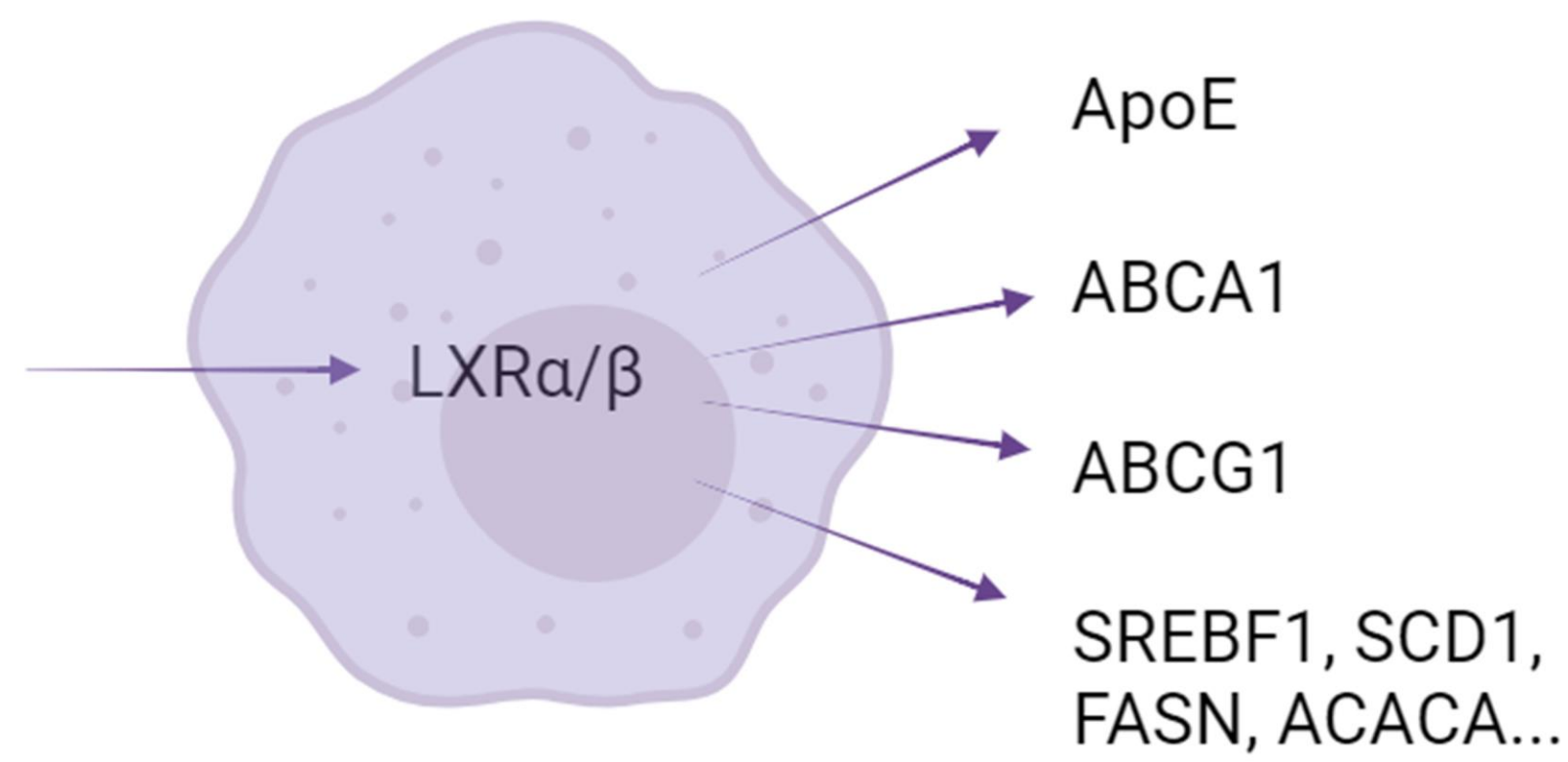
