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Immune Cell Recruitment Affects Metabolic-related Gene Expression

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Abstract: Brown adipose tissue (BAT) plays essential role in metabolic- and thermoregulation and displays morphological and functional plasticity in response to environmental and metabolic challenges. BAT is a heterogeneous tissue containing adipocytes and various immune-related cells, however, their interaction in regulation of BAT function is not fully elucidated. Fractalkine is a chemokine synthesized by adipocytes, which recruits fractalkine receptor (CX3CR1)-expressing leukocytes into the adipose tissue. Using transgenic mice, in which the fractalkine receptor, *Cx3cr1* gene was replaced by *Gfp*, we evaluated whether deficiency in fractalkine signaling affects BAT remodeling and function in high-fat-diet - induced obesity. Homo- and heterozygote male CX3CR1-GFP mice were fed with normal or fat enriched (FatED) diet for 10 weeks. Interscapular BAT was collected for histological and qPCR analysis. Heterozygous animals in which fractalkine signaling remains intact, gain more weight during FatED than CX3CR1 deficient *gfp/gfp* homozygotes. FatED in controls resulted in macrophage recruitment in the BAT with increased expression of proinflammatory mediators (*Il1a,b*, *Tnfa* and *Ccl2*). Local BAT inflammation was accompanied by increased expression of lipogenic enzymes and resulted in BAT "whitening". By contrast, fractalkine receptor deficiency prevented accumulation of tissue macrophages, selectively attenuated the expression of *Tnfa*, *Il1a* and *Ccl2*, increased BAT expression of lipolytic enzymes (*Atgl*, *Hsl* and *Mgl1*) and upregulated genes involved thermo-metabolism (*Ucp1*, *Pparg* *Pgcl1a*) in response to FatED. These results highlight the importance of fractalkine-CX3CR1 interaction in recruitment of macrophages into the BAT of obese mice which might contribute to local tissue inflammation, adipose tissue remodeling and regulation of metabolic-related genes.

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His group identified for the first time that fractalkin is an adipokine.

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Editors-in Chief
BBA/ Molecular and Cell Biology of Lipids

March, 07, 2016

Dear Editors,

Enclosed please find our manuscript entitled: "Brown Adipose Tissue in Obesity: Fractalkine-receptor Dependent Immune Cell Recruitment and Metabolic-related Gene Expression" to be published in BBA Molecular and Cell Biology of Lipids.

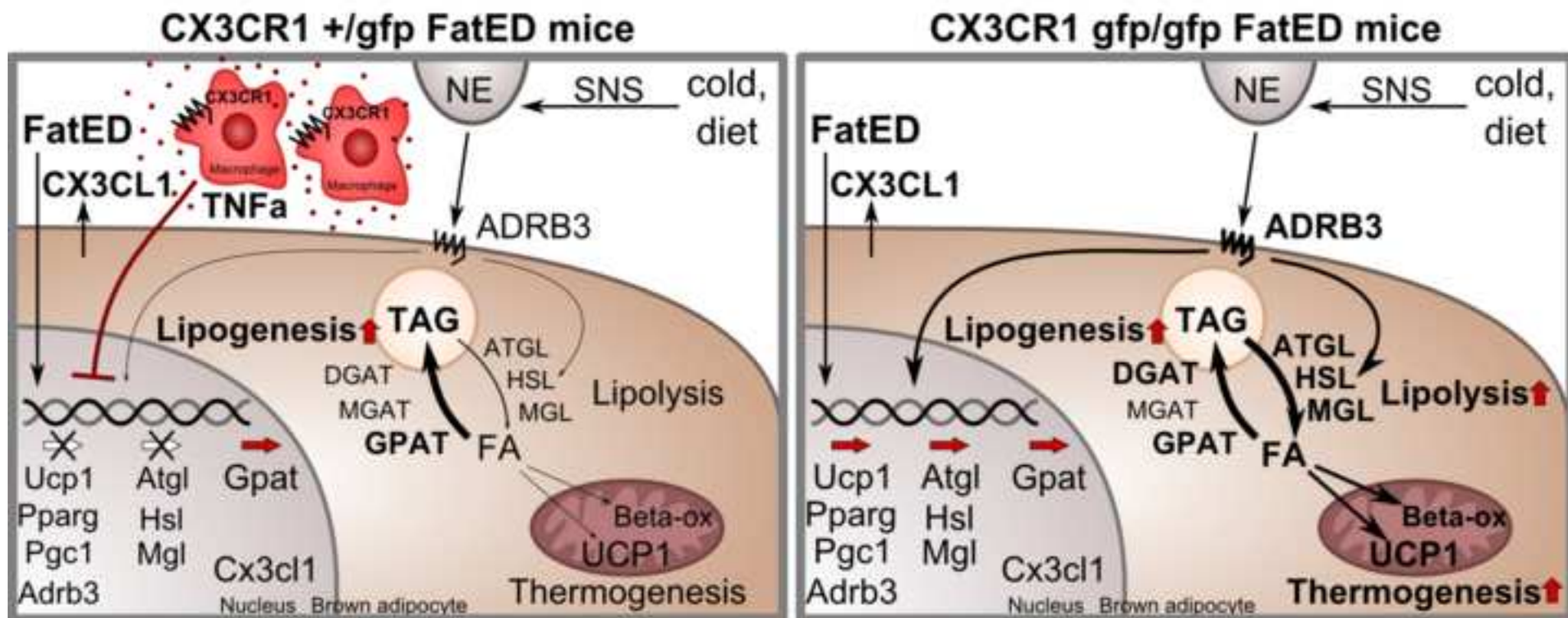
In this study, we aimed to identify the chemokine mechanism which is responsible for recruitment of macrophages in the brown adipose tissue in response to high fat diet. We have found that fat-enriched diet results in "whitening" the BAT of obese animals, upregulation of local production of proinflammatory cytokines, chemokines, impaired induction of lipolytic enzymes and thermogenic genes. Among the chemokines upregulated in obese mice BAT, fractalkine might play a crucial role in diet-induced morphological and functional rearrangements, because mice with targeted deletion of fractalkine receptor gain less weight and have reduced proinflammatory cytokine expression in the BAT. Mice with deficient fractalkine signaling were able to mount significant upregulation of thermometabolic and lipolytic enzymes's genes in the BAT when exposed to fat enriched diet.

These results are novel and important, because the role of fractalkine in BAT function has not been addressed before. On the basis of our present and previous findings, fractalkine signaling might be a potential target to fight obesity and metabolic inflammation.

This study is a continuation of our previously published paper (Polyák et al. BBI 38:25-35,2014) in which we studied the role of fractalkine in the hypothalamus, **WAT** and liver. Here we confirm that this present manuscript describes our original findings on **BAT** and has not been and will not be submitted for publication elsewhere.

Sincerely,

Krisztina J. Kovács, PhD



Highlights (for review)

- Fat enriched diet results in accumulation of leukocytes into the BAT and local inflammation.
- BAT accumulation of macrophages depends on fractalkine signaling.
- Thermogenic and lipolytic genes are induced in FatED mice with impaired fractalkine signaling.
- Fractalkine receptor deficient mice are protected from FatED-induced obesity.

