

ORION-1 Primary efficacy & safety outcomes

LDL-C reduction from 6 to 9 months following single or second injections of inclisiran, a novel siRNA compound

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On behalf of the ORION-1 investigators

Background Major progress is being made in ASCVD



PCSK9 inhibition is now a validated target for reducing LDL-C and ASCVD¹

PCSK9 mAb therapy requires 12-26 injections per year

Adherence data with PCSK9 mAbs show no substantial improvement over statins²

Poor adherence and LDL-C variability are associated with poor outcomes³

These limitations are most relevant in high risk patients with high LDL-C

- Sabatine M et al NEJM 2017
- Hines D et al ACC 2017 abstract #1203-313
- Bangalore S et al JACC 2015;65:1539-48











Rationale and objective Inclisiran: a novel agent to address unmet needs



Harnessing RNAi offers an alternative treatment for PCSK9 and LDL-C1

- Inclisiran, a synthetic siRNA molecule, inhibits PCSK9 synthesis in the liver²
- In Phase I, 300 mg inclisiran lowered LDL-C 50-60% for 84 days (n=69)³

Objective of ORION-1

Evaluate optimal dosing regimens in patients with elevated LDL-C and high CV risk

- Wittrup A & Lieberman J Nature Rev Gen 2015;16: 543-52
- 2. Fitzgerald K et al. Lancet 2013;9911:60-8
- 3. Fitzgerald K et al. N Engl J Med 2017;376 (1):41-51





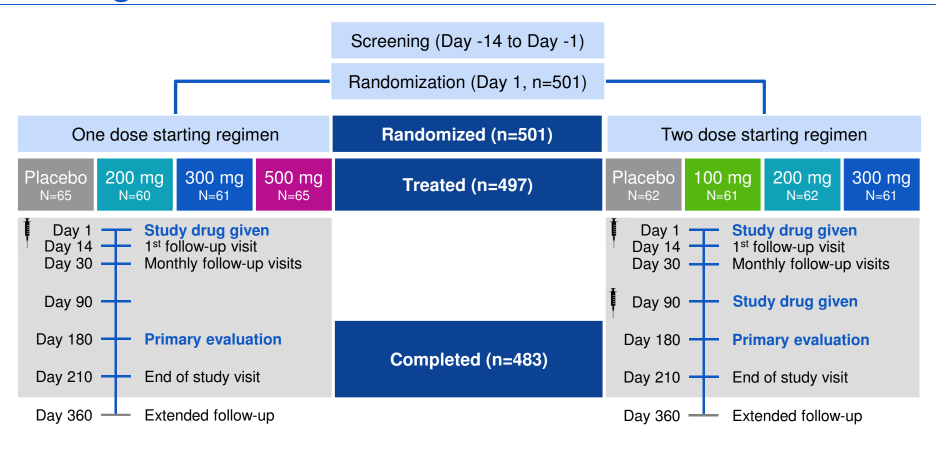






Methods Trial design











Patients High-risk CV patients, balanced by randomization



		One dose starting regimen		Two dose starting regimen	
		Placebo	Inclisiran	Placebo	Inclisiran
		N=65	N=186	N=62	N=184
Age	Mean years	62	63	63	64
Male sex	%	64.6	67.7	53.2	66.3
Prior ASCVD	%	69.2	67.9	74.2	68.3
Statin Rx	%	70.3	74.4	77.0	70.2
LDL-C	Mean mg/dL	128.5	125.9	125.2	133.0
Non-HDL-C	Mean mg/dL	157.8	156.5	157.1	165.6
Аро-В	Mean mg/dL	102.4	103.2	104.6	107.7
Lipoprotein(a)	Median nmol/L	27.0	34.0	50.5	40.0
PCSK9	Mean ng/mL	404.7	428.7	431.3	416.2