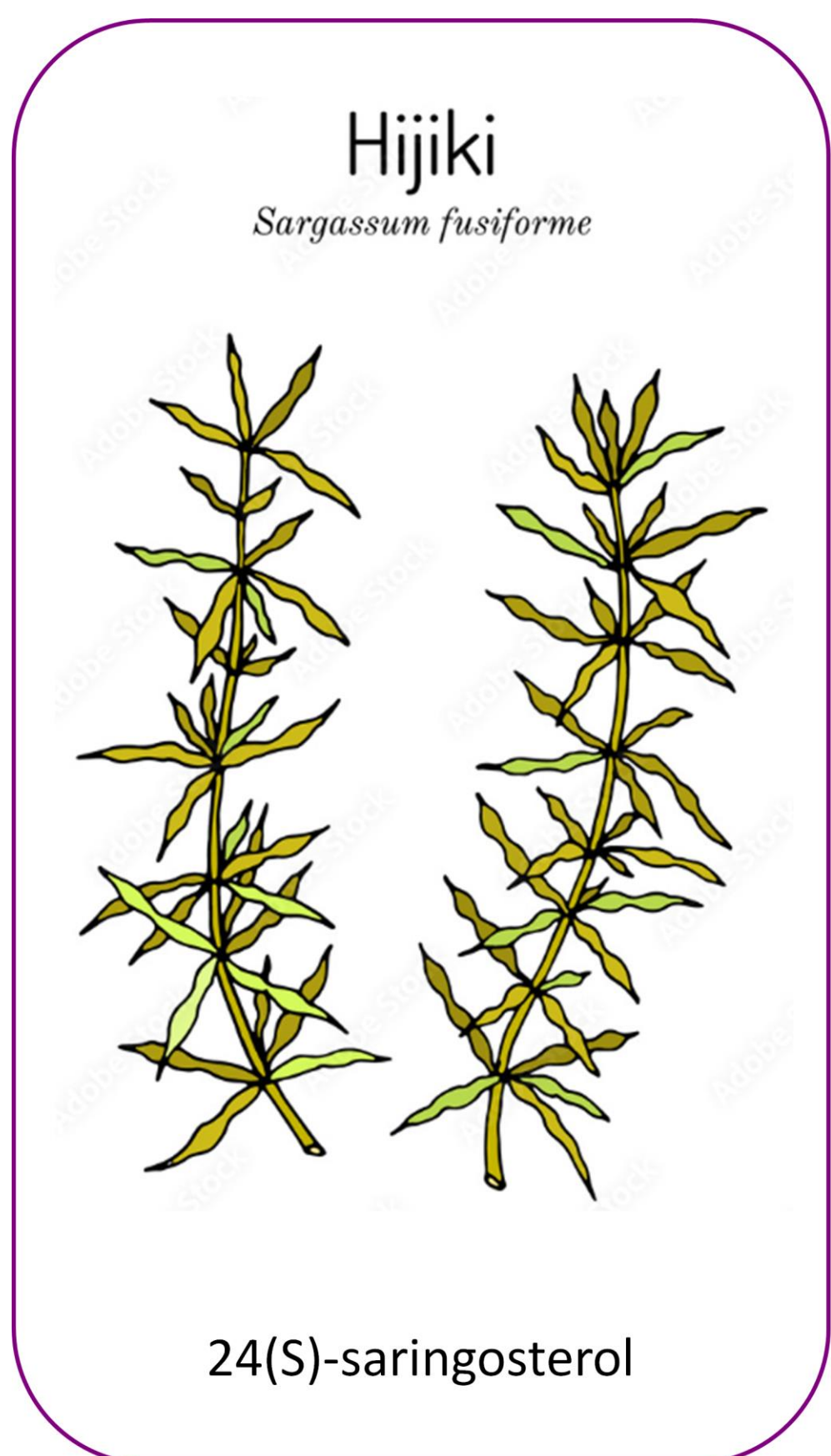


