Associations Between Cannabis Use, Abdominal Fat Phenotypes and Insulin Traits

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Abstract

Background: General obesity has been linked to dysregulation of the endocannabinoid system in humans. However, there is a lack of studies on the relationship between cannabis use and specific abdominal fat phenotypes. The aim was to investigate the associations between cannabis use and magnetic resonance imaging-derived fat phenotypes, as well as indices of insulin sensitivity and insulin secretion.

Methods: In this cross-sectional study, magnetic resonance imaging was used to quantify subcutaneous fat volume (SFV), visceral fat volume (VFV), intra-hepatic fat deposition (IHFD), intra-pancreatic fat deposition (IPFD) and skeletal muscle fat deposition (SMFD) by two independent observers. Insulin sensitivity was determined based on HOMA-IS, Raynaud index and Matsuda index, whereas insulin secretion was determined based on HOMA-β, insulinogenic index 30' and insulinogenic index 60'. A validated questionnaire was used to ascertain participants’ cannabis use. Linear regression models were constructed, adjusting for demographics, glycated hemoglobin, physical activity, tobacco smoking and alcohol consumption.

Results: A total of 120 individuals were included. Cannabis use explained 9.2% of variance in IHFD, 4.4% in SMFD, 3.4% in VFV, 0.4% in SFV and 0.2% in IPFD. Regular cannabis users had significantly greater IHFD compared with never users, in both the unadjusted (P = 0.002) and all adjusted (P = 0.002; P = 0.008) analyses. The other fat phenotypes did not differ significantly between either regular or non-regular users compared with never users. Regular cannabis users had significantly greater insulin secretion (as defined by the insulinogenic index 60') compared with never users, in both the unadjusted (P = 0.049) and all adjusted (P = 0.003; P = 0.004) analyses. The other indices of insulin sensitivity in either regular or non-regular cannabis users compared with never users.

Conclusion: Regular cannabis use may be a risk factor for non-alcoholic fatty liver disease (but not IPFD) and may alter the neuromodulation of insulin secretion. Further investigations are now warranted to elucidate the mechanisms underlying these associations.

Keywords: Cannabis use; Intra-hepatic fat; Intra-pancreatic fat; Skeletal muscle fat; Visceral fat; Subcutaneous fat; Magnetic resonance imaging; Insulin traits

Introduction

The Cannabis sativa plant is among the earliest crops cultivated by man and its use dates back to 2737 BC [1]. In recent decades, with the discovery of the endocannabinoid system (ECS), there has been an increasing trend towards legalizing medicinal cannabis use. To date, 33 US states have legalized cannabis for medicinal use, with 11 of these also allowing recreational use [2]. The US Food and Drug Administration has approved the use of cannabis-based drugs for the treatment of seizures associated with Lennox-Gastaut syndrome and Dravet syndrome, for nausea and vomiting associated with cancer chemotherapy in patients who have been unresponsive to first-line treatments, and for anorexia associated with weight loss in acquired immunodeficiency syndrome patients [3]. Given the numerous health claims and the increasing trend towards decriminalization of cannabis, there has been a call to better understand the acute and chronic risks of cannabis use. Mechoulam and colleagues [2] reviewed the current literature on cannabis use-related health effects. Of the 68 studies analyzed, they found 62 reported various adverse outcomes [2]. However, these effects are not completely understood and research is far from sufficient to recommend a “safe” therapeutic dose of cannabis-based drugs.

In the context of metabolic health, the relationship between cannabis use and metabolic sequelae has been speculated. “Marijuana”, the most commonly used recreational form of cannabis, contains high concentrations of Δ-9-THC (the main psychoactive constituent of the plant), which is known to have appetite-inducing properties [4]. Cannabinoid
receptors (CB$_1$R and CB$_2$R) are present both centrally and pe-
ripically, and their endogenous ligands, endocannabinoids, play a central role in appetite regulation, energy balance, li-
pid metabolism, insulin sensitivity and β-cell function [5]. A
clear association exists between ECS dysregulation and the
development of obesity and type 2 diabetes. Animal stud-
ies have demonstrated that chronic CB$_1$R stimulation favors
adiposity, independent of calorie intake [4]. Several experimen-
tal models have shown metabolic effects from peripheral
CB$_1$R activation, including increased lipogenesis in hepatocy-
cyes and adipocytes, reduced insulin responsiveness of myo-
cytes, as well as impaired insulin sensitivity and secretion
in the pancreatic β cells [4]. Consistent with these findings,
human clinical trials using CB$_1$R antagonists have proven to
be effective in reducing body weight as well as improving
dyslipidemia and insulin sensitivity [4]. These findings from
human and animal studies suggest that cannabis use may be
not innocuous and may be privy to the pathophysiology of
metabolic syndrome.

