UNITED STATES SECURITIES AND EXCHANGE COMMISSION

Washington, D.C. 20549

FORM 6-K

REPORT OF FOREIGN PRIVATE ISSUER PURSUANT TO RULE 13a-16 OR 15d-16 UNDER THE SECURITIES EXCHANGE ACT OF 1934

For the month of March 2023

Commission File Number: 001-41562

NewAmsterdam Pharma Company N.V. (Exact name of registrant as specified in its charter)

Gooimeer 2-35 1411 DC Naarden The Netherlands (Address of principal executive office)

Indicate by check mark whether the registrant files or will file annual reports under cover of Form 20-F or Form 40-F:	
Form 20-F ⊠ Form 40-F □	
Indicate by check mark if the registrant is submitting the Form 6-K in paper as permitted by Regulation S-T Rule 101(b)(1): \Box	
Indicate by check mark if the registrant is submitting the Form 6-K in paper as permitted by Regulation S-T Rule 101(b)(7): \Box	

On March 7, 2023, 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 Report on Form 6-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 Report on Form 6-K.

EXHIBIT INDEX

Exhibit No. Description

99.1 NewAmsterdam Pharma Company N.V. Corporate Presentation

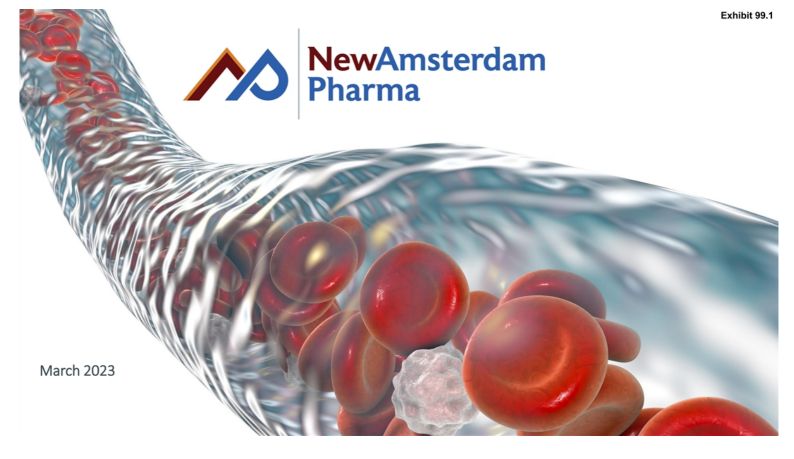
SIGNATURE

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, thereunto duly authorized.

NewAmsterdam Pharma Company N.V.

March 7, 2023

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





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," "anticipate," "intend," "expect," "should," "polan," "predict," "potential," "seem," "seek," "future," "outlook" and similar expressions that predict or indicate future events or that are not statements of historical matters. These forward-looking 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, under a company of initiation, completion and data readouts for clinical trials and the potential approval of the Company's product candidate; 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; the third product candidate; 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; 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; the therapeutic and curative potential of the Company's product candidate; 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 assurantee, an assurantee and are not predictions of actual performance. These forward

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 Date

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 is developing oral obicetrapib as CETPi to address major unmet need in hypercholesterolemia and other cardiometabolic diseases

Company history:

- Mid-'20: Founded after spin-out from Amgen
- Jan '21: \$196M Series A
- June '22: >\$1B European license with Menarini Group (including €142.5M committed capital + significant back-end economics)
- Nov '22: NASDAQ listing (NAMS) with \$328M total proceeds (\$93M from FLAC trust in deSPAC, plus \$235M PIPE colead by Frazier and Bain)
- Jan '23: ~\$1B market cap and \$450M cash





- 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
- Simple, oral, once-daily, low dose CETP inhibitor with strong LDL-lowering efficacy and safety observed through Phase 2b:
 - ~50% LDL-lowering as monotherapy, ~59% in combination with ezetimibe, observed on top of high-intensity statins
 - Strong safety and tolerability in >700 pts
 - Robust effects on ApoB, non-HDL-C, HDL-C and Lp(a)
- Convenient oral format + low cost of goods sold (COGS) enables broad market access strategy to address existing unmet need

>35M addressable patient population

~59% LDL-lowering observed in Phase 2b \$4B+ global market opportunity

Achievements and milestones:

- Fully-funded through 2026 including CVOT readout and potential global launch
- Significant number of patients randomized in Phase 3 to-date, first DSMB meeting completed
- Late-stage inflection points expected from 2023-2025 (including Phase 2 & 3 clinical readouts and registrational filings)
- Pipeline expansion potential in Alzheimer's disease and diabetes



Expert cardiometabolic leadership supported by top investors

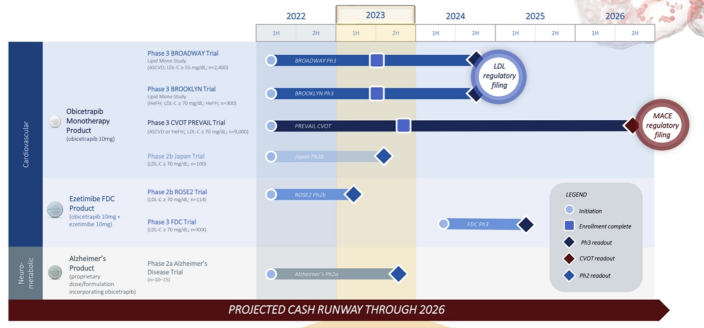


Michael Davidson, M.D. CEO	John Kastelein, M.D. CSO	Do	ouglas Kling COO	Lir	na Gugucheva CBO	Lou	ise Kooij CFO		narsch, M.D. DO
© CORVIDIA	uniQure	•	CORVIDIA		SCORPION'		I GENZYME 🗳	_	eneca Doche
SEASONED BOA OF DIRECTORS:	ARD SANDER SLOOTWEG Managing Partner, Forbion	JULIET AUDET Partner, Forbion	JAMIE TOPPER Managing Partner, Frazier Ufe Sciences	NICHOLAS DOWNING Partner, Bain Capital	LOU LANGE Partner, Asset Management Ventures	MICHAEL DAVIDSON CEO, NewAmsterdam Pharma	JOHN KASTELEIN CSO, NewAmsterdam Pharma	JOHN W. SMITHER Independent	
BACKED I	BY TOP TIER BVF RS: medicxi	Forbion.	FRAZIER	_	ainCapital R		TAL VIVING	LSP	





Net proceeds expected to fund obicetrapib development through several value-creating milestones



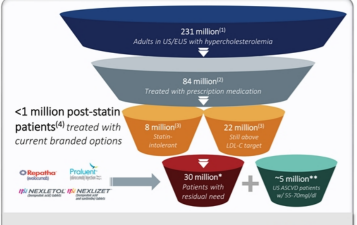


Numerous catalysts throughout 2023

ote: Other than as noted, the pipeline represents trials that are currently ongoing. Projections are subject to inherent limitations. Actual results may differ from expectations. The timing of regulatory submissions is subject to additional discussions in the contract of the pipeline represents trials that are currently ongoing. Projections are subject to inherent limitations. Actual results may differ from expectations. The timing of regulatory submissions is subject to additional discussions.

