UNITED STATES SECURITIES AND EXCHANGE COMMISSION

Washington, D.C. 20549

FORM 8-K

CURRENT REPORT
Pursuant to Section 13 OR 15(d)
of The Securities Exchange Act of 1934

Date of Report (Date of earliest event reported): January 22, 2024

NewAmsterdam Pharma Company N.V.

(Exact name of registrant as specified in its charter)

The Netherlands (State or other jurisdiction of incorporation) 001-41562 (Commission File Number) N/A (I.R.S. Employer

Gooimeer 2-35 Naarden The Netherlands (Address of principal executive offices)

1411 DC (Zip Code)

+31 (0) 35 206 2971
(Registrant's telephone number, including area code)

Not Applicable (Former name or former address, if changed since last report)

Check the appropriate box below if the Form 8-K is intended to simultaneously satisfy the filing obligation of the registrant under any of the following provisions:

- ☐ Written communication pursuant to Rule 425 under the Securities Act (17 CFR 230.425)
- ☐ Soliciting material pursuant to Rule 14a-12 under the Exchange Act (17 CFR 240.14a-12)
- ☐ Pre-commencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))
- ☐ Pre-commencements communications pursuant to Rule 13e-4(c) under the Exchange Act (17 CFR 240.13e-4(c))

Securities registered pursuant to Section 12(b) of the Act:

Title of each class	Trading Symbols	Name of each exchange on which registered
Ordinary Shares, nominal value €0.12 per share	NAMS	The Nasdaq Stock Market LLC
Warrants to purchase Ordinary Shares	NAMSW	The Nasdaq Stock Market LLC

- Indicate by check mark whether the registrant is an emerging growth company as defined in Rule 405 of the Securities Act of 1933 (§230.405 of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (§240.12b-2 of this chapter).
- If an emerging growth company, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any new or revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act.

tem 7.01 Regulation FD Disclosure.

On January 22, 2024, NewAmsterdam Pharma Company N.V. (the "Company") posted an updated corporate investor presentation on its website (https://www.newamsterdampharma.com/). A copy of the corporate investor presentation is furnished as Exhibit 99.1 to this Current Report on Form 8-K. The information contained on, or that can be accessed from, the Company's website is not incorporated into, and does not constitute a part of, this Current Report on Form 8-K.

The information contained in this Item 7.01, including Exhibit 99.1, is being "furnished" and shall not be deemed "filed" for purposes of Section 18 of the Securities Exchange Act of 1934, as amended (the "Exchange Act"), or otherwise subject to the liability of that Section or Sections 11 and 12(a)(2) of the Securities Act of 1933, as amended (the "Securities Act"). The information contained in this Item 7.01, including Exhibit 99.1, shall not be incorporated by reference into any registration statement or other document pursuant to the Securities Act or into any filing or other document pursuant to the Exchange Act, except as otherwise expressly stated in any such filing.

Item 9.01 Financial Statements and Exhibits.

(d) Exhibits.

EXHIBIT NUMBER

EXHIBIT DESCRIPTION

99.1 NewAmsterdam Pharma Company N.V. Corporate Presentation

104 Cover Page Interactive Data File (embedded within the Inline XBRL document)

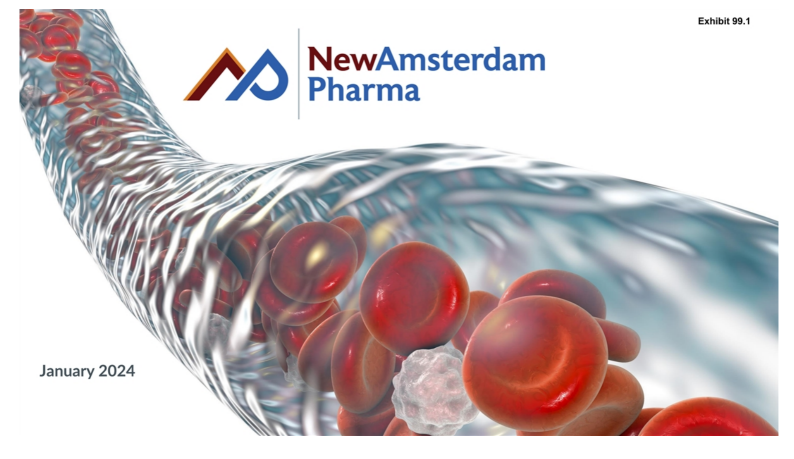
SIGNATURE

Pursuant to the requirements of the Securities Exchange Act of 1934, the registrant has duly caused this report to be signed on its behalf by the undersigned hereunto duly authorized.

NewAmsterdam Pharma Company N.V.

By: /s/ Michael Davidson Michael Davidson Chief Executive Officer

Dated: January 22, 2024





This presentation (together with oral statements made in connection herewith, this "Presentation") is for informational purposes only. This Presentation shall not constitute an offer to sell, or the solicitation of an offer to buy, any securities, nor shall there be any sale of securities in any states or jurisdictions in which such offer, solicitation or sale would be unlawful.

Forward Looking Statements

Certain statements included in this Presentation that are not historical facts are forward-looking statements for purposes of the safe harbor provisions under the United States Private Securities Litigation Reform Act of 1995. Forward looking statements generally are accompanied by words such as "believe," "may," "will," "estimate," "continue," "intend," "expect," "should," "would," "plan," "predict," "potential," "seem," "seek," "future," "outlook" and simila expressions that predict or indicate future events or trends or that are not statements include, but are not limited to, statements by NewAmsterdam Pharma Company N.V.
("NewAmsterdam" or the "Company") regarding estimates and forecasts of other financial and performance metrics and projections of market opportunity; expectations and timing related to the success, cost and timing of product development activities, including timing of initiation, completion and data readouts for clinical trials and the potential approval of the Company's product candidate; the timing for enrolling patients; the timing and forums for announcing the state of the Company's product candidate; the timing for enrolling patients; the timing and forums for announcing the state of the Company's product candidate; the timing for enrolling patients; the timing and forums for announcing the state of the Company's product candidate; the timing for enrolling patients; the state of the Company's product candidate; the timing and forums for announcing the state of the Company's product candidate; the timing for enrolling patients; the state of the Company's product candidate; the timing for enrolling patients; the state of the Company's product candidate; the timing for enrolling patients; the state of the Company's product candidate; the timing for enrolling patients; the state of the Company's product candidate; the state of the Company's product candi data; the size and growth potential of the markets for the Company's product candidate; the therapeutic and curative potential of the Company's product candidate; financing and other business milestones; the Company's expected acash runway; and the Company's plans for commercialization. These statements are based on various assumptions, whether or not identified in this Presentation, and on the current expectations of the Company's management and are not predictions of actual performance. These forward-looking statements are provided for illustrative purposes only and are not intended to serve as and must not be relied on as a guarantee, an assurance, a prediction, or a definitive statement of fact or probability. Actual events and circumstances are difficult or impossible to predict and may differ from assumptions. Many actual events and circumstances are beyond the control of the Company. These forwardlooking statements are subject to a number of risks and uncertainties, including changes in domestic and foreign business, market, financial, political, and legal conditions; risks related to the approval of NewAmsterdam's product candidate and the timing of expected regulatory and business milestones; whether topline, initial or preliminary results from a particular clinical trial will be predictive of the final results of that trial and whether results of early clinical trials will be indicative of the results of later clinical trials; ability to negotiate definitive contractual arrangements with potential customers; the impact of competitive product candidates; ability to obtain sufficient supply of materials; global economic and political conditions, including the Russia-Ukraine conflict, and the war in Israel; the effects of competition on NewAmsterdam's future business; and those factors discussed in documents filed by the Company with the SEC. Additional risks related to NewAmsterdam's business include, but are not limited to: uncertainty regarding outcomes of the company's ongoing clinical trials, particularly as they relate to regulatory review and potential approval for its product candidate; risks associated with the Company's efforts to commercialize a product candidate; the Company's ability to negotiate and enter into definitive agreements on favorable terms, if at all; the impact of competing product candidates on the Company's ability to continue to source the raw materials for its product candidate, together with the risks described in the Company's filings made with the U.S. Securities and Exchange Commission from time to time.

