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Resting-state brain connectivity following extra virgin olive oil intake in healthy adults: a randomised crossover pilot neuroimaging substudy

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This pilot randomised crossover neuroimaging substudy ($n = 9$) identified increased resting-state occipital functional connectivity following daily consumption of (poly)phenol-rich extra virgin olive oil compared with regular olive oil. Urinary hydroxytyrosol-glucuronide excretion increased significantly, and a significant interaction between intervention and metabolite levels was observed on occipital connectivity. These preliminary findings require replication in larger, adequately powered studies.

Introduction

Brain function and connectivity can be influenced by dietary factors, including foods rich in (poly)phenols. Previous evidence suggests that (poly)phenol-rich diets, such as the Mediterranean diet (MD), are associated with enhanced cognitive performance and brain connectivity, particularly in older adults.^{1,2}

Extra-Virgin Olive Oil (EVOO) is one of the main dietary sources of (poly)phenols within the MD. Unlike regular olive oil (OO), EVOO retains its antioxidant phenolic compounds due to natural minimal processing.^{3,4} Previous research has shown that greater adherence to MD is associated with stronger functional brain connectivity, especially in older adults.⁵

EVOO phenolic compounds as a whole exhibit strong antioxidants and anti-inflammatory properties. These bioactives, including oleuropein, hydroxytyrosol (HT), and related secoiridoids, neutralize reactive oxygen species, reduce oxidative damage to lipids, proteins, and DNA, and modulate signalling pathways involved in neuroinflammation and synaptic plasticity.^{6–8} Moreover, recent epidemiological evidence indicates that regular EVOO consumption is associated with reduced risk of cognitive decline and dementia, reinforcing its role in brain protection within the MD framework.^{9,10}

Among these compounds, HT is particularly relevant due to its potent antioxidant and neuroprotective activity. Importantly, HT is rapidly metabolised into HT-glucuronide, which is widely recognised as a validated biomarker of EVOO intake.¹¹

Despite growing evidence linking (poly)phenol-rich dietary patterns with cognitive and neurovascular outcomes, no previous study, to our knowledge, has evaluated the effects of EVOO consumption on human brain connectivity using resting-state functional magnetic resonance imaging (rs-fMRI). While prior research has primarily focused on older popu-

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lations, examining EVOO intake in healthy young adults may help detect early alterations in functional connectivity before age-related decline occurs.

This pilot randomised crossover trial was designed to explore whether daily EVOO intake is associated with differences in resting-state functional connectivity in healthy young adults. We combined rs-fMRI with urinary biomarker analysis to assess changes in brain connectivity in parallel with (poly)phenol metabolites. We hypothesised that short-term consumption of high-(poly)phenol EVOO, compared with low-(poly)phenol olive oil, would induce measurable differences in resting-state connectivity patterns.

Materials and methods

Study design and participants

This study is based on data from the HEVOOC trial (NCT05898113), a randomised, crossover clinical study aimed to evaluate the effects of daily consumption of 'Corbella' EVOO compared to regular OO in healthy young adults ($n = 28$).

Sample size was calculated based on the brain-derived neurotrophic factor (BDNF) variable using <https://www.surveysystem.com/sscalc.htm> for paired comparisons, with a significance level of 5% and 80% power, considering an expected dropout rate of 20%.

Participants were randomly allocated to one of two intervention sequences: (1) EVOO followed by regular olive oil (OO), or (2) the reverse sequence, with a 4-week washout period between phases. Each intervention phase lasted 4 weeks, during which participants consumed either EVOO or OO at a dose of 0.7 g per kg body weight per day, containing 227.7 mg kg^{-1} and 12.3 mg kg^{-1} total (poly)phenols, respectively. Participants were instructed to maintain their habitual diet and incorporate the assigned oil, either raw and/or cooked, at any time of the day.

Prior to the study, a 7-day run-in period was conducted, during which participants replaced EVOO with low-polyphenol olive oil and followed standardised dietary restrictions, avoiding olives, high-(poly)phenol foods (such as coffee, citrus fruits, apples, and berries), and limiting alcohol to a maximum of one glass of wine or beer per week. Participants were recruited in Barcelona, Spain, from June to September 2023, through online and institutional advertisements at Hospital Clínic and the University of Barcelona.