Identification of two new Side Chain Oxidized Sterols as Novel Liver X Receptor Agonists with Therapeutic Potential in the Treatment of Cardiovascular and Neurodegenerative Diseases

Marcella Palumbo 1, Na Zhan 2 3, Boyang Wang 2, Nikita Martens 3 4, Yankai Liu 2, Shangge Zhao 2, Gardi Voortman 3, Jeroen van Rooij 3, Frank Leijten 3, Tim Vanmierlo 4 5, Folkert Kuipers 6 7, Johan W Jonker 6, Vincent W Bloks 6, Dieter Lütjohann 8, Francesca Zimetti 1, Maria Pia Adorni 9, Hongbing Liu 2, Monique T Mulder 3

1Department of Food and Drug, University of Parma, 43124 Parma, Italy; 2Key Laboratory of Marine Drugs, Ministry of Education, School of Medicine and Pharmacy, Ocean University of China, Qingdao 266003, China; 3Department of Internal Medicine, Erasmus Medical Center, 3015 CN Rotterdam, The Netherlands; 4Department of Neuroscience, Biomedical Research Institute, Hasselt University, 3500 Hasselt, Belgium; 5School for Mental Health and Neuroscience, Maastricht University, 6229 ER Maastricht, The Netherlands; 6Department of Pediatrics, University Medical Center Groningen, University of Groningen, 9713 GZ Groningen, The Netherlands; 7European Research Institute for the Biology of Ageing (ERIBA), University Medical Center Groningen, University of Groningen, 9713 GZ Groningen, The Netherlands; 8Institute of Clinical Chemistry and Clinical Pharmacology, University Hospital Bonn, 53105 Bonn, Germany; 9Unit of Neurosciences, Department of Medicine and Surgery, University of Parma, 43125 Parma, Italy.

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Background

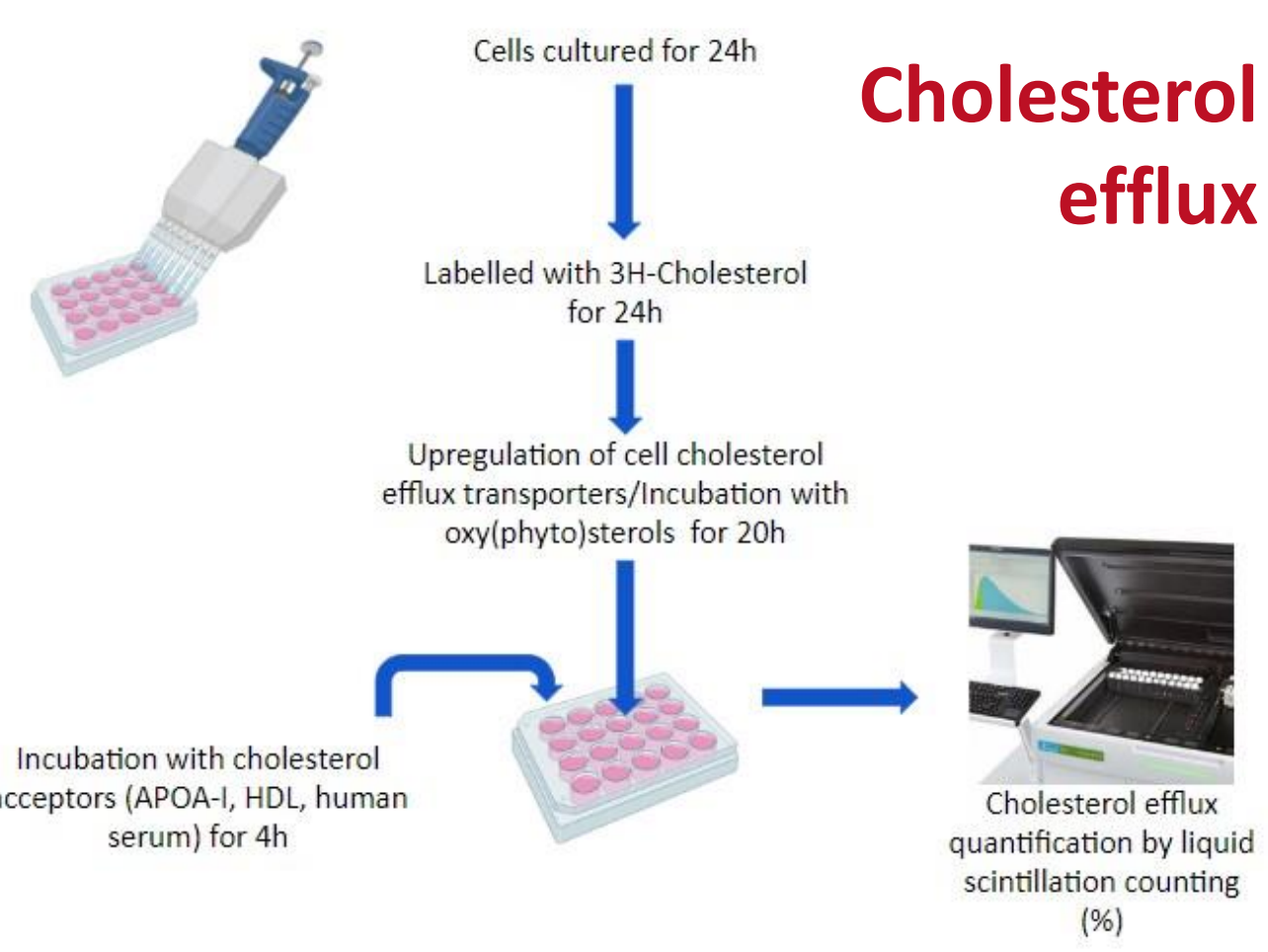
The **nuclear liver X receptors (LXR) α and β** , may be potential **therapeutic targets** in cardiovascular and neurodegenerative diseases because of their key role in the regulation of lipid homeostasis and inflammatory processes.