Evidence for cannabis use and obesity has been derived
largely from experimental models and epidemiological data
using body mass index (BMI) [6, 7]. More recently, focus has
shifted to the distribution of fat, including visceral and ec-
topic, as a greater predictor of metabolic risk [8]. Muniyappa
and colleagues [4] have demonstrated a relative visceral adi-
posity and adipose tissue insulin resistance (IR) in chronic
cannabis users, suggesting differential and tissue-specific ef-
effects of regular cannabis use. In the context of ectopic fat
deposition, only preliminary studies have linked cannabis
use with the risk of fatty liver, which have predominantly
been animal studies [9, 10]. Magnetic resonance (MR) im-
aging has emerged as the "gold standard" for differentiating
the abdominal fat compartments and quantifying ectopic fat,
such as in the liver and pancreas. To date, no attempts have
been made to comprehensively characterize the phenotype of
fat deposition (using validated MR-derived measurements)
associated with cannabis use. Further, it has not been inves-
tigated how the distribution of fat (both visceral and ectopic)
associated with cannabis use affects insulin secretion and
sensitivity.

The primary aim of this study was to investigate the asso-
ciation between cannabis use and MR-derived fat phenotypes.
The secondary aim was to evaluate the association between
cannabis use and insulin traits.

Materials and Methods

Study design and study population

This was a cross-sectional study, which represented a sub-
project within the main project. The main project focused on
intra-pancreatic fat deposition (IPFD) after an attack of pan-
creatitis and its results were published elsewhere [11]. Partici-
pants with available data on cannabis use were included in the
present study if they were of at least 18 years of age, provided
written informed consent, and had a history of pancreatitis. Ex-
clusion criteria were other diseases of the exocrine pancreas,
interventions involving the pancreas, malignancy, pregnancy,
celiac disease, cystic fibrosis, history of steroid use, or medi-
cal evaluation or treatment for acute infectious or inflammatory
conditions within the preceding 6 months [11]. Individuals with
contraindications to MR imaging (e.g. electronic device im-
plantations, metallic foreign bodies) were also excluded from
this study. The study was approved by the Health and Disability
Ethics Committee (13/STH/182) and conducted in compliance
with the ethical standards of the responsible institution on hu-
man subjects as well as with the Helsinki Declaration.

Ascertainment of cannabis use

At the time of MR acquisition, participants completed a
standardized questionnaire, which asked about their history
of cannabis use. They were first asked “Have you ever used
marijuana?”. Those who answered no to this question were cat-
ergORIZED as “never” cannabis users. Those who answered yes to
this question were “ever” cannabis users, and were then asked
“How often did/do you use marijuana?”. The options for this
question were: 1) Occasionally; 2) Less than once per week; 3)
Once per week; 4) 2-3 times per week; and 5) Daily. Answers
to this question were used to categorize participants into “non-
regular” and “regular” cannabis users, where “non-regular” us-
ers were defined as less than once per week (answers 1-2) and
“regular” users defined as one or more times per week (answers
3-5), respectively, in line with published literature [12].

MR imaging protocol

All participants underwent abdominal MR scans, performed
at the Centre of Advanced Magnetic Resonance Imaging (The
University of Auckland) using a 3.0 Tesla MAGNETOM
Skyra MR scanner (Siemens, Erlangen, Germany). The de-
tailed protocol was described elsewhere [11]. In brief, while
in the supine position, participants were asked to breath-hold
during end expiration. Axial T1-weighted volumetric inter-
polated breath-hold examination Dixon sequence was ap-
plied with the following parameters: true form abdomen shim
mode; field of view (FOV), 420 mm; base resolution, 320;
echo time (TE), 1.27 ms, 2.5 ms; repetition time (TR), 3.85
ms; flip angle, 9; pixel bandwidth, 920 Hz; slice thickness,
5 mm. Four types of images were generated: in-phase, out-
of-phase, fat and water images. These images were retrieved
from the MR scanner and exported as DICOM files after
which they were analyzed using ImageJ software (National
Institutes of Health, USA).

Subcutaneous fat volume (SFV) and visceral fat volume (VFV)

Quantification of SFV and VFV was conducted from the L2
to the L5 levels by segmentation of the subcutaneous and vis-
ceral fat compartments using the free-hand tool of ImageJ.
Care was taken to exclude all non-adipose tissue, abdominal
organs and blood vessels from the measurement of visceral
Fat. The calculation of volumes required summation of the pixel contents (from the complete series of slices) and multiplying by the total pixel area and slice thickness, as described elsewhere [13].