Inclusion criteria were age between 18 and 35 years, BMI < 30 kg m^{-2} , absence of chronic diseases, and non-smokers. Exclusion criteria included food allergies or intolerances, adherence to extreme or restrictive diets, excessive alcohol intake (>30 g day^{-1} for men, >20 g day^{-1} for women), pregnancy, menopause, or breastfeeding. For the present neuroimaging substudy ($n = 9$), participants from the HEVOOC cohort underwent rs-fMRI assessments. Participant flow throughout the study is presented in the SI (Fig. S1). All participants provided written informed consent. The study was conducted in accordance with the Declaration of Helsinki and was approved

by the Ethics Committee of Hospital Clínic (HCB/2023/0074) and the University of Barcelona (IRB00003099).

Neuroimaging acquisition and analysis

Image acquisition and analysis were performed at the IDIBAPS Magnetic Resonance Imaging Core Facility. MRI scans were conducted using a 3T Siemens PRISMA scanner and included both T_1 -weighted structural images (TR = 2.5 s, TE = 4.37 ms, voxel size: $1 \times 1 \times 1 \text{ mm}^3$) and rs-fMRI acquisitions (TR = 800 ms, TE = 37 ms, total volumes = 450; voxel size: $2 \times 2 \times 2 \text{ mm}^3$).

The neuroimaging protocol included structural and rs-fMRI, analysed *via* independent component analysis (ICA) and permutation testing. Preprocessing steps comprised: (1) slice timing correction using FSL (v.6.0.7.1),¹² (2) motion correction and detection of motion outliers: motion parameters were estimated using SPM12,¹³ together with frame displacement (FD) and DVARS computation.¹⁴ Motion parameters, FD, and DVARS were included as confound regressors in the BOLD signal estimation; (3) detection of motions outliers: a quality criteria of mean FD <0.5 mm was applied or if more than 10% of volumes with FD >0.5 mm; (4) registration of functional images to the T_1 -weighted structural image to mitigate EPI distortion using ANTs Registration (v.2.4.4);¹⁵ and (5) spatial smoothing with sigma = 3 mm and band-pass filtering between (0.01 and 0.15 Hz) using Nilearn (v.0.10.1).¹⁶ After pre-processing, images were registered into the standard MNI space using elastic registration by ANTs. Visual inspection of raw and pre-processed images was performed during the pipeline to ensure image quality.

Urinary (poly)phenol metabolites

Urinary metabolites were measured in 24 hour urine samples using liquid chromatography (Agilent 1290 Infinity II) coupled to high-resolution mass spectrometry (Agilent 6560 Ion Mobility QTOF LC/MS) operated in negative ionization mode. Analyses were performed at the Separation Techniques Unit of the Scientific and Technological Centers (CCiTUB), Universitat de Barcelona. Sample preparation and analysis followed a previously described protocol developed by our research group.¹⁷ Based on prior literature, the analysis specifically targeted (poly)phenolic compounds characteristic of EVOO: HT, HT-glucuronide, and HT-sulfate.^{18,19}

Covariate assessment

Covariates were selected based on prior knowledge of their potential influence on both (poly)phenol metabolites and resting state brain connectivity.^{20–24} These included total energy intake, assessed as a continuous variable (kcal day^{-1}) using a validated 151-item food frequency questionnaire (FFQ);²⁵ and adherence to the MD, evaluated using the 17-item MD Adherence Screener (MEDAS),²⁶ which provides a score reflecting higher or lower MD adherence.

Objective physical activity was measured with ActiGraph® wGT3X-BT accelerometers (ActiGraph LLC, Pensacola, FL). Each device was programmed with the participant's study ID



and worn on the non-dominant wrist continuously for 7 consecutive days prior to each study visit. Data were analysed with ActiLife software (v6.13.4). Moderate-to-vigorous physical activity (MVPA) was defined as ≥ 1952 counts per minute (cpm), using Freedson adult cut-points,²⁷ and expressed as the average daily minutes above this threshold. All covariates were measured at baseline and adjusted for in statistical models.

