The intake of high-fat diets induces an obesogenic-like gene expression profile in peripheral blood mononuclear cells, which is reverted by dieting

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Abstract
Peripheral blood mononuclear cells (PBMC) are increasingly used for nutrigenomic studies. In this study, we aimed to identify whether these cells could reflect the development of an obesogenic profile associated with the intake of high-fat (HF) diets. We analysed, by real-time RT-PCR, the dietary response of key genes related to lipid metabolism, obesity and inflammation in PBMC of control rats, rats fed a cafeteria or a commercial HF diet and rats fed a control diet after the intake of a cafeteria diet (post-cafeeteria model). Cafeteria diet intake, which resulted in important overweight and related complications, altered the expressions of most of the studied genes in PBMC, evidencing the development of an obesogenic profile. Commercial HF diet, which produced metabolic alterations but in the absence of noticeably increased body weight, also altered PBMC gene expression, inducing a similar regulatory pattern as that observed for the cafeteria diet. Regulation of carnitine palmitoyltransferase I (Cpt1a) mRNA expression was of special interest; its expression reflected metabolic alterations related to the intake of both obesogenic diets (independently of increased body weight) even at an early stage as well as metabolic recovery in post-cafeeteria animals. Thus, PBMC constitute an important source of biomarkers that reflect the increased adiposity and metabolic deregulation associated with the intake of HF diets. In particular, we propose an analysis of Cpt1a expression as a good biomarker to detect the early metabolic alterations caused by the consumption of hyperlipidic diets, and also as a marker of metabolic recovery associated to weight loss.

Key words: Blood cells: Biomarkers: Obesity: High-fat diets

Peripheral blood mononuclear cells (PBMC) represent a fraction of WBC composed of monocytes and lymphocytes. In addition to their role in immunity, PBMC possess specific characteristics that make them interesting biological material to be used in the search for molecular biomarkers, including early biomarkers of disease. PBMC can express a large proportion of the genome(1) and can be easily and repeatedly collected in sufficient quantities to perform transcriptomic analysis, in contrast to the more invasive sampling of the liver, adipose tissue or the other tissues.(2) They have a fast turnover and are in continuous interaction with most parts of the body, and thus they can reflect gene expression profiles of certain pathologies and can be used for clinical diagnosis(3–5). Moreover, PBMC can respond to hormonal changes reflecting the metabolic response of different organs, as they possess receptors for hormones such as insulin, glucagon or leptin(6–9).

Recently, gene expression analysis in PBMC has become a key target for nutrigenomic studies. It is well known that PBMC can reflect the metabolic effect of different diets such as hyperlipidic diets on gene expression, especially on lipid and cholesterol metabolism and on inflammatory response(10–12). These cells can also respond to specific nutrients such as PUFA, polyphenols or L-carnitine(13–14) and to changes in feeding conditions (fasting/feeding)(10,11,15,16). PBMC are also good biological material to perform obesity studies. Obesity is linked to a pro-inflammatory state(17), and in humans obesity and diabetes affect PBMC gene expression of different cytokines such as resistin, Tnfα and IL-6(18,19). In addition, we have previously described in rodents that these cells reflect alterations related to diet-induced obesity, such as impairment in the response to feeding conditions, even in early stages of body weight gain(10,11), as well as metabolic recovery associated with weight loss based on the recovery of fasting response(20). In fact, we have previously presented PBMC as a potential source of obesity biomarkers(21).

Our previous studies analysing the effect of high-fat (HF) diets on PBMC gene expression have mainly been based on the study of genes involved in lipid metabolism. In this study, we aimed to study a pool of obesity biomarkers, linked to different processes related to obesity development (lipid metabolism, obesity regulation and inflammation), which could be altered in response to the intake of HF diets since an early stage as well as

Abbreviations: Cpt1a, carnitine palmitoyltransferase I; HF, high-fat; PBMC, peripheral blood mononuclear cells; PCAF, post-cafeeteria.

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to describe their reversibility after withdrawal to a control balanced diet. Moreover, different types of HF diets were used – a commercial HF and a cafeteria diet – inducing different degree of overweight and metabolic alterations. Description of early and flexible/reversible obesity biomarkers is extremely useful in terms of development of prevention strategies aimed at maintaining proper body weight.