Safety No safety concerns: Adverse events similar to placebo



Safety population	One dose star	ting regimen	Two dose starting regimen	
	Placebo	Inclisiran	Placebo	Inclisiran
	N=65	N=186	N=62	N=184
	n (%)	n (%)	n (%)	n (%)
Any TEAE	46 (70.8)	140 (75.3)	50 (80.6)	142 (77.2)
Serious	3 (4.6)	17 (9.1)	6 (9.7)	24 (13.0)
Severe	2 (3.1)	11 (5.9)	7 (11.3)	19 (10.3)
Related	12 (18.5)	39 (21.0)	18 (29.0)	51 (27.7)
Injection site reaction	0	7 (3.8)	0	12 (6.5)

TEAEs (treatment emergent adverse events) - similar incidence placebo vs inclisiran:

One dose starting regimen: Nasopharyngitis, myalgia, back pain, cough, arthralgia, headache Two dose starting regimen: Myalgia, headache, diarrhea, nasopharyngitis, arthralgia, back pain









Safety

No safety concerns: Liver and muscle enzymes



No LFT elevations related to drug

- Transient transaminase increases no differences between randomized groups
 - -0.8% placebo
 - 0.8% inclisiran

No difference in incidence of myalgias or CPK enzyme elevation

One clinically relevant case of myonecrosis on placebo

No deaths related to drug administration

- Two deaths¹ > 100 days beyond injection and clearly related to underlying disease
 - 1: Patient A: History of CHD, MI and PCI died of a fatal MI on Day 104 of the study. (500mg x1 dose)

Patient B: Developed complications of aortic aneurysm surgery including an aorto-esophageal fistula requiring esophagectomy, leading to infection of the prosthesis, sepsis, and stroke, culminating in death on Day 198 of the study. Patient also had AF, chronic renal failure, emphysema, HT and obesity. (200mg x2 doses)











Safety

No safety concerns: Other relevant parameters



No thrombocytopenia

No neuropathy

No immunogenicity (no anti-drug antibodies)