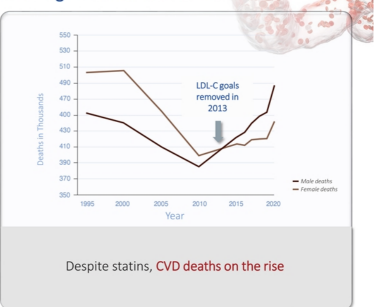


Despite availability of statins, CVD remains the leading cause of death worldwide



Key factors limiting penetration include **product limitations** and **market** access hurdles

**Factoring in new U.S. guidance lowering LDL-C goals to 55 mg/dL for very high-risk patients, which will grow addressable patient population





[1] Uterature review suggesting hypercholesterolemia prevalence of ~94mm in the US (average of He et al. 2020, Mercado et al. 2015, Munther et al. 2013) and ~137mm in EUS (average of Gomes-Huelgas et al. 2010, Guallar-Castillon et al. 2012, Trage et al. 2012, Grau et al. 2013).

| 2012 Grau et al. 2013, Crau et al. 2013, Crau et al. 2014, Grau et al. 2

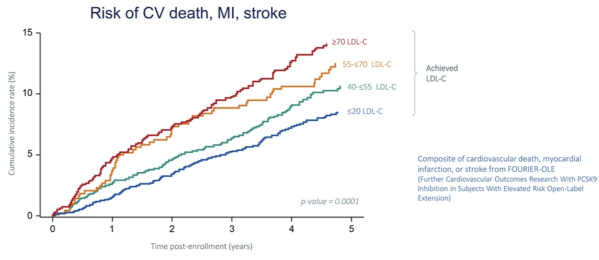
and 59% compliance for statins & Nex/Nex (based on statin literature) treated patient volume estimates were derived from the prescription data and extrapolated to the EUS.

3) simm statin-individent & Zarma shove UILC tragger Everentage of patients in each category estimated from Trinity quantitative market research and the — percentages were then applied to the estimated 8Amm treated number above.
4) carma branded patients: 2020 US prescription data for Repatha, Praluent, and Nexietol/Nexilizet were guiled from the Bloomberg Prescription Data Portal that Trinity subscribes to; assuming 12 scripts/year and 70% compliance for PCSR9s (based 6754) Biotechnois of 65% compliance for Abov/Nex (based on statin literature) analysis of solvening activates users derived from the negoriation data and extragolated in the EUS.



Cardiovascular field is returning to the mantra of "Lower is Better", as supported by recently published FOURIER-OLE analysis

In patients with atherosclerotic cardiovascular disease, long-term achievement of lower LDL-C levels, down to <20 mg/dL (<0.5 mmol/L), was associated with lowest of cardiovascular outcomes compared to patients who achieved less robust LDL-C reduction

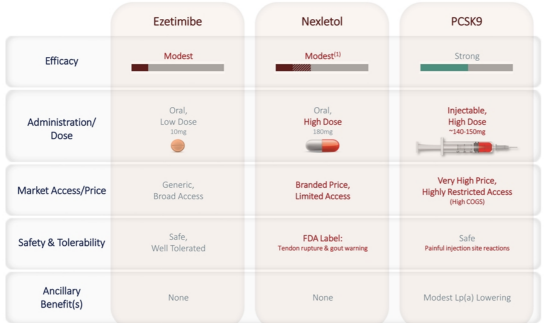




Note: Gaba P, et. al., Association Between Achieved Low-Density Lipoprotein Cholesterol Levels and Long-Term Cardiovascular and Safety Outcomes: An Analysis of FOURIER-OLE. Circulation. 2023 Feb 13. doi: 10.1161/CIRCULATIONAHA.122.063399



Currently approved post-statin LDL-lowering products all fall short of the profile patients



Patients need a drug that fits the following profile:

Strong⁽¹⁾

Oral, Low Dose

Disruptive Price, **Broad Access** (Low COGS)

Safe, Well-Tolerated

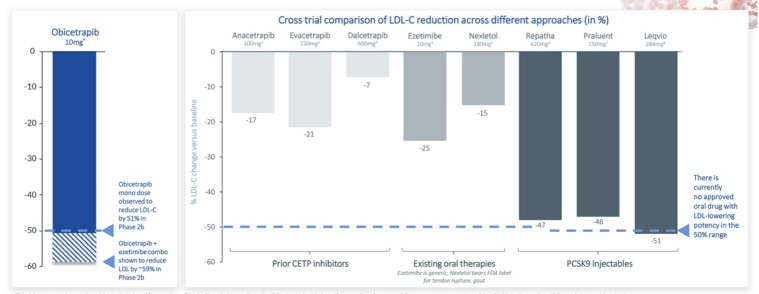
Strong Lp(a) Lowering, **HDL** Raising has Potential Benefits in Alzheimer's and Diabetes



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Note: Novartis' stated price of Inclisiran to be approximate to net prices of PCSK9 mAt
(1) Lighter shaded region of bar chart represents the additional potential efficacy for



~50% LDL-C reduction efficacy would be virtually identical to PCSK9 injectables and is substantially better than other oral therapies



The trials represented were selected due to their shared features that reflect the Phase 3 objecterable 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) patients population — ASCVD in skequivalent patients (including primary hypercholesterolemia and Heffl) and where possible, selected studies where LDL-C measured by preparative ultracentrifusation (PLC) as opposed to Friedewald: noted below are those instances where PLUC was not used—this is important because at low IDL-C levels (+S DI me/dL). Acclusted IDL-C by Friedewald is coverestimated: certain significant deviations from these parameters are provided in the footnotes.



