# Characterization of dietary intervention in the non-obese CDAA-HFD mouse model of advanced NASH with progressive fibrosis

### Authors

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### BACKGROUND & AIM

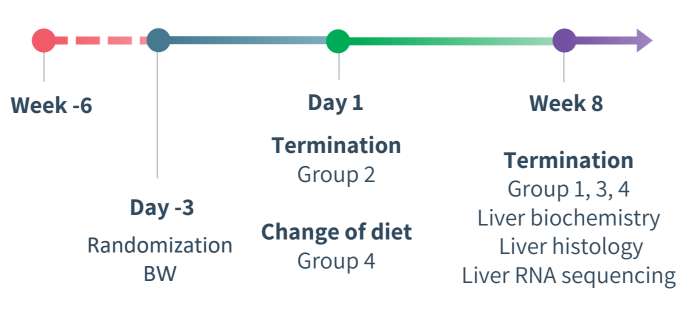
Current standard of care for non-alcoholic steatohepatitis (NASH) involves lifestyle modification, notably dietary intervention, aiming to promote regression or resolution of NASH and liver fibrosis.

We have recently characterized dietary intervention (chow reversal) in the translational GAN diet-induced obese (DIO) mouse model of fibrosing NASH (Møllerhøj et al. Clin Transl Sci, 2022). The present study aimed to evaluate chow-reversal in the non-obese cholinedeficient L-amino-acid defined high-fat diet (CDAA-HFD) mouse model of advanced NASH with progressive fibrosis.