**Methods**

**Animals, diets and experimental design**

The animal protocol followed in this study was reviewed and approved by the Bioethical Committee of the University of the Balearic Islands, following its guidelines for the use and care of laboratory animals. We analysed gene expressions in PBMC of male Wistar rats (Charles River Laboratories España, SA) from a previous study(22). Male Wistar rats aged 2 months were distributed into four groups fed different diets for 4 months: a control group (control; n 6) fed a control diet (with 10 kcal% from fat, ref. D12450B; Research Diets, Brogaarden); a HF group (n 7) fed a commercial hyperlipidic diet (with 60 kcal% from fat, ref. D12492; Research Diets, Brogaarden); a cafeteria group (CAF; n 6) fed cafeteria diet plus control diet to ensure nutritional requirements of the animals (CAF group ingested 62 kcal% from fat); and a post-cafeteria group (PCAF; n 6) fed for 2 months with cafeteria diet plus control diet and then for another 2 months exclusively with the control diet. The gross composition of the diets is described in Table 1. Animals were housed in pairs at 22°C with a 12 h light–12 h dark cycle. Food and water were provided ad libitum and body weight was measured twice a week. At the end of the experimental period, animals were killed at the beginning of the light cycle by decapitation, and different white adipose tissue depots, both visceral (epididymal, mesenteric and retroperitoneal) and subcutaneous (inguinal), were harvested and weighed to determine adiposity.

**Peripheral blood mononuclear cells isolation**

From the age of 2–6 months, blood samples (1.5–2–5 mL) were collected monthly in ad libitum-fed conditions from the saphena vein at the beginning of the light cycle (08.00–10.00 hours), using heparin in NaCl (0.9%) as anticoagulant. After blood collection, PBMC were isolated by Ficoll gradient separation, according to the instructions indicated by the manufacturer (GE Healthcare Bio Sciences), with some modifications. In brief, the anticoagulant-treated blood was diluted with an equal volume of balanced salt solution, which was prepared by mixing two stock solutions (1:10): solution A (5·5 ms-anhydroxy d-glucose, 5 ms-CaCl₂.2H₂O, 0·98 ms-MgCl₂.6H₂O, 5·4 ms-KCl, 1·5 ms-Tris) and solution B (140 ms-NaCl). Next, the blood was layered carefully over Ficoll without intermixing (1·5 mL of Ficoll for 2 mL of blood mixed with balanced salt solution) in a centrifuge tube and centrifuged at 900 g for 40 min at 20°C. PBMC, together with platelets, were harvested from the interface between Ficoll and sample layers. This layer was then centrifuged in balanced salt solution at 400 g for 10 min at 20°C to wash PBMC and to remove the platelets. Every month, blood samples were collected from the saphena vein, stored at room temperature for 1 h and then centrifuged at 1000 g for 10 min at 4°C to collect the serum.

**Measurement of adiposity, circulating parameters and homeostatic model assessment of insulin resistance**

Fat and lean mass were determined using an EchoMRI-700™ (Echo Medical Systems LLC). Lipid content in the liver was determined using the Folch method(23). Circulating levels of

| Table 1. Diet composition: (a) macronutrient proportion and (b) diet fatty acid profile* |
|---------------------------------|----------------|----------------|----------------|----------------|----------------|----------------|
|                                 | Control diet  | HF diet        | CAF diet       | CAF diet + control diet |
| (a) Macronutrient proportion    |                |                |                |                |
| Carbohydrates                   | 67·3           | 70             | 26·3           | 20             | 21·3           | 26             | 22·5           | 27             |
| Fats                            | 4·3            | 10             | 34·9           | 60             | 23·2           | 64             | 22·5           | 62             |
| Proteins                        | 19·2           | 20             | 26·2           | 20             | 8·2            | 10             | 8·8            | 11             |
| Total                           | 82·8           | 100            | 72·1           | 100            | 57·7           | 100            | 49·2           | 100            |
| kJ/g                            | 1.612          | 2.192          | 1.368          | 1.368          |
| kcal/g                          | 3.85           | 5.24           | 3.27           | 3.27           |
| (b) Diet fatty acid profile     |                |                |                |                |
| Fatty acid profile              |                |                |                |                |
| SFA (%)                         | 25·1           | 37·1           | 46·9           |                |
| MUFA (%)                        | 34·7           | 48·0           | 43·4           |                |
| PUFA (%)                        | 40·2           | 16·9           | 9·40           |                |
| Linoleic:linolenic ratio        | 6·86           | 7·84           | 11·7           |                |
| Total n-6:n-3 ratio             | 7·00           | 8·77           | 4·44           |                |

HF, high fat; CAF, cafeteria.

