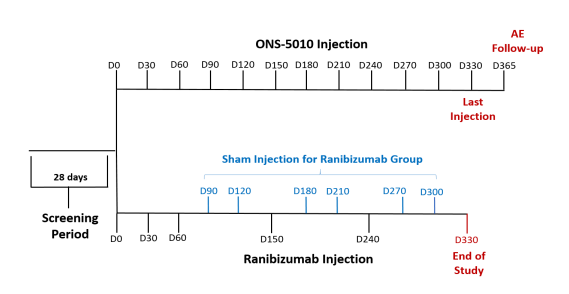
Trial Design Highlights:

- Randomized Masked Controlled Trial
- ONS-5010 vs LUCENTIS® (ranibizumab)
- 61 subjects enrolled
- Trial conducted in Australia
- Expected to support planned new U.S. BLA filing in 2021



ONS-5010 Demonstrated Safety and Efficacy in Clinical Experience Trial

Title: A clinical effectiveness, multicenter, randomized, double-masked, controlled trial of the efficacy and safety of ONS-5010 in subjects with subfoveal choroidal neovascularization (CNV) secondary to age-related macular degeneration



Trial Design:

- 30 treatment-naïve or previously treated wet AMD patients per arm
- Baseline visual acuity 20/40 to 20/320
- ONS-5010 dosed monthly vs ranibizumab dosed 3 initial monthly injections, followed by quarterly dosing
- Efficacy read-out at the Month 11 visit

Proof-of-Concept Achieved

- Desired proportion of 3-line visual acuity gainers achieved
- Desired mean gain in visual acuity achieved
- Zero ocular inflammation observed
- In this trial, safety was comparable to published bevacizumab studies, such as CATT



Positive Results From Clinical Experience Trial

- ONS-5010 demonstrated anticipated safety and efficacy signals consistent with previously published ophthalmic bevacizumab research
- No significant statistical differences in efficacy and safety
- Results provide support for the established design and protocol for ongoing U.S.-based Phase 3 pivotal trial
- No ocular adverse events of intraocular inflammation, vasculitis or retinal artery occlusion such as those recently reported for other anti-VEGFs in treating retinal diseases

| Trial Enrollment | | ONS-5010 (N=31) | Ranibizumab (N=30) | Overall (N=61) |
|---------------------------------|-----|--------------------|-----------------------|-------------------|
| Prior Anti-VEGF Treatment | Yes | 25 (80.6%) | 15 (50.0%) | 40 (65.6%) |
| | No | 6 (19.4%) | 15 (50.0%) | 21 (34.4%) |

| Subgroup Analysis of Treatment- Naïve Subjects | ONS-5010 | Ranibizumab |
|---|--------------|------------------|
| Subjects achieving > 15 letters BCVA at Month 11 | 2/6 (33%) | 4/14 (28.6%) |

| Subgroup Analysis | ONS-5010 | Ranibizumab |
|--|--------------|---------------|
| Proportion of treatment-naïve Subjects with baseline visual acuity of <67 Letters (20/50 or worse) | 2/4 (50%) | 4/10 (40%) |

- ONS-5010 ITT 3-line Visual Acuity Gainers Subgroup Summary
 - Treatment-naïve ONS-5010: **2/6 33.3**% (historical **CATT 31**% bevacizumab monthly historical PIER **13.1**% ranibizumab quarterly historical **EXCITE 14.2**% ranibizumab quarterly)
 - Treatment-naïve & 20/50 or worse ONS-5010: **2/4 50%** (historical **CATT 31% bevacizumab monthly** historical **PIER 13.1% ranibizumab quarterly** historical **EXCITE 14.2% ranibizumab quarterly**)
- ONS-5010 ITT BCVA Subgroup Summary
 - Treatment-naïve +7.3 letters (historical CATT +8.0)
 - Treatment-naïve & 20/50 or worse +8.3 letters (historical CATT +8.0)



Ongoing Pivotal Trial

2nd Registration Trial



Enrollment completed

Topline data expected mid-2021

Trial Highlights:

- Randomized Masked Controlled Trial
- ONS-5010 vs LUCENTIS® (ranibizumab)
- 227 patients enrolled
- Trial conducted in the United States
- Both trial arms include predominantly treatment-naïve patients with baseline VA less than 20/50 at trial start
- Safety & efficacy data expected to support planned new U.S. BLA filing in 2021



Ongoing Pivotal Trial Design Informed by Clinical Experience Trial – With Larger Sample Size



Randomized masked controlled trial with 227 subjects



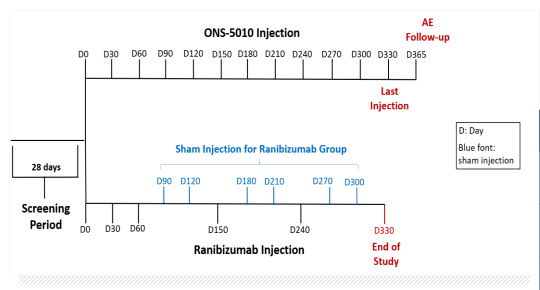
ONS-5010 administered monthly X 12



LUCENTIS dosing arm (PIER dosing) – Three initial monthly injections followed by fixed quarterly dosing