If any of these risks materialize or NewAmsterdam's assumptions prove incorrect, actual results could differ materially from the results implied by these forward-looking statements. There may be additional risks that are presently unknown by the Company or that NewAmsterdam currently believes are immaterial that could also cause actual results to differ from those contained in the forward-looking statements. In addition, forward-looking statements reflect NewAmsterdam's expectations, plans, or forecasts of future events and views as of the date of this Presentation and are qualified in their entirety by reference to the cautionary statements herein. NewAmsterdam anticipates that subsequent events and developments will cause the Company's assessments to change. These forward-looking statements should not be relied upon as representing NewAmsterdam's assessments as of any date subsequent to the date of this Presentation. Accordingly, undue reliance should not be placed upon the forward-looking statements. Neither NewAmsterdam nor any of its affiliates undertakes any obligation to update these forward-looking statements, except as required by law.

Market Data

Certain information contained in this Presentation relates to or is based on third-party studies, publications, surveys and NewAmsterdam's own internal estimates and research. In addition, all of the market data included in this Presentation involves a number of assumptions and limitations, and there can be no guarantee as to the accuracy or reliability of such assumptions. Finally, while NewAmsterdam believes its internal research is reliable, such research has not been verified by any independent source and NewAmsterdam cannot guarantee and makes no representation or warranty, express or implied, as to its accuracy and completeness.

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NewAmsterdam
Pharma





Obicetrapib in multiple Phase 3 trials for hypercholesterolemia – Key valuedriving data expected in 2024

Significant unmet need for oral LDL-lowering therapy as adjunct to statins:

- 35mm+ patients in US/EU5 are not achieving LDL-lowering goals despite standard-of-care
- \$3-4B+ global market opportunity

Simple, oral, once-daily, low dose CETP inhibitor with strong LDL-lowering observed through five Phase 2 trials:

- 43% mean LDL-lowering as monotherapy, 59% mean in combination with ezetimibe, observed on top of highintensity statins
- Tolerability data in >800 pts, with blinded data in >10,000 pts
- Robust effects on ApoB, non-HDL-C, HDL-C and Lp(a)

Convenient oral format potentially enables broad market access to address unmet need

Cash at year end 2023: ~\$340 million(1)

Multiple pivotal data readouts expected from 2024-2026

- 1Q 2024: Complete Phase 3 enrollment for PREVAIL
- 1Q 2024: Initiate Phase 3 fixed-dose combination ("FDC") trial

Anticipated Phase 3 data readouts:

- 3Q 2024: BROOKLYN
- 4Q 2024: BROADWAY
- 1Q 2025: TANDEM Fixed-Dose Combination
- 2026: PREVAIL CVOT

Additional pipeline expansion potential in Alzheimer's disease and diabetes

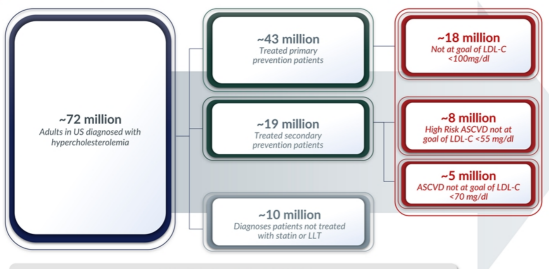
Upcoming catalysts build on 2023 progress:

Enrollment complete in BROOKLYN & BROADWAY

Positive data in ROSE2, Phase 2b Trial in Japanese Patients Initial data from Phase 2a Trial in Early Alzheimer's



Obicetrapib designed to address the ~30M patients in US on drug but not at goal



Of the ~30M treated patients not at goal, ~18M were "far from goal" (greater than 20%) and 6M were not taking statins

US Branded Lipid Lowering Market

Potential key factors limiting penetration include product limitations and market access hurdles:

Low prescriber enthusiasm for existing TPPs

Payors restrict access

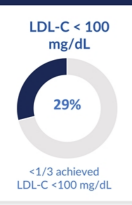


ASCVD=atherosclerotic cardiovascular disease; HeFH=heterozygous familial hypercholesterolemia; LDL-C=low-density lipoprotein-cholesterol; LLT=lipid lowering treatment.
Source: Merative Marketscan Claims Linked with Lab Data. 2019 - 2022. 12 months continuous data for each patient (6 months LB and 6 months LF from 1st observed statin treatmen



Majority of ASCVD/HeFH patients are not achieving LDL-C targets

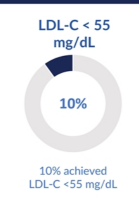
Primary prevention HeFH
patients with
an LDL-C target <100 mg/dL
(2011-2017)¹

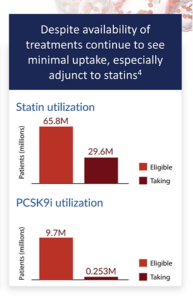


ASCVD patients with an LDL-C target of LDL<70 or <55 mg/dL (2017-2018)²



Very high risk ASCVD patients with an LDL-C target <55 mg/dL (2020-2021)³





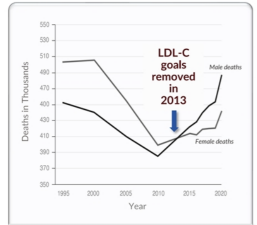


EVDP-atherosclerotic cardiovascular diseases: HeFH-heteroargous familial hypercholesterolemia; LDL-Colow-density lipoportorie-in-Oolesterol.