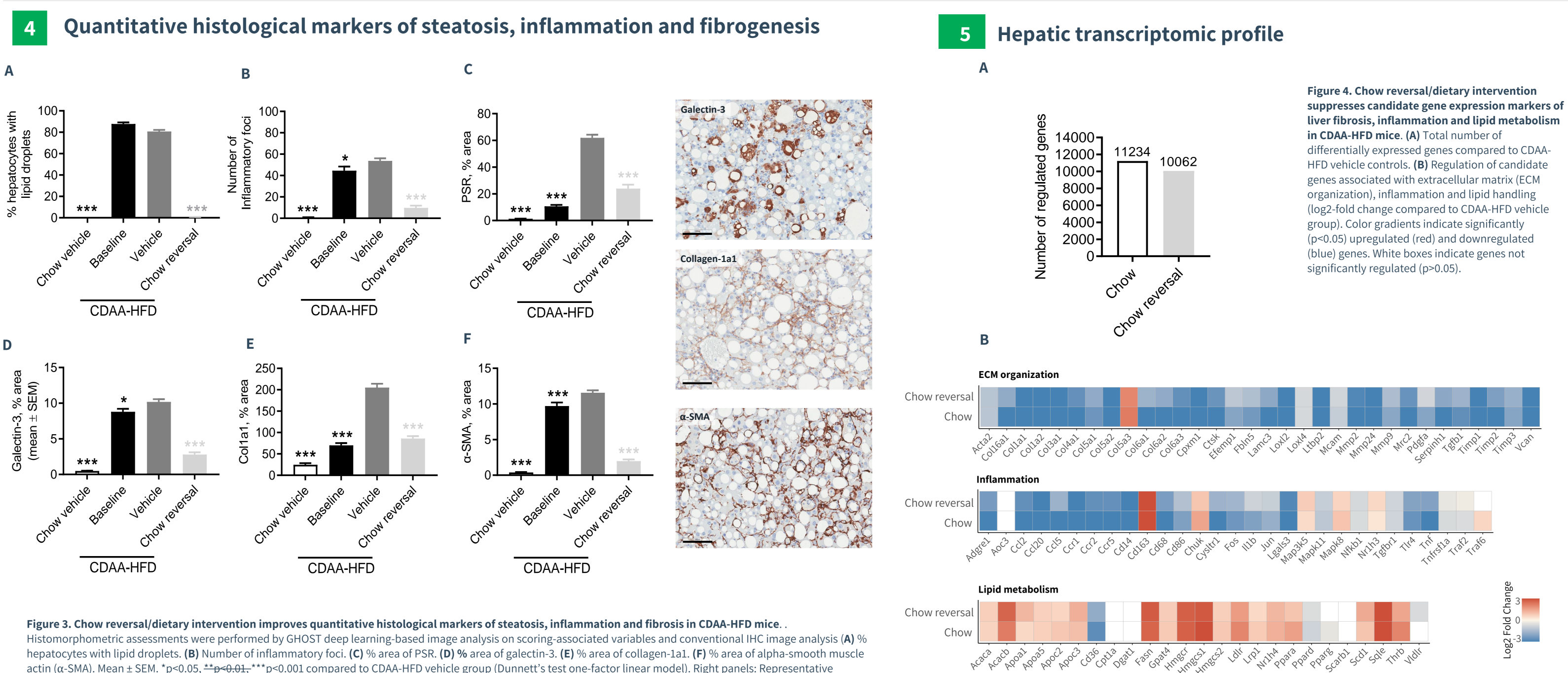
## METHODS

C57BL/6JRj mice were fed chow or choline-deficient high-fat diet (CDAA-HFD, 45 kcal% fat, 0.1% methionine, 1% cholesterol, 28 kcal% fructose) for 6 weeks before treatment start (i.e. after induction of fibrosis). Prior to treatment, animals were randomized into treatment groups based on body weight. A baseline group (n=12) was terminated at study start. Dietary intervention was performed by switching from CDAA-HFD to chow feeding on treatment day 1 (n=12). Chow-fed mice (n=8) served as normal controls. Terminal endpoints included plasma and liver biochemistry, NAFLD Activity Score (NAS), fibrosis stage quantitative liver histology and transcriptome signatures.





Group no.	Group	Name	Number of animals
1	Chow	Chow	8
2	Baseline CDAA-HFD	Baseline CDAA-HFD	12
3	Vehicle CDAA-HFD	Vehicle CDAA-HFD	12
4	Chow reversal	Chow reversal	12



actin (α-SMA). Mean ± SEM. \*p<0.05, \*\*p<0.01, \*\*\*p<0.001 compared to CDAA-HFD vehicle group (Dunnett's test one-factor linear model). Right panels: Representative galectin-3, collagen 1a1 and  $\alpha$ -SMA photomicrographs (scale bar, 100  $\mu$ m).

