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Trials of obeticholic acid for non-alcoholic steatohepatitis

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(Article begins on next page)



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Obeticholic acid: an effective treatment for diabetic patients with non-alcoholic steatohepatitis?

RUNNING TITLE: OCA and NASH

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To the Editor:

we read with interest the results of the FLINT trial, evaluating the effect of obeticholic acid (OCA) on liver histology in non-alcoholic steatohepatitis (NASH)¹.

A remarkable yet overlooked result of this trial was the different histological response between diabetic and non-diabetic participants: among the former, liver histology improved in 53% patients with OCA vs. 19% on placebo (OR for improvement with OCA: 4.6, 95%CI=2.0-10.6, p=0.0003), while in non-diabetic patients, liver histology improved in 37% patients with OCA vs. 23% with placebo (OR: 2.0, 95%CI=0.8-4.7, p=0.12).

The impact of glucose intolerance on histological response to OCA was evident also across progressive stages of pancreatic β -cell dysfunction, as estimated by HOMA B-cell index (online Appendix Table S2)¹

Beside the FLINT, only another randomized trial, enrolling diabetic patients, evaluated OCA in non-alcoholic fatty liver(NAFLD)². Therefore current evidence for effectiveness of farnesoid X receptor(FXR) agonists is lacking in non-diabetic individuals, a substantial proportion of NASH population.

Additionally, consistent with other randomized trials in NASH, the percentage of responders approached 50% in the FLINT, leaving a substantial proportion of patients without an effective treatment³. NASH is an heterogeneous conditions and diverse mechanisms of liver injury likely operate in different patient populations to promote liver disease progression. In humans, altered bile acid metabolism, which represents the rationale for using semi-synthetic bile acids, has been more convincingly demonstrated in diabetic than in non-diabetic individuals⁴, thereby potentially explaining the lack of response to OCA among non-diabetic individuals in the FLINT. Hence, further assessment of FXR agonists in nondiabetic NASH is mandatory, and an approach tailored to individual metabolic profile may be required to tackle NASH epidemic.

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Author Contributions.

Giovanni Musso: conceived the article, undertook literature search and acquired data, critically analyzed the results, drafted the article, gave final approval

Giovanni Musso, as the corresponding author, had full access to all the data and takes final responsibility for the decision to submit for publication.

Maurizio Cassader: undertook literature search and acquired data, critically analyzed the results, contributed to draft of the article, gave final approval

Roberto Gambino: undertook literature search and acquired data, critically analyzed the results, contributed to draft of the article, gave final approval

Conflicts of interest.

Giovanni Musso has no present or past conflict of interest or financial relationship to disclose. No funding bodies had any role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

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