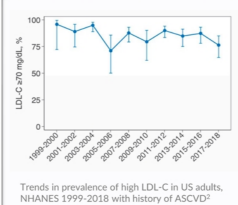
Schreuder MM, et al. LDL cholesterol targets rarely achieved in familial hypercholesterolemia patients: A sex and gooder-specific namalysis. Atherosclerosis. 2023 2. Gao Y, Shah LM, Ding J, Martin SS. US trends in cholesterol screening, lipid levels, and sid-lowering medication use in US adults, 1999 to 2018. J Am Heart Assoc. 2023;12(3):e028205; 3. Katzmann IL, et al. Simulation study on LDL cholesterol target attainment, treatment costs, and ASCVD events with bempedoic acid in patients at high ord name high patients rate in Color (2018). The color of the Co



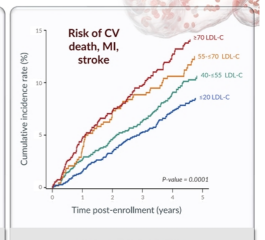
Increased CV events following removal of LDL-C guidelines in 2013



Despite statins, CVD deaths are on the rise



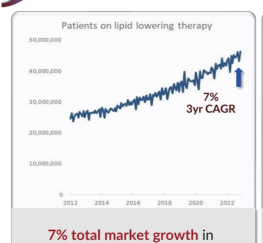
~75% of ASCVD patients are NOT at their risk-based LDL-C goal



Numerous studies demonstrate resurgence of paradigm "lower is better"



Resurgence of the "lower is better" paradigm leading to significant US market growth



the US





18% non-statin patient growth

Driven by generic ezetimibe, given lack of convenient and efficacious alternatives

Recent guideline and label changes driving renewed acceleration

June 2023: ACC updated guidelines to target LDL-C <55 mg/dl in high risk patients in line with ESC/EAS November 2023: FDA highlights need to reduce access restrictions for LLTs. Labels updated from "on top of maximally tolerated statins" to "treatment of primary hyperlipidemia" for some LLTs



Source: Symphony Health data through October 2023



Few approved post-statin LDL lowering products, which are limited by efficacy, convenience and/or payor access

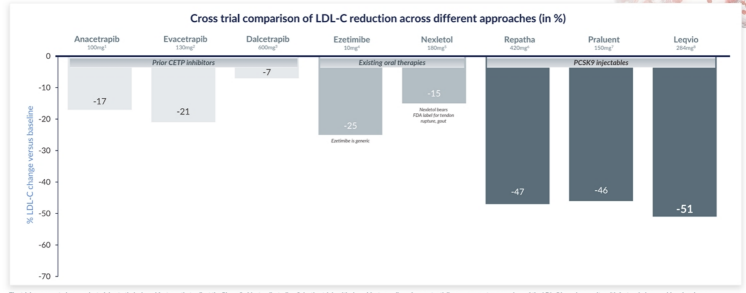
						THE WEATHER
	Ezetimibe(1)	Nexletol ⁽²⁾	PCSK9i ⁽³⁾	Oral PCSK9 ⁽⁴⁾	Obicetrapib ⁽⁵⁾	Obi + Eze ⁽⁵⁾
Approval	Approved	Approved	Approved	LDL data 2026E (CVOT data 2029E)	LDL data 2024E (CVOT data 2026E)	LDL data 2025E
MACE Benefit	7%	13%	15%	TBD	TBD	TBD
Observed LDL-C Reduction	25%	15%	45-50%	50-59%	43-51%	63%
Administration	Oral (small molecule)	Oral (small molecule)	Injectable (mAb)	Oral (peptide)	Oral (small molecule)	Oral (small molecule)
Dosing	10mg	180mg	140-150mg	380mg (20mg API + 360mg SNAC)	10mg	20mg (10mg Obi + 10mg Eze)
Food Effect	No	No	No	Yes (8hr fast & 30min wait)	No	No
Safety & Tolerability	Safe, Well-Tolerated	Tendon rupture & gout warning on label	Safe, injection site reactions	SNAC technology has previously been observed to have tolerability concerns ⁽⁶⁾	Well-Tolerated compared to placebo	Well-Tolerated compared to placebo
Lp(a) lowering	Raises	None	15-30%	20-25%	47-57%	40%



ote: The above data do not represent head-to-head comparisons. Actual results may differ from expectations. Obicetrapib mono and Exetimibe combo, along with the Oral PCSK9 have not been approved by any regulatory authority. Ee estimated dates. Jurises: 1. Pl Zetia table 7. refers to; Gagne, C et al. Am J Cardiol 2002. LDL: C measured only using Friedewald 2. Pl Needledol 2. Pl Needledol 2. Pl Needledol 2. Pl Needledol 2. So mg/ds. 3. AMA 2019;322(18):1780-1788. LDL: C measured using Friedewald 2. Pl Needledol 2. Pl Needledol 2. Pl Needledol 2. So mg/ds. 3. AMA 2019;322(18):1780-1788. LDL: C measured using Friedewald 2. Pl Needledol 2. Pl Needled



40-50% LDL-C reduction comparable to high efficacy PCSK9 injectables



The trials represented were selected due to their shared features that reflect the Phase 3 obicetrapib studies. Selecting trials with shared features allows for a potentially more accurate comparison of the LDL-C lowering results, with factors being considered such as:
a) presence of intensive LDL-lowering therapy including (high intensity) statins and PCSK9 inhibitors, b) patient population – ASCVD or ASCVD risk equivalent patients (including primary hypercholesterolemia and HefH-1) and c) where possible, selected studies where LDL-C measured by preparative ultracentrifugation (PUC) as opposed to Friedewald; noted below are those instances where PUC was not used – this is important because at low LDL-C levels (< 50 mg/dL), calculated LDL-C by Friedewald is overestimated; certain significant devia from these parameters are provided in the footnotes.

Note: The above trials and data do not represent head-to-head comparisons. Actual results may differ from expectations.