Statistical analysis

The neuroimaging analyses were conducted in a small subset of participants as part of a pilot, exploratory substudy. Accordingly, these analyses were intended to generate preliminary insights rather than to provide definitive or generalisable whole-brain inferences.

ICA was performed using the MELODIC toolbox within the FSL to identify resting-state brain networks. Dual regression was applied to obtain subject-specific spatial maps and time series. Group-level comparisons were conducted with the FSL randomize tool and threshold-free cluster enhancement (TFCE) to evaluate differences in brain activation across conditions (baseline *vs.* washout, pre-OO *vs.* post-OO, and pre-EVOO *vs.* post-EVOO). Statistical significance for imaging data was set at $p < 0.005$. Cohen's *d*-effect was computed voxel-wise and average in the cluster showing significant differences identified using randomise. Bootstrapping was then applied to the mean cluster connectivity in each subject to estimate the corresponding confidence interval.

To evaluate the effects of the dietary interventions on brain connectivity and urinary biomarkers, linear mixed-effects models (LMMs) were used to account for the repeated-measures crossover design. Each model included a random intercept for participant ID to account for within-subject correlation. Model 1 included intervention type, period, and the baseline value of the outcome variable. Model 2 additionally adjusted for baseline total energy intake, Mediterranean diet adherence score, and moderate-to-vigorous physical activity. Associations between urinary HT-glucuronide levels and brain connectivity were examined using models that included an interaction term between metabolite levels and intervention. Model assumptions were assessed by visual inspection of residual Q-Q plots and residuals *versus* fitted value plots to evaluate normality and homoscedasticity. When normality was not satisfied, variables were log-transformed prior to analysis. Standardised effect sizes (Cohen's *d*) were calculated for the main intervention effects by dividing the estimated fixed-effect coefficient by the residual standard deviation of the model. All analyses were performed using Stata 17.0 (StataCorp, College Station, TX) and statistical significance was set at $p < 0.05$.

Results

Participant characteristics

Of the 28 participants enrolled in the HEVOOC randomised clinical trial, nine healthy adults consented to undergo additional neuroimaging assessments and were therefore

Table 1 Baseline characteristics of the study participants included in the pilot study

Characteristics	Participants (<i>n</i> = 9)
Sex, male : female (<i>n</i>)	5 : 4
Age (years)	31 [27;32]
Blood pressure measures	
SBP (mmHg)	112.5 [104; 121]
DBP (mmHg)	73.7 [69; 73]
Anthropometric and metabolic parameters	
Weight (kg)	64.1 [58.1; 75.1]
BMI (kg m^{-2})	23 [21.2; 25.1]
Body fat (%)	24.7 [22.5; 29.4]
Muscle mass (%)	34.1 [29.5; 37.7]
Visceral fat (%)	6 [3; 7]
Waist circumference (cm)	76.5 [73; 84]
TC (mg dL^{-1})	165 [153; 216]
LDL-C (mg dL^{-1})	95 [78; 122]
HDL-C (mg dL^{-1})	52 [48; 56]
Questionnaire of healthy habits	
MVPA min day^{-1}	303 [276.9; 321.4]
MedDiet score	10 [9; 11]
Total energy intake (Kcal day^{-1})	2210.6 [2128.2; 2725.1]

Data are presented as median [IQR]. Abbreviations: SBP – systolic blood pressure; DBP – diastolic blood pressure; BMI – body mass index; TC – total cholesterol; HDL-C – high-density lipoprotein cholesterol; LDL-C – low-density lipoprotein cholesterol; MVPA – moderate-to-vigorous physical activity; MedDiet – Mediterranean diet adherence score.

included in the present exploratory analysis. Baseline characteristics are presented in Table 1. Participants were normotensive, had no cardiometabolic risk factors, and exhibited a moderate adherence to the MD (median score: 10/17).