1 Brown Adipose Tissue in Obesity: Fractalkine-receptor Dependent Immune Cell
2 Recruitment Affects Metabolic-related Gene Expression
3
4

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14

15

16

17 **Abstract**

18 Brown adipose tissue (BAT) plays essential role in metabolic- and thermoregulation and
19 displays morphological and functional plasticity in response to environmental and metabolic
20 challenges. BAT is a heterogeneous tissue containing adipocytes and various immune-related
21 cells, however, their interaction in regulation of BAT function is not fully elucidated. Fractalkine
22 is a chemokine synthesized by adipocytes, which recruits fractalkine receptor (CX3CR1)-
23 expressing leukocytes into the adipose tissue. Using transgenic mice, in which the fractalkine
24 receptor, *Cx3cr1* gene was replaced by *Gfp*, we evaluated whether deficiency in fractalkine
25 signaling affects BAT remodeling and function in high-fat-diet - induced obesity. Homo- and
26 heterozygote male CX3CR1-GFP mice were fed with normal or fat enriched (FatED) diet for 10
27 weeks. Interscapular BAT was collected for histological and qPCR analysis. Heterozygous
28 animals in which fractalkine signaling remains intact, gain more weight during FatED than
29 CX3CR1 deficient *gfp/gfp* homozygotes. FatED in controls resulted in macrophage recruitment
30 in the BAT with increased expression of proinflammatory mediators (*Il1a,b, Tnfa and Ccl2*).
31 Local BAT inflammation was accompanied by increased expression of lipogenic enzymes and
32 resulted in BAT “whitening”. By contrast, fractalkine receptor deficiency prevented
33 accumulation of tissue macrophages, selectively attenuated the expression of *Tnfa, Il1a and Ccl2*,
34 increased BAT expression of lipolytic enzymes (*Atgl, Hsl and Mgl1*) and upregulated genes
35 involved thermo-metabolism (*Ucp1, Pparg Pgc1a*) in response to FatED. These results highlight
36 the importance of fractalkine-CX3CR1 interaction in recruitment of macrophages into the BAT
37 of obese mice which might contribute to local tissue inflammation, adipose tissue remodeling and
38 regulation of metabolic-related genes.

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42 **Keywords:** fractalkine, macrophage, inflammation, triglyceride metabolism, thermogenesis,
43 obesity, BAT

44

45 **Abbreviations:** BAT – brown adipose tissue, WAT – white adipose tissue, CX3CL1 –
46 fractalkine, CX3CR1 – fractalkine receptor, GFP – green fluorescent protein, HFD – high fat
47 diet, FatED – fat enriched diet, ND – normal diet, PND – postnatal day, qPCR – quantitative real
48 time polymerase chain reaction, IL1A – interleukin 1 alpha, IL1B – interleukin 1 beta, IL6 –
49 interleukin 6, TNFa – tumor necrosis factor alpha, CCL2 (MCP1) - chemokine (C-C motif)
50 ligand 2, UCP1 – uncoupling protein 1, PPARG - peroxisome proliferator-activated receptor
51 gamma, PGC1A (PPARGC1A) - peroxisome proliferator-activated receptor gamma, coactivator
52 1 alpha, TH - tyrosine hydroxylase, ADRB3 - adrenoceptor beta 3, DIO2 - Type 2 Iodothyronine
53 Deiodinase, GLUT4 - Glucose transporter type 4, DGAT1 - diacylglycerol O-acyltransferase 1,
54 MGAT - mannosyl (alpha-1,3-)-glycoprotein beta-1,2-N-acetylglucosaminyltransferase, GPAT -
55 glycerol-3-phosphate acyltransferase, ATGL (PNPLA2) - adipose triglyceride lipase, HSL
56 (LIPE) - lipase, hormone-sensitive, MGL - monoglyceride lipase, FA – fatty acid.

57

58

59 **1. Introduction**

60 Obesity and diabetes are worldwide epidemics driven by the disruption in energy balance [1].

61 Brown adipose tissue (BAT) is the major site for cold- and diet-induced thermogenesis with
62 which BAT significantly affects systemic glucose and lipid metabolism [2-4]. In 2007

63 Nedergaard et al. published that adult humans possess active BAT [5]. The amount of BAT is
64 inversely correlated with body-mass index, especially in older people [6]. Metabolically active
65 BAT seems to be particularly low in patients with obesity or diabetes [7]. These results suggest a
66 significant role of brown adipose tissue in adult human metabolism and opens new opportunities
67 to develop therapeutic interventions to treat obesity.

68 Brown adipocytes and inducible brown-in-white (brite, beige) adipocytes are multilocular and
69 contain significantly higher number of mitochondria than other adipocytes in the body [8]. These
70 cells are specialized to dissipate energy in the form of heat by uncoupled thermogenesis,
71 mediated by the dissociation of mitochondrial respiratory chain electron transport from ATP
72 synthesis via the action of uncoupling protein UCP1. In addition to adipocytes, adipose tissues
73 contains various immune-related cells including resident macrophages, eosinophils, mast cells
74 and T cells, which significantly contribute to their function via release (adipo)cytokines and
75 transmitters in paracrine or endocrine fashion. [9-12]. Both types of adipose tissues (BAT and
76 WAT) are sensitive to environmental (temperature) - hormonal (T3, leptin, insulin,
77 corticosteroid) - and metabolic (high fat diet) cues and display significant cellular and functional
78 remodeling in response to these challenges. For instance, high fat diet results in hypertrophy and
79 hyperplasia of white adipocytes and recruitment of monocytes into the WAT [13]. Furthermore,
80 in obese animals and humans there is a shift from alternatively (anti-inflammatory) polarized
81 macrophages to those that produce predominantly proinflammatory mediators [14, 15]. However,

82 the accumulation of macrophages to BAT, the mechanisms that recruit and activate them, and
83 their effect on thermometabolic genes has not been fully elucidated. Because these changes
84 contribute to insulin resistance and low grade systemic metabolic inflammation which is seen in a
85 subset of obese patients with metabolic X [16], it is important to understand the mechanisms that
86 recruit and activate adipose tissue macrophages and the means with which local inflammation
87 affects lipid metabolism and thermoregulation.

88 Fractalkine (CX3CL1) is a chemokine expressed in endothelial cells, vascular smooth muscle
89 cells, hepatocytes, adipocytes and neurons as a transmembrane protein and involved in trafficking
90 and capturing various leukocytes (monocytes, macrophages, microglia) expressing its' cognate
91 receptor, CX3CR1 [17, 18]. Fractalkine -released from the cell surface by proteolytic cleavage-
92 acts in paracrine and endocrine manner and has been identified in the WAT as a novel
93 adipocytokine with increased expression in obese individuals [19]. It has been shown previously
94 that lack of CX3CL-CX3CR1 signaling results in reduced macrophage accumulation into white
95 adipose tissue and reduced body weight gain during the development of obesity [20].

96 The aim of the present study was to identify the role of fractalkine/CX3CR1 signaling in the
97 recruitment of monocytes into the brown adipose tissue and to reveal the role of local
98 inflammation in regulation of genes involved in triglyceride- and thermo-metabolism in obese
99 mice.