Sources: Forculation. 2021;144:e564-e939 17065. 1. Bowman, Let al. N Engl J Med 2017. 2. Amirhossein, Set al. Curr Pharmaceutical Design 2016. Meta-analysis - Also included hyperlipidaemia patients. LDL-C measured using direct assays and Friedewald. 3. de Grooth et al. Circulation 2022; LDL-C measured only using Friedewald and did not require subjects to be on prior statin therapy or present with ASCVD. 4. PIZetia table 7. refers to; Gagne, C et al. Am J Cardiol 2002. LDL-C measured only using Friedewald and did not require subjects to be on prior statin therapy or present with ASCVD. 4. PIZetia table 7. refers to; Gagne, C et al. Am J Cardiol 2002. LDL-C measured only using Friedewald and did not require subjects to be on prior statin therapy or present with ASCVD. 4. PIZetia table 7. refers to; Gagne, C et al. Am J Cardiol 2002. LDL-C measured using refedewald and did not require subjects to be on prior statin therapy or present with ASCVD. 4. PIZetia table 7. refers to; Gagne, C et al. Am J Cardiol 2002. LDL-C measured using refedemal and did





Obicetrapib program designed to overcome limitations of prior CETP inhibitors

	Torcetrapib
Observed LDL-C reduction	20%
CETP inhibition	80%
Dosing	60mg
Blood pressure increase	Yes
Aldosterone increase	Yes
Lp(a) lowering	unknown
ApoB lowering	10%
OUTCOMES STUDIES	
Name	ILLUMINATI
Patients	15,067
Baseline LDL-C (mg/dl)	79.7

Dalcetrapib ⁽²⁾
7%
40%
600mg
No
No
unknown
None
Dal-OUTCOMES
15.871
76.4
NS
31 mo
1.04
No LDL-C benefit

Evacetrapib ⁽³⁾
21%
65%
100mg
No
No
20-25%
15%
ACCELERATE
12,092
81.1
25
26 mo
1.01
Short follow-up but mortality benefit (HR 0.84)

Anacetrapib ⁽⁴⁾)
17%	
90%	
100mg	
No	
No	
20-25%	
18%	
REVEAL	
30,449	
61	
11	
49 mo	
0.91	
As expected, love baseline and LD reduction	

98%
10mg
No
No
47-57%
25%-35%
PREVAIL
>9,000 (expected)
~105 (expected)
TBD
42 mo (expected)
TBD
TBD

Obicetrapib⁽⁵⁾
43%



Explanation

Median follow-up Result (HR)

LDL-C reduction (mg/dl)

Note: The above trials and data do not represent head-to-head comparisons. Actual results may differ from expectations.

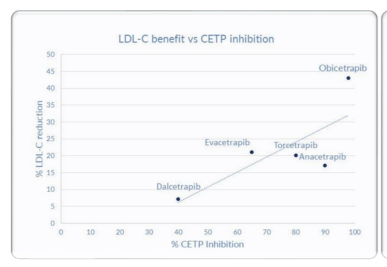
20 18 mo

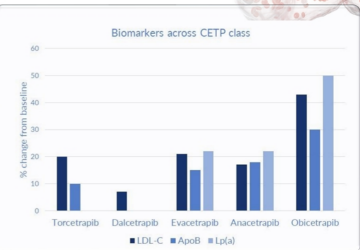
1.25

Off target tox



Enhanced LDL-C reduction with Obicetrapib's greater potency

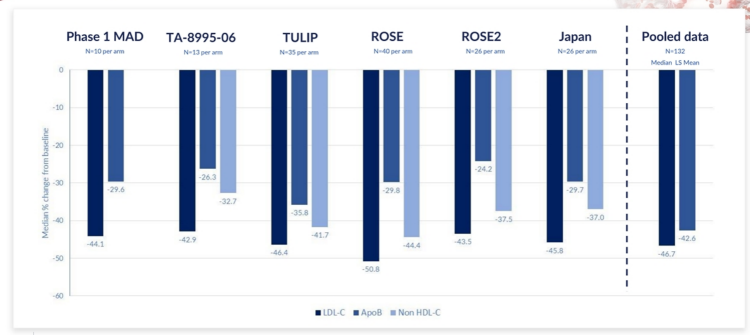






lote: The above trials and data do not represent head-to-head comparisons. Actual results may differ from expectations.

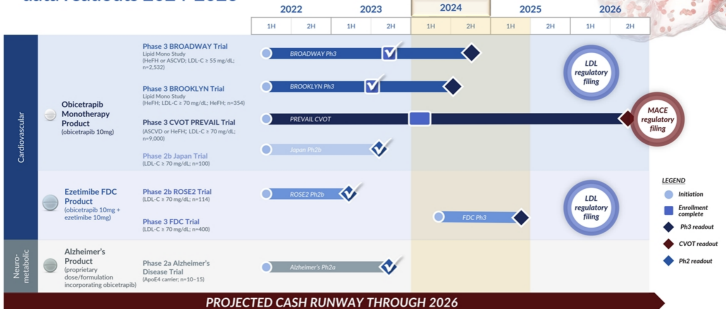
Obicetrapib Phase 1/2 studies: Consistent benefits observed in lipid biomarkers





burce: Company data for obicetrapib 10mg monotherapy, Pooled data includes TULIP, ROSE, ROSE2, and Japan Phase 2 data set

Sufficient cash expected to fund the Company through multiple potential pivotal data readouts 2024-2026



NewAmsterdam Pharma

Numerous catalysts expected throughout 2024-2026



BROOKLYN study design

Objective: To evaluate the effects of obicetrapib in patients with heterozygous familial hypercholesterolemia (HeFH)

Inclusion criteria

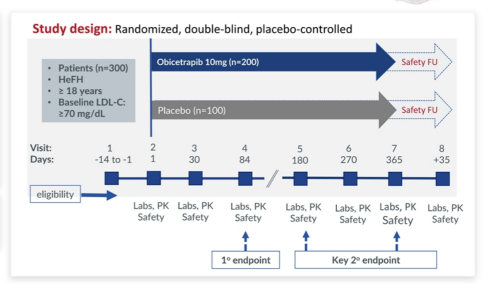
- HeFH by genetic confirmation, and/or WHO Criteria/Dutch Clinical Network, and/or Simon Broome criteria
- 70% of patients on HS
- 10% Statin Intolerant
- Stable lipid lowering therapies with an LDL ≥ 70 mg/dL and TG ≤ 400 mg/dL

Exclusion criteria

- CV disease < 3 months
- HofH
- · Uncontrolled hypertension

Primary efficacy endpoint

Percent change from baseline in LDL-C compared to the placebo group







BROADWAY study design

Objective: To evaluate the effect of obicetrapib on top of max tolerated lipid-modifying therapy in patients with HeFH and or ASCVD



Have a fasting serum LDL-C at Screening (Visit 1) as follows:

Have a fasting serum LDL-C at 55 mg/dL (≥ 1.4 mmol/L) to <100 mg/dL (< 1.4 mmol/L) 0 Ron-HDL-C = 85 mg/dL (≥ 2.2 mmol/L) to <130 mg/dL (< 2.6 mmol/L) with at least 2 risk enhancers

OR

Have a fasting serum LDL-C ≥ 100 mg/dL (≥ 2.6 mmol/L) OR non-HDL-C ≥ 130 mg/dL (≥ 3.4 mmol/L).