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

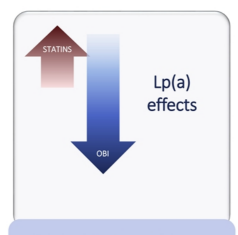
verse: *Circulation. 2021;144:e564—e593 17065. 1. Bowman, L et al., N Engl J Med 2017. 2. Amirhossein, S et al. Curr Pharmaceutical Design 2016. Meta-analysis - Also included hyperlipidaemia patients. LDL-C measured using direct assays and indexwald. 3. de Grooth et al. Circulation. 2002. LDL-C measured only using Friedewald and did not require subjects to be on prior statin therapy or present with ASCVD. 4. Pl Zetia table 7. refers to; Gagne, C et al. Am J Cardiol 2002. LDL-C measured villy using Friedewald and direct assay for LDL-C 50 mg/dL. 6. Pl Repatha; study 2. refers to; Goldberg, A et al. JAMA 2019;32(18):1780-1788. LDL-C measured using Friedewald and direct assay for LDL-C 50 mg/dL. 6. Pl Repatha; study 3. refers to; Blom, D et al. N Engl J Med 2014. Also challed hashed interpretation and the properties of the path of the pa



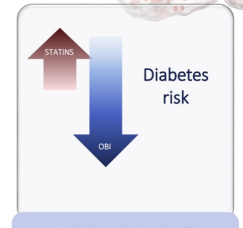
Obicetrapib has potential to be ideal complement to statin therapy



Obicetrapib's LDL and ApoB lowering potency may finally take statin patients to goal



Obicetrapib's robust Lp(a)-lowering effect (~57% lowering seen in P2) counter-acts Lp(a)-raising effect observed with statins

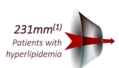


CETPi has been shown clinically to reverse diabetes risk, potentially ameliorating key patient concerns about high dose statins





Obicetrapib has potential to solve a substantial unmet medical need in dyslipidemia



>35mm patients not sufficiently addressed by available treatment options

Fewer than 1mm patients treated with current branded options: PCSK9s Bempedoic Acid Payors highly restrict access THE Low prescriber enthusiasm **PROBLEM** Relatively low patient compliance

OBICETRAPIB



- ~50% LDL-lowering observed for monotherapy, ~59% in combination with ezetimibe in Phase 2b
- ✓ Potential for attractive pricing to unlock broad
- ✓ Strong safety and tolerability profile observed
- ✓ Convenient once-daily oral tablet
- √ High prescriber and payor enthusiasm

>\$4B⁽²⁾ global market opportunity



THE SOLUTION



US market research indicates strong prescriber and payor enthusiasm, wide window of pricing opportunity to enable broad market access strategy

Strong enthusiasm from prescribers and payors:

PRESCRIBER ENTHUSIAM:

	Nexletol	PCSK9i	Obicetrapib
Efficacy	Moderate/Low*	High	High
Safety/ Tolerability	Moderate	High	High
Route of Administration	High	Low	High

PAYOR ENTHUSIAM:

"Biggest need (is) a product that works like the PCSK9s but with a more convenient ROA and no diabetes risk"

"Fulfills an important unmet need"





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Key: Blue cells denote high ratings (6.5 – 9) / Yellow cells denote high ratings (6.5 – 9)



Developing monotherapy and FDC in parallel builds in optionality for patients and prescribers, optimizes chances of broad market access

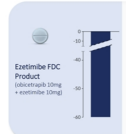


- o Expected to be preferred:
 - Initially, by less aggressive prescribers and those seeking more experience with CETPi class
 - By prescribers with lingering misperceptions about ezetimibe
 - Over time, expect that this will be the product that moves into earlier lines of therapy and primary care utilization

Obicetrapib mono utilization

Ezetimibe FDC utilization

Note: Nexletol/Nexlizet scripts are currently generating 1:1 sales, however ESPR has signaled expectations for Nexlizet to become preferred with time



o Expected to be preferred:

- By more aggressive prescribers & KOLs
- · For highest risk patients
- By more physicians over time as treatment paradigms shift, CETPi experience is gained

Parity pricing with monotherapy should ensure parity access CVOT data from both obicetrapib and ezetimibe monotherapies + Phase 3 FDC data would be included in FDC label MORE EXPERIENCE WITH CETPI

plus

BROADER ADOPTION OF NEW TREATMENT PARADIGMS:

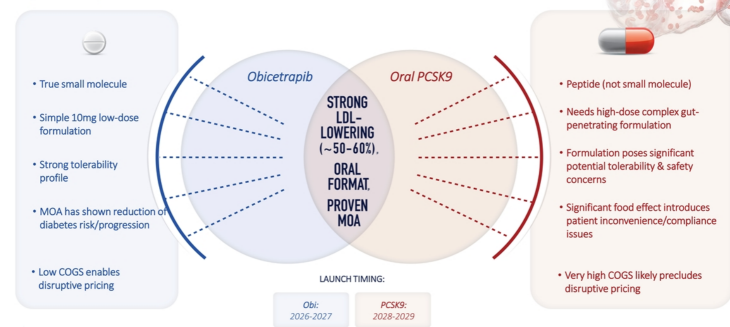
- / "Lower is better"
- ✓ Decreased LDL target goals
 - Earlier-line use of combination therapy

Expected to drive heavier FDC utilization over time





Obicetrapib will launch at least 2-3 years ahead of any oral PSCK9 with a better optimized product profile



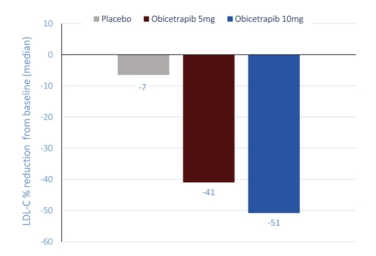






In ROSE Phase 2b clinical trial, obicetrapib demonstrated robust LDL-C lowering as adjunct to high intensity statins¹

Preparative ultra-centrifugation (PUC) is "gold-standard" for LDL-C quantification



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

Time	Placebo	Obicetrapib 5mg	Obicetrapib 10mg
Baseline Median	90.0	95.0	88.0
	(63, 204)	(54, 236)	(39, 207)
	N=40	N=39	N=40
	86.0	53.0	49.5
EoT Median	(43, 137)	(13, 126)	(23, 83)
	N=39	N=39	N=40
% Change	-6.5	-41.45	-50.75
from Baseline	(-53.9, 31.6)	(-71.2, 62.3)	(-76.9, 15.6)
(median)	N=39	N=38*	N=40
% Change from Baseline	-4.76	-37.98	-44.15
LS mean (95% CI)	(-11.74, 2.22)	(44.80, -31.17)	(-50.95, -37.35)
P-value	0.1814	<0.0001	<0.0001

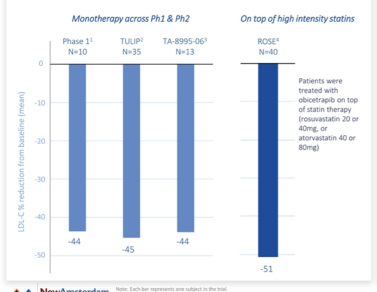


In obicetrapib 5mg arm, 40 patients were randomized. N-value at end-of-treatment decreased to 38 because one patient was missing an LDL value at baseline and a second patient was missing an LDL value at end-of-treatment readir



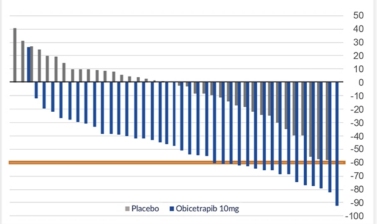
Robust consistency of efficacy profile across multiple studies and within ROSE

LDL-C lowering activity of obicetrapib 10mg across all efficacy studies to-date





- · Almost all patients experienced meaningful LDL-C reductions
- LDL-C reduction of >60% was observed in 40% of obicetrapib 10mg patients





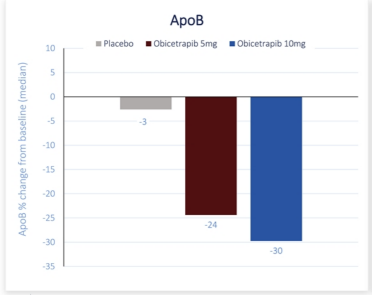
NewAmsterdam Note: Each bar represents one subject in the trial.

