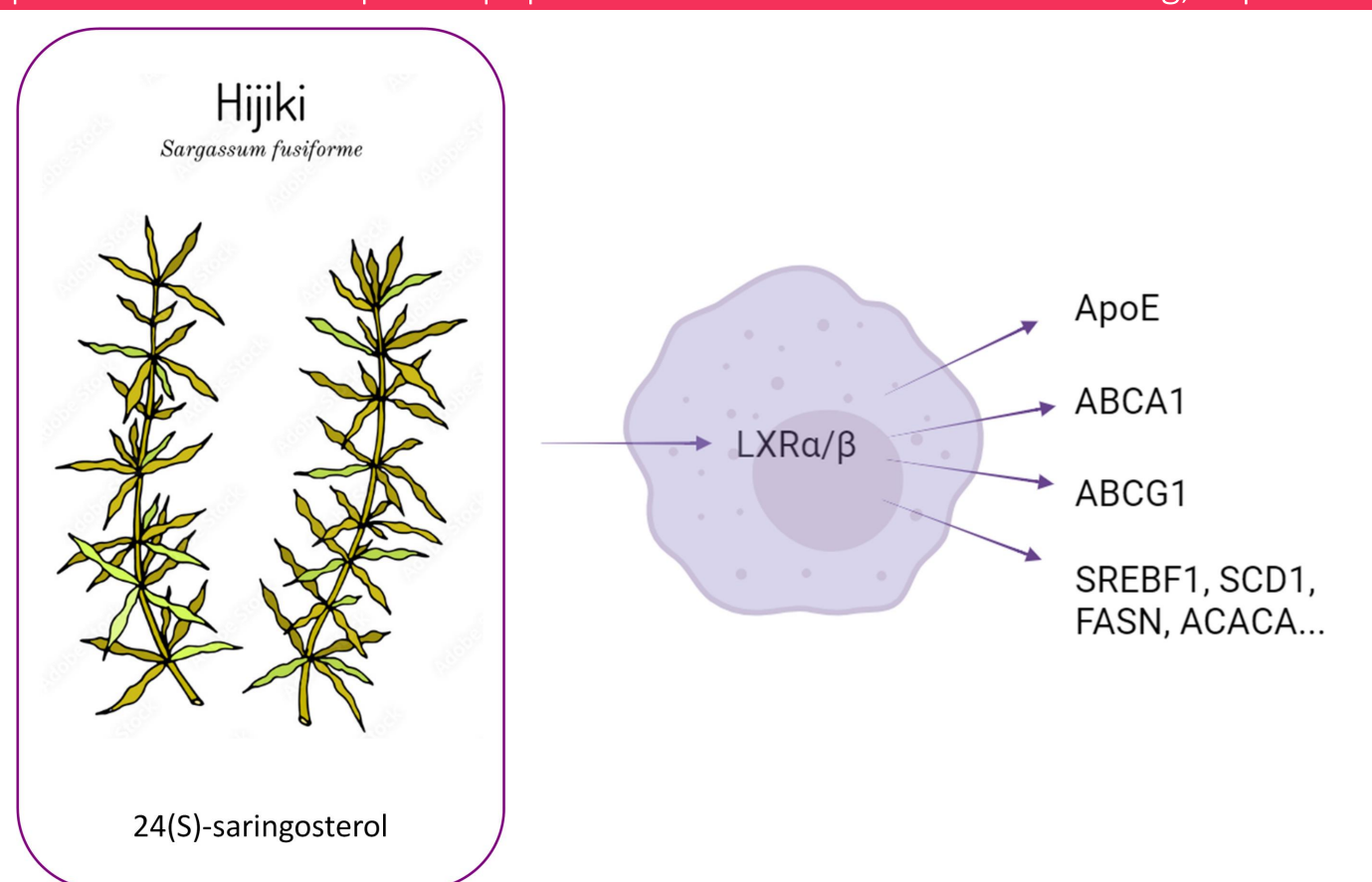
Identification of two new Side Chain Oxidized Sterols as Novel Liver X Receptor Agonists with Therapeutic Potential in the Treatment of Cardiovascular and Neurodegenerative Diseases