Risk enhancers:

Age of >60 years:

Recent MI (>3 and <24m prior to Randomization);

Type 2 diabetes mellitus;

Current cigarette smoking;

hsCRP ≥ 2.0 mg/L;

TG >150 mg/dL (>1.7 mmol/L);

Lp(a) >30 mg/dL (<70 nmol/L);

HDL-C <40 mg/dL (<1.0 mmol/L);

Exclusion criteria

- HoFH
 Uncontrolled hypertension

Primary efficacy endpoint

% change from BL to Day 84 in LDL-C for obicetrapib vs placebo

Key secondary efficacy endpoint

CV death, MI, stroke, or non-elective coronary







PREVAIL trial design leverages lessons learned



- · Study design:
 - o n = 9000
 - o Inclusion: ASCVD patients on maximally tolerated statins with risk enhancers and LDL-C > 70mg/dl
 - Minimum follow up 2.5 years
- · Primary endpoint: 4-point MACE
- First secondary: 3-point MACE
- Prespecified endpoints:
 - o Conversion of pre-diabetes to diabetes
 - o A1c levels in diabetes patients
- Patient populations of interest
 - o Patients on PCSK9
 - o Patients on GLP-1
 - o Patients on SGLT-2

Applying lessons from prior CVOTs

Greater LDL-lowering activity anticipated

42.6% observed in Phase 2

plus

Targeting higher baseline LDL patients

~100mg/dl anticipated

Higher absolute LDL-C reduction expected to lead to greater MACE benefit

Longer duration of follow up

Median of 42 months vs. only 2.1 years in ACCELERATE

plus

Targeting higher-risk patient population
ASCVD patients further enriched with with risk enhancers shown in REVEAL long-term follow up to have stronger relative risk reduction (high LDL/ApoB, diabetes, high triglycerides, recent MI)



More time + higher patient risk potentially maximizes opportunity for MACE reduction

Differentiated secondary endpoints

Lp(a)-lowering, HDL-raising, diabetes, and Alzheimer's benefits



Potentially enhanced commercial profile vs. other LDLlowering agents + potential therapeutic area expansion





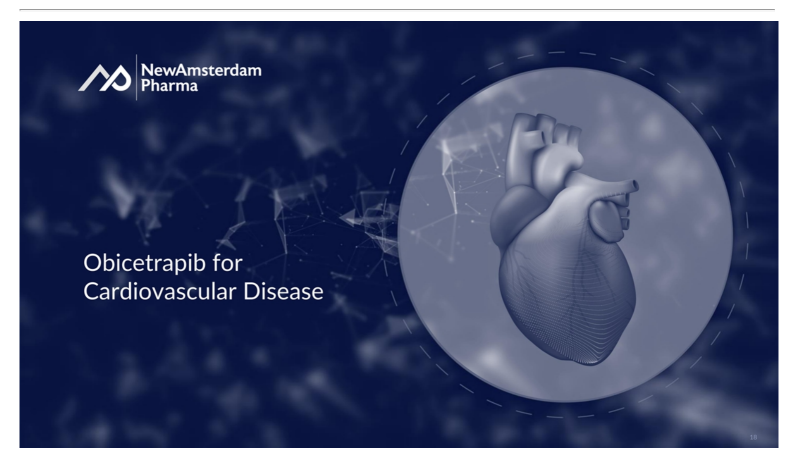
2023 achievements pave the way for potential 2024 value inflection milestones



	1Q 202	<u>!</u> 4	3Q 2024	4Q 2024		1Q 2025
2024	Complete enrollment for PREVAIL CVOT	Initiate FDC Phase 3 trial	BROOKLYN Phase 3 topline	BROADWAY Phase 3 topline	2025	TANDEM FDC Phase 3 topline
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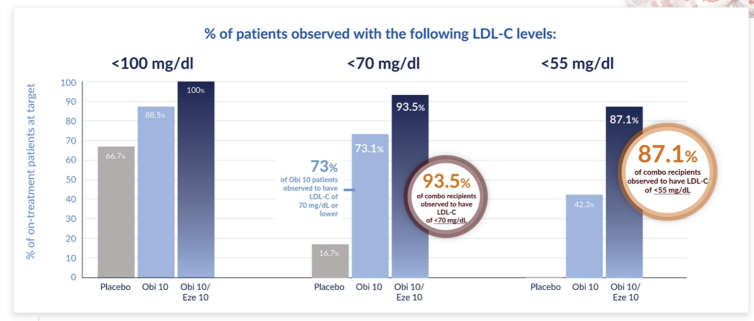


: Projections are subject to inherent limitations. Actual results may differ from expectations. The timing of regulatory submissions is subject to additional discussions with regulator





Exceptional LDL goal attainment observed with ezetimibe + obicetrapib combination, including >87% of patients observed to attain <55 mg/dl LDL-C levels

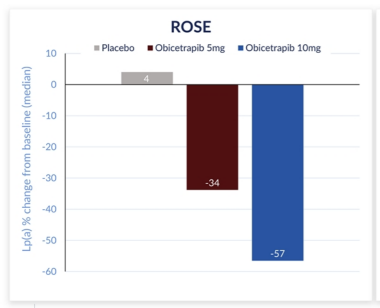


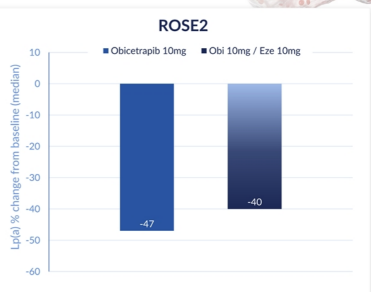




Lp(a) percent reduction from baseline in ROSE¹ and ROSE2²

• Lp(a) is emerging as a strong and independent marker of CVD risk and an exciting new CVD drug target



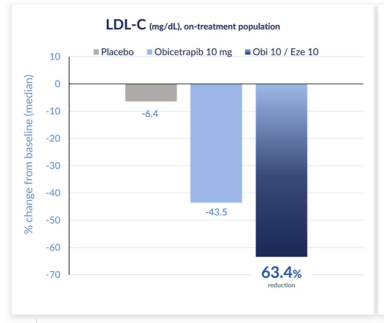




Source: 1. Nicholls SJ, et al. Nat Med 2022;28:1672-1678. 2. Ballantyne CM, et al. J. of Clinical Lipidology 20