100

101
102
103 **2. Materials and methods**
104 2.1. *Animals and diet*
105 Experiments were performed in male CX3CR1 +/gfp (+/gfp), and CX3CR1 gfp/gfp (gfp/gfp)
106 mice [17]. Animals were obtained from the European Mouse Mutant Archive (EMMA
107 Cx3cr1^{tm1Litt} MGI:2670351). The background C57Bl/6J strain has been shown to be genetically
108 vulnerable to diet-induced obesity [21]. In these mice, the *Cx3cr1* gene was replaced by a *Gfp*
109 reporter gene such that heterozygote CX3CR1 +/gfp mice express GFP in cells of the myeloid
110 lineage and retain receptor function, whereas monocytes in homozygote CX3CR1 gfp/gfp mice are
111 labeled with GFP and lack functional CX3CR1. Genotype of the animals has been verified by
112 PCR using combination of three different primers as described by Jung et al [17].

113 Animals were housed in groups of 4-5/cage at the minimal disease (MD) level of the Medical
 114 Gene Technology Unit of our Institute, had free access to food and water and were maintained
 115 under controlled conditions: temperature, 21 °C ± 1 °C; humidity, 65%; light-dark cycle, 12-h
 116 light/12-h dark cycle, lights on at 07:00. At 35 days of age, both CX3CR1 +/gfp (n=25) and
 117 CX3CR1 gfp/gfp (n=25) mice were randomly distributed into two groups. The first group,
 118 normal diet (ND), received standard chow (VRF1 (P), Special Diets Services (SDS), Witham,
 119 Essex, UK.). The second group received fat-enriched diet (FatED), by providing a 2:1 mixture of
 120 standard chow and lard (Spar Budget, Budapest, Hungary). The energy content and macronutrient
 121 compositions of the two diets are given in Table 1. All procedures were conducted in accordance
 122 with the guidelines set by the European Communities Council Directive (86/609 EEC) and
 123 approved by the Institutional Animal Care and Use Committee of the Institute of Experimental
 124 Medicine (permit number: 22.1/3347/003/2007).

125 ***Table 1. Energy content and macronutrient composition of diets***
 126

	ND - standard chow		FatED - mixed chow	
	g%	kcal%	g%	kcal%
Protein	19,1	22,5	12,7	9,7
Carbohydrate	55,3	65,0	36,9	28,0
Fat	4,8	12,6	36,5	62,3
kcal/g	3,40		5,27	

127
 128

129 **2.2. *Experimental design***

130 Mice were fed with normal diet (ND) or fat enriched diet (FatED) starting at age of 35 days.
 131 Mice were decapitated ten weeks later, interscapular brown adipose tissues were collected,
 132 sampled and stored at -70°C for qPCR, tissue samples were also obtained for histology. A set of
 133 animals underwent cold tolerance test.

134

135 2.3. *Histology*

136 BAT tissue samples were immersion fixed in 4% w/v paraformaldehyde in 0.1 mol l⁻¹ phosphate
137 buffer, pH 7.4 (PB) for 3 days and stored in 1% w/v paraformaldehyde in 0.1 mol l⁻¹ PB at 4°C
138 then were embedded in paraffin, sectioned and stained with hematoxylin-eosin (H&E).
139 Microscopic slides were digitalized with Panoramic Digital Slide Scanner (3DHISTECH Kft.,
140 Hungary). Lipid droplet areas of brown adipose cells were counted under 40x magnification in
141 one field of view with ImageJ software (NIH, USA).

142

143 2.4. *Core body temperature measurement and cold challenge*

144 Rectal temperature was measured with Multithermo thermometer (Seiwa Me Laboratories Inc.,
145 Tokyo, Japan). To assess cold tolerance, set of animals (n = 30) from both genotypes were fasted
146 for 5 hours, then placed into new individual cages with minimal bedding and transferred to cold
147 room (4°C). Rectal temperature was measured before and 60, 120, 180 and 240 min after cold
148 exposure.

149

150 2.5. *Gene expression analysis by quantitative real-time PCR*

151 Total RNA was isolated from brown adipose tissue samples with QIAGEN RNeasyMiniKit
152 (Qiagen, Valencia, CA, USA) according the manufacturer's instruction. To eliminate genomic
153 DNA contamination, DNase I (Fermentas) treatment was used. Sample quality control and the
154 quantitative analysis were carried out by NanoDrop (Thermo Scientific). Amplification was not
155 detected in the RT-minus controls. cDNA synthesis was performed with the High Capacity
156 cDNA Reverse Transcription Kit (Applied Biosystems, Foster City, CA, USA). The designed
157 primers (Invitrogen) were used in real-time PCR reaction with Power SYBR Green PCR master
158 mix (Applied Biosystems, Foster City, CA, USA) on ABI StepOnePlus instrument. The gene

159 expression was analyzed by ABI StepOne 2.3 program. The amplicon was tested by Melt Curve
 160 Analysis. Measurements were normalized to ribosomal protein S18 (*Rps18*) expression [22].

161

162 **2.6. Primer design**

163 Primers used for the comparative CT (threshold cycle) experiments were designed by the Primer
 164 Express 3.0 program. Primer sequences are shown in Table 2.

165

166 **Table 2. Mouse specific primer sequences used for rtPCR**

167

Gene	Forward sequence	Reverse sequence
<i>Adrb3</i>	ACCGCTCAACAGGTTTGA	GGGGCAACCAGTCAAGAAGAT
<i>Atgl</i>	GCCATGATGGTGCCCTATACT	TCTTGGCCCTCATCACCAGAT
<i>Ccl2 (Mcp1)</i>	CCAGCACCAGCACCAGCCAA	TGGATGCTCCAGCCGGCAAC
<i>Cx3cl1</i>	CCGCGTTCTTCCATTTGTGT	GGTCATCTTGTCGCACATGATT
<i>Dgat1</i>	GTTCCCGTCCAGGGTGGTAGT	CGCACCTCGTCCTCTTCTAC
<i>Dio2</i>	ACAAACAGGTTAAACTGGGTGAAG	CGTGCACCACACTGGAATTG
<i>Gfp</i>	GGACGACGGCAACTACAAGA	AAGTCGATGCCCTTCAGCTC
<i>Glut4</i>	AGGAACTGGAGGGTGTGCAA	GGATGAAGTGCAAAGGGTGAG
<i>Gpat</i>	AGTGAGGACTGGGTTGACTG	GCCTCTCCGGCTCATAAGG
<i>Hsl</i>	AGCCTCATGGACCCTCTTCTA	TCTGCCTCTGTCCCTGAATAG
<i>Il1a</i>	CCATAACCCATGATCTGGAAGAG	GCTTCATCAGTTTGTATCTCAAATCAC
<i>Il1b</i>	CTCGTGGTGTCTGGACCCATATGA	TGAGGCCCAAGGCCACAGGT
<i>Il6</i>	CTCTGCAAGAGACTTCCATCC	AGTCTCCTCTCCGGACTTGT

<i>Mgat</i>	TGGTTCTGTTTCCCGTTGTTC	GAAACCGGCCCGTTACTCAT
<i>Mgl</i>	CTTGCTGCCAAACTGCTCAA	GGTCAACCTCCGACTTGTTC
<i>Pgc1a (Pparg1a)</i>	ATGTGCAGCCAAGACTCTGT	TTCCGATTGGTCGCTACACC
<i>Pparg2</i>	CTCCTGTTGACCCAGAGCAT	TGGTAATTTCTTGTGAAGTGCTCA
<i>Rps18</i>	TCCAGCACATTTTGCAGTA	TTGGTGAGGTCGATGTCTGC
<i>Th</i>	TCTCAGAGCAGGATACCAAGCA	GCATCCTCGATGAGACTCTGC
<i>Tnfa</i>	CAGCCGATGGGTTGTACCTT	GGCAGCCTTGTGCCTTGA
<i>Ucp1</i>	GGTCAAGATCTTCTCAGCCG	AGGCAGACCGCTGTACAGTT

168

169 2.7. *Statistical analysis*

170

171 Statistical analysis was performed by factorial ANOVA with Newman–Keuls post-hoc test in
 172 Statistica 11 (StatSoft Inc.). The results are shown as means ± SEM. In all cases $p < 0.05$ was
 173 considered significant.