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

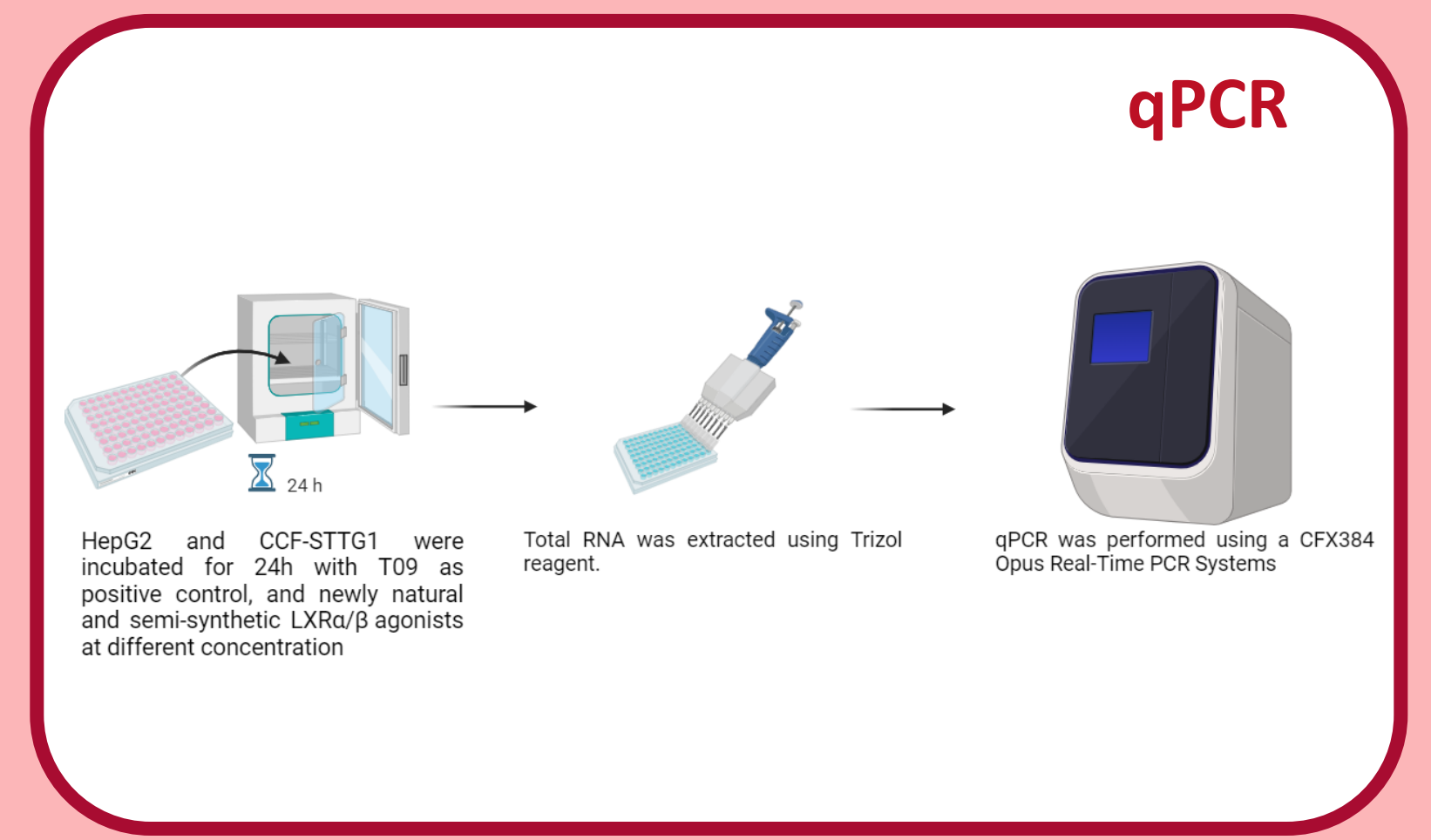
Specific **oxy(phyto)sterols** differentially modulate the transcriptional activity of **LXRs**, providing opportunities to develop **new therapies**. However, this development is precluded by unwanted side effects, such as hypertriglyceridemia and hepatic steatosis due to hepatic LXR α activation.