Obicetrapib/ezetimibe observed to lower LDL-C by 63.4% on top of HIS in ROSE2



Median (min, max) LDL-C levels (mg/dL) at baseline & EoT

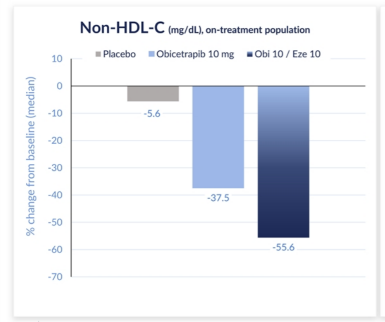
Time	Placebo	Obi 10 mg	Obi 10 / Eze 10
	95.5	100.0	87.0
Baseline Median	(60, 211)	(35, 189)	(62, 152)
	(N=40)	(N=26)	(N=31)
	88.0	55.5	39.0
EoT Median	(55, 188)	(21, 148)	(15, 96)
	(N=36)	(N=26)	(N=31)
% Change	-6.4	-43.5	-63.4
from Baseline	(-36.4, 96.7)	(-78.4, 22.6)	(-83.7, -29.7)
(Median)	(N=36)	(N=26)	(N=31)
% Change from Baseline	-0.85	-39.20	-59.23
LS mean (95% CI)	(-7.75, 6.05)	(-47.41, -30.99)	(-66.75, -51.71)
P-value	-	<0.0001	<0.0001

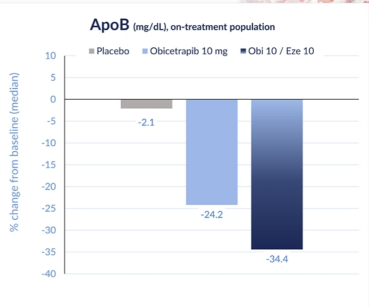


Source: Ballantyne CM, et al. J. of Clinical Lipidology 202:



ROSE2: Non-HDL-C and ApoB percent change from baseline (Day 84)

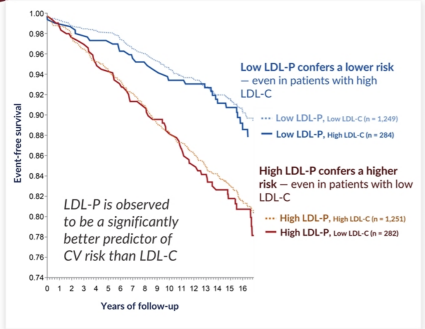






Source: Ballantyne CM, et al. J. of Clinical Lipidology 2023

LDL-P believed to be one of the most robust predictors of cardiovascular risk



- Small dense LDL particles are more likely to be trapped in arterial wall than larger-sized LDL particles
- High LDL-P levels typically signify that a patient has a higher proportion of small dense LDL particles vs. larger-sized LDL particles



Even though all LDL particles contain only one ApoB protein, small dense LDL particles have a less massive ApoB protein

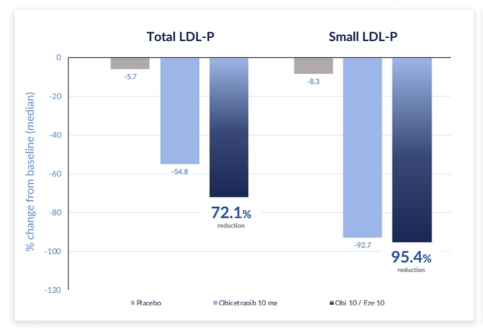




Source: Cromwell WC, et al. Clin Lipidol. 2007 December 1; 1(6): 583-59



ROSE2 showed significant reduction in total and small LDL particles, bringing patients who had baseline elevated LDL-P to optimal parameters⁽¹⁾



Patients taking the Obi/Eze combo observed to achieve optimal LDL-P profiles

ROSE2 placebo	ROSE2 Obi / Obi + Eze
1012.8	495 / 300
717.5	73.4 / 47.5
20.26	21.0 / 21.0
	1012.8 717.5

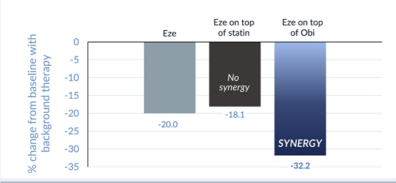
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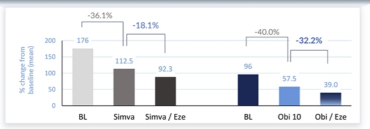
	High	Moderate	
LDL-P (nmol/L)	>1816	935-1816	<935
Small LDL- P (nmol/L)	>820	467-820	<467
LDL size (nm)	≤20.5	N/A	>20.5

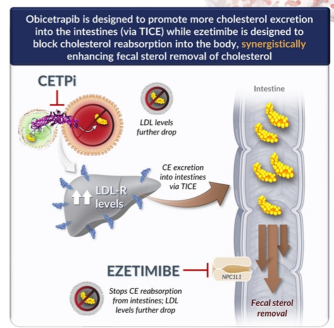


Sources: 1. Ballantyne CM, et al. J. of Clinical Lipidology 2023, 2. LipoFraction NMR practitioner Guideline

Stronger LDL-lowering observed with ezetimibe in obicetrapib combo vs. ezetimibe with statins, potentially due to a synergistic mechanism of action for obi/eze combo(1)









Source: Ballantyne CM, et al. J. of Clinical Lipidology 2023 Note: The calculations included in the graphs here recresent the Company's hypothetical calculation assuming one patient was treated with each drug independently



Favorable safety profile observed in all LDL Phase 1 & 2 clinical studies

	Comparator ⁽¹⁾ (N=231)	Pooled Obicetrapib (5, 10mg) ⁽²⁾ (N=309)
TEAEs (%)	(11 202)	(11 307)
TEAEs, total	136 (58.9)	173 (55.9)
TEAEs, related	45 (19.5)	49 (15.8)
TEAEs, severe	5 (2.2)	7 (2.3)
TESAEs		
*TESAEs, total	6 (2.6)	4 (1.3)
TESAEs, related	0	0
Deaths	0	0
Withdrawals study / medication		
TEAEs leading to discontinuation of study drug	13 (5.6)	13 (4.2)



^{*} There were three additional TESAEs in other obicetrapib dose arms: two in the TULIP 2.5mg arm, and one in the Lp(a) 2.5mg arm; none were considered to be related to study drug.

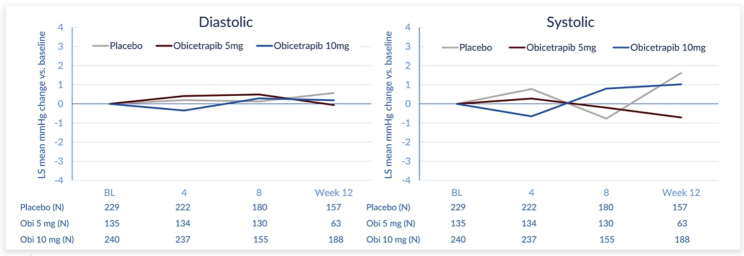
(1) The Comparator group included patients receiving placebo and non-obicetrapib monotherapy.