No pro-inflammatory symptoms or elevated markers



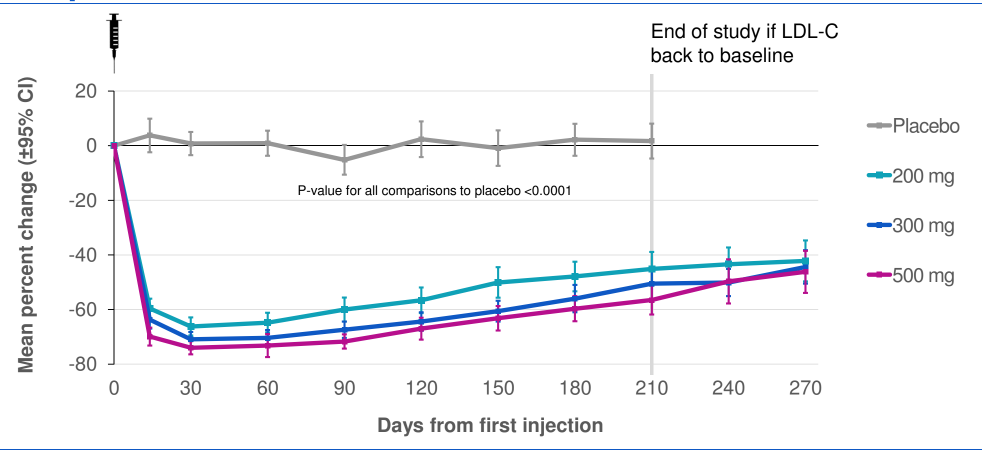






Efficacy: One dose starting regimen Clamped PCSK9 knockdown







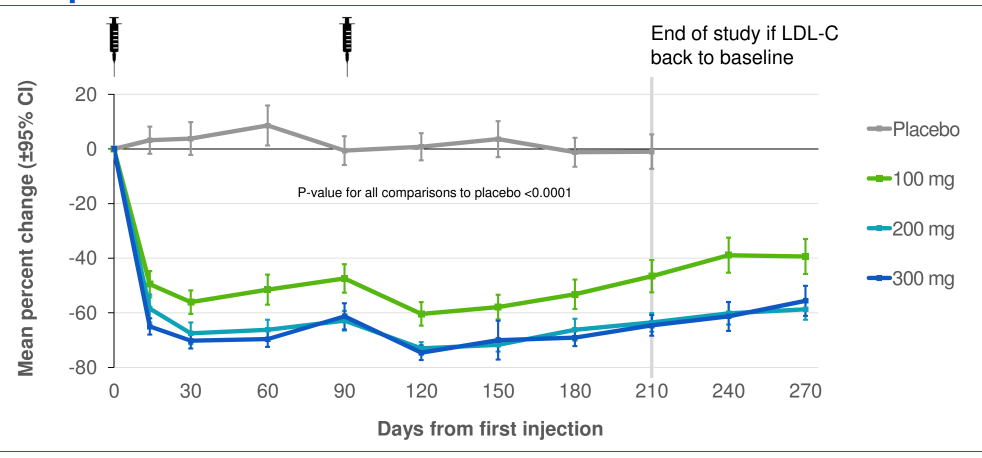






Efficacy: Two dose starting regimen Clamped PCSK9 knockdown







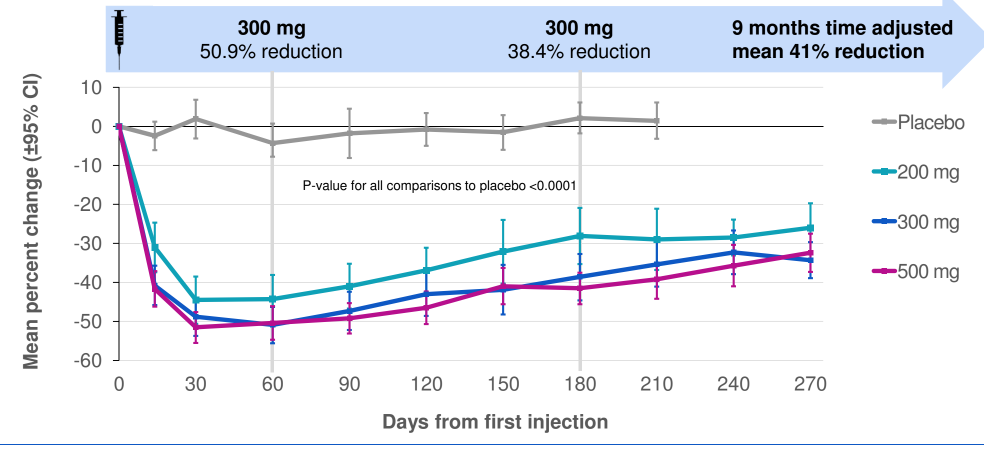






Efficacy: One dose starting regimen Robust, sustained LDL-C reductions – 300 mg optimal





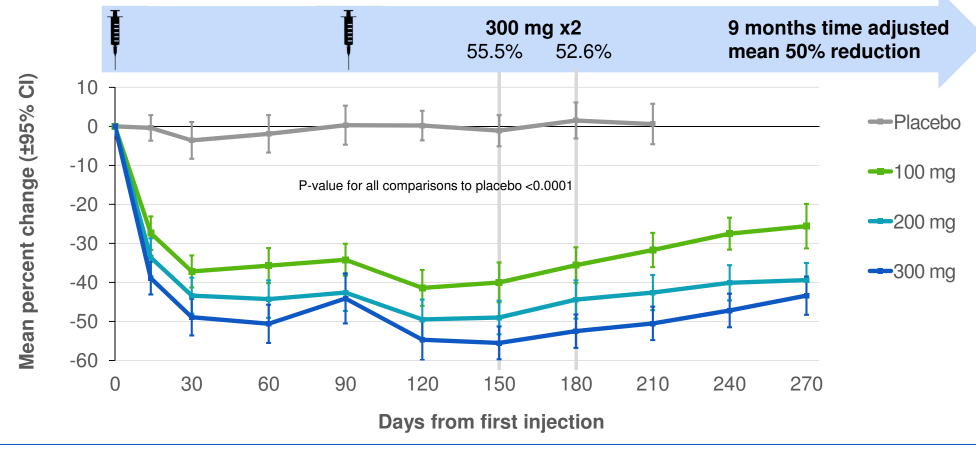








Efficacy: Two dose starting regimen Robust, sustained LDL-C reductions – optimal start regimen