**Intra-pancreatic fat deposition (IPFD)**

IPFD (%) was measured using the “MR-opsy” technique, as per previously published protocol [14]. In brief, two candidate slices (of 5 mm thickness), with clear visualization of the pancreas, were selected from the series of abdominal MR scans. Three regions of interest were placed in the head, body and tail region of the pancreas for estimation of IPFD (Fig. 1a, b). A thresholding range of 1-20% was applied, to eliminate potential inclusion of the non-parenchymal tissue within the selected region of interest (including blood vessels, visceral fat and the main pancreatic duct), as previously described [14]. IPFD was calculated as the average pancreatic fat fraction of both slices.

Figure 1. Fat phenotypes measurements for a 78-year-old European man who was never smoker and had low-to-moderate average alcohol consumption. (a, b) Intra-pancreatic fat deposition measurement by region of interest placement in the head, body and tail of the pancreas on out-of-phase and fat images. (c, d) Intra-hepatic fat deposition measurement with placement of the voxel within the liver and spectroscopy data using fat and water signals collected at the time of scan acquisition. (e, f) Skeletal muscle fat deposition measurement with selection of total erector spinae muscle area and quantification of fat pixels using the thresholding function of ImageJ.
Participants underwent a mixed meal test at the COSMOS Clinic at the University of Auckland (New Zealand). Venous blood samples were taken at baseline following at least 8 h overnight fast. All participants consumed a standardized mixed meal of BOOST™ drink and venous blood samples were collected at fasting and post-prandially [17]. Glycated hemoglobin (HbA1c) and plasma glucose were analyzed at the tertiary referral medical laboratory, LabPlus (Auckland, New Zealand). Participants’ diabetes status was determined based on their HbA1c level at the time of MR acquisition. Insulin and C-peptide were measured using the MILLIPLEX MAP Human metabolic hormone magnetic bead panel based on the Luminex xMAP technology (Luminex Corporation, Austin). Results were calculated based on the fluorescent reporter signals recorded by the Luminex xPONENT software (MILLIPLEX Analyst 5.1).

Intra-hepatic fat deposition (IHFD)

IHFD (%) was determined using single-voxel MR-spectroscopy. A 20 × 20 × 20 mm voxel was positioned in the right lobe of the liver ≥ 10 mm away from the edge, bile ducts and blood vessels (Fig. 1c). Automated shimming was performed prior to signal acquisition to improve B0 homogeneity. Spectra were acquired using a free-breathing navigator-triggered spin echo acquisition with TR = 3,000 ms, TE = 33 ms, 50 averages. Acquisition time for each spectrum was 5 min. Both water-suppressed and non-water-suppressed spectra were acquired, with the non-water-suppressed spectrum acting as a reference for IHFD quantification (Fig. 1d). Spectra were processed and analyzed using SIVIC (San Francisco, CA, USA) [15]. MR spectroscopy-derived IHFD was defined as follows: fat fraction = area under fat peak/(area under combined fat and water peaks) × 100%.

Skeletal muscle fat deposition (SMFD)

Total muscle area and intra-muscular fat area of erector spinae muscles were measured using a single axial slice at the lower endplate of L3 vertebra, as it had previous been demonstrated that the L3 level is a reliable marker for total skeletal muscle fat [16]. The free-hand tool of ImageJ was used to outline the left and right erector spinae muscles followed by measurement of total pixel content (Fig. 1e) [16]. Further, to calculate the intra-muscular fat area, the threshold-function of ImageJ was used to convert grayscale pixels into binary images, using global histogram-derived method (Fig. 1f). Care was taken not to include extra-muscular fat (i.e. beyond the fascial layer of the erector spinae muscles). Total muscle area and intra-muscular fat area were calculated by multiplying the selected total pixel content with pixel surface area. The ratio of fat-free cross-sectional muscle area to total cross-sectional muscle area was determined by subtracting intra-muscular fat area from the total muscle area and dividing this value by the total muscle area. SMFD was defined as follows: fat fraction = (1 - fat-free cross-sectional muscle area to total cross-sectional muscle area ratio) × 100%.

Inter-observer reliability

Two observers, blinded to participant characteristics and group allocation, measured SFV, VFV, IPFD and SMFD independently. Average measurement values of the two independent measurements were used for all statistical analyses. Inter-observer reliability of measurements was evaluated using intraclass correlation coefficients (ICCs), with <0.5, 0.5 - 0.75, 0.75 - 0.9 and >0.9 being indicative of poor, moderate, good and excellent inter-rater reliability, respectively [11].

Laboratory analyses

Participants underwent a mixed meal test at the COSMOS Laboratory analyses and excellent inter-rater reliability, respectively [11].
(0 g/day), light to moderate (1 - 40 g/day) and excess (> 40 g/day). A binary variable was generated for physical activity status, defined as less or more than 2.5 h of moderate-to-vigorous physical activity per week.