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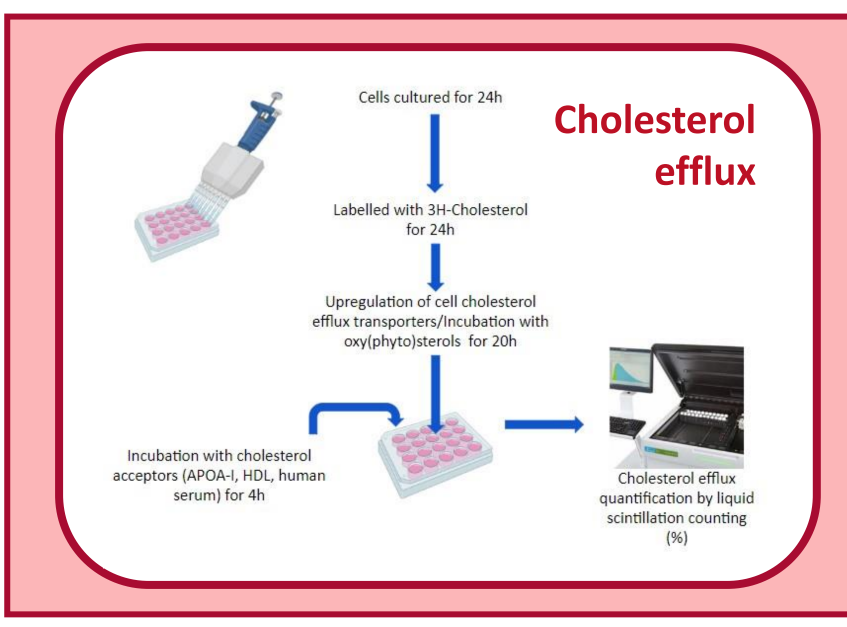
Background

The nuclear liver X receptors (LXR) α and β , may be potential therapeutic targets in cardiovascular and neurodegenerative diseases because of their key role in the regulation of lipid homeostasis and inflammatory processes.

Among the mechanisms involved in the maintenance of macrophage cholesterol homeostasis, cholesterol efflux plays a crucial role.

Specific oxy(phyto)sterols differentially modulate the transcriptional activity of LXRs, providing opportunities to develop new therapies. However, this development is precluded by unwanted side effects, such as hypertriglyceridemia and hepatic steatosis due to hepatic LXR α activation.

The aim of this study was to investigate the effect of newly isolated oxyphytosterols from Sargassum Fusiforme and new synthesized side chain oxidized sterols analogs on cholesterol efflux and on genes involved in this process.



Methods

Cellular cholesterol efflux was evaluated with a radioisotopic cell-based assay on human hepatocellular carcinoma cell line (HepG2) after treatment with n=13 new LXR α/β agonists at different concentrations. Gene expression was assessed in HepG2 and in human astrocytoma cells (CCF-STTG1) by qPCR. T0901317 $1\mu M$ was used as positive control.