174

175 **3. Results**

176

177 3.1. *Fractalkine receptor deficiency prevents FatED-induced obesity*

178 In agreement with our previous findings [20], 10 weeks on FatED increased body weight of mice
 179 (diet effect: $F(1,14) = 20.84$, $p < 0.001$), but the weights were significantly lower in fractalkine
 180 receptor deficient, gfp/gfp FatED group (genotype*diet effect: $F(1,14) = 6.59$, $p < 0.05$) (Fig.
 181 1A).

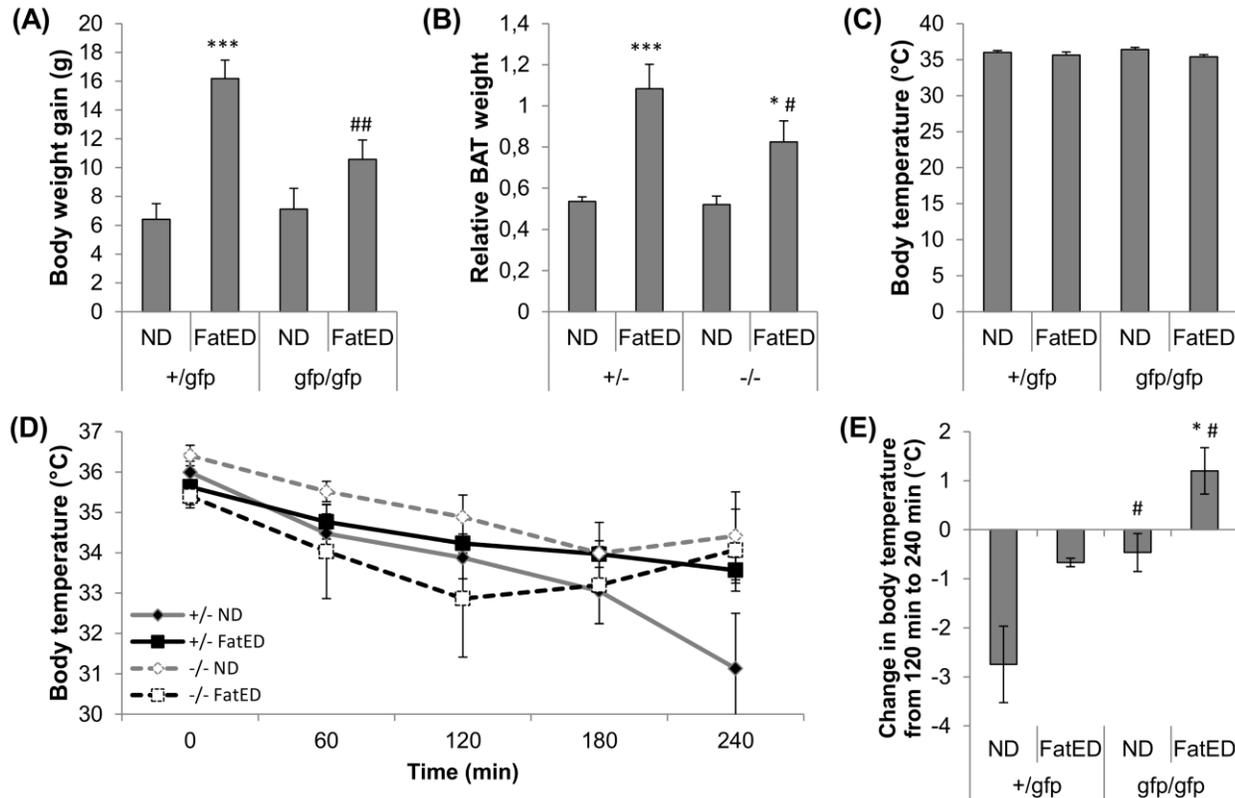
182 Although the daily food consumption of all FatED mice was lower, the daily energy intake was
183 comparable to those on normal diet. Because we did not detect significant genotype effect in
184 cumulative food- and energy intake and fecal output, these factors may not be responsible for the
185 differences seen in body weight gain (Supplementary Table 1).

186 However, significant diet- and genotype effects have been revealed in the relative BAT weight.
187 There was an increase in response to FatED (diet effect: $F(1,14) = 11.5$, $p < 0.01$; genotype
188 effect: $F(1,14) = 4.63$, $p < 0.05$), but it was significantly lower in *gfp/gfp* group (Fig.1B).

189

190 3.2. *Fractalkine deficient mice activate thermogenesis in response to acute cold.*

191 The core temperature of *ad libitum* fed mice was not different (Fig. 1C). When fasted mice were
192 placed to cold, the rectal temperature of all mice gradually decreased. However, after 2 hours in
193 cold, the temperature of homozygous animals started to increase back to the normal and the
194 increase in FatED mice was significantly higher than that seen in heterozygous animals fed by
195 control- or FatED (Fig. 1D-E).



196

197 **Figure 1. Body weight gain, BAT weight and cold tolerance test.**

198 Mean±SEM values of body weight gain (A) and relative BAT weight (B) of mice kept on normal
 199 (ND) or fat enriched diet (FatED) for 10 weeks. C) Core body temperature at room temperature.
 200 D-E) Changes in body temperature during cold tolerance test * $p < 0.05$, ** $p < 0.01$, *** $p <$
 201 0.001 vs. ND, # $p < 0.05$, ## $p < 0.01$ vs. +/gfp (Newman–Keuls post hoc comparison). FatED –
 202 fat enriched diet, ND – normal diet.

203

204 **3.3. Lack of fractalkine receptor attenuates diet-induced accumulation of macrophages**