Statistical analyses

All statistical analyses were performed using SPSS for Windows Version 25 (SPSS Inc., Chicago, IL, USA). Extreme values in the data (defined as cases with values/standardized residuals greater than ± 3 standard deviations) were considered outliers and were excluded from the analyses [13]. Data were expressed as median and interquartile range (IQR) or frequency count and percentage (%) for continuous and categorical variables, respectively. Differences in baseline characteristics between never, non-regular and regular cannabis users were examined using analysis of variance (for continuous variables) and Chi-squared test (for categorical variables).

Statistical analyses were performed in two stages. First, to investigate the variation in each fat phenotype (SFV, VFV, IPFD, IHFD and SMFD) explained by cannabis use, the most robust R² metric from univariate linear regression analysis was yielded for each association. Regular and non-regular cannabis users were compared with never cannabis users (reference group). Further, multiple variable linear regression models were used to adjust for potential confounders. For each of the five fat phenotypes as separate dependent variables, three models were constructed: 1) Unadjusted; 2) Adjusted for age, sex and ethnicity; and 3) Adjusted for age, sex, ethnicity, HbA1c, physical activity, tobacco smoking status and alcohol consumption. Values were reported as the B-coefficients, 95% confidence intervals (CIs) and associated P-values.

Second, to investigate the associations between cannabis use and the six indices of insulin sensitivity (HOMA-IS, Raynaud index and Matsuda index) and insulin secretion (HOMA-β, insulinogenic index 30', and insulinogenic index 60'), univariate and multiple variable linear regression models were used. As in the first stage of analysis, regular and non-regular cannabis users were compared with never cannabis users (reference group). To investigate the variation in each insulin sensitivity/secretion index explained by cannabis use, the most robust R² metric from the univariate analysis was yielded for each association. The same three models were constructed as above. A two-sided P-value of less than 0.05 was considered statistically significant in all analyses.

Results

Characteristics of participants

A total of 120 individuals were recruited into the study, of whom 91 were never cannabis users, 21 were non-regular cannabis users and 8 were regular cannabis users. For the fat phenotypes of the total cohort, mean ± standard deviation SFV was 3,090 ± 1,432 cm³, VFV was 2,158 ± 1,140 cm³, IPFD was 9.52±1.79%, IHFD was 10.14±9.40% and SMFD was 15.15±6.81%. For the insulin sensitivity indices of the total cohort, mean ± standard deviation HOMA-IS was 0.67 ± 0.40, Raynaud index was 4.18 ± 8.17 and Matsuda index was 0.72 ± 0.51. For the insulin secretion indices of the total cohort, mean ± standard deviation HOMA-β was 151.28 ± 82.52, insulinogenic index 30' was 0.62 ± 0.52 and insulinogenic index 60' was 0.65 ± 0.44. Characteristics of the study participants according to cannabis use group are presented in Table 1.

Associations between cannabis use and the fat phenotypes in the study groups

The inter-rater reliability of the measurements of the MRI-derived fat phenotypes was as follows: ICC of 0.998 for SFV, 0.997 for VFV, 0.970 for IPFD and 0.984 for SMFD.

Cannabis use contributed the most to variance in IHFD (R² = 9.2%), followed by SMFD (R² = 4.4%), then VFV (R² = 3.4%). Cannabis use contributed the least to variance in IPFD (R² = 0.2%) (Fig. 2). IHFD was significantly greater in regular cannabis users when compared with never cannabis users, in both the unadjusted (P = 0.002) and two adjusted (P = 0.002; P = 0.008) models. Non-regular cannabis users did not have significantly greater IHFD compared with never cannabis users (Table 2). SFV was significantly greater in regular cannabis users when compared with never cannabis users, in the unadjusted analysis (P = 0.024). This was not significant in the two adjusted models (Table 2). There were no significant differences in VFV, IPFD or SMFD when compared with never cannabis users, regardless of their level of cannabis use.

Associations between cannabis use and the insulin traits in the study groups

Cannabis use contributed the most to variance in the Raynaud index (R² = 8.2%) out of all the insulin sensitivity indices. Cannabis use explained 0.6% of variance in both HOMA-IS and the Matsuda index. There were no significant differences in HOMA-IS, Matsuda index, or Raynaud index when compared with never cannabis users, in both the unadjusted and two adjusted models (Table 3).