* Control (D12450B) and HF (D12492) diets were from Research Diets and were purchased from Brogaarden and served as dry pellets. Cafeteria HF diet containing different fatty acids was prepared as previously described(11,21), and was administered while allowing free access to control diet. (a) Percentage of the macronutrients in grams and kcal as well as kJ/g (kcal/g) of foods of the commercial HF and the cafeteria diet; in addition, we provide macronutrient proportions ingested by the cafeteria group considering the intake of the cafeteria diet plus control diet. (b) Fatty acid profile of the diets (SFA, MUFA and PUFA) expressed as % of total amount of fat, linoleic:linolenic acid and total n-3:n-6 dietary fatty acid ratios. Cafeteria diet composition was calculated based on the foodstuffs ingested by the animals(27). Of note, both HF diets had similar sucrose content (approximately 9 g/100 g of food).
glucose, leptin and TAG were measured in serum samples obtained at the end of the experimental period. Glucose concentration was measured using an Accu-Chek Glucometer (Roche Diagnostics), leptin concentration was measured using an enzymatic colorimetric kit (Mercodia AB), and glucose concentration was determined using the formula of Matthews et al.\textsuperscript{[23]} to assess insulin resistance.

**Total RNA isolation**

Total RNA from PBMC samples was extracted using Tripure Reagent (Roche Diagnostics) and then purified with E.Z.N.A. MicroElute RNA Clean Up (Omega Bio-tek). RNA yield was quantified in a NanoDrop ND 1000 spectrophotometer (Nanodrop Technologies), and its integrity was confirmed using agarose gel electrophoresis.

**Real-time RT-PCR analysis**

Real-time RT-PCR was carried out to determine mRNA expression levels of obesity-related genes in PBMC samples. Total RNA of 80 ng was denatured at 90°C for 1 min and then reverse transcribed to complementary DNA (cDNA) using M-MuLV RT (Applied Biosystems) at 42°C for 60 min and a final step of 5 min at 99°C in a 2720 Thermal Cycler (Applied Biosystems). Each PCR was performed from a diluted (1/5) cDNA template, forward and reverse primers (5 µM) and Power SYBER Green PCR Master Mix (Applied Biosystems) in a total volume of 11 µl, with the following profile: 10 min at 95°C, followed by a total of forty temperature cycles (15 s at 95°C and 1 min at 60°C) with a final cycle of 15 s at 95°C, 1 min at 60°C and 15 s at 95°C. In order to verify the purity of the products, a melting curve was produced after each run according to the manufacturer’s instructions. The threshold cycle (C\textsubscript{T}) was calculated by the instrument’s software (StepOne Software version 2.0; Applied Biosystems), and the relative expression of each mRNA was calculated as a percentage of control rats, using the 2\textsuperscript{−ΔΔCt} method\textsuperscript{[25]}. Data were normalised against the reference gene guanosine diphosphate dissociation inhibitor 1 (Gdi1). Gdi1 was chosen based on microarray data, which showed its stability in control and diet-induced obese animals\textsuperscript{[16]}. Primers for the different genes are described in Table 2. All primers were obtained from Sigma Genosys (Sigma-Aldrich Quimica SA).

**Statistical analysis**

All data are expressed as means with their standard errors. Differences between groups were analysed using ANOVA, and least significance difference (LSD) post hoc was used when differences were statistically significant (P<0.05). The effect of fasting was analysed using Student’s t test. Analyses were performed with SPSS for windows (SPSS). Threshold of significance was defined at P<0.05, and is indicated when different.