205 *into the BAT.*

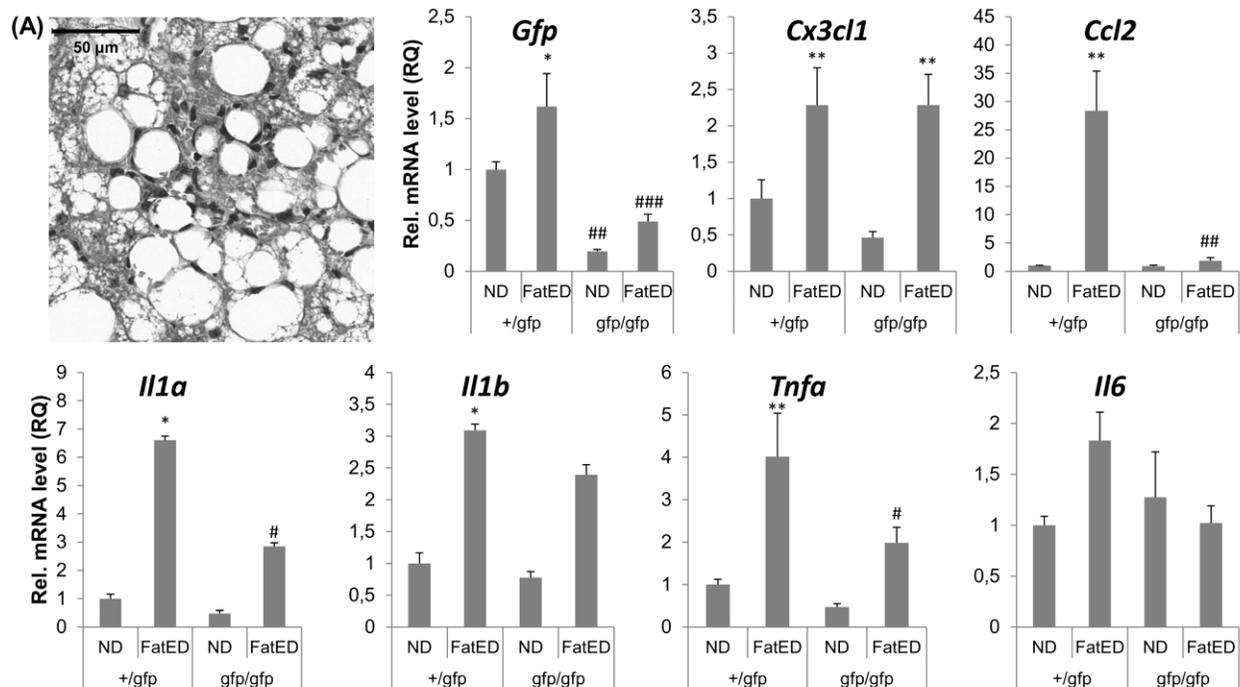
206 To reveal leukocyte recruitment into the BAT, we relied on GFP transgene expression, which
 207 occurs in myeloid cells of CX3CR1+/gfp and gfp/gfp mice. Increased number of leukocytes
 208 were observed in the BAT sections of FatED mice. Furthermore, “crown like structures” (CLS) -
 209 similar to those found in WAT of obese animals [23] - were observed in BAT: enlarged
 210 adipocytes filled with single lipid droplet were surrounded by numerous immune cells in FatED

211 +/gfp mice (Fig. 2A). No similar cellular scenario has been detected in gfp/gfp FatED and all
212 ND groups.

213 Because *Gfp* expression in the tissue is proportional to the number of macrophages, we calculated
214 the normalized *Gfp* mRNA levels to compare the number of macrophages within the BAT of our
215 experimental groups. In CX3CR1 +/gfp mice, FatED resulted in an increase of *Gfp* expression,
216 however, in *Cx3cr1* homozygotes the relative quantity of *Gfp* did not change in response to
217 FatED, suggesting that lack of fractalkine receptor prevents the accumulation of CX3CR1+
218 monocytes into BAT (Fig.2).

219 CCL2 (MCP-1) and CX3CL1, among others, are monocyte attracting chemokines, which serve
220 as signals for monocytes to accumulate to the sites of inflammation. CCL2 also contributes to the
221 local proliferation of tissue macrophages. To reveal the importance of these chemokines in
222 accumulation of GFP-positive immune cells to the BAT, we have compared their relative
223 expression in mice exposed to ND or FatED.

224 As shown in Figure 2 fractalkine (*Cx3cl1*) mRNA level was elevated in response to FatED in
225 both genotypes, however, diet-induced expression of *Ccl2* was detected only in CX3CR1 +/gfp
226 mice (*Gfp*: diet effect: $F(1,11) = 8.68$, $p < 0.05$; genotype effect: $F(1,11) = 38.97$, $p < 0.001$.
227 *Cx3cl1*: diet effect: $F(1,11) = 33.13$, $p < 0.001$. *Ccl2*: diet effect: $F(1,11) = 9.08$, $p < 0.05$; genotype
228 effect: $F(1,11) = 7.99$, $p < 0.05$; genotype * diet: $F(1,11) = 7.87$, $p < 0.05$).



229

230 **Figure 2. Macrophage accumulation to BAT and expression of proinflammatory cytokines**
 231 A) Representative image of CX3CR1 +/gfp FatED BAT. Adipocytes with enlarged lipid droplets
 232 in the BAT of CX3CR1 +/gfp mice are surrounded by leukocytes. Scale bar = 50 μm. Mean ±
 233 SEM values for relative mRNA levels in BAT: Gfp, chemokines: Cx3cl1, Ccl2, pro-inflammatory
 234 cytokines: Il1a, Il1b, Tnfa, Il6. * p < 0.05, ** p < 0.01 vs. ND, # p < 0.05, ## p < 0.01 vs. +/gfp
 235 (Newman–Keuls post hoc comparison). FatED – fat enriched diet, ND – normal diet.

236

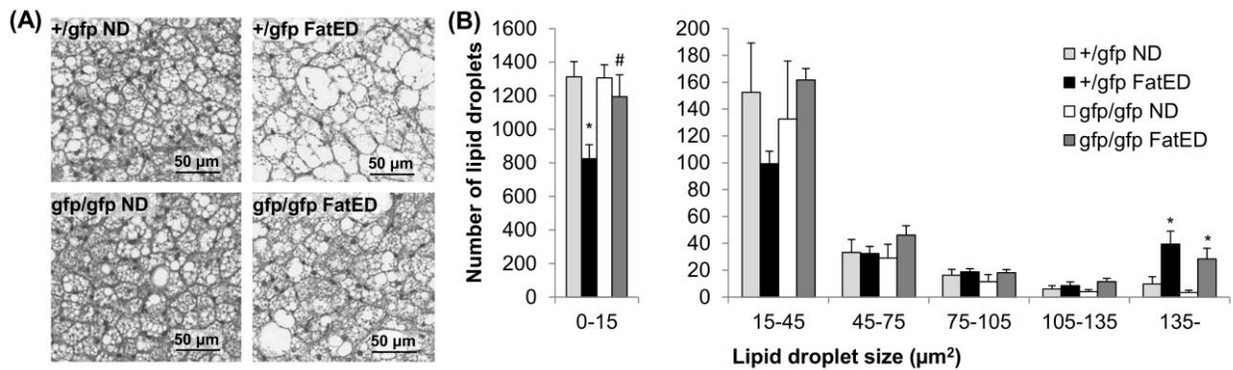
237 3.4. Recruitment of macrophages in the BAT is accompanied by increased expression
 238 of proinflammatory cytokines

239 As we have detected significant differences in the number of Gfp-positive profiles in the BAT of
 240 mice exposed to FatED, next, we compared the mRNA levels of pro-inflammatory cytokines
 241 produced by macrophages in the BAT. Il1a, Il1b and Tnfa were higher in +/gfp FatED group
 242 (Il1a: diet effect: F(1,11) = 11.93, p < 0.01; Il1b: diet effect: F(1,11) = 19.09, p < 0.01; Tnfa: diet
 243 effect: F(1,11) = 23.75, p < 0.001; genotype effect: F(1,11) = 7.61, p < 0.05), but in gfp/gfp
 244 FatED mice only Il1b expression was elevated, although the increase was not significant (Fig. 2).