Cannabis use contributed the most to variance in the insulinogenic index 60' (R² = 20.3%) out of all the insulin secretion indices. Cannabis use explained 4.4% of variance in the HOMA-β index and 1.7% of variance in the insulinogenic index 30'. Insulinogenic index 60' was significantly greater in regular cannabis users when compared with never cannabis users, in both the unadjusted (P = 0.049) and two adjusted (P = 0.003; P = 0.004) models. Non-regular cannabis users did not have significantly higher insulinogenic index 60' compared with never cannabis users (Table 3). There were no significant differences in the insulinogenic index 30' or HOMA-β when compared with never cannabis users, in both the unadjusted...
and two adjusted models (Table 3).

**Discussion**

This study has investigated the association between cannabis use and comprehensively characterized fat phenotypes derived from MR imaging, including visceral and subcutaneous fat as well as ectopic fat in the liver, pancreas and skeletal muscle. Further, this study has presented data on the association between cannabis use and several indices of insulin sensitivity and secretion (both fasting and post-prandial). All MR-derived measurements completed by two independent observers had excellent inter-rater reliability (ICC > 0.9). Further, the used statistical models accounted for potential covariates in both analyses of cannabis use with fat phenotypes and cannabis use with insulin traits. We report two main findings from this study. First, regular cannabis users had significantly greater IHFD (but not SFV, VFV, IPFD or SMFD) compared with never cannabis users. This remained significant after adjusting for age, sex, ethnicity, HbA1c, tobacco smoking, alcohol consumption and physical activity. Moreover, the amount of variance explained by cannabis use was greatest for IHFD (R² = 9.2%). Second, regular cannabis users had significantly greater insulin secretion (as defined by the insulinogenic index 60') when compared with never cannabis users, whereas insulin...
sensitivity was not significantly altered in cannabis users. This study presents novel findings and there are many unknowns regarding the mechanisms that underlie them. We speculate that, at least in part, our results may be attributed to upregulated neurohumoral stimulation from the central ECS, which is supported by evidence of the effect of cannabis use on autonomic regulation of β-cell secretion [18, 19]. The ECS is a neuromodulatory system that acts on both the central and peripheral nervous systems, as well as non-neural peripheral sites [18]. CB1Rs are present in all sites involved in the control of energy homeostasis. While CB1Rs are predominantly found in the central nervous system, lower levels are detected in skeletal muscle, liver, pancreas and gastrointestinal tract [18]. CB1Rs are among the most abundant G-protein coupled receptors expressed in the brain and are present in the forebrain, basal ganglia, cerebellum, hippocampus and cerebral cortex [20]. The ECS is the main central system involved in appetite-regulation and food intake. Pharmacologic administration of CB1 agonists increases food consumption and, in the case of Δ-9-THC, has been observed even in satiated animals [4, 5, 20]. Moreover, systemic CB1R blockade by SR141716A (rimonabant) reduces food intake in both fed and fasted animals [20]. Its modulation of food intake is believed to be two-fold. First, through mesolimbic pathways and dopaminergic release, it activates the reward mechanisms for food consumption (usually of highly palatable foods). Second, through hypothalamic regulation of the level and/or activity of anorectic and orexigenic mediators that induce appetite. CB1R expression in the hypothalamus, a key area for the regulation of energy balance, is relatively low but highly efficient, and activation stimulates the release of key neurotransmitters (including but not limited to noradrenaline and serotonin). Both noradrenaline and serotonin are key neurohumoral regulators of insulin secretion, and they can exert both stimulatory and inhibitory effects on the pancreatic β cells [21]. Acute increases in sympathetic neurotransmission initially inhibit insulin secretion but over longer periods elevate its plasma levels. While it is assumed that noradrenaline mediates the inhibitory actions of sympathetic stimulation on glucose-stimulated insulin secretion, it is accepted that it also has stimulatory actions through its direct activation of β2-adrenoreceptors and α2-adrenoreceptors on both the β and α cells [22]. Serotonin can also influence insulin output, either by stimulating cellular receptors to enhance release of noradrenaline from islet sympathetic terminals, or through direct intracellular regulation of insulin release [21]. It could be speculated then that stimulation of the central ECS enhances autonomic neurotransmission to the pancreatic islet cells and induces supraphysiologic insulin secretion. Insulin drives glucose uptake into cells where it is either stored or used directly as fuel. The predominant sites of glucose uptake are the liver, skeletal muscle and adipocytes. Ectopic fat deposition typically reflects the failure of intracellular lipid homeostasis to prevent lipotoxicity as a consequence of accumulation of saturated lipid-storing white adipocytes [23]. In a hyperinsulinemic state, glucose uptake may exceed the energy requirements of the cells and lead to excess glucose storage, driving fatty infiltration into organs such as the liver. The above arguments suggest that significantly increased IHFD in regular cannabis users observed in the present study might be due to chronic stimulation of CB1Rs promoting excess insulin secretion. Cannabinoids are also known to activate PPARγ, the isoform of PPARs that mediates some of the metabolic effects seen with both endo- and phyto-cannabinoids. Palomba and colleagues [24] recently recognized a potential role for PPARγ in the modulation of leptin activity by hypothalamic CB1Rs, which further supports the effect of cannabis through CB1R mediation.