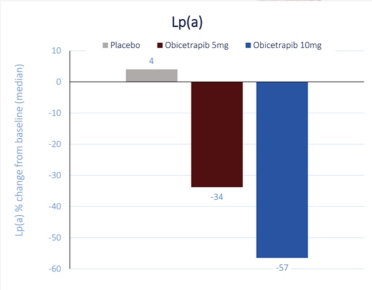
Pharma Source: 1. Ford J, et al. Br J Clin Pharmacol 2014;78:498-508; 2. Hovingh GK, et al. Lancet 2015;386:452-460; 3. Data on file; 4. Nicholis 5J, et al. Nat Med 2022;28:1672-1678



ApoB & Lp(a) percent reduction from baseline in ROSE¹

- ApoB is currently considered the most important biomarker for CVD risk reduction (in addition to non-HDL and LDL-C)
- Elevated Lp(a) is emerging as a strong and independent marker of cardiovascular risk and an exciting new CVD drug target







ource: 1. Nicholls SJ, et al. Nat Med 2022;28:1672-1678

ROPRIFTARY 19



Obicetrapib safety profile in ROSE and across multiple studies: overview of AEs, SAEs and withdrawals

 ${\it Clean safety profile; Similar \, rates \, of \, drug \, discontinuation \, were \, observed \, for \, obicetrapib \, and \, placebo}$

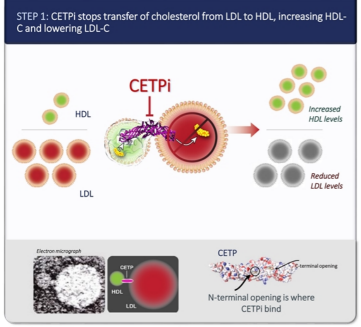
ROSE Safety	Placebo (N=40)	Obicetrapib, 5mg (N=40)	Obicetrapib, 10mg (N=40)
AEs (%)			
AEs, total	19 (47.5)	15 (37.5)	8 (20.0)
AEs, related	4 (10.0)	2 (5.0)	1 (2.5)
AEs, severe	1 (2.5)	0	0
SAEs			
SAEs, total	2 (5.0)	0	0
SAEs, related	0	0	0
Deaths	0	0	0
Withdrawals study / medication			
TEAEs leading to discontinuation of study drug	1 (2.5)	0	0
TESAEs leading to discontinuation of study	0	0	0
TESAEs leading to discontinuation of	0	0	0

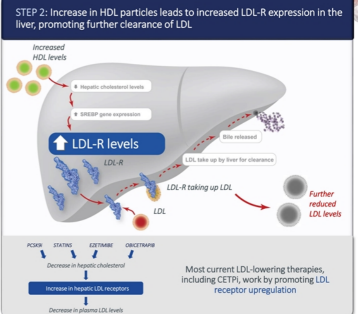
Pooled Safety	Comparator ⁽¹⁾ (N=231)	Obicetrapib Pooled 5mg + 10mg ⁽²⁾ (N=309)
TEAEs (%)		
TEAEs, total	136 (58.9)	173 (55.9)
TEAEs, related	45 (19.5)	49 (15.8)
TEAEs, severe	5 (2.2)	7 (2.3)
TESAEs		
*TESAEs, total	6 (2.6)	4 (1.3)
TESAEs, related	0	0
Deaths	0	0
Withdrawals study / medication		
TEAEs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





CETP inhibition works to lower LDL via a two-step mechanism of action





PROPRIETARY 22

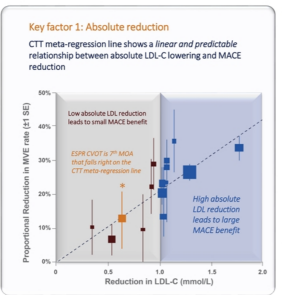


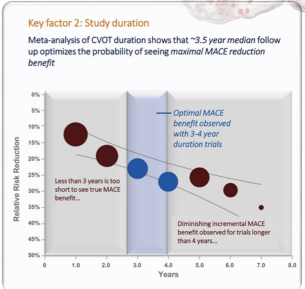
ote: Figures adapted from Meng Zhang, at al., Assessing the mechanisms of cholesteryl ester transfer protein inhibitors, Blochimica et Blophysica Acta (BBA) - Molecular and Cell Biology of Upids, 1862(12), 2017, 1606-1617, and from Lei D, et al.,



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









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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.
Lancet 2005;366-1267-78; Silverman MG, et al. JAMA 2016;27:1289-1297. * ESPR CVOT depiction based on ACC 2023 presentation, drawn as approximation to existing meta



Obicetrapib program designed to overcome limitations of all prior CETP inhibitors

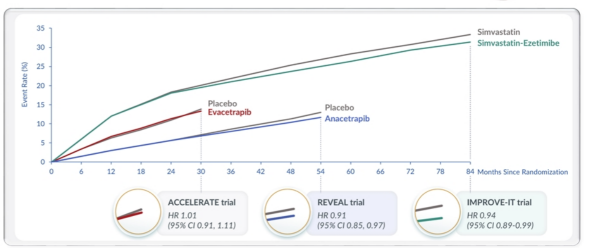




ACCELERATE, REVEAL and IMPROVE-IT demonstrate that CVOT study duration should be long enough to see optimal MACE benefit

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







Curves 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 admission or coronary 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 unstable angina, and in REVEAL as the composite of coronary death, myocardial infarction, or coronary revascularization.

annon 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;377:1217-122



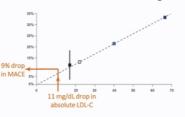
MACE benefits in CVOT of anacetrapib (REVEAL) observed to be exactly as expected, informing NewAmsterdam's CVOT design for objectrapib



At 4.1 years, two important learnings:

Learning 1: Predictable MACE benefit

- 9% drop in MACE is exactly predicted by the CTT metaregression line
- · Indicates CETPi behaves like statins in reducing MACE