Effects of EVOO on resting-state functional connectivity

Analysis of rs-fMRI data (TFCE-corrected, FSL Randomize; $p < 0.005$) revealed increased activation in a visual network localised to the left occipital cortex, including its superior and inferior lateral areas, following EVOO intake (Fig. 1a). The average voxel-wise Cohen's *d* within the significant cluster was 1.058 with a bootstrapped 95% CI [0.147, 0.464] for the mean cluster connectivity change. Fig. 1b displays the average visual network connectivity before and after each intervention, showing notably higher post-EVOO connectivity levels compared to regular OO. Extracted mean connectivity values were analysed using LMMs, functional connectivity was significantly greater following EVOO compared with regular OO consumption in the fully adjusted model (Model 2: $\beta = 0.20$; 95% CI: 0.03, 0.37; $p = 0.016$). Cohen's *d* = 1.46. Results were consistent in the minimally adjusted model (Table S1).

Together, these findings support an association between EVOO intake and increased intrinsic functional connectivity in occipital regions involved in visual processing.

Effect of EVOO on urinary HT-glucuronide and its association with occipital connectivity

Urinary log-transformed HT-glucuronide excretion were significantly higher following EVOO compared with regular OO consumption in the fully adjusted model ($\beta = 1.24$; 95% CI: 0.13,



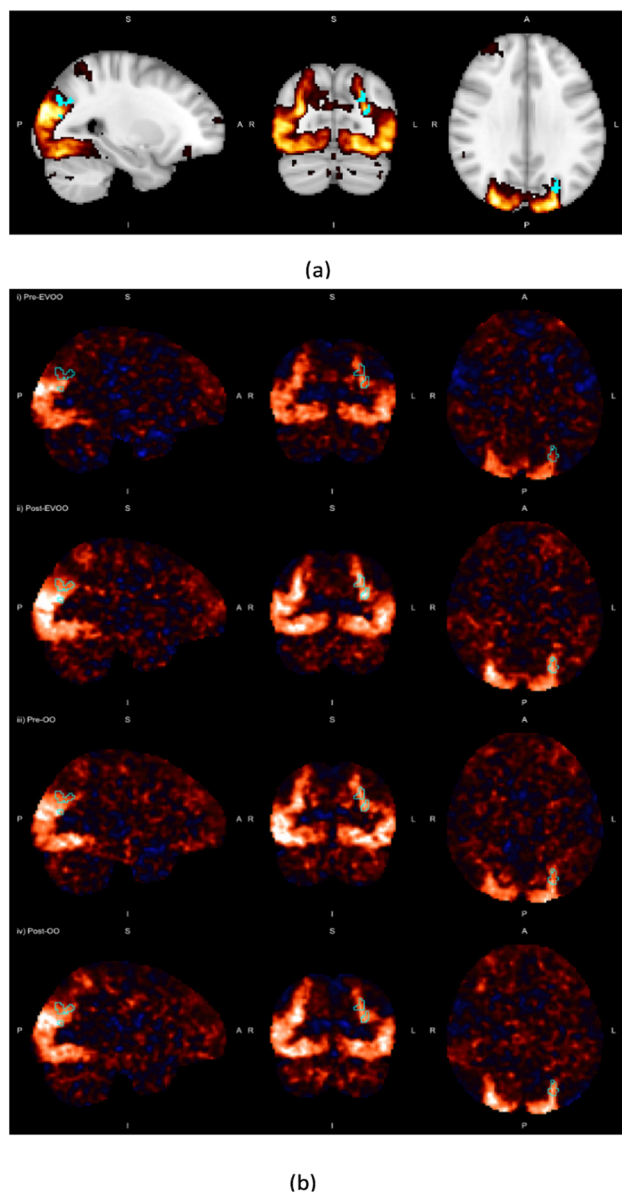


Fig. 1 Changes in Brain Activation following Dietary Interventions. (a) Effect of the EVOO intervention in an occipital functional network. Red-yellow colours indicate the spatial map of the network obtained from ICA. Light blue indicates the points where significant differences in activation between pre- and post-EVOO intervention. (b) Mean functional connectivity maps of the significant ICA component: (i) pre-EVOO, (ii) post-EVOO, (iii) pre-OO and (iv) post-OO. Significant differences were detected in the areas outlined in light blue. Orientation: S (superior), I (inferior), A (anterior), P (posterior), L (left), R (right).