We found cannabis use was significantly associated with
Table 2. Associations Between Cannabis Use and the Studied Fat Phenotypes

| Fat phenotype                          | Cannabis use | Mean ± SD   | Model 1 | Model 2 | Model 3 |
|----------------------------------------|--------------|-------------|---------|---------|---------|
|                                        |              | B           | 95% CI  | P-value | B       | 95% CI  | P-value | B       | 95% CI  | P-value |
|                                        |              | Lower       | Upper   |         | Lower   | Upper   |         | Lower   | Upper   |         |
| Subcutaneous fat volume, cm³           | Never        | 2,769 ± 1,421 | Ref     | Ref     | Ref     | Ref     | Ref     | Ref     | Ref     | Ref     |
|                                        | Non-regular  | 2,928 ± 266  | -0.008  | -759.8  | 706.9   | 0.943   | -0.043  | -911.6  | 617.3   | 0.702   |
|                                        | Regular      | 4,061 ± 1,644 | 0.246   | 174.3   | 2467.1  | 0.024   | 0.180   | -190.4  | 2118.9  | 0.100   |
| Visceral fat volume, cm³               | Never        | 2,390 ± 1,134 | Ref     | Ref     | Ref     | Ref     | Ref     | Ref     | Ref     | Ref     |
|                                        | Non-regular  | 2,190 ± 350  | 0.006   | 596.7   | 631.7   | 0.955   | -0.032  | -674.7  | 495.9   | 0.762   |
|                                        | Regular      | 2,522 ± 1,653 | 0.072   | 647.4   | 1272.9  | 0.519   | 0.048   | -674.8  | 1093.4  | 0.639   |
| Intra-pancreatic fat deposition, %     | Never        | 9.7 ± 1.7    | Ref     | Ref     | Ref     | Ref     | Ref     | Ref     | Ref     | Ref     |
|                                        | Non-regular  | 9.9 ± 1.7    | -0.070  | 1.37    | 0.71    | 0.529   | -0.028  | 1.24    | 0.97    | 0.811   |
|                                        | Regular      | 9.1 ± 1.8    | 0.007   | 1.53    | 1.63    | 0.952   | 0.034   | 1.38    | 1.87    | 0.764   |
| Intra-hepatic fat deposition, %        | Never        | 9.2 ± 8.4    | Ref     | Ref     | Ref     | Ref     | Ref     | Ref     | Ref     | Ref     |
|                                        | Non-regular  | 7.2 ± 2.0    | 0.097   | 2.00    | 5.42    | 0.361   | 0.096   | 2.43    | 5.83    | 0.415   |
|                                        | Regular      | 12.6 ± 7.8   | 0.336   | 3.45    | 15.03   | 0.002   | 0.370   | 3.96    | 16.39   | 0.002   |
| Skeletal muscle fat deposition, %      | Never        | 16.1 ± 5.0   | Ref     | Ref     | Ref     | Ref     | Ref     | Ref     | Ref     | Ref     |
|                                        | Non-regular  | 8.4 ± 1.9    | -0.147  | 6.20    | 1.19    | 0.181   | 0.070   | 1.93    | 4.32    | 0.448   |
|                                        | Regular      | 11.3 ± 5.5   | -0.135  | 9.38    | 6.16    | 0.217   | 0.017   | -4.28   | 5.16    | 0.853   |

Model 1 was unadjusted. Model 2 was adjusted for age, sex and ethnicity. Model 3 was adjusted for age, sex, ethnicity, HbA1c level, tobacco smoking, alcohol consumption and physical activity. SD: standard deviation; B: beta coefficient; CI: confidence interval.
Table 3. Associations Between Cannabis Use and the Studied Insulin Traits