Learning 2: Baseline levels were too low

- Baseline 60 mg/dL already below U.S. guideline goals
- Modest drug LDL-lowering potency (17%) resulted in very small absolute reduction (only 11 mg/dL)



At 6.4 years: 20% additional MACE risk reduction

 $An a cetrapib's \ long \ half-life \ causes \ it \ to \ continue \ to \ have \ effects \ in \ patients \ (patients \ remained \ randomized)$

At both time readouts, REVEAL showed statistically significant drop across **all composites** of MACE*



ource: The HPS3/TIMISS-REVEAL Collaborative Group. European Heart Journal (2021) 00, 1–9; The HPS3/TIMISS-REVEAL Collaborative Group. N Engl J Med 2017; 377:1217-1227



REVEAL supports translation from <u>absolute</u> LDL reduction to MACE benefit

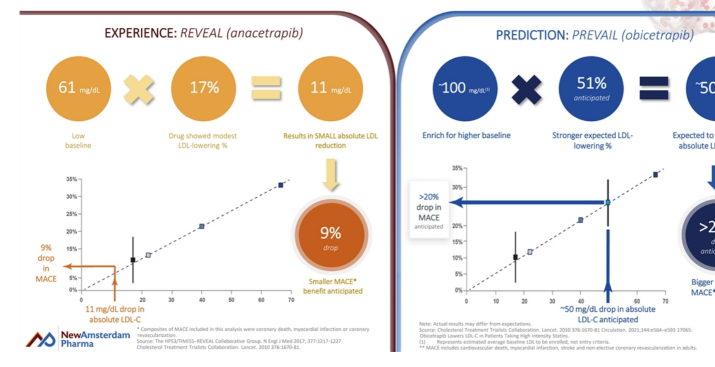
~50 mg/dL

Expected to result in more

absolute LDL reduction

>20%

Bigger expected MACE** benefit





PREVAIL trial design & power calculations leverage lessons learned



· Study design:

- o n = 9000
- o Inclusion: ASCVD patients on maximally tolerated statins with risk enhancers and LDL-C > 70mg/dl
- o Minimum follow up 2.5 years
- Primary endpoint: 4-point MACE
- First secondary: 3-point MACE
- Prespecified endpoints:
 - Conversion of pre-diabetes to diabetes
 - o A1c levels in diabetes patients

· Power assumptions:

- o 4.4% annual event rate
- o 45mg/dl reduction in LDL-C
- o 90% power to achieve a 20% relative risk reduction (p < 0.01)

Differentiation over prior CVOTs

Superior LDL-lowering activity

51% observed in Phase 2b

plus

Targeting higher baseline LDL patients



Higher absolute LDL-c reduction leads to greater MACE benefit

Longer duration of follow up

Median of 48 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)



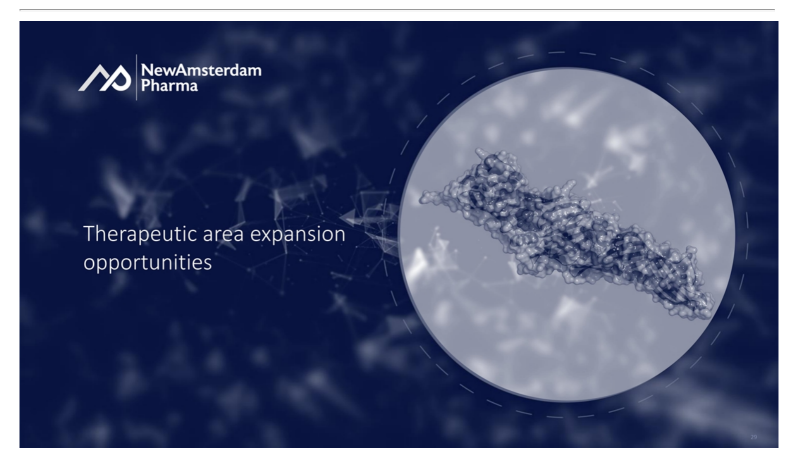
More time + higher patient risk maximizes opportunity for MACE reduction

Differentiated secondary endpoints



Enhanced commercial profile vs. other LDL-lowering agents + potential therapeutic area expansion



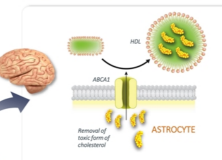




HDL and apoA1 effects potentially offer pipeline expansion opportunities in Alzheimer's , AMD and diabetes

HDL is the "vacuum cleaner" of the body, effluxing free cholesterol out of peripheral tissues to promote healthy cell function &

 NewAmsterdam is exploring potential HDL- and apoA1- raising benefits in other indications such as Alzheimer's disease (AD) and diabetes

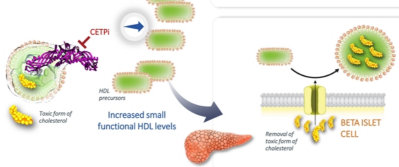


POSITIVE TISSUE EFFECTS IN THE BRAIN

- Increasing HDL and apoA1 is expected to promote healthy tissue survival in the brain²
- Administration of CETPi to APP/CETP knock-in mice observed to promote cholesterol removal from the brain and improve cognition³
- We are testing obicetrapib in Alzheimer's patients in a Phase 2a biomarker study

Obicetrapib observed to increase HDL by >160%

survival1

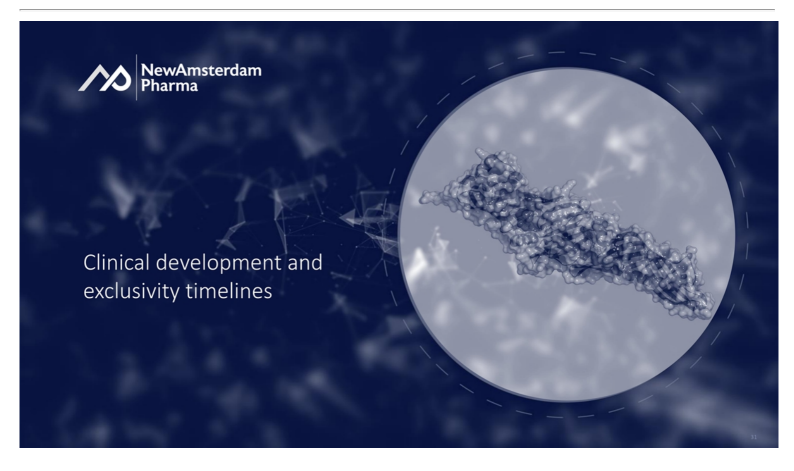


POSITIVE BETA ISLET CELL EFFECTS IN THE PANCREAS

- Increasing HDL is expected to promote beta islet cell survival in the pancreas, potentially improving insulin production^{4,5}
- All four prior CETPi CVOTs demonstrated statistically significant reduction of diabetes risk or reversal of diabetes progression⁶⁻⁹
- We are measuring diabetes progression as an endpoint in PREVAIL and are exploring regulatory paths for a diabetes prevention indication