2.35; $p = 0.028$) Similar estimates were observed in the minimally adjusted model (Table S1). In exploratory analyses, changes in urinary log-transformed HT-glucuronide levels were positively associated with changes in occipital functional connectivity in the fully adjusted model ($\beta = 3.34$; 95% CI: $-0.02, 6.72$; $p = 0.052$). Estimates from the minimally adjusted model were comparable (Table S1).

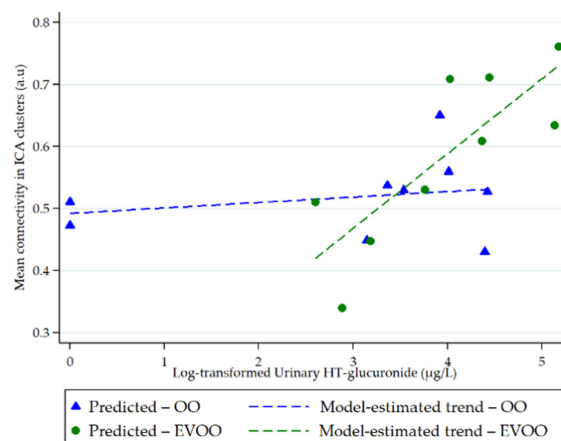


Fig. 2 Brain activation and urinary HT-glucuronide by intervention. Individual occipital network activation values are plotted against log-transformed urinary HT-glucuronide levels for OO (lavender triangles) and EVOO (green circles) groups ($n = 9$). Dashed lines represent model-estimated trend lines for each intervention. Linear mixed-effects modelling revealed a significant positive association between HT-glucuronide levels and brain activation in the EVOO group compared to OO ($\beta = 0.13$, $p = 0.002$).

Interaction between EVOO Intake and HT-glucuronide on brain activity

Notably, a significant interaction was observed between urinary HT-glucuronide levels and intervention group on occipital network activation in the fully adjusted model ($\beta = 0.13$; 95% CI: $0.05, 0.21$; $p = 0.002$). Similar estimates were obtained in the minimally adjusted model (Model 1: $\beta = 0.14$; 95% CI: $0.06, 0.21$; $p < 0.001$), indicating that the interaction effect was robust to additional adjustment. Fig. 2 plots occipital network activation against log-transformed urinary HT-glucuronide levels for each intervention, illustrating that the association between HT-glucuronide and brain activation differed by intervention.

Discussion

This exploratory pilot study provides initial evidence that daily intake of 0.7 g per kg body weight of EVOO for one month was associated with increased resting-state functional connectivity within an occipital network, compared with regular, low-(poly) phenol OO. Activation increases were localised to the left occipital cortex, areas involved in visuospatial attention, motion perception (superior), and object recognition (inferior). EVOO consumption also significantly increased urinary HT-glucuronide excretion. Notably, a significant interaction between HT-glucuronide excretion and intervention was observed on occipital functional connectivity, indicating that the relationship between phenolic metabolite levels and visual network connectivity differed according to the intervention.

These results complement prior studies associating MD adherence with improved brain connectivity²⁸ but uniquely



isolate EVOO's contribution. While previous work identified interesting effects of EVOO consumption on frontoparietal brain regions involved in working memory,²⁰ our findings point to specific modulation of visual cortical networks even in the absence of external stimuli. Activation was specifically localised in the left occipital cortex, a region implicated in visuospatial attention and motion perception in its superior portion, and object recognition in its inferior segment.^{29,30}

Urinary HT-glucuronide increased significantly following EVOO consumption. The observed interaction between HT-glucuronide levels and intervention further supports the hypothesis that EVOO-specific phenolics may play a relevant role in mediating its neurological benefits.