| Insulin traits                        | Cannabis use | Mean ± SD | Model 1 | Model 2 | Model 3 |
|---------------------------------------|--------------|-----------|---------|---------|---------|
|                                       |              |           | B       | 95% CI  | P-value |
|                                       |              |           | Lower   | Upper   |         |
|                                       |              |           |         |         |         |
| Indices of insulin sensitivity        |              |           |         |         |         |
| HOMA-IS (fasting)                     | Never        | 0.68 ± 0.42 | Ref     | Ref     | Ref     |
|                                       | Non-regular  | 0.61 ± 0.35 | -0.020  | -0.536  | 0.911   |
|                                       | Regular      | 0.45 ± 0.27 | -0.079  | -0.745  | 0.480   |
| Raynaud index (fasting)               | Never        | 3.30 ± 2.56 | Ref     | Ref     | Ref     |
|                                       | Non-regular  | 9.28 ± 20.77 | 0.281   | -0.79   | 15.38   |
|                                       | Regular      | 1.99 ± 1.71 | -0.054  | -26.76  | 12.33   |
| Matsuda index (post-prandial)         | Never        | 0.72 ± 0.53 | Ref     | Ref     | Ref     |
|                                       | Non-regular  | 0.74 ± 0.35 | -0.032  | -0.732  | 0.609   |
|                                       | Regular      | 0.46 ± 0.38 | -0.072  | -0.976  | 0.643   |
| Indices of insulin secretion          |              |           |         |         |         |
| HOMA-β (fasting)                      | Never        | 142.6 ± 75.2 | Ref     | Ref     | Ref     |
|                                       | Non-regular  | 223.8 ± 134.6 | 0.212   | -31.78  | 126.42  |
|                                       | Regular      | 155.9 ± 44.3 | 0.167   | -50.49  | 140.20  |
| Insulinogenic index 30' (post-prandial) | Never     | 0.66 ± 1.27 | Ref     | Ref     | Ref     |
|                                       | Non-regular  | -0.03 ± 0.25 | -0.128  | -2.823  | 1.475   |
|                                       | Regular      | 1.91 ± 0.76 | -0.026  | -2.284  | 2.015   |
| Insulinogenic index 60' (post-prandial) | Never     | 0.53 ± 1.12 | Ref     | Ref     | Ref     |
|                                       | Non-regular  | 0.28 ± 0.72 | -0.156  | -2.203  | 0.980   |
|                                       | Regular      | 1.76 ± 1.49 | 0.408   | 0.010   | 3.187   |

Model 1 was unadjusted. Model 2 was adjusted for age, sex and ethnicity. Model 3 was adjusted for age, sex, ethnicity, HbA1c level, tobacco smoking, alcohol consumption and physical activity. SD: standard deviation; B: beta coefficient; CI: confidence interval.
the insulinogenic index 60’ (reflecting increased insulin secretion at 60 min following mixed meal) but no significant changes in insulin sensitivity. The lack of observed difference in insulin sensitivity is unsurprising, given the overall studied population did not have marked abnormalities of glucose metabolism. Moreover, the group of regular cannabis users had lower median HbA1c (35 mmol/mol) than both nonregular (37 mmol/mol) and never (39 mmol/mol) users (Table 1). Increased insulin secretion may reflect an augmented insulin response following cannabis use and suggests the potential role of CB1Rs in the gastrointestinal tract. DiPatrizio and colleagues [25] found that a diet high in fat altered intestinal endocannabinoid levels, which upregulated CB1Rs in vagal nerve fibres. Investigations of β-cell function historically have used intravenous glucose. As far as the oral route is concerned, the relationship becomes more complex and is influenced by the rate of glucose absorption, neural activation, and incretin action. Incretins, such as glucose-dependent insulinotropic polypeptide (GIP) and glucagon like peptide-1 (GLP-1), enhance insulin secretion in response to nutrients and are responsible for the augmented response to glucose when given orally rather than intravenously [26]. Several animal models have isolated CB1Rs on enteroendocrine cells in rodents and have suggested these play a role in the regulation of incretin secretion [27]. Chia and colleagues [26] demonstrated a significant increase in fasting and 60 min post-oral glucose tolerance test GIP levels following administration of a CB1 agonist, but no change in GLP-1 levels. They also found an increase in GIP levels in obese individuals, suggesting increased GIP is likely a consequence of increased endocannabinoid levels. Further, rodents fed dietary linoleic acid (which increased weight gain) were found to have elevated endocannabinoid levels in both the liver and the gut [28]. Linoleic acid is also an endogenous activator of PPARs, which have further been implicated in the regulation of metabolism and energy homeostasis. The above arguments may explain why we found increased insulin secretion, but no significant change in insulin sensitivity.