**Results and discussion**

**Body weight, adiposity and circulating parameters**

As we have previously described using the same set of animals\textsuperscript{[22]}, cafeteria diet intake resulted in a progressive increase in body weight; rats were 18% (1st month), 27%, (2nd month), 33% (3rd month) and 37% overweight by the end of the experimental period (4th month). This increased body weight was accompanied by progressively increased adiposity every month (27, 35, 43 and 46% of body weight as fat), with increased serum leptin levels and decreased circulating TAG due to a reduction in VLDL production rate and an increase in TAG removal from the blood\textsuperscript{[26]}. These animals had drastic alterations related to the metabolic syndrome, including hyperglycaemia, insulin resistance (evidenced by increased HOMA-IR index) and increased liver lipid content (Table 3). However, animals fed a commercial HF diet presented an initial increased body weight, which was significant after the 1st month of diet (data not shown), but was normalised thereafter, and therefore these animals did not present increased body weight at the end of the experimental period. Nevertheless, HF animals had increased adiposity throughout the experimental period (22, 28, 32 and 35% of body weight as fat after 1, 2, 3 and 4 months of dietary treatment, respectively) and metabolic alterations, although not so marked as those observed with cafeteria diet feeding. In these animals, as in cafeteria diet-fed animals, we observed increased circulating leptin, lower circulating TAG levels and increased fat deposition in the liver (Table 3). In spite of increased adiposity (because of increased

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### Table 2. Nucleotide sequences of primers and amplicon size used for real-time RT-PCR amplification

<table>
<thead>
<tr>
<th>Genes</th>
<th>Forward primer (5′–3′)</th>
<th>Reverse primer (5′–3′)</th>
<th>Amplicon size (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Apob48r</td>
<td>GGGCTACATCAGGCTTGTAG</td>
<td>TTCTCCCTCTACACCTTC</td>
<td>150</td>
</tr>
<tr>
<td>Cebpβ</td>
<td>GAGCGACAGATACAGATGCCG</td>
<td>TGGTGCTCAGTCAGGTTG</td>
<td>96</td>
</tr>
<tr>
<td>Cpt1a</td>
<td>GTCGGACACATTACAGGACAT</td>
<td>TGGACACCATAGAGGACAG</td>
<td>250</td>
</tr>
<tr>
<td>Fasn</td>
<td>CGCGGATCTGATGGCATTAT</td>
<td>ACACAGGGACGGATAATGC</td>
<td>222</td>
</tr>
<tr>
<td>Il-6</td>
<td>ATATTTTCAGGAGGAGATTTAG</td>
<td>GTGCAATCATGGCTGTATCA</td>
<td>79</td>
</tr>
<tr>
<td>Slc27a2</td>
<td>TTTCCGAGCCAGCAGTTG</td>
<td>TCTCTCAGTAGGATTTCC</td>
<td>157</td>
</tr>
<tr>
<td>Sreb1a</td>
<td>CCGACCACCGTTACACC</td>
<td>GCGCTGGGTCTTCTTGT</td>
<td>198</td>
</tr>
<tr>
<td>Tnfo</td>
<td>CCGATTCTGCGATTCTACCC</td>
<td>TGCGTTCCAGAAGCAATGAC</td>
<td>237</td>
</tr>
<tr>
<td>Gdi1 (reference gene)</td>
<td>CCGCACAAGGCGAATACATC</td>
<td>GACTCTCTGAAAGCCTACAA</td>
<td>159</td>
</tr>
</tbody>
</table>

\textsuperscript{Apob48r, apoprotein B48 receptor; Cebpβ, CCAAT/enhancer-binding protein β; Cpt1a, carnitine palmitoyltransferase 1; Fasn, fatty acid synthase; Slc27a2, solute carrier family 27 member 2; Sreb1a, sterol regulatory element-binding protein 1a; Gdi1, guanosine diphosphate dissociation inhibitor 1.}
size of both visceral and subcutaneous adipose depots analysed, data not shown), animals in the commercial HF group maintained their body weight because of a decrease in lean mass, evidencing a loss in muscle mass. The more harmful effect of the cafeteria diet in comparison with the commercial HF diet seems not to be related to a different fatty acid profile or to a different sucrose content of the diets (as seen in Table 1), but rather to increased energy intake resulting from voluntary and persistent hyperphagia, which we have previously reported for highly palatable cafeteria dietary feeding(27) in comparison with what occurred in the commercial HF group, in which we observed an increased energy intake during the 1st month, but this hyperphagia disappeared thereafter and even decreased during the last month of dietary treatment(22). Animals in the PCAF had an important decrease in body weight and adiposity, but this was not completely reverted (Table 3). PCAF animals displayed an important improvement in their metabolic parameters, normalisation in serum leptin, glucose and TAG levels, and normalisation of liver lipid content. This experimental design is of special interest for the identification of biomarkers as we can compare the effect of two hyperlipidic diets, both causing health problems, but one of them, the commercial HF diet, was not associated with increased body weight but with increased body fat. Moreover, the post-cafeteria model gives the opportunity to analyse body weight loss and its related metabolic recovery.

