

Metabolic dysfunction-associated steatotic liver disease (MASLD) encompasses a heterogenous series of disorders ranging from fatty liver to more severe metabolic dysfunction-associated steatohepatitis (MASH). Thyroid hormone receptor beta (THRβ) is a clinically validated target for the treatment of MASH, with THRβ agonists able to selectively reduce fat deposits in the liver and potentially prevent the downstream consequences of MASLD (inflammation, fibrosis, cirrhosis, etc.). ALG-055009 (Fig. 1) is a THRB agonist that has demonstrated significant reductions in liver fat (placebo-adjusted median relative reductions up to 46.2%) and atherogenic lipids in patients with presumed MASH and stage 1-3 liver fibrosis. Here, we present the effects of ALG-055009 in a diet-induced obese (DIO) mouse model and human liver cells.

Phase 2a HERALD study highlights¹

(see 2025 EASL presentation SAT-451 & posters SAT -430, -450, -451) – Primary endpoint achieved with robust reductions in liver fat content at Week 12 • 70% of patients achieved >30% at 0.7 mg dose - Significant reductions in atherogenic lipids (e.g. LDL-C, lipoprotein (a) & apolipoprotein b) – Dose-dependent increases in SHBG (marker of THR-β activation in liver) – Well-tolerated, with rates of GI-related AEs similar to placebo



METHODS

High Fat Diet-Induced Obese Mouse Study: C57BL/6J mice were fed with a high fat diet (HFD; D12492) for 14 weeks, followed by drug treatment for 4 weeks. ALG-055009 treatment groups included two oral QD dose levels of 0.5 and 1.5 mg/kg, and four oral BID dose levels ranging from 0.075 to 0.35 mg/kg/dose. Pharmacodynamic endpoints included total and low-density lipoprotein (LDL) cholesterol. Liver gene expression was monitored by RT-qPCR. Previously reported reductions in serum lipid levels are shown below (Table 1).

Table	1. High	Fat Diet	-Induced	Obese	Mouse	Study
		1				

Group	Diet	Treatment	Dose (mg/kg, PO)	Dose Frequency	TC (mean % change)	LDL-C (mean % change)
1	ND	Vahiela	_	BID	-28.2****	-34.4**
2	HFD	venicie			0.0	0.0
3	HFD	ALG-055009	0.5	QD	-25.1****	-37.5***
4			1.5		-37.8****	-40.6***
5			0.075	BID	-9.6	-9.4
6			0.15		-17.2**	-34.4**
7			0.25		-34.4****	-59.4****
8			0.35		-44.3****	-56.3****

ND =normal chow diet (D12450J); HFD =high fat diet (D12492); PO =oral dosing; BID =twice daily; QD =once daily; n =6 animals per group; TC =total cholesterol; LDL-C =low-density lipoprotein cholesterol; statistical analysis: ordinary one-way ANOVA with Dunnet's multiple comparisons test (compared to HFD-Vehicle group at 28 days post-dose); ** =p-value <0.01; *** =p-value <0.001; **** =p-value <0.0001

In Vitro Gene Expression Assays: Huh-7 cells were cultured in media supplemented with 10% charcoalstripped FBS and treated with vehicle or increasing concentrations of ALG-055009 or MGL-3196 for 24 hours. RNA was extracted and the resulting cDNA was used in RT-qPCR. Primary human hepatocytes (PHH) were plated, serum-starved for 24 hours, and then treated with vehicle, ALG-055009, or MGL-3196 for 24 hours at the indicated doses. RNA was extracted and was either used for cDNA library preparation and subsequent RNA-Seq or cDNA was prepared for use in downstream RT-qPCR analysis.

REFERENCES

- 1) NCT06342947
- 2) doi: 10.1016/j.metabol.2019.153994
- 3) doi: 10.1186/s11658-024-00675-6
- 4) doi: 10.1038/s41598-017-11212-1
- 5) doi: 10.1530/ETJ-22-0211
- 6) doi: 10.1152/ajpendo.90736.2008
- 7) doi: 10.1016/j.atherosclerosis.2022.04.006
- 8) doi: 10.1016/j.jcmgh.2019.10.010
- 9) doi: 10.1194/jlr.M700378-JLR200 10) doi: 10.1016/j.jhep.2006.02.011
- 11) doi: 10.3892/ijmm.2020.4479 12) doi: 10.1016/j.livres.2022.09.003
- 13) doi: 10.1016/j.jcmgh.2022.03.011
- 14) doi: 10.3390/ijms21062061 15) doi: 10.1038/srep45049

ALG-055009, a potent and selective THR^β agonist for the treatment of MASH, induces pro-metabolic and anti-fibrotic gene expression in the liver of DIO mice ALIGOS

P. Althoff¹, J. Song¹, L. Adame¹, T. Lin², K. Gupta¹, K. Vandyck², D. McGowan², S. Stevens¹, A. Stoycheva¹, L.M. Blatt¹, L. Beigelman¹, J.A. Symons¹, P. Raboisson², J. Deval¹, and Xuan (Susan) G. Luong^{1²} ¹Aligos Therapeutics, Inc., South San Francisco, CA; ²Aligos Belgium BV, Leuven, Belgium, *Corresponding author: xluong@aligos.com



FGF21 (fibroblast growth factor 21): liver glucose and lipid metabolism²



Enpp2 (autotaxin): stellate cell activation and liver fibrosis³



Dio1 (deiodinase, iodothyronine, type I): thyroid hormone activation and lipid metabolism⁵

Figure 5a. Values reported indicate fold-change relative to Vehicle group () at 24 hours post-dose; statistical analysis: nonlinear regression fit with variable slope

ABCD2 (ATP binding cassette subfamily D member 2): fatty acid transport⁶ LPA (lipoprotein (a)): cholesterol transport⁷



reflecting its superior potency reported in preclinical rodent models and in patients in the clinic. Financial disclosure: all authors are current employees of Aligos Therapeutics Inc.