Pharmacokinetic studies show that over 60% of phenolic metabolites detected post-EVOO consumption circulate as conjugates, mainly glucuronides and sulfates.³¹ While urinary HT-glucuronide is a validated biomarker of EVOO consumption,³² and may contribute to observed effects, it is likely that other co-occurring bioactive compounds in EVOO also play a role in modulating brain connectivity.

In addition to human studies, preclinical evidence supports the neuromodulatory role of EVOO. Recent work in obese rats demonstrated that EVOO supplementation reduced hypothalamic inflammation (arcuate nucleus) and improved metabolic parameters compared to controls suggesting broader neuromodulatory effects across distinct brain regions.³³ Although these findings concern different brain regions and mechanistic pathways, they provide biological plausibility for the cerebral effects of EVOO.

Strengths of this study include its nutritional randomised crossover design with a control, fMRI-based neural assessment, phenolic profiling, and within-subject control. The inclusion of healthy young participants helps control age-related variability in brain function. However, the study has limitations. Most notably, the small sample size ($n = 9$) may limit statistical power and the generalizability of the findings, particularly with respect to the neuroimaging outcomes. Additionally, the relatively short duration of the intervention (one month) may be insufficient to capture long-term neuroplastic changes. While the crossover design improves internal validity by allowing each participant to serve as their own control, replication of these results in larger, adequately powered samples, ideally at least double the current size, is necessary to confirm the robustness and reproducibility of the observed effects.

Conclusions

One month of consuming EVOO (0.7 g per kg body weight per day) was associated with preliminary differences in resting-state functional connectivity and higher urinary excretion of HT-glucuronide in healthy young adults, compared to one month of consuming regular, highly refined OO. The observed interaction between olive oil-derived phenolic metabolites and functional connectivity suggests a potential neuro-modulatory

effect of EVOO. However, these findings should be interpreted with caution due to the exploratory nature and small sample size of the neuroimaging substudy. These novel findings warrant further investigation in larger, adequately powered, and long-term studies to confirm the observed associations and to explore the underlying mechanisms of the (poly) phenols present in EVOO.

Author contributions

Conceptualization, M. P., A. V.-Q., R. C., R. E. and R. M. L.-R.; methodology, R. M. G.-R., E. M.-M., A. H., P. G., C. A.-R. and I. D.-L.; software, E. M.-M. and A. H.; validation, C. D.-V., I. D.-L. and R. M. L.-R.; formal analysis, R. M. G.-R., C. D.-V., E. M.-M., A. H., P. G., C. A.-R. and I. D.-L.; investigation, R. M. G.-R., E. M.-M., A. H., P. G., C. A.-R. and S. H.-B.; resources, R. C. and R. E.; data curation, R. M. G.-R. and C. D.-V.; writing – original draft, R. M. G.-R.; writing – review & editing, C. D.-V., E. M.-M., A. H., P. G., C. A.-R., S. H.-B., I. D.-L., M. P., A. V.-Q., R. C., R. E. and R. M. L.-R.; visualization, R. M. G.-R.; supervision, R. M. L.-R.; project administration, R. M. L.-R.; funding acquisition, R. E. and R. M. L.-R. All authors have read and agreed to the published version of the manuscript.

Conflicts of interest

R. M. L.-R. reported personal fees from Cerveceros de España, UNIDECO, Adventia, Wine in Moderation, and Ecoveritas S. A., all outside the submitted work. R. E. reported grants from the Fundación Dieta Mediterránea and Fundación Cerveza y Salud, as well as personal fees for lectures from Brewers of Europe, Fundación Cerveza y Salud, Instituto Cervantes (Albuquerque, Milan, Tokyo), Pernod Ricard, and the Wine and Culinary International Forum. He also received non-financial support for the organization of a national nutrition congress and for feeding trials with products from Grand Fountain and Uriach Laboratories (Spain). The funders had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript; or in the decision to publish the results.

Data availability

The data supporting this article have been included as part of the supplementary information (SI), including Supplementary Dataset 1. Supplementary information is available. See DOI: <https://doi.org/10.1039/d5fo05016b>.

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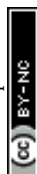
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