Regulation of mRNA expression by the intake of hyperlipidic diets in peripheral blood mononuclear cells

To identify early markers of obesity, we screened for alterations in the expressions of a set of selected genes (involved in lipid metabolism, obesity and inflammation) in PBMC, after chronic intake of the two studied HF diets, at different times of diet exposure. Parts of the studied genes were selected on the basis of their potential usefulness according to a previous microarray analysis performed by our group in PBMC of obese animals(21) – that is, the case of the lipid metabolism genes studied and of the obesity-related gene Slc27a2.

Lipid metabolism genes. Proper lipid metabolism regulation is crucial in order to handle the increased fat intake provided by hyperlipidic diets. Coincident to what was observed in key energy homeostatic tissues such as adipose tissue or liver in the same animals(22) at the end of the experimental period, the intake of the HF diets studied increased PBMC expression of a key β-oxidation gene, Cpt1a, but inhibited the expression of the lipogenic genes Srebp and Fasn (Fig. 1), reflecting a physiological adaptation to increased fat intake. For Cpt1a, a clear regulatory effect was also observed at different times of dietary treatment. Cpt1a codes for carnitine palmitoyltransferase I, an enzyme that enables the entrance of long-chain fatty acids across the inner mitochondrial membrane for subsequent β-oxidation. Its mRNA expression turned out to be of special interest to be considered as a marker of adiposity as it was equally increased as a result of the intake of both hyperlipidic diets (commercial HF and cafeteria diet) at the age of 4 months (after 2 months of diet), and this increased expression was maintained 1 month later. Unexpectedly, at the end of the experimental period, Cpt1a expression was only observed in the PCAF group. We also assessed the expression of two lipogenic genes – Fasn and Srebp1a. Fasn codes for fatty acid synthase, an enzyme directly involved in fatty acid synthesis, and Srebp1a codes for a transcription factor that is critical for the expression of numerous genes involved in fatty acid synthesis. It is well described that the expressions of lipogenic genes are decreased by HF diets in order to decrease de novo lipogenesis, thereby promoting the use of fatty acids from the diet(29). The two HF diets studied decreased Fasn and Srebp1a mRNA expressions; however, this decrease required long-term exposure, as it was not observed until the end of the experimental period (after 4 months of dietary treatment). Moreover, the expressions of these genes were not recovered after withdrawal of the

Table 3. Characterisation of animals of the different groups
(Mean values with their standard errors; n = 6–7)*

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>HF</th>
<th>CAF</th>
<th>PCAF</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weight (g)</td>
<td>486a,b</td>
<td>502a,b</td>
<td>668a,b</td>
<td>534a,b</td>
</tr>
<tr>
<td>Fat mass (g)</td>
<td>110a,b</td>
<td>177a,b</td>
<td>296a,b</td>
<td>142a,b</td>
</tr>
<tr>
<td>Lean mass (g)</td>
<td>325a,b</td>
<td>260a,b</td>
<td>315a,b</td>
<td>339a,b</td>
</tr>
<tr>
<td>Leptin (μg/l)</td>
<td>14.4±0.26</td>
<td>19.2±0.88</td>
<td>43.2±2.5</td>
<td>12.3±2.1</td>
</tr>
<tr>
<td>Glucose (mmol/l)</td>
<td>6.00±0.17</td>
<td>6.33±0.22</td>
<td>6.67±0.11</td>
<td>6.28±0.33</td>
</tr>
<tr>
<td>TAG (mg/ml)</td>
<td>2.05±0.15</td>
<td>1.56±0.19</td>
<td>1.06±0.11</td>
<td>3.07±0.21</td>
</tr>
<tr>
<td>HOMA-IR</td>
<td>3.90±0.76</td>
<td>4.60±1.02</td>
<td>7.42±1.48</td>
<td>6.98±1.07</td>
</tr>
<tr>
<td>Liver lipid content (mg/g tissue)</td>
<td>12.9±3.7</td>
<td>36.8±4.5</td>
<td>101.2±24</td>
<td>24.3±8.2</td>
</tr>
</tbody>
</table>

* HF animals were fed a commercial HF diet and CAF animals a CAF diet plus control diet for 4 months; PCAF animals were fed a CAF diet plus control diet for 2 months and a control diet for 2 months more. LSD post hoc was used after ANOVA.