245

246 3.5. Fat-enriched diet results in “whitening” of BAT

247 As shown in Fig. 3A, fat enriched diet resulted in “whitening” of interscapular brown adipose
 248 tissue. Histological analysis of BAT revealed enlarged brown adipose cells with few large lipid
 249 droplets in +/gfp FatED mice, reminiscent of white adipocytes filled with a single lipid droplet
 250 were also present. In gfp/gfp mice kept on FatED, multilocular brown adipocytes were more
 251 abundant than in +/gfp FatED mice and comparable to those BAT cells in ND mice. Frequency
 252 distribution analysis of lipid droplet areas in BAT revealed that FatED shifted the droplet areas
 253 to larger sizes, less droplets were under 15 μm^2 and more over 135 μm^2 ($F(1,14) = 8.62, p < 0.05$;
 254 $F(1,14) = 16.76, p < 0.01$, respectively). In gfp/gfp FatED mice significantly more small lipid
 255 droplets were present than in +gfp heterozygotes (Fig. 3B).



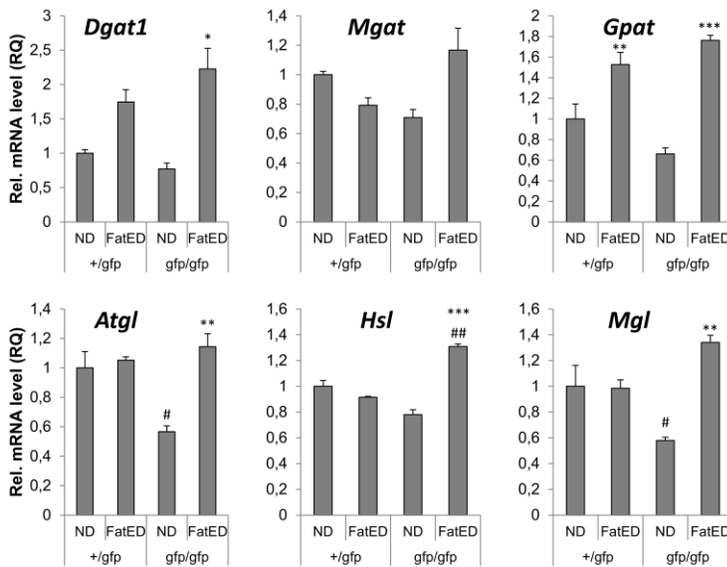
256
 257 **Figure 3. Quantitative histological analysis of BAT**

258 A) Representative histological images of hematoxylin-eosin stained BAT sections. FatED fed
 259 CX3CR1 +/gfp mice have larger lipid droplets. Scale bars = 50 μm . B) Frequency distribution of
 260 lipid droplet areas in one field of view. * $p < 0.05$ vs. ND, ** $p < 0.01$ vs. ND, # $p < 0.05$ vs.
 261 +/gfp, ## $p < 0.01$ vs. +/gfp (Newman–Keuls post hoc comparison). FatED – fat enriched diet,
 262 ND – normal diet.

263
 264 3.6. Fat storage and lipolytic/lipogenic enzymes in BAT

265 To analyze whether differences in lipid metabolism contribute to diet-induced phenotypic- and
 266 morphological changes in BAT of +/gfp and gfp/gfp animals, we have measured expression of
 267 enzymes involved in lipid synthesis and lipolysis in the BAT.

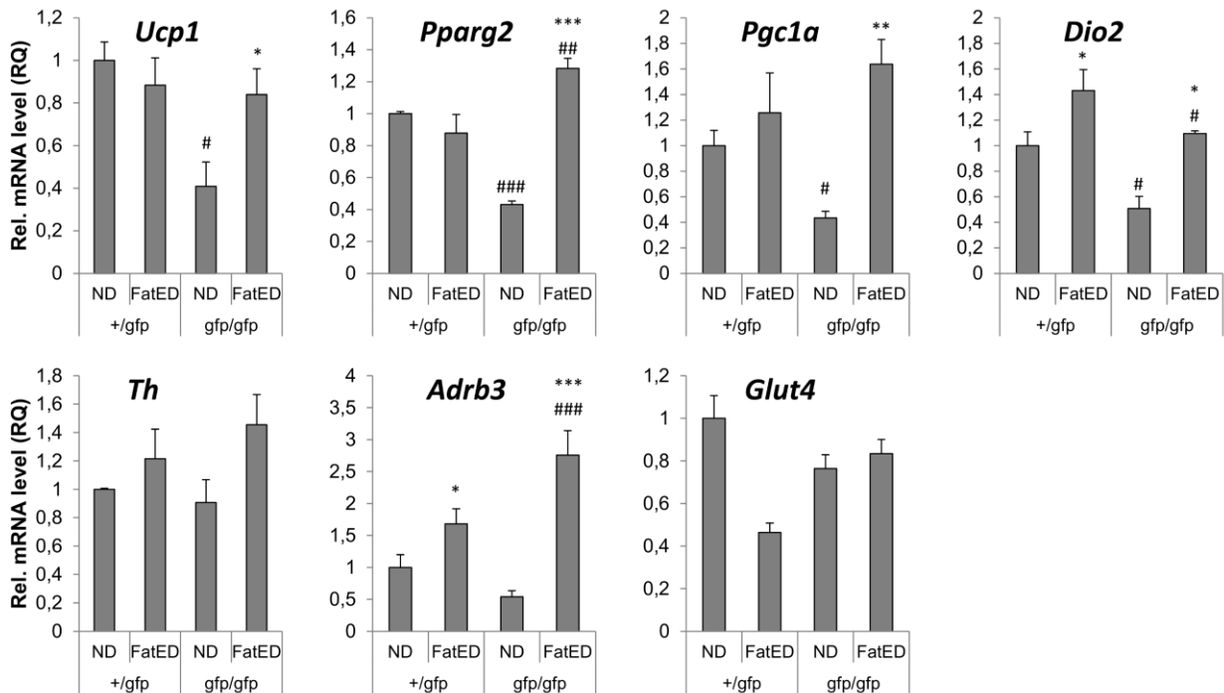
268 FatED upregulated lipogenic enzymes, *Dgat1* and *Gpat* mRNA expression in both genotypes.
 269 Lipolytic enzyme expression did not change in response to FatED in +/gfp mice. In gfp/gfp ND
 270 fed mice express lower levels of *Atgl* and *Mgl*, but FatED upregulated all lipolytic enzymes'
 271 mRNA expression (Fig. 4) (Lipogenic enzymes: *Dgat1* (diet effect: $F(1,13) = 76.94, p < 0.001$;
 272 diet * genotype: $F(1,13) = 6.63, p < 0.05$); *Mgat* (diet effect: $F(1,13) = 9.79, p < 0.01$); *Gpat*
 273 (diet effect: $F(1,13) = 129.54, p < 0.001$; genotype effect: $F(1,13) = 9.36, p < 0.01$; diet *
 274 genotype: $F(1,13) = 12.44, p < 0.01$). Lipolytic enzymes: *Atgl* (diet effect: $F(1,13) = 22.12, p <$
 275 0.001); *Hsl* (diet effect: $F(1,13) = 18.02, p < 0.001$); *Mgl* (diet effect: $F(1,13) = 32.10, p <$
 276 0.001)).



277
 278 **Figure 4. Gene expression of lipogenic and lipolytic enzymes in the BAT.** Mean \pm SEM values
 279 for relative mRNA levels in BAT. * $p < 0.05$, ** $p < 0.01$, *** $p < 0.0001$ vs. ND, # $p < 0.05$, ##
 280 $p < 0.01$ vs. +/gfp (Newman–Keuls post hoc comparison). FatED – fat enriched diet, ND – normal
 281 diet.