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

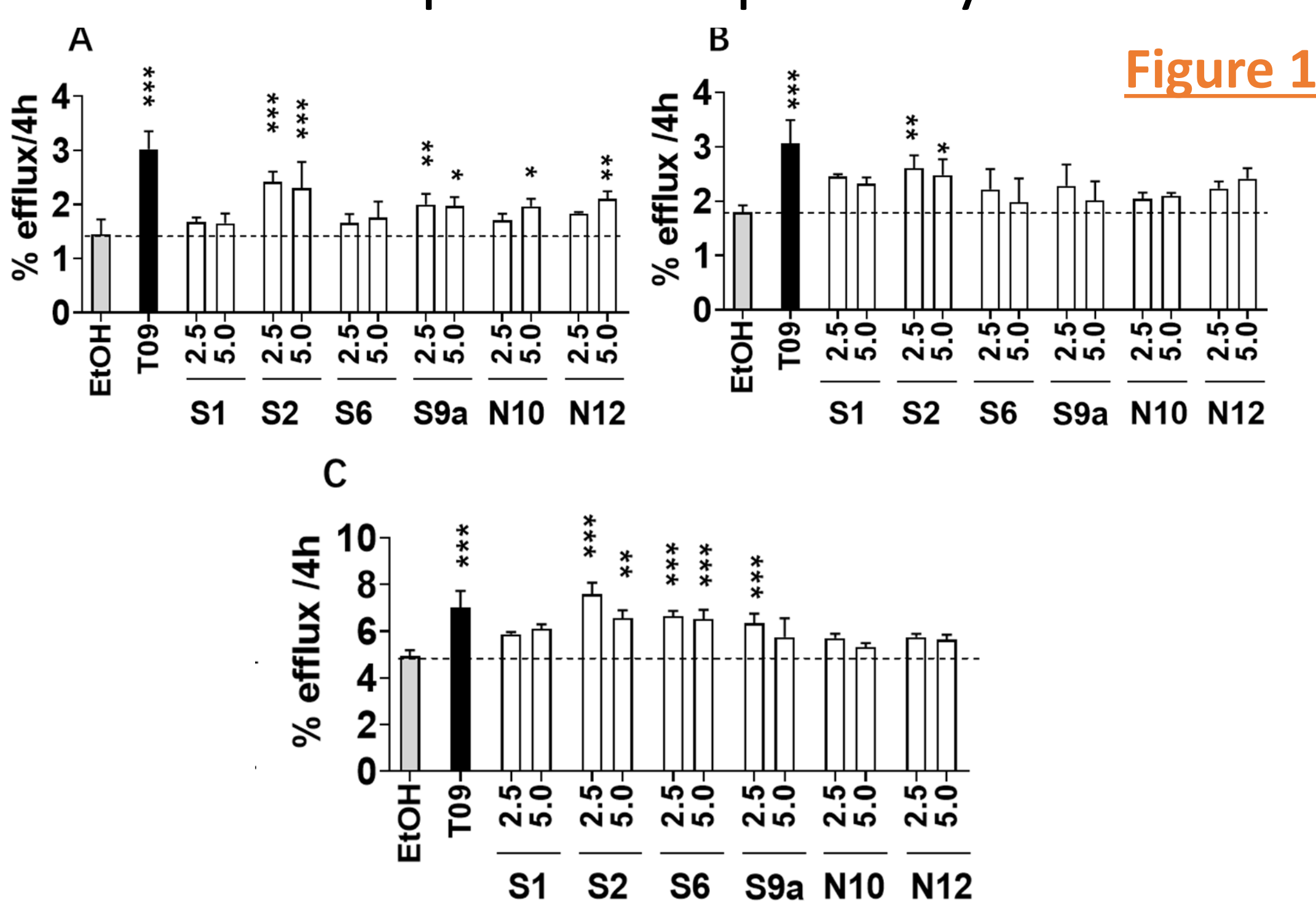


Cellular cholesterol efflux was evaluated with a radioisotopic cell-based assay on human hepatocellular carcinoma cell line (HepG2) after treatment with n=13 new LXR α / β agonists at different concentrations. Gene expression was assessed in HepG2 and in human astrocytoma cells (CCF-STTG1) by qPCR. T0901317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



Results

Among the 5 natural and semi-synthetic 24-oxidized 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 serum (**C**) (as cholesterol acceptor), respectively, while **S6** increased cholesterol efflux by 9.3% and 6.7% in presence of APOA-I (**A**), and human serum (**C**) (**Figure 1**).

Regarding the impact of these compounds on gene expression, in HepG2 cells **S2** and **S6** did not 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. In astrocytes, **S2** and **S6** slightly increased **APOE**, **ABCA1**, and **ABCG1** mRNA levels (**Figure 3**).