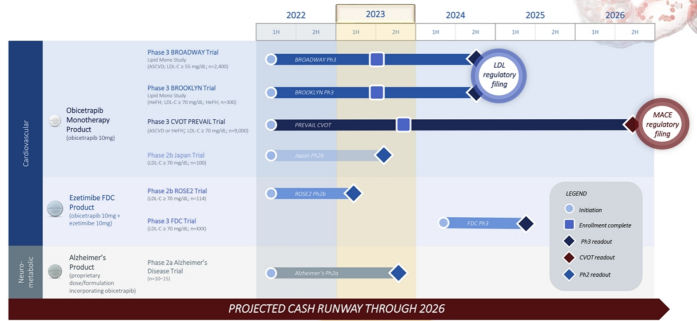


Sources: 1. Tall AR, et al. Nat Rev Immunol 2015;15:104-116; 2. Hottman DA, et al. Neurobiol Dis 2014 2014;72:22-36; 3. Phenix J, et al. Alzheimer's Dement. 2021;17[Suppl. 3]:e051134; 4. Fryirs MA, et al. Arteriosclerosis, Thrombosis, and Vascular Biology, 2010;30:1642–1648; 5. Petremand J, et al. Diabetes 2012;61:100-1111; 6. Barter PJ, et al. N Engl J Med 2007;357:2109-2122; 7. Schwartz GG, et al. N Engl J Med 2012;367:2089-2099; 8. Lincoff AM, et al. N Engl J Med 2017;376:1933-1942; 9. The HPS3/TIMI55–REVEAL Collaborative Group. N Engl J Med 2017; 377:1217-1227





Net proceeds expected to fund obicetrapib development through several value-creating milestones





Numerous catalysts throughout 2023

te: Other than as noted, the pipeline represents trials that are currently ongoing. Projections are subject to inherent limitations. Actual results may differ from expectations. The timing of regulatory submissions is subject to additional discussion to represent the regulators.



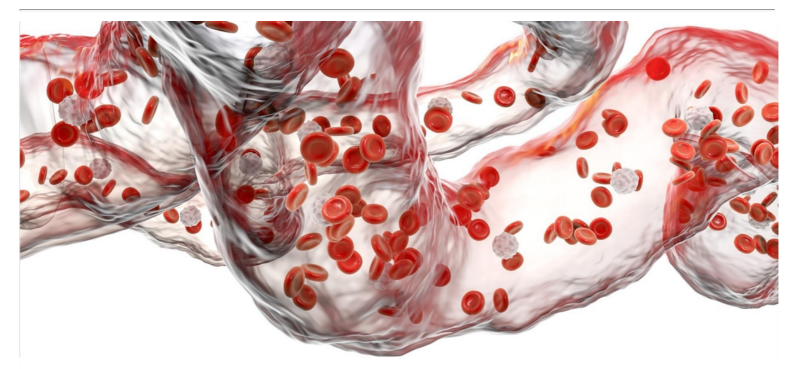
Projected exclusivity timelines in the EU and US

Assumes EU approval on June 1, 2025 and US approval on July 1, 2027

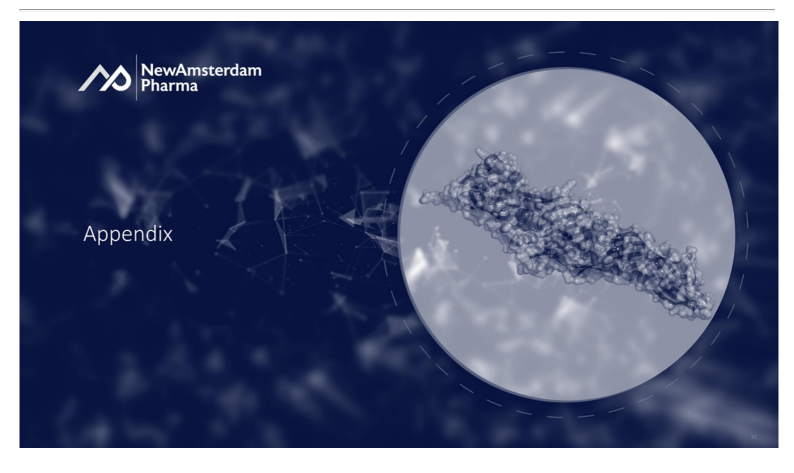




tes: Filled colors = granted patents & dotted lines = pending patents; one patent only to be selected for SPC/PTE; an earlier US approval leads to earlier regulatory expiry & shorter PTE; *including pediatric extension 6m; ** will be pending once a "T sendication; in 18 disk) and the second patents will offer the second patents and the second patents are second patents.









Obicetrapib development history

Preclinical—Phase 1b (2008-2012)







Manufacturing scale-up (2015-2016)

"Paused" clinical development to scale-up manufacturing to support Phase 3/commercial supply
Discovered manufacturing process that led to current \$3/month projected peak COGS
In 2015/2016, a "perfect storm" of external factors (incl. Merck and Lilly CETPi CVOT read-outs) led to a re-prioritization within Amgen's CVD portfolio, and obicetrapib was de-prioritized to 4th-in-line for a CVOT

AMGEN

• Dr. Kastelein & Forbion spun obicetrapib from Amgen to NAMS in 2020, resulting in full transfer of IP ownership & full global rights

Amgen and MTPC are shareholders of NAMS, hold no other



Phase 2b through CVOT and commercialization (2020 through present)





Menarini partnership overview

Partnership with Menarini brings in significant non-dilutive capital and could enable NewAmsterdam to simultaneously launch obicetrapib in different markets with the ideal partner to optimize the commercial opportunity in the EU

RIGHT PARTNER

Leading presence in cardiovascular disease:

• 18 marketed products in cardiometabolic diseases

• #1 share of voice among cardiologists, internists

· >2,500 sales reps and >280 specialist field force

• Successfully secured access for >500 products

Strong partnering track record, with 60+ partnerships



RIGHT TIME



RIGHT DEAL

MENARINI IS A LEADING EUROPEAN PHARMACEUTICAL ~3 YEARS FROM LAUNCH OPTIMIZES EUROPEAN COMPANY DEVELOPMENT WITH AN EXCELLENT EUROPEAN

· Pricing and access in Europe is critical for

- obicetrapib's success
- · Local expertise is needed to jumpstart P&MA strategy, including evidence generation and proactive HTA engagement
- Strong relationships with EU KOLs will support obicetrapib market entry while more effectively disseminating the obicetrapib value story