HF, high fat; CAF, CAF, cafeteria; PCAF, post-cafeteria; HOMA-IR, homoeostatic model assessment of insulin resistance; LSD, least significance difference.
cafeteria diet. Thus, according to our data, only Cpt1a gene expression is revealed as a potential early marker of increased adiposity and obesity-like features regardless of body weight, as animals fed a cafeteria diet presented increased body weight but not those fed a commercial HF diet, although both diets produced increased Cpt1a expression. This is of interest as the number of subjects who are not obese but present metabolic features characteristic of the obese status is increasing in our society\(^{(29)}\), evidencing the need for proper biomarkers for preventive strategies. In addition, Cpt1a could also be useful as a marker of metabolic recovery related to weight loss, based on the recovery of its mRNA expression in the PCAF group. In a previous study, we also reported the usefulness of Cpt1a expression in PBMC as an indicator of metabolic recovery based on recovery of its fasting response in PCAF animals in comparison with the fasting insensitivity observed in cafeteria diet-fed obese animals\(^{(20)}\). Thus, an analysis of Cpt1a expression levels in PBMC could serve as a sensitive indicator of metabolic risk factors directly related to lipid metabolism/adiposity.

Fig. 1. Expressions of genes involved in lipid metabolism (carnitine palmitoyltransferase I (Cpt1a), fatty acid synthase (Fasn) and sterol regulatory element-binding protein 1a (Srebp1a) in peripheral blood mononuclear cells of control (■), high-fat (□), cafeteria (△) and post-cafeteria (★) animals at different ages of diet administration (from 2 to 6 months), measured by real-time RT-PCR. Values are means (n 6–7), with standard errors of ratios of specific mRNA levels relative to guanosine diphosphate dissociation inhibitor 1, expressed as a percentage of the value of control animals that was set to 100 %. \(a, b, c\) Mean values with unlike letters were significantly different (\(P < 0.05\)). One-way ANOVA was performed for each age group, least significance difference (LSD) post hoc test was used after ANOVA.
Obesity-related genes. There is a wide number of genes available in the literature whose altered expressions are related to obesity and are considered as obesity markers. To perform our study, we selected three obesity-related genes, which have been shown to be expressed in PBMC (Fig. 2). One of the selected genes was apoprotein B48 receptor (\textit{Apob48r}) whose expression has been described to increase in monocytes after HF feeding, increasing lipid accumulation in these cells\textsuperscript{(30,31)}. Our results show increased \textit{Apob48r} expression in response to the two tested hyperlipidic diets after 2 months of diet exposure (4-month-old animals), but this increased expression disappeared after longer exposure to the diets. Increased \textit{Apob48r} expression after a HF meal has been shown to be time-dependent: it increases in the postprandial state and is recovered thereafter\textsuperscript{(31)}. On the basis of this, our results would indicate a response to the increased fat proportion of the diet after 2 months of dietary treatment, followed by an adaptation and normalisation of \textit{Apob48r} expression thereafter. Another gene defined as an obesity marker is \textit{Cebp\textbeta}, which encodes for a CCAAT/enhancer-binding protein (C/EBP)\textbeta\ transcription factor, an important regulator in controlling lipid metabolism and in the development of diet-induced obesity\textsuperscript{(32)}. C/EBP\textbeta\ expression has been shown to be increased in the liver, adipocytes and also in macrophages in response to HF diets, being responsible for dietary-induced inflammation in mice\textsuperscript{(33)}. According to this, our results show how 3 and 4 months of cafeteria diet feeding increased PBMC \textit{Cebp\textbeta} gene expression, although mRNA returned to basal levels in the PCAF. Commercial HF diet also induced increased \textit{Cebp\textbeta} expression, but only after 4 months of diet administration. Our group has