(2) The pooled obicetrapib group includes patients treated with obicetrapib as a monotherapy and in combination with atorvastatin, rosuvastatin and ezetimibe.



Obicetrapib does not show an effect on systolic and diastolic blood pressure

- A dedicated meta-analysis of the obicetrapib ROSE2, ROSE, TULIP, OCEAN, and TA-8995-203 study did not reveal any signal in systolic and diastolic blood pressure
- By contrast, in the cardiovascular outcome trial ILLUMINATE, torcetrapib showed a significant 5.4 and 2.0mm Hg
 increase in systolic blood and diastolic pressure and was associated with a significant decrease in serum potassium, and
 increases in serum sodium, bicarbonate and aldosterone





Sources: Circulation 2021;144:e564–e593 17065: Obicetrapib Lowers LDL-C in Patients Taking High Intensity Statins: Results From Rose Clinical Trial.





Obicetrapib program designed to overcome limitations of all prior CETP inhibitors



SAFETY OFF-TARGET TOXICITY. INCREASED BLOOD PRESSURE. ALDOSTERONE (seen early in Phase 2) Safe & well-tolerated Strong safety Safe & profile across ~59k well-tolerated Safe & well-tolerated

We believe that all prior CETPi were developed with a misguided focus on HDL increase (rather than LDL decrease) as the primary MoA for CVD risk reduction, leading to inappropriate compound selection or inappropriate CVOT design



INSUFFICIENT TRIAL DURATION (only 2 years) Sufficient duration (4.1 years, with 6.3 year follow up) Baseline LDL too low (60 mg/dL)

COMMERCIALLY UNVIABLE - HIGH LIPOPHILICITY AND FAT TISSUE ACCUMULATION LED TO 4+ YEAR

HALF-LIFE

OBICETRAPIB⁵

observed - but drug accumulated in fat tissue

Meaningful MACE benefit



Tolerability profile observed in >800 patients through Phase 2b

No concerns seen in biomarker safety data, including blood pressure-associated biomarkers √ ~43% LDL-LOWERING **OBSERVED IN PHASE 2B**

✓ ~59% LDL-LOWERING **OBSERVED IN FDC PHASE 2**

√ Longer trial duration (4 yrs)

√ High baseline LDL (100 mg/dL)⁽¹⁾

= PREVAIL CVOT design expected to translate into 15-20% MACE benefit

√ Favorable PK/PD profile

√ No accumulation in fat tissue observed



NewAmsterdam Note: The above trials and data do not represent head-to-head comparisons. Obicetrapib has not been approved for marketing by any regulatory authority.

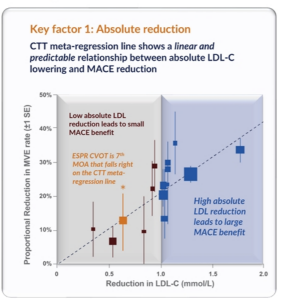
(1) Represents estimated average baseline IDL to be enrolled, not entry criteria.

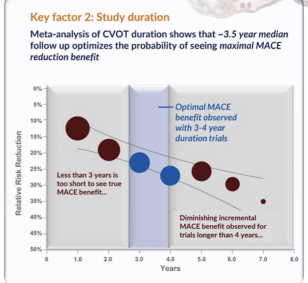
Sources: 1. Barter Pl, et al. K regij 1 Med 2017; 377:1217-1227; 5. Data on file



Absolute reduction of LDL-C and ApoB, and duration of that reduction are believed to be key to reducing cardiovascular risk









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Pharma

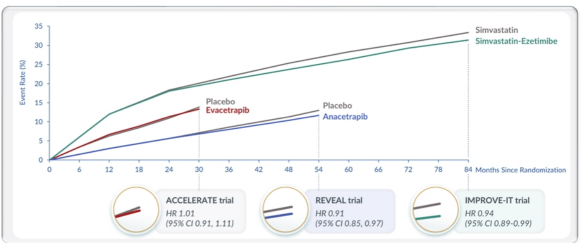
Sources: American Heart Association, CDC, Mayo clinic, Global Health estimates 2016: Deaths by Cause, Age, Sex, by Country and by Regio, 2000-2016, Geneva, WHO; 2018. Inacet 2005;966-1267-78; Silverman MG, et al. JAMA 2016;27:1289-1297. * ISPR CVOT depiction based on ACC 2023 presentation, drawn as approximation to existing meta



ACCELERATE, REVEAL and IMPROVE-IT support our belief that CVOT study duration should be long enough to see optimal MACE benefit

Kaplan-Meier curves for these trial, with very similar absolute ApoB reductions, show separation later than 2 years, which is the point in time that ACCELERATE stopped





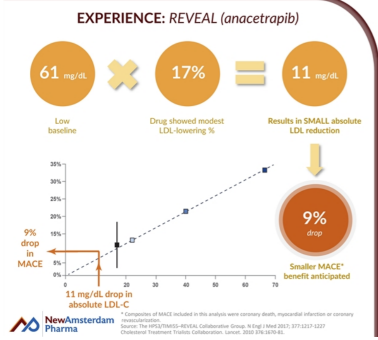


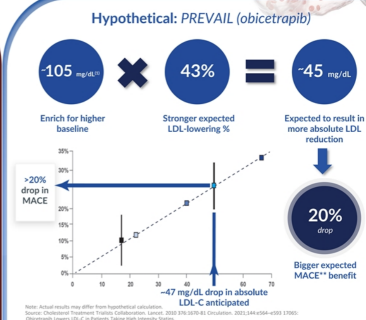
rives are for the primary efficacy endpoint, which in IMPROVE-IT was defined as the composite of death from cardiovascular disease, a major coronary event (nonfatal myocardial infarction, documented unstable angina requiring hospital admissior recoronary revascularization occurring at least 30 days after randomization), or nonfatal stroke, in ACCELERATE as the composite of death from cardiovascular causes, myocardial infarction, stroke, coronary revascularization for instable angina, and in REVEAL as the composite of coronary death, myocardial infarction, or coronary revascularization.

innon CP, et al. N Engl J Med 2015:372:2387-2397. Lincoff AM, et al. N Engl J Med 2017:376:1933-1942. Bowman L, et al. N Engl J Med 2017:377:1217-1222