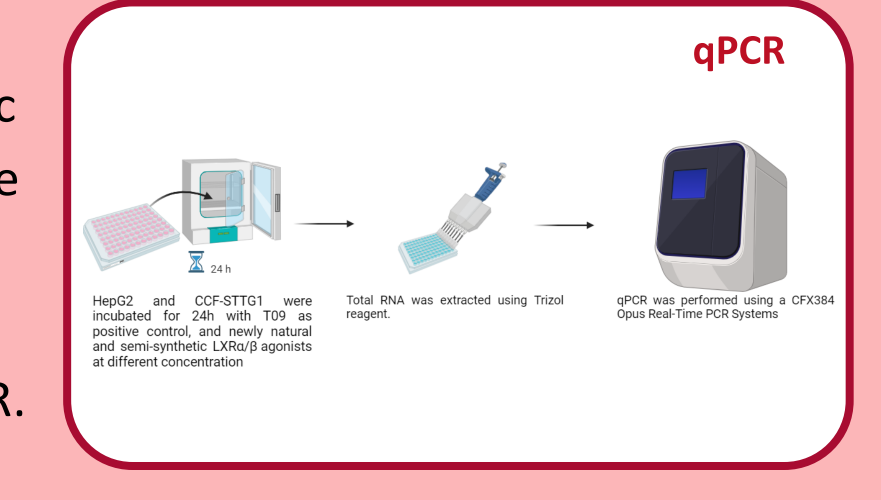
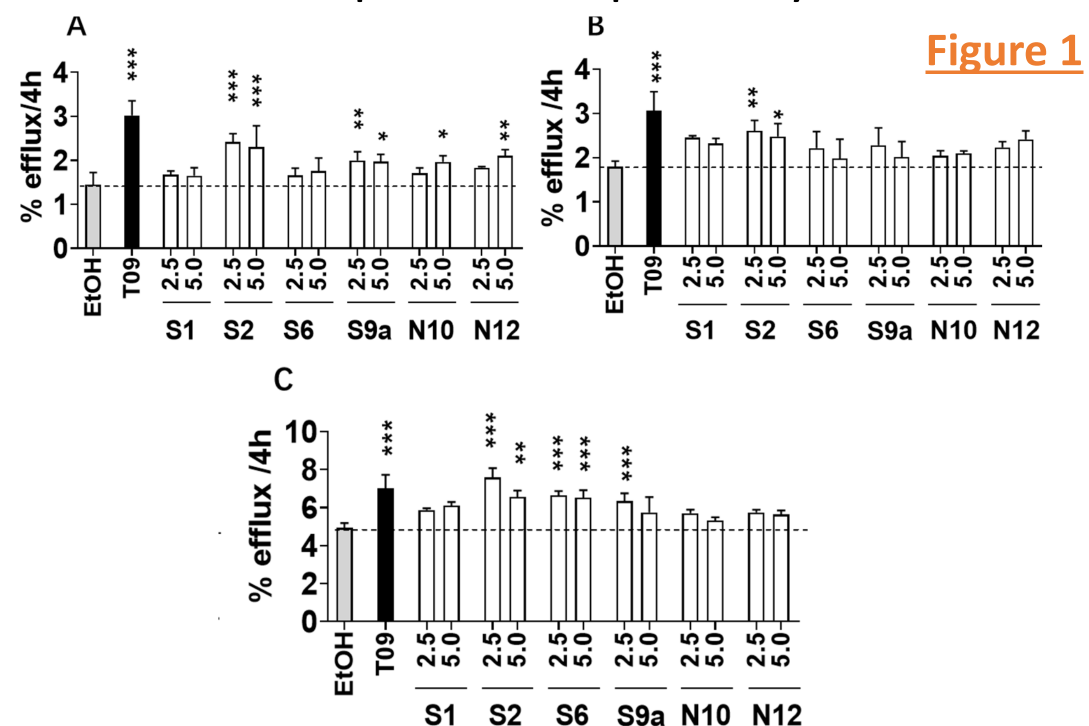


Figure 2

HepG2

Among the 5 natural and semi-synthetic 24oxidized sterols with high potency for LXR α/β activation, we identified two synthetic side chain compounds (S2 and S6) that also promote several protective pathways.



In detail, S2 increased cholesterol efflux from HepG2 by 54.3%, 15.2%, and 24.5% using isolated APOA-I (A) and HDL (B), and human cholesterol acceptor), serum respectively, while **S6** increased cholesterol efflux by 9.3% and 6.7% in presence of APOA-I (A), and human serum (C) (Figure 1).

impact Regarding the ot compounds on gene expression, in HepG2 cells **S2 S6** did and upregulate the expression of the main genes involved in cholesterol efflux ABCA1 and ABCG1 (data not shown), in addition they did not affect the expression of SREBF1, SCD1, FASN, or ACC1 (Figure 2), responsible for the hepatic side effects which are usual for synthetic pan-LXR agonist.

astrocytes, **S2** and slightly **S6** increased APOE, ABCA1, and ABCG1 mRNA levels (Figure 3).