Fig. 2. Expression of obesity-related genes (apoprotein B48 receptor (\textit{Apob48r}), CCAAT/enhancer-binding protein \textbeta\ (\textit{Cebp\textbeta}) and solute carrier family 27 member (\textit{Slc27a2}) in peripheral blood mononuclear cells of the same animals described in Fig. 1, measured by real-time RT-PCR. Statistics: same as in Fig. 1. □, Control; ■, high fat; ■, cafeteria; ■, post-cafeeteria.
previously presented an analysis of \textit{Slc27a2} expression in PBMC as a potential overweight biomarker associated with the intake of a cafeteria diet\(^{21}\). This gene codes for solute carrier family 27 (fatty acid transporter), member 2, a free long-chain fatty acid transporter whose overexpression in the liver is related to hepatic steatosis\(^{34}\). As we observed in our previous experiment with a different set of animals\(^{21}\), \textit{Slc27a2} expression dramatically increased in PBMC of cafeteria diet-fed rats. We observed how these highly increased expression levels were present from the 1st month of dietary treatment (3-month-old animals), when animals only presented 18\% increased body weight and were maintained throughout the experimental period (when animals reached 37\% overweight). On the other hand, surprisingly, no change in \textit{Slc27a2} mRNA levels was observed in animals fed a commercial HF diet, which did not develop overweight. These data suggest that increased PBMC \textit{Slc27a2} expression would be reflecting increased body weight but not the effect of the intake of a HF diet \textit{per se}. However, as increased fat content in the diet produced metabolic alterations even in the absence of obesity, this gene would not be considered as a precise marker of disease or metabolic alterations as the previously mentioned \textit{Cpt1a}, whose expression was altered (increased) in response to the intake of both hyperlipidic diets. Interestingly, increased \textit{Slc27a2} expression was not reverted in the PCAF group, maybe because of the fact that post-cafeteria animals still maintain a significant part of increased body weight. Further studies are required to understand the important increase in the expression of this gene, exclusively in cafeteria diet-fed animals.

\textbf{Inflammatory genes.} Obesity represents an expansion of adipose tissue mass characterised by a state of chronic inflammation\(^{35–37}\). Moreover, a pro-inflammatory state is positively correlated with obesity-related complications such as type 2 diabetes and CVD\(^{37,38}\). Thus, gene expression analysis of inflammatory cytokines in immune cells can be used to identify a pro-inflammatory pattern related to obesity. We have analysed the expression of two well-known pro-inflammatory markers, IL-6 and TNF\(_{\alpha}\), which are commonly involved in the aetiology of obesity (Fig. 3). There is a lot of controversy regarding IL-6 gene expression and obesity development. Although IL-6 mRNA levels increase in adipose tissue in an obesogenic state\(^{39}\), different studies have not detected changes in IL-6 gene expression in the liver or in blood samples of obese subjects\(^{35,40}\). Our results in PBMC at different times of the intake of HF diets did not show any conclusive evidence. Cafeteria diet feeding produced no change in IL-6 expression, and the same was true for the commercial HF diet, except at the end of the experimental period when IL-6 mRNA levels decreased in the HF group. As expected\(^{37}\), \textit{Tnfa} gene expression increased in PBMC of our cafeteria-obese animals; increased expression was observed from the age of 4–6 months, and was recovered in PCAF animals. Commercial HF diet had a lower impact, and increased \textit{Tnfa} expression was only observed in 4-month-old animals (after 2 months of diet). According to these results, \textit{Tnfa} gene expression reflects increased adiposity associated with the intake of a cafeteria diet, but not that related to the commercial HF diet, which
produces lower increased adiposity and lower metabolic complications.

**Conclusion**

Our results clearly indicate that PBMC gene expression can reflect the development of an obesogenic profile related to the intake of hyperlipidic diets, with alterations in selected lipid metabolism, obesity-related, and inflammatory genes. Long-term cafeteria diet intake, associated with increased body weight and important metabolic deregulations, produced a higher impact in the studied genes in comparison with a commercial HF diet, which is related to increased adiposity and metabolic complications, but not to increased body weight. Remarkably, we were able to observe a time-dependent effect on the studied biomarkers, some of them being altered at the beginning and others at the end of the dietary treatment – the latter being representative of cumulative adiposity or metabolic damage. Altogether, Qpfla gene expression was of special interest as an early predictive biomarker of increased adiposity and metabolic damage associated with HF diet feeding, as well as of certain metabolic recovery associated with weight loss. Identification of new molecular biomarkers in blood samples is extremely useful to develop minimally invasive health preventive strategies.

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