# 1 Study outline

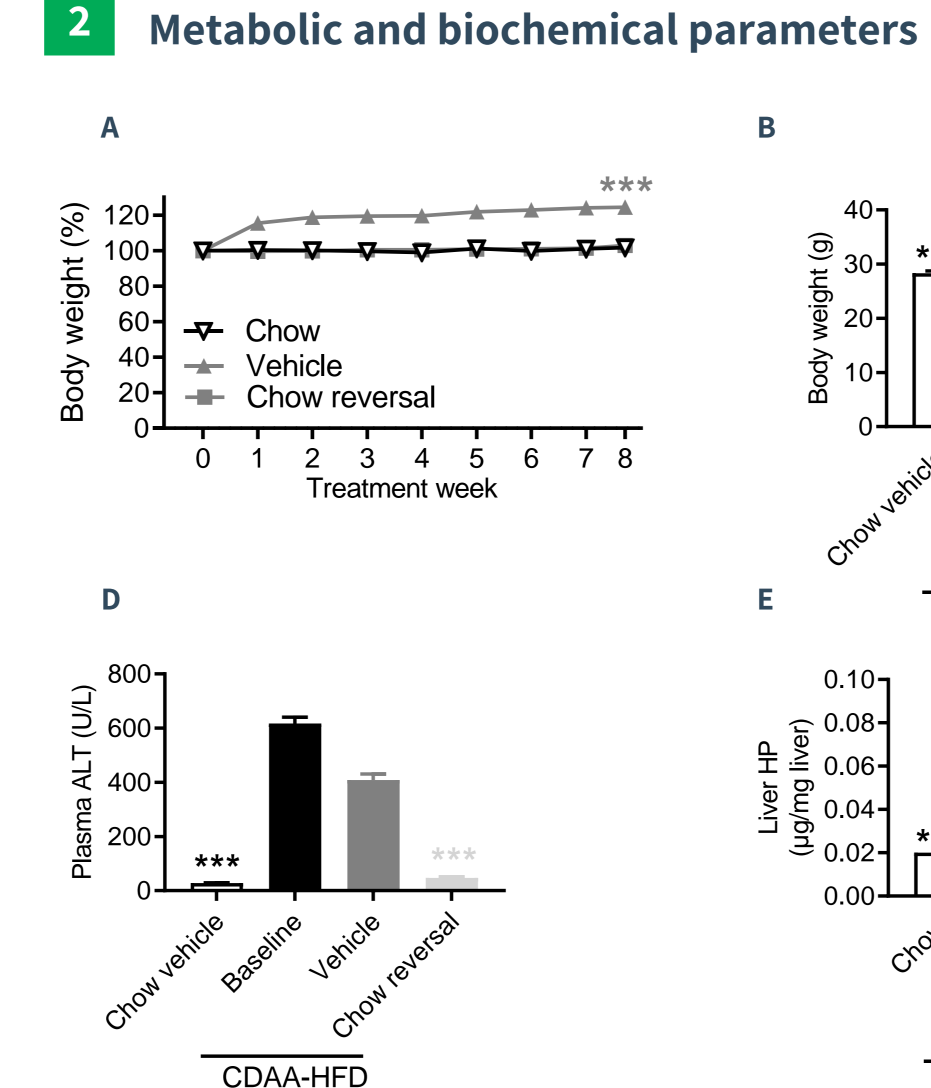
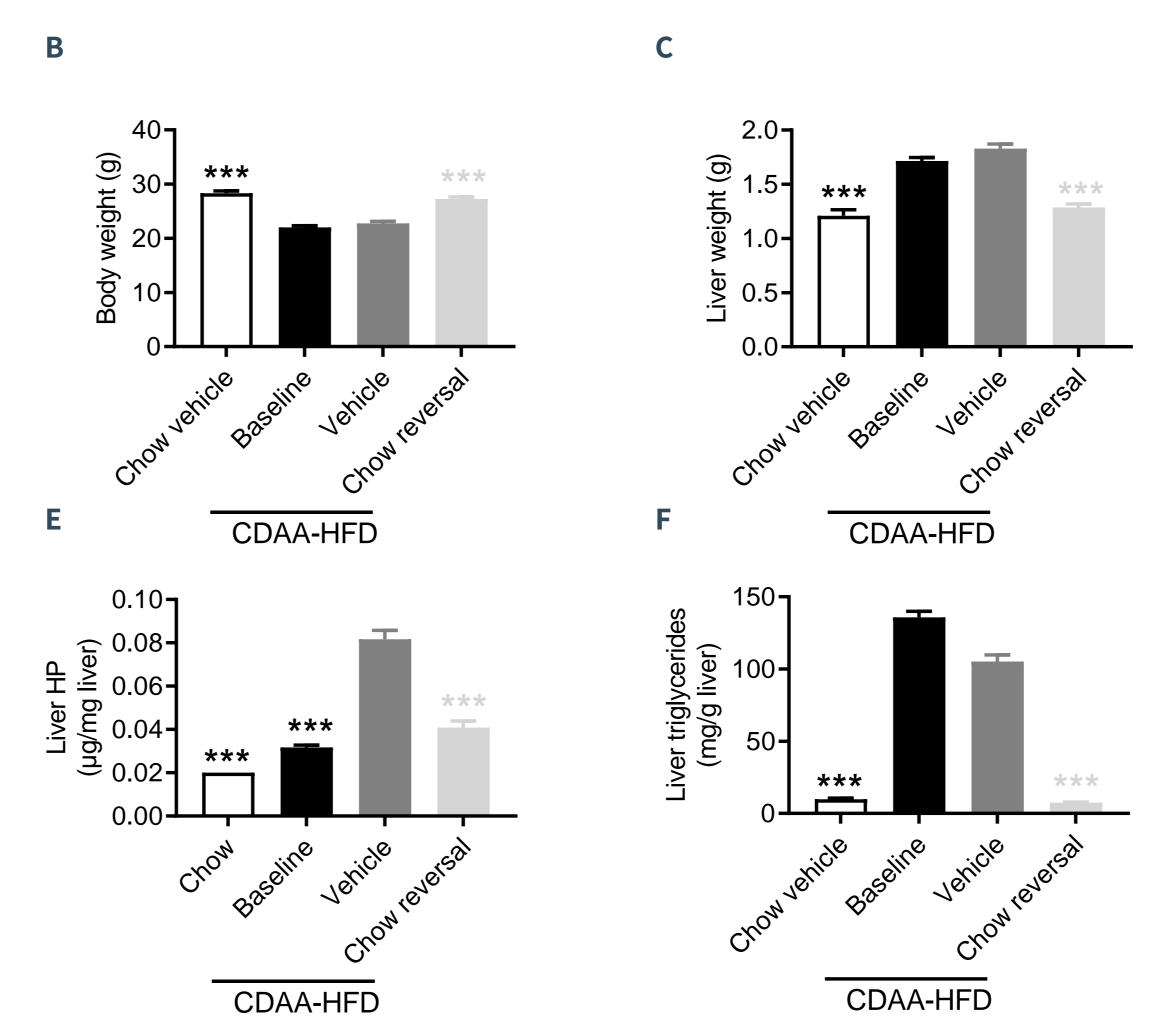


Figure 1. Chow reversal/dietary intervention reduces body weight, improves hepatomegaly plasma ALT, liver hydroxyproline and triglycerides levels in CDAA-HFD mice. (A) Body weight change relative (%) to day 0. (B) Terminal body weight (g). (C) Terminal liver weight (g). (D) Terminal plasma alanine aminotransferase (ALT, U/L). (E) Terminal liver hydroxyproline (HP, µg/mg). (F) Terminal liver triglycerides (. \*p<0.05, \*\*p<0.01, \*\*\*p<0.001 compared to corresponding CDAA-HFD vehicle group (Dunnett's test one-factor linear model).