Figure 2 HepG2

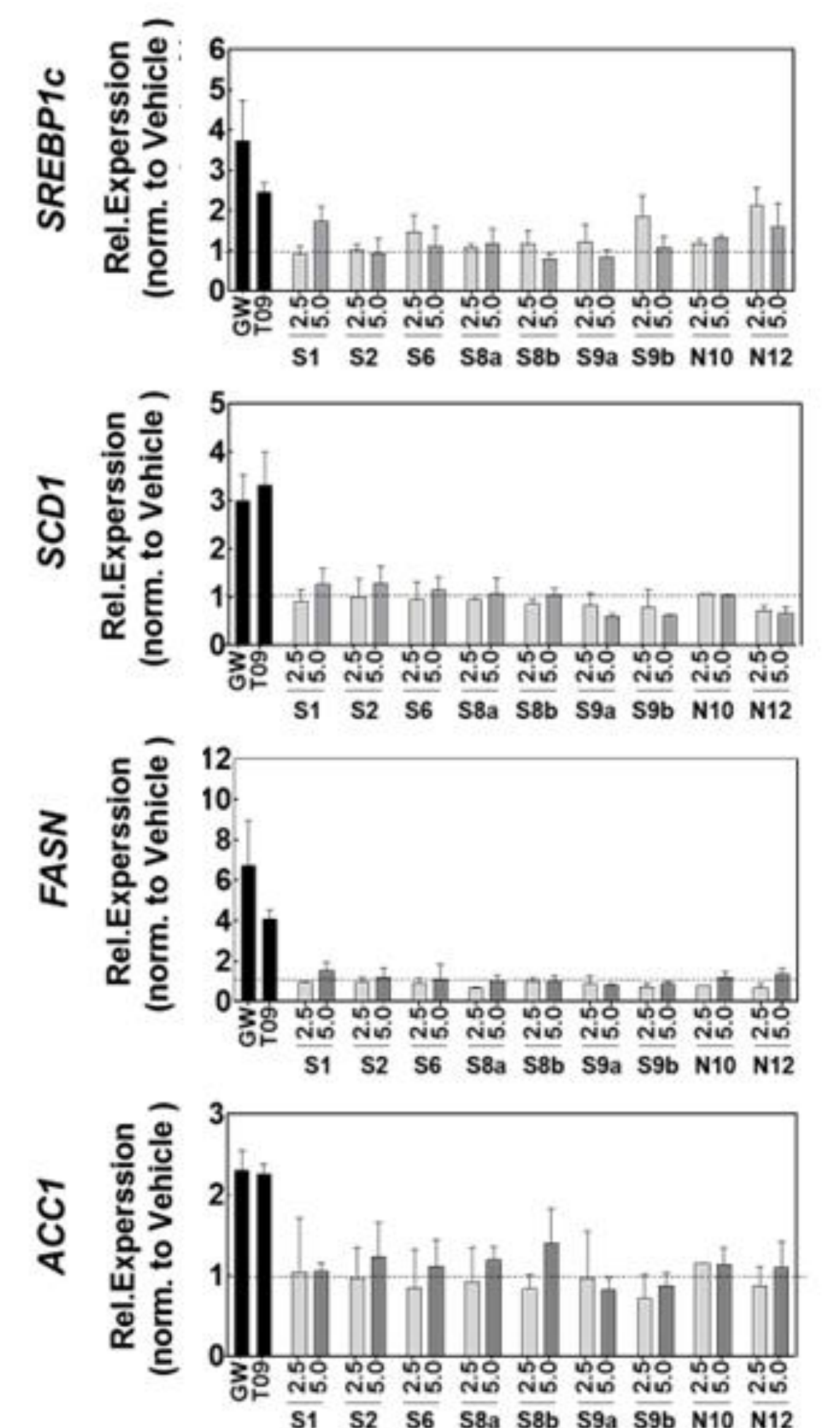
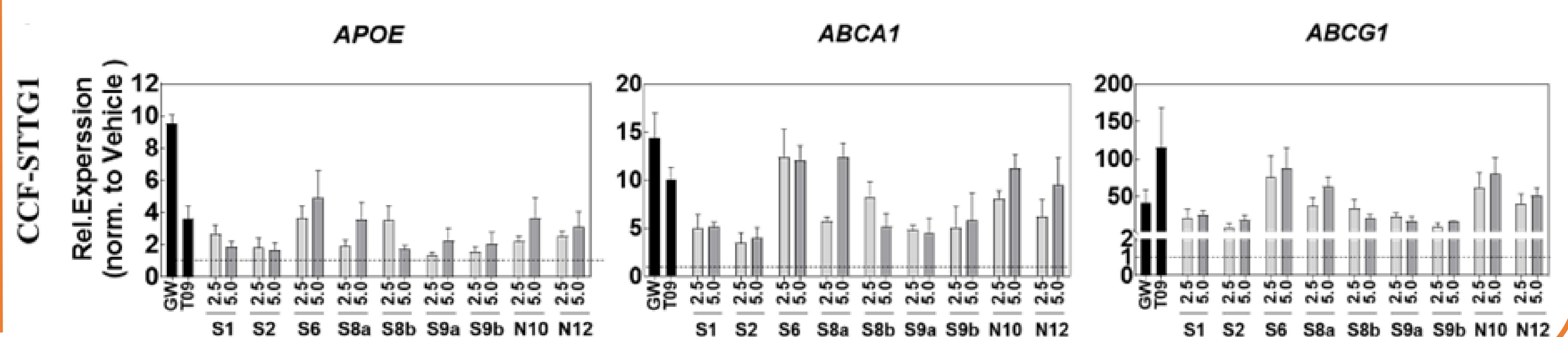


Figure 3



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.