Another possible explanation for our findings is that peripheral CB1Rs may be preferentially distributed in particular sites and result in augmented physiological effects from stimulation of the ECS. Peripheral CB1Rs play an important role in glucose homeostasis by modulating lipogenesis in the liver and adipose tissue. CB1R activation enhances de novo lipogenesis and triglyceride accumulation in hepatocytes and adipocytes, while minimizing lipolysis and fatty acid oxidation [28]. Liu and colleagues [29] demonstrated a reversal of diet-induced IHFD through peripheral CB1R blockade in rodents. They also found increased fatty acid oxidation with CB1R blockade in mice both with and without diet-induced obesity. This suggests a possible propensity for cannabis-related effects on hepatocytes over other peripheral cell types, due to more concentrated CB1Rs than in other peripheral sites. Alternatively, the differing effects of Δ-9-THC on fat deposition may be due to variable expression of CB1 and CB2 receptors in the peripheral ECS. While CBR expression has been shown in the islets of Langerhans, evidence is conflicting on the receptor subtypes in each of the different islet cells [19]. Previous in vivo experiments on rodents demonstrated metabolic derangement with reduced glucose uptake following administration of CB1 agonists, but the opposite with CB2 agonists [30]. Given the lack of research conducted on CB2Rs, both in their relative distribution and functions, we can only speculate their function in both metabolic dysfunction and ectopic fat deposition. Greater understanding of the interplay between the two types of cannabinoid receptors is needed to understand how cannabis differentially affects various organs and systems in human studies. Given that the relative expressions of CB1 to CB2 receptors in the peripheral tissues are currently unknown, variability in the level of expression in the liver compared with the pancreas or skeletal muscle may explain the increased IHFD compared with other fat phenotypes.

The present study has several limitations. First, the questionnaire asked about cannabis use did not specify the type of cannabis product or the mode of administration. Second, the relative amounts of Δ-9-THC versus non-Δ-9-THC cannabinoids in the used cannabis preparations are unknown. However, our question referred to “marijuana”, which is most frequently referred in the literature as recreational cannabis (with much greater Δ-9-THC than non-Δ-9-THC) [4]. Therefore, we assumed the effects seen were more the consequence of Δ-9-THC and not the CBD components (which are frequently used for therapeutic purposes). Third, there is a risk of under-reporting when asking about substance use. This would have resulted in our study yielding more conservative findings. Fourth, this was a post-hoc analysis of data on participants recruited into a study on sequelae of pancreatitis. Although all the 120 individuals had a history of mild pancreatitis, no individual had signs of acute inflammation on their MR scans and all had complete clinical resolution of their pancreatitis prior to MR acquisition [31, 32]. This also provided a relatively homogenous group of individuals who had a detailed clinical phenotype [33, 34]. They were also potentially a population with greater pain concerns due to their history of pancreatitis and subsequent impaired quality of life [35]. Hence, they could be considered an at-risk group for cannabis use or dependence, or potentially could be considered a group that may benefit from the therapeutic use of cannabis if delivered in controlled conditions [36]. Fifth, some individuals with diabetes received insulin, which may have influenced abdominal body fat composition. However, the effect is unlikely to be material as none of the participants in the regular cannabis use group and only three participants in the other two groups received insulin. Sixth, the number of regular cannabis users was rather limited. This study is considered pilot and hypothesis-generating. Power calculations of future studies in the field will benefit from the data presented here. Seventh, the analyses did not adjust for BMI or waist circumference. Whilst it is acknowledged that these parameters might have influenced the studied associations, it is also acknowledged that these are less accurate than MR-derived visceral and subcutaneous fat compartments (used as dependent variable in the present study) for metabolic health. Adjusting the studied associations for BMI or waist circumference would have introduced multicollinearity into the regression analyses and therefore undermined the statistical assumptions [13]. Last, given the cross-sectional nature of the study, inference of causality between cannabis use and fat phenotypes or insulin traits cannot be made. However, to the best of our knowledge, no longitudinal study has been conducted in humans on cannabis use. Future studies are now war-
ranted to track temporal changes in fat phenotypes with regular cannabis use, both recreationally and therapeutically.

In conclusion, our study presents first data on the differential association between cannabis use and MR-derived fat phenotypes, including SFV, VFV, IPFD, IHFD and SMFD. Our results show significantly greater IHFD (but not SFV, VFV, IPFD or SMFD) in regular cannabis users compared with never cannabis users, both in the unadjusted and adjusted analyses. Cannabis use contributes most to variance in IHFD, and least to IPFD. Our results show significantly altered insulin secretion (but not insulin sensitivity) in regular cannabis users compared with never cannabis users. These findings provide novel insights into the relationship between cannabis use, different abdominal fat phenotypes, and insulin traits, the knowledge on which will likely increase in the future.

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Conflict of Interest

None to declare.

Informed Consent

Informed consent was given by all study participants.

Author Contributions

Study concept and design: MSP. Patient recruitment: GAR, JC, JK, CES, AM. Acquisition of data: CES, JK, AM. Analysis and interpretation of data: CES and JC. Statistical analysis: CES and JC. Drafting of manuscript: CES. Critical revision of the manuscript: JC, GAR, AM, MSP. Study supervision: MSP.

Data Availability

The authors declare that data supporting the findings of this study are available within the article.

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