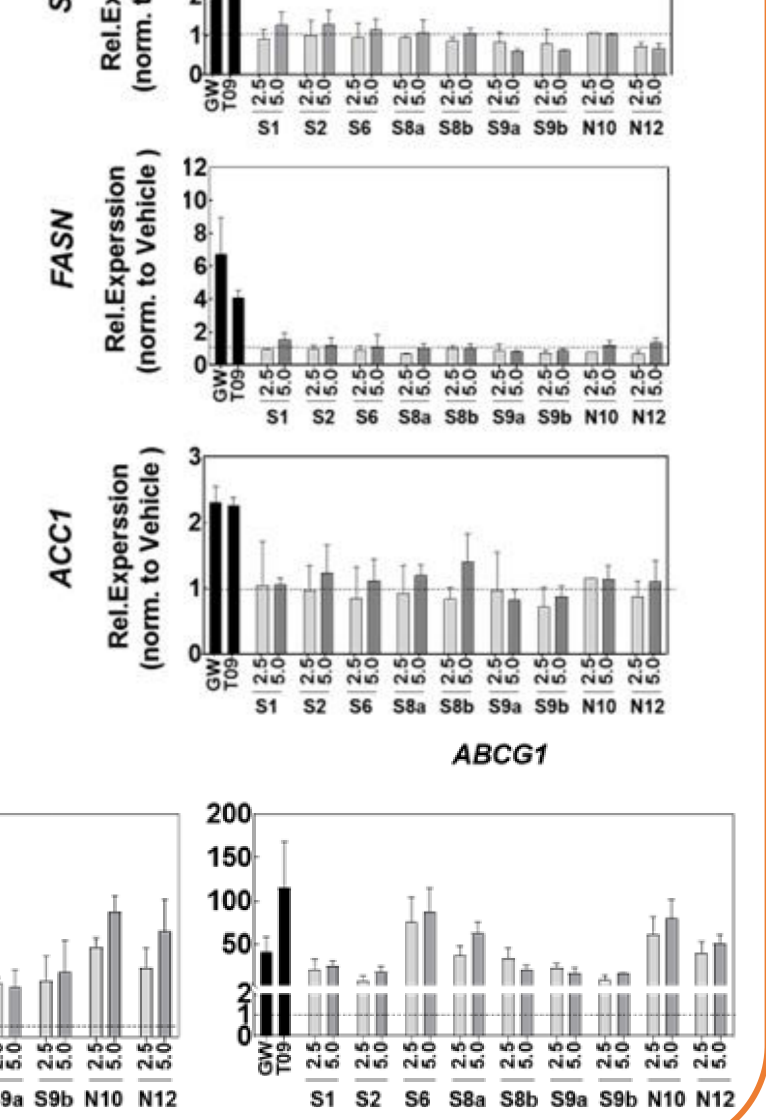
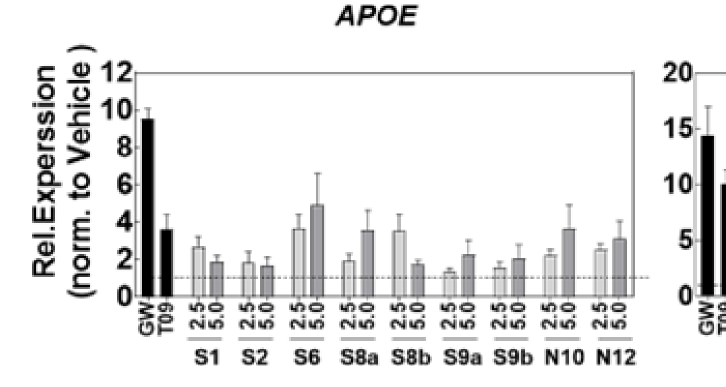
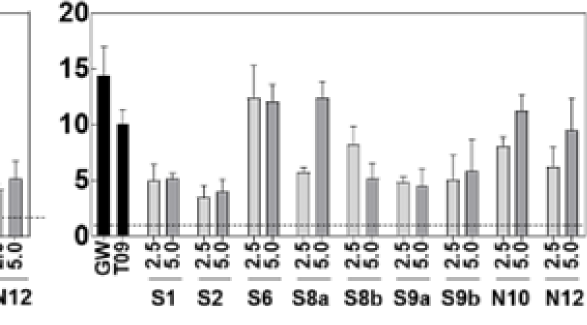


Figure 3





ABCA1

Conclusions

These results put the premises to identify and develop novel LXR-activating 24-oxidized sterols as potential therapeutic options in neurodegenerative and cardiovascular diseases.