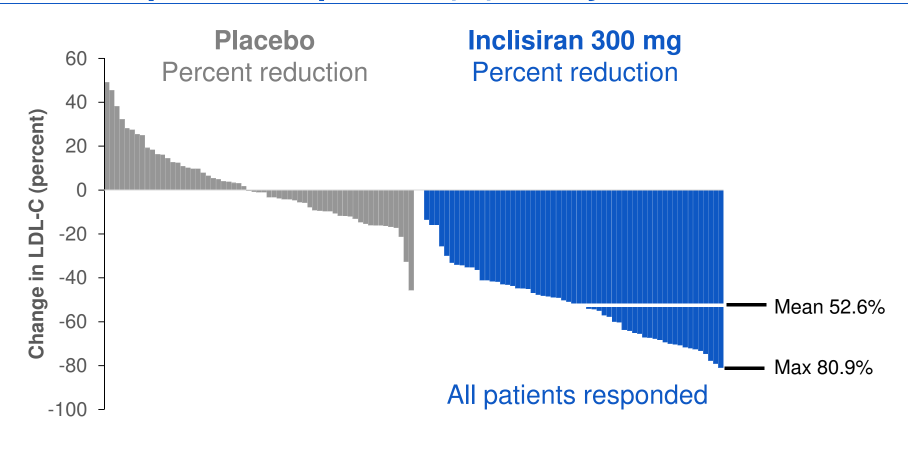






Efficacy: Two dose starting regimen Individual patient responses (%) at day 180







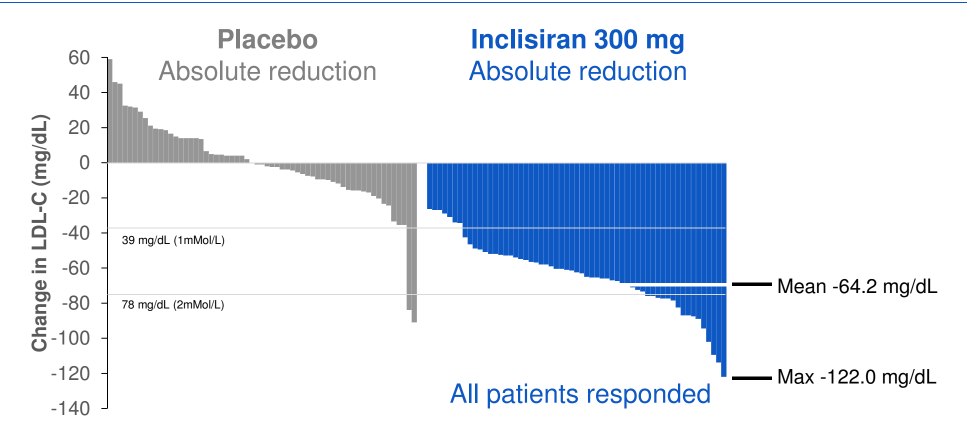






Efficacy: Two dose starting regimen Individual patient responses (mg/dL) at day 180















Conclusions Two 300 mg starting dose regimen for inclisiran selected



No safety concerns

Optimal dosage 300 mg given twice as starting regimen then Q6 monthly

- All patients responded with significant LDL-C lowering
- At 6 months, mean LDL-C↓ of 52.6% (64 mg/dL), and up to 81% (122 mg/dL)

Unique attributes of inclisiran address multiple unmet needs

- LDL-C variability within individuals is practically eliminated
- Injection burden reduced substantially
- Sustained effect between infrequent injections
- Opportunity to improve patient adherence









Implications Inclisiran will move into phase III trials



In an ORION-1-like population, inclisiran 300 mg delivers sustained LDL-C lowering of 60-65 mg/dL

In a CVOT, this is likely to confer substantial reductions in MACE

ORION-4 will study CV outcomes with inclisiran in high risk primary and secondary prevention patients with average LDL-C ~130 mg/dL









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ORIGINAL ARTICLE

Inclisiran in Patients at High Cardiovascular Risk with Elevated LDL Cholesterol

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