282
 283 3.7. Fractalkine receptor deficiency affects the expression of BAT thermogenic and
 284 metabolic-related markers

285 Because BAT significantly contributes to energy expenditure via non-shivering thermogenesis by
286 using fatty acids and glucose as fuels, next we investigated the diet-induced expression of
287 thermogenic related markers in the BAT of mice with or without fractalkine signaling.
288 In +/gfp mice, fat enriched diet did not affect expression of *Ucp1*, *Pparg* and *Pgc1a*, however,
289 *Dio2* and *Adrb3* mRNA levels were elevated. Gfp/gfp mice express less *Ucp1*, *Pparg2* and
290 *Pgc1a*, than +/gfp mice during normal dieting, exposure to FatED resulted in significantly
291 elevated expression of these mRNAs in the BAT. *Pparg2* and *Adrb3* mRNA levels in gfp/gfp
292 FatED mice was higher than in +/gfp FatED mice (Fig. 5) (*Ucp1*: diet effect: $F(1,11) = 23.68$, p
293 < 0.001 ; genotype * diet: $F(1,11) = 13.74$, $p < 0.01$; *Pparg2*: diet effect: $F(1,11) = 26.88$, $p <$
294 0.001 , genotype * diet: $F(1,11) = 43.91$, $p < 0.001$; *Pgc1a*: diet effect: $F(1,11) = 15.74$, $p < 0.01$;
295 genotype * diet: $F(1,11) = 7.75$, $p < 0.05$. *Dio2*: diet effect: $F(1,11) = 24.70$, $p < 0.001$;
296 genotype effect: $F(1,11) = 17.35$, $p < 0.01$. *Adrb3*: diet effect: $F(1,11) = 89.78$, $p < 0.001$;
297 genotype effect: $F(1,11) = 7.56$, $p < 0.05$; genotype * diet: $F(1,11) = 28.23$, $p < 0.001$).
298 Because adipose tissue macrophages synthesize and release catecholamines locally in response to
299 cold [24], we have been interested how *Th*, the key enzyme in catecholamine synthesis, varies in
300 the BAT in response to diet. Neither the genotype nor the diet affected *Th* expression in the BAT.



301
 302 **Figure 5. Gene expression of BAT thermogenic and metabolic-related markers.** Mean \pm
 303 SEM values for relative mRNA levels in BAT * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$ vs. ND; # p
 304 < 0.05 , ## $p < 0.01$ ###, $p < 0.001$ vs. +/gfp (Newman–Keuls post hoc comparison). FatED – fat
 305 enriched diet, ND – normal diet.

306 **4. Discussion**

307 Present results reveal significant morpho-functional immune- and metabolic rearrangements in
 308 the brown adipose tissue of mice kept on fat enriched diet. We have found “whitening” of BAT
 309 in FatED fed mice with increased accumulation and recruitment, of mononuclear cells in the
 310 BAT, differential overexpression of pro-inflammatory mediators that results in local metabolic
 311 inflammation. Fractalkine/fractalkine receptor system is involved in the recruitment of
 312 mononuclear cells into the BAT and regulation of adipose inflammation, since mice lacking
 313 functional fractalkine receptor display reduced expression of proinflammatory cytokines and
 314 improved profile of diet-induced genes involved in lipid metabolism and thermogenesis. These
 315 changes in the BAT might contribute to the obesity-resistant phenotype of mice lacking
 316 fractalkine signaling.

317 Both type of adipose tissue displays significant morphological and functional plasticity driven by
318 metabolic-, environmental- and hormonal cues [25]. “Browning” of the white adipose tissue is
319 well recognized. For instance, clusters of UCP1 expressing cells referred to as “brite” (brown in
320 white or beige) adipocytes appear in the white adipose tissue in response to cold, while there is a
321 downregulation of *Ucp1* mRNA levels together with phenotypic appearance of white adipocytes
322 in the BAT at thermoneutral conditions [26]. Here we have shown „whitening” of BAT in
323 response to fat-enriched diet in mice, which is due to coalescence of lipid droplets. Similar,
324 distorted lipid droplet architecture has also been reported in mice kept on high fat diet for 13
325 weeks [27, 28]. Increase of the size of lipid droplets might indicate an imbalance between lipid
326 synthesis and lipolysis. Indeed, our present results show that obese CX3CR1 +/gfp mice were
327 unable to induce lipolytic enzyme expression in the BAT, which might be responsible for fat
328 deposition in BAT of these animals.

329 In addition to morphological changes of adipocytes we found recruitment/accumulation of
330 mononuclear cells into the BAT of +/gfp mice kept on fat-enriched diet. These data are consistent
331 with previous results showing that genetic and diet-induced obesity results in chronic
332 inflammation in the BAT [29-31]. By contrast, Fitzgibbons et al. found very low level of immune
333 cell enriched transcripts in the BAT from C57BL6/J mice fed a high-fat diet for 13 weeks [27].
334 Thus the extent of BAT inflammation is largely depends on the strain and conditions used. Our
335 estimation of leukocyte accumulation is based on the normalized expression of *Gfp* mRNA in
336 CX3CR1-gfp transgenic mice [20] in which monocytes (except eosinophils and neutrophils), a
337 special subset of NK cells and dendritic cells express *Gfp* as described previously [17].
338 Recruitment of leukocytes into the white adipose tissue and their role in metabolic inflammation
339 is well recognized both in genetic- and diet-induced rodent models as well as in human obesity
340 [32]. Feeding a high fat diet to C57Bl6 mice has been shown to promote large increases of

341 various leukocytes, among those T cells and neutrophils are the first on the scene, followed by
342 monocytes by 8-10 weeks on diet [33]. Adipose tissue macrophages have been mechanistically
343 implicated in low grade, long lasting, metabolic inflammation and glucose intolerance seen in
344 diet-induced obesity [34, 35]. It has been hypothesized that dying adipocytes initiate macrophage
345 recruitment to the adipose tissue, however, recent findings emphasize the role of various
346 chemokines originating from adipocytes and/or from the stromal vascular fraction. In this respect
347 the monocyte attractant protein, CCL2 (MCP1) and its receptor CCR2 have been the most
348 intensively studied [36]. Although *Mcp1* mRNA level in the adipose tissue is elevated within 7
349 days and plasma MCP1 concentration increased 4 weeks after starting high fat diet, genetic
350 disruption of MCP1 signaling did not confer resistance to diet-induced obesity in mice or reduce
351 adipose tissue macrophage infiltration in the WAT [36], indicating involvement of additional
352 monocyte attractants.

353 Fractalkine has been recently identified as an adipo-chemokine, which is elevated in obese people
354 and patients with type2 diabetes [19]. CX3CL1 is implicated in recruitment of leukocytes in
355 clinical syndromes of adipose tissue inflammation and atherosclerosis. Indeed, we have recently
356 shown increased expression of fractalkine in the epididymal white fat pad of obese mice to be
357 accompanied with increased number of tissue macrophages and upregulated expression of
358 proinflammatory cytokines [20]. Here we report, for the first time, that fat enriched diet induces
359 fractalkine expression in brown adipose tissue as well. Based on the facts that expression of
360 fractalkine (CX3CL1) has been significantly elevated in all groups fed with FatED, while mice
361 lacking the fractalkine receptor accumulated significantly less GFP+ cells into the BAT and
362 display less severe local tissue inflammation than controls, we propose a role of
363 fractalkine/fractalkine receptor system in recruitment of macrophages into the BAT.