MENARINI TO RECEIVE EXCLUSIVE RIGHTS TO OBICETRAPIB MONOTHERAPY AND EZETIMIBE FDC FOR CVD IN EUROPE

NewAmsterdam retains all other global rights and is eligible for significant non-dilutive financial terms:

- €142.5mm committed capital (Consisting of €115mm upfront + €27.5mm committed R&D funding)
- Up to €863mm payable upon achievement of certain clinical, regulatory and commercial milestones
- · Tiered royalties from teens to mid-twenties



NewAmsterdam Pharma

and GPs in EU5

Deep commercial expertise:

members in Europe

· 540 launches in 50 countries

spanning small biotech to large pharma

UP TO €1,005.5mm OF TOTAL CASH CONSIDERATION



LDL-C, ApoB and Lp(a) are causal in the development of atherosclerosis and heart disease

ApoB is a molecule that envelopes LDL in a 1:1 ratio

LDL-C



ApoB: Each LDL-C particle contains one ApoB molecule

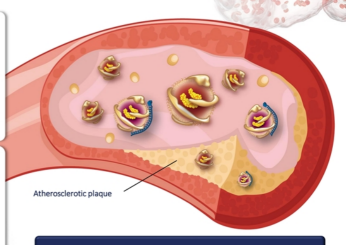
- ApoB-containing particles can become trapped in the arterial wall
- High ApoB levels lead to atherosclerotic plaque formation and buildup over time



Each Lp(a) particle has a LDL particle that contains an ApoB wrapped with an Apo(a) molecule



- Genetic mutations can lead to higher Lp(a) levels in certain individuals
- These individuals are at a higher risk of developing arterial plaques and experiencing adverse cardiovascular events



Reducing the number of ApoB and Lp(a) particles in circulation is critical to halting plaque build-up and reducing CV risk





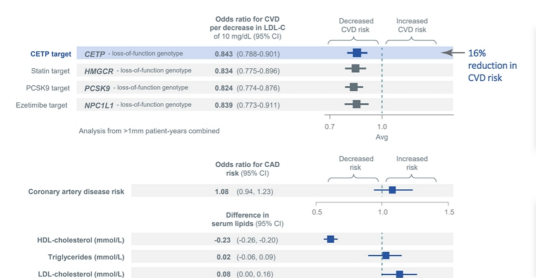
Genetic support that CETPi drives CVD benefit through LDL reduction

Analysis of >1mm patient-years' shows loss-of-function equivalent to targets of other LDL-lowering drugs

-0.3

0.0

0.3



- A 16% reduction in CVD risk is observed for every 10 mg/dL decrease in LDL levels
- This is ~equivalent to the effect seen in loss-of-function genotypes for statins, PCSK9 inhibitors and ezetimibe

More CETP = more CAD risk, less HDL, more LDL and more ApoB

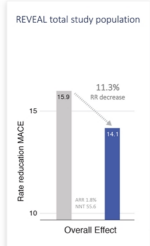


Sources: Ference BA et al. JAMA 2017; Blauw et al; Genome-Wide Association Study on Circulating CEI



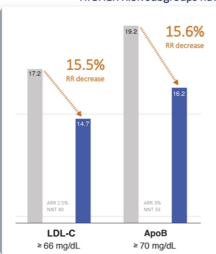
REVEAL long-term follow-up identified risk enhancers important for PREVAIL

HIGHER RISK subgroups have higher event rates & larger treatment effects





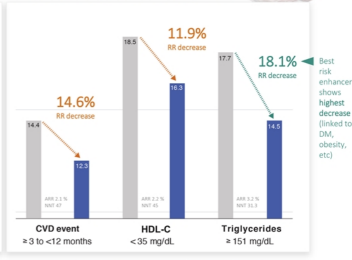
NewAmsterdam Pharma



PREVAIL study inclusion criteria requires high baseline LDL-C (also translates to high ApoB)

Inclusion criteria: LDL-C ≥70 mg/dL

Source: The HPS3/TIMI55-REVEAL Collaborative Group. European Heart Journal (2021) 00, 1–9



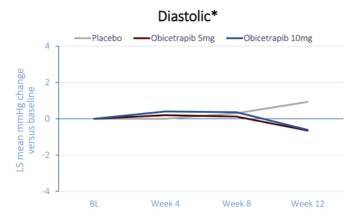
PREVAIL study <u>risk enhancers</u> will increase high-risk patient populations

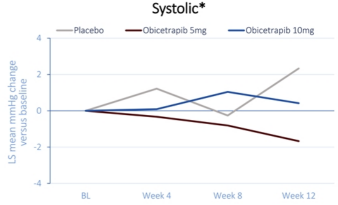
Inclusion criteria: Lp(a) >75 mg/dL, HDL-C <40 mg/dL, triglycerides >150 mg/dL



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

- · In the TULIP study, obicetrapib did not show any effect on blood pressure, aldosterone and electrolytes
- A dedicated meta-analysis of the obicetrapib ROSE, TULIP and OCEAN 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



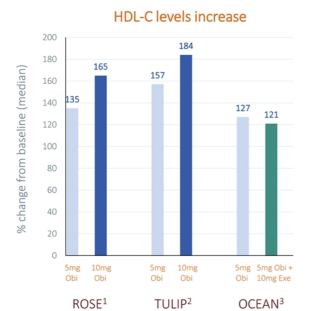


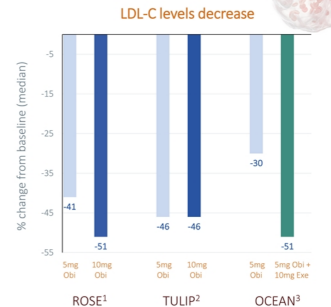


Sources: Circulation 2021;144:e564–e593 17065: Obicetrapib Lowers LDL-C in Patients Taking High Intensity Statins: Results From Rose Clinical Tria



Consistent LDL-C lowering and HDL-C increase observed across three Phase 2 studies of obicetrapib







Source: 1. Nicholls SJ, et al. Nat Med 2022;28:1672-1678; 2. Hovingh GK, et al. Lancet 2015;386:452-460; 3. Data on File



OCEAN: Phase 2b evaluating 5mg obicetrapib in combination with 10mg ezetimibe

Objective To evaluate the effect of obicetrapib in combination with ezetimibe compared to placebo on LDL-C

Inclusion criteria

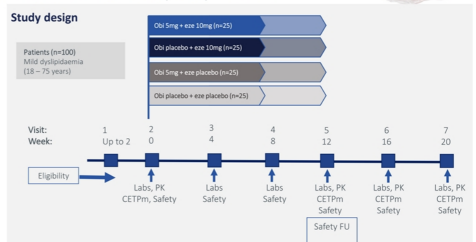
- · Mild dyslipidaemia
- Fasting LDL-C levels >2.5 mmol/L and <4.5 mmol/L