Primary endpoint difference in proportion of subjects gaining 15 letters of BCVA at Day 330





| Comparison of trial Parameters | Clinical Experience Trial | Pivotal Trial | Rationale for Change from Clinical Experience Trial to Pivotal Trial Parameters |
|-----------------------------------|---|--|---|
| Prior Treatment | Both treatment-naïve and previously treated | Treatment-naïve, only | Treatment-naïve subjects have more active disease (leakage on fluorescein angiography) and worse vision; more room to improve |
| Baseline Visual Acuity | 20/40 to 20/320 BCVA (73 to 25 letters) | 20/50 to 20/320 BCVA (67 to 25 letters) | Better baseline VA (20/40 or better) is associated with less gain in VA and a lower proportion gaining ≥3-lines compared to worse VA (20/50 or worse) |
| Planned Sample Size | 25 per am | 110 per arm | To support 90% power to detect a difference between arms in the proportion of responders |



Ongoing Open-Label Safety Study Supports BLA Requirements







Full enrollment of 195 subjects achieved in less than one month, significantly ahead of planned four-month schedule

Data from study to be included in complete data package to support BLA for wet AMD

Trial Highlights:

- Open-label safety study
- Enrolled 195 subjects with wet age-related macular degeneration (wet AMD), diabetic macular edema (DME) or branch retinal vein occlusion (BRVO)
- Subjects to receive three doses of ONS-5010 over a threemonth period
- Conducted to ensure adequate number of safety exposures to ONS-5010 / LYTENAVA™ (bevacizumab-vikg)



Commercial Planning Activities Underway



With an enhanced safety and cost-effectiveness profile, Outlook Therapeutics expects ONS-5010 to be widely adopted by payors and clinicians worldwide and to become the first-line drug of choice for payor-mandated "step edit" in the United States for retinal indications



Physician and Patient Outreach



Aligning Key Opinion Leaders



Payor Community Engagement

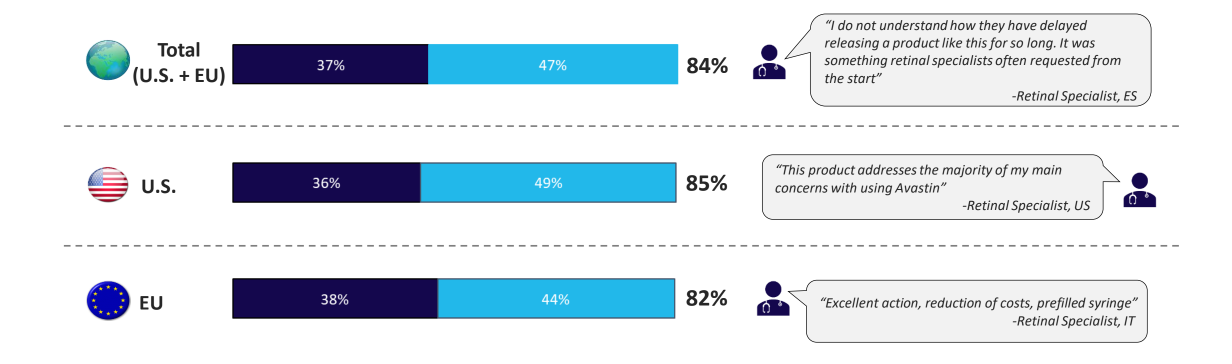
Discussions with Potential Strategic Partners Progressing

- Engaged with several life sciences companies that could result in a strategic partnership and definitive agreement for ONS-5010 as soon as the end of 2020
- Established joint venture with Syntone Technologies for commercializing ONS-5010 in Greater China



Physicians Want Approved Bevacizumab

>80% of retinal specialists express interest in an FDA-approved ophthalmic bevacizumab to treat wet AMD, DME and BRVO





Manufacturing and Regulatory Progress Towards Commercialization







Manufacturing

Best-in-class cGMP manufacturing partners



Pre-Filled Syringes

Supply agreement for a best-inclass pre-filled ophthalmic syringe



Regulatory

Tentatively granted ATC code for ophthalmic bevacizumab





• Lead product candidate ONS-5010 / LYTENAVATM has potential to be first FDA-approved ophthalmic formulation of bevacizumab for use in multiple retinal indications

Potential FDA Approval in 2022

Targeting \$13.1 Billion Global Anti-VEGF Market¹

Potential for 12 Years of Market Exclusivity

Management Team with Extensive Clinical/Regulatory
 Ophthalmology & Drug Development Experience