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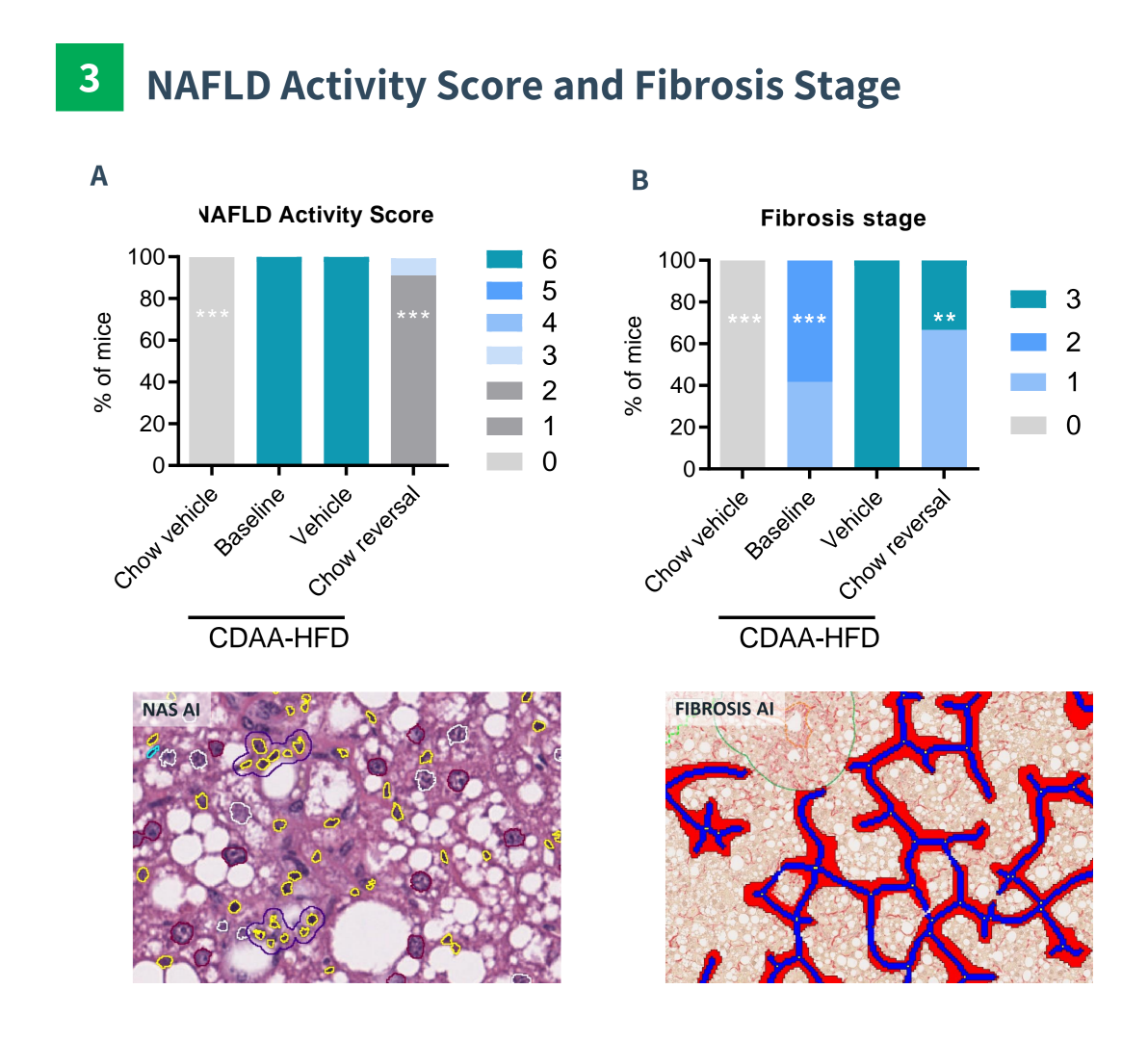


Figure 2. Chow reversal/dietary intervention improves NAFLD activity score and Fibrosis Stage in CDAA-HFD mice. Histopathological scores were determined by Gubra Histopathological Objective Scoring Technique (GHOST) deep learning-based image analysis. (A) NAFLD Activity Score (NAS). (B) Fibrosis Stage. \*\*p<0.01, \*\*\*p<0.001 compared to CDAA-HFD vehicle group (One-sided Fisher's exact test with Bonferroni correction). Bottom panels: Representative HE and PSR photomicrographs used for GHOST evaluation.

# CONCLUSION

Chow reversal/dietary intervention in CDAA-HFD mice:

- Normalizes body and liver weight as well as plasma, liver biochemistry and liver hydroxyproline levels.
- + Improves both NAFLD Activity Score and fibrosis stage.
- + Reduces quantitative histological markers of steatosis, inflammation and fibrosis
- + Suppresses hepatic genes linked to inflammation and fibrosis

Effects of dietary intervention in the non-obese CDAA-HFD mouse model of NASH with progressive fibrosis are in good agreement with clinical outcomes in NASH patients.