364 By contrast, data from Morris et al. indicate that CX3CR1 is not required for trafficking of
365 macrophages to- and their retention in, the epididymal WAT in mice with diet-induced obesity
366 [37].

367 Here we report significant increase of mRNA expression of proinflammatory cytokines *Il1a*, *Il1b*
368 and *Tnfa* in the BAT of FatED mice, the pattern and rate of increase of cytokines were
369 comparable with those observed in WAT. It remains unknown however, if these cytokines
370 originate in resident or recruited population of immune cells within the BAT. When compared to
371 heterozygote controls, we have found reduced expression of proinflammatory mediators in the
372 BAT of CX3CR1 *gfp/gfp* mice, indicating the role of fractalkine signaling in activation of
373 monocytes in the BAT as well. Accumulation of proinflammatory adipose tissue macrophages in
374 obese BAT shows similarities to foam cell formation in atherosclerotic plaques, which is also
375 dependent on the presence of CX3CR1 [38]. It should be recognized, however, that different
376 subsets of monocytes use different chemokine patterns with which to accumulate in various
377 inflammatory targets [39].

378 One interesting finding of the present study is the lack of diet-induced elevation of CCL2
379 (MCP1) in fractalkine receptor deficient mice, suggesting some mechanistic relationship between
380 these chemokines.

381 Among the cytokines induced by FatED in the BAT, TNFa might play a prominent role in
382 morphofunctional rearrangements. For instance, TNFa decreased the expression of functionally
383 active ADRB3 receptors in brown adipocytes and consequently attenuated the thermogenic and
384 lipolytic actions of SNS activity [40]. Studies on 3T3-L1 adipocytes revealed that TNFa
385 decreases the expression of lipolytic enzymes *Atgl* and *Hsl* [41] Conversely, TNFa deficiency in
386 genetically obese (*ob/ob*) mice resulted in less severe obesity, decrease in brown adipocyte
387 apoptosis, and increased expression of *Adrb3* and *Ucp1* with significant improvement in

388 thermogenetic capacity [40]. Fractalkine receptor deficient mice (gfp/gfp), in which FatED- did
389 not induce local *Tnfa* expression, are protected from excessive weight gain, display improved
390 glucose tolerance and induction of *Adrb3*, *Atgl* and *Hsl* mRNA in the BAT.

391 The next obvious question was how FatED-induced proinflammatory environment affects energy
392 expenditure/cold tolerance and BAT expression of metabolic-related and thermogenic genes.
393 CX3CR1 +/gfp mice were unable to increase BAT expression of *Ucp1*, *Pparg2*, and *Pgc1a* in
394 response to FatED, which might explain their obesity prone phenotype and impaired cold
395 tolerance during fat enriched diet. Indeed, it has been recently shown that macrophage derived
396 proinflammatory cytokines in general-, and TNFa in particular, suppress the induction of *Ucp1*
397 promoter activity and mRNA expression [42, 43]. Nevertheless, the interaction between adipose
398 tissue macrophages, proinflammatory cytokines and adipocytes is quite complex and occurs at
399 several functionally distinct loci of obesity.

400
401 These results clearly suggest that diet-induced recruitment of macrophages in the BAT of +/gfp
402 mice through the release of proinflammatory cytokines like TNFa, results in local inflammation
403 and may attenuate the sympathetic nervous system (SNS) induced thermogenesis and lipolysis in
404 BAT, leading to fat accumulation, driving a vicious circle. However, impaired fractalkine
405 signaling (in gfp/gfp mice) breaks this circle by attenuating the accumulation of brown adipose
406 tissue macrophages and their cytokine production, which results in diet-induced upregulation of
407 *Atgl*, *Hsl* and *Mgl* lipogenic enzymes and *Ucp1* mRNA in the BAT, which changes are likely to
408 contribute to the improved thermoadaptive response and the leaner phenotype seen in fractalkine
409 receptor deficient mice.

410

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414

415

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Food intake (g)

Genotype	Diet	Weeks									
		1	2	3	4	5	6	7	8	9	10
+/gfp	ND	3,41 ± 0,58	4,63 ± 0,28	3,97 ± 0,37	4,52 ± 0,14	4,45 ± 0,15	3,69 ± 0,15	7,33 ± 0,37	3,99 ± 0,34	4,9 ± 0,26	3,31 ± 0,36
	FatED	2,88 ± 0,11	3,38 ± 0,11	2,58 ± 0,1	2,83 ± 0,08	2,84 ± 0,18	3,24 ± 0,16	3,17 ± 0,15	3,03 ± 0,1	3,02 ± 0,16	3,53 ± 0,24
gfp/gfp	ND	3,53 ± 0,53	5,75 ± 0,79	4,14 ± 0,15	4,69 ± 0,36	4,44 ± 0,27	4,23 ± 0,35	6,62 ± 0,11	4,38 ± 0,8	5,15 ± 0,16	4,1 ± 0,26
	FatED	2,7 ± 0,09	3,25 ± 0,19	2,68 ± 0,19	2,88 ± 0,22	2,74 ± 0,08	3,32 ± 0,23	3,07 ± 0,19	2,9 ± 0,13	2,94 ± 0,16	3,37 ± 0,14

Energy intake (kcal)

Genotype	Diet	Weeks									
		1	2	3	4	5	6	7	8	9	10
+/gfp	ND	11,59 ± 1,99	15,74 ± 0,96	13,5 ± 1,26	15,4 ± 0,5	15,13 ± 0,53	12,55 ± 0,51	24,93 ± 1,26	13,59 ± 1,15	16,66 ± 0,89	11,27 ± 1,25
	FatED	15,14 ± 0,6	17,82 ± 0,62	13,6 ± 0,53	14,92 ± 0,46	14,95 ± 0,98	17,07 ± 0,85	16,69 ± 0,83***	15,95 ± 0,53	15,92 ± 0,86	18,56 ± 1,27**
gfp/gfp	ND	12,01 ± 1,82	19,56 ± 2,68	14,08 ± 0,53	15,95 ± 1,23	15,09 ± 0,94	14,38 ± 1,22	22,5 ± 0,37	14,91 ± 2,74	17,52 ± 0,55	13,94 ± 0,89
	FatED	14,2 ± 0,52	17,13 ± 1	14,09 ± 1,05	15,14 ± 1,16	14,45 ± 0,43	17,5 ± 1,21	16,17 ± 1,03**	15,28 ± 0,7	15,5 ± 0,88	17,73 ± 0,77

Faeces (g)

Genotype	Diet	Weeks					
		1	3	4	6	7	8
+/gfp	ND	0,93 ± 0,04	0,93 ± 0,04	0,85 ± 0,04		0,89 ± 0,02	0,99 ± 0,07
	FatED	0,52 ± 0,04***	0,54 ± 0,02***	0,47 ± 0,03***	0,51 ± 0,03	0,42 ± 0,01***	0,44 ± 0,02***
gfp/gfp	ND	0,95 ± 0,03	0,91 ± 0,08	0,81 ± 0,03		0,96 ± 0,16###	1,02 ± 0,03
	FatED	0,51 ± 0,01***	0,55 ± 0,02***	0,49 ± 0,02***	0,51 ± 0,02	0,43 ± 0,01***	0,48 ± 0,01***

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