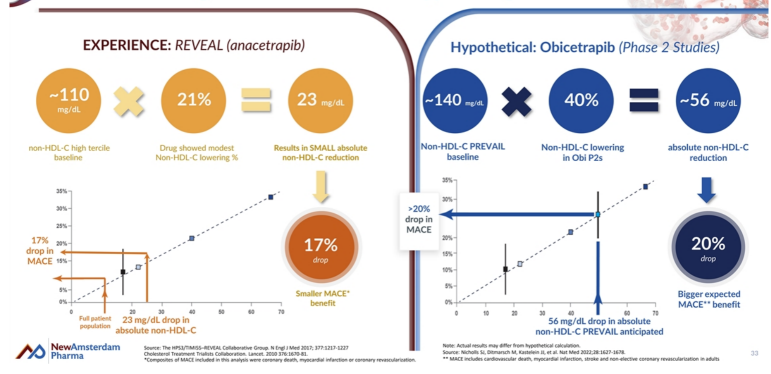


REVEAL data supports translation from absolute LDL reduction to MACE benefit





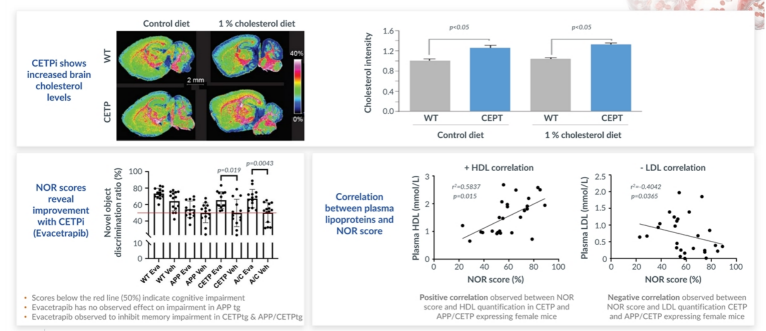
REVEAL data in high tercile in non-HDL-C supports larger MACE benefit







CETP knock-in mice observed to increase brain cholesterol levels and CETPi rescues cognition in preclinical models of CETP-induced AD



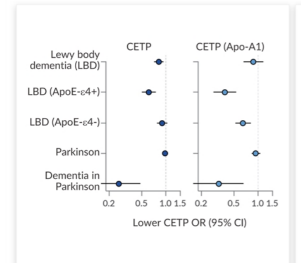


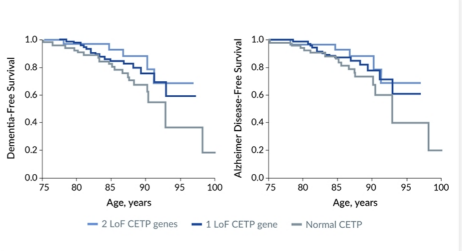
NewAmsterdam Pharma Source: Felix Oestereich, et al., The Cholesteryl Ester Transfer Protein (CETP) raises Cholesterol Levels in the Brain and affects Presenilin-mediated Gene Regulation, Journal of Lipid Research, vol. 63, no.9, 2022.



CETP loss-of-function (LoF) genotype may be associated with slower memory decline and lower AD risk

- CETP's potential involvement in CNS cholesterol homeostasis is supported by genetic data
- CETP LoF genotype may be associated with lower CETP activity & a corresponding increase in HDL levels





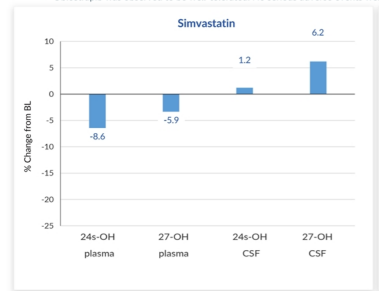


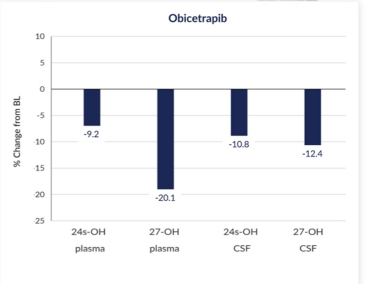
Source: JAMA, January 13, 2010-Vol 303, No. 2



Initial data for Obicetrapib 10mg observed to decrease 24s- & 27-hydroxycholesterol ("OH") in both plasma and cerebrospinal fluid ("CSF")

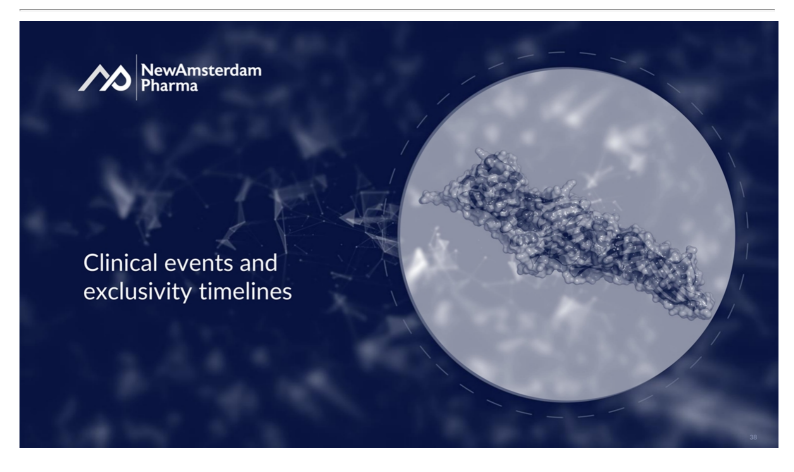
- In separate trials with different protocols and endpoints, Simvastatin was observed to only reduce 24s- and 27-OH in plasma
- Obicetrapib was observed to be well-tolerated. No serious adverse events were reported, nor were any adverse events considered to be related to the study drug.







Source: Alzheimer Dis Assoc Disord. 2010; 24(3): 220–226 and Company data
Note: The results shown above do not represent head-to-head comparisons. The data was obtained from clinical trials with different objectives, designs and patients. Actual results may differ from expectations.





Projected exclusivity timelines in the EU and US Assumes EU approval 4Q 2025 and US approval 1Q 2027

		2023	2024	2025	2026	2027	2028	2029	2030	2031	2032	2033	2034	2035	2036	2037	2038	2039	2040	2041	2042	1
Regulatory exclusivity	EU																	C max 0/40				
	US	US approval 1/1/27 NCE exclusivity (5 years) 1/32 30 mo. stay 7/34 6 mo. pedi. (1/35)														PTE ma 1/41						
1 st gen		Origin	Original genus patent family 8/27 (U.S.); 4/25 (EU)																			
2 nd gen ⁽¹⁾	EU	Species selection/low dose patent family SPC 8/39*												/39*								
	US	Speci	es selec	ction/lo	w dose	patent	family					2/3	4	••••	P	TE	••••	8	/39*			
3 rd gen	EU	Proprietary form (COM) patent family**																7,	/43			
	US			•••	Prop	rietary 1	orm (C	OM) pa	itent fai	mily	••••	••••	••••	•••	••••	••••	••••	••••	• • • •		7/	/43





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