Exclusion criteria

- Currently taking any lipid-lowering therapy Any clinical manifestation of atherosclerotic CVD
- Diagnosis of type 1 or type 2 diabetes mellitus
- Uncontrolled hypertension

Primary efficacy endpoint

• Percent change from baseline in LDL-C for the combination therapy group compared to the placebo group

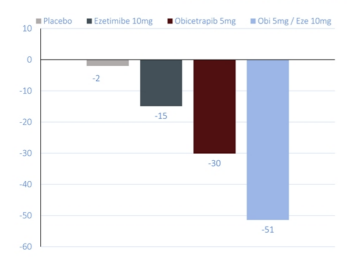






In OCEAN, obicetrapib was observed to have LDL-lowering efficacy that is additive with ezetimibe

LDL-C in mg/dL by PUC & Percent change from baseline



Median (min, max) LDL-C levels (mg/dL) at baseline and EOT

Time	Placebo	Ezetimibe 10mg	Obicetrapib 5mg	Obi 5 + Eze 10mg
Baseline Median	136.0	127.0	121.0	123.0
	(101, 177)	(76, 189)	(82, 153)	(89, 186)
	(N=24)	(N=27)	(N=27)	(N=27)
EoT Median	138.0	105.0	86.5	63.5
	(88, 193)	(66, 142)	(38, 137)	(34, 133)
	(N=25)	(N=24)	(N=26)	(N=24)
% change from BL median	-2.0	-14.90	-30.10	-51.40
	(-24.5, 35.9)	(-46.8, 46.9)	(-56.7, 19.1)	(-69.6, 8.1)
	(N=27)	(N=25)	(N=25)	(N=24)
% change from BL	1.40	-12.86	-30.70	-40.95
LS Mean (95% CI)	(-6.03, 8.84)	(-20.29, -5.42)	(-38.21, -23.19)	(-48.73, -33.16)
p-value	0.7116	0.0007	<0.0001	<0.0001

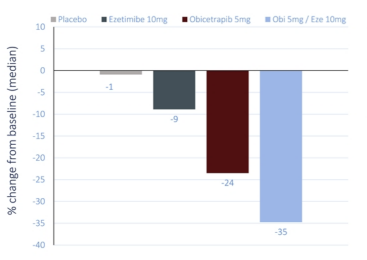






OCEAN: ApoB in mg/dL & percent change from baseline

Median (min, max) ApoB levels (mg/dL) at baseline and EOT



Time	Placebo	Ezetimibe 10mg	Obicetrapib 5mg	Obi 5mg+ Eze 10mg
	105.5	103.0	102.0	105.0
Baseline Median	(74, 141)	(79, 133)	(74, 124)	(77, 158)
	(N=28)	(N=28)	(N=28)	(N=27)
	107.0	94.0	75.0	73.0
EoT Median	(69, 153)	(59, 137)	(45, 103)	(49, 105)
	(N=27)	(N=25)	(N=26)	(N=24)
% change	-0.9	-8.9	-23.5	-34.8
from BL	(-19.8, 25.4)	(-45.4, 32.3)	(-39.3, 21.2)	(-53.0, 8.9)
Median	(N=27)	(N=25)	(N=26)	(N=24)



Data on File



OCEAN safety: TEAEs, TESAEs and withdrawals overview

Obicetrapib 5mg well tolerated compared to placebo

				0.0
	Placebo	Ezetimibe 10mg	Obicetrapib 5mg	Obi 5mg / Eze 10mg
TEAEs (%)				
TEAEs, total	6 (21.4)	8 (28.6)	4 (14.3)	9 (33.3)
TEAEs, related	4 (14.3)	3 (10.7)	1 (3.6)	3 (11.1)
TEAEs, severe	0	1 (3.6)	0	0
TESAEs				
TESAEs, total	0	2 (7.1)	0	0
TESAEs, related	0	0	0	0
Deaths	0	0	0	0
Withdrawal's study / medication				
TEAEs leading to discon of study drug	1 (3.6)	1 (3.6)	0	2 (7.4)
TESAEs leading to discon of study	0	1 (3.6)	0	0



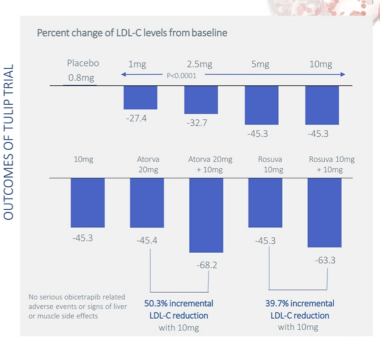
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TULIP Phase 2a trial results

TULIP clinical trial design

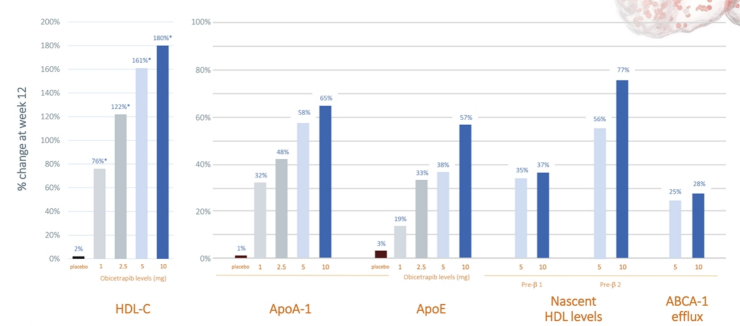
- Randomized double-blind placebo-controlled trial, once daily dosing
- Endpoints:
 - Percent change in LDL-C at week 12 compared to baseline
 - Safety
 - Tolerability
- Duration: 12 weeks treatment
- Patients:
 - 364 patients (aged 18-75 years)
 - Fasting LDL-C levels: 2.5 4.5 mmol/L







TULIP: HDL-C, ApoA-1, ApoE and nascent HDL levels increases observed







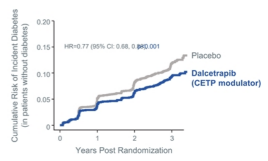
Obicetrapib bears significant potential to also beneficially impact diabetes risk

CETP validated both genetically and clinically to reduce/reverse diabetes risk

- While statins increase diabetes risk by 11-15%1, CETP inhibition is clinically demonstrated to reverse existing diabetes, decrease new-onset diabetes, and improve glucose control
- Genetic CETP deficiency lowers glucose levels²



Dalcetrapib reduces the risk of diabetes by 26% and increases the conversion to non-diabetes in high risk CV patients



Sources: ¹ Sattar. Lancet. 2010 Feb 27; ² Brian ference, data on file; ³ REVEAL trial. N Engl J Med. 2017 Sep 28; ⁴ Barter. Circulation. 2011 Aug 2



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