Research Report

Obesity-mediated inflammation may damage the brain circuit that regulates food intake

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ABSTRACT

Adiposity is associated with chronic low-grade systemic inflammation and increased inflammation in the hypothalamus, a key structure in feeding behavior. It remains unknown whether inflammation impacts other brain structures that regulate feeding behavior. We studied 44 overweight/obese and 19 lean individuals with MRI and plasma fibrinogen levels (marker of inflammation). We performed MRI-based segmentations of the medial and lateral orbitofrontal cortex (OFC) and hippocampal volumes. Gray matter (GM) volumes were adjusted for head size variability. We conducted logistic and hierarchical regressions to assess the association between fibrinogen levels and brain volumetric data. Using diffusion tensor imaging (DTI), we created apparent diffusion coefficient (ADC) maps and conducted voxelwise correlational analyses. Fibrinogen concentrations were higher among the overweight/obese (t[61]=−2.33, \(P=0.023\)). Lateral OFC associated together with fibrinogen correctly classified those with excess of weight (accuracy=76.2\%, sensitivity=95.5\%, and specificity=31.6\%). The lateral OFC volumes of overweight/obese were negatively associated with fibrinogen (r=−0.37, \(P=0.016\)) and after accounting for age, hypertension, waist/hip ratio and lipid and sugar levels, fibrinogen significantly explained an additional 9\% of the variance in the lateral OFC volume (\(\Delta R^2=0.093, \Delta F P=0.046\)). Among overweight/obese the associations between GM ADC and fibrinogen were significantly positive (P<0.001) in the left and right amygdala and the right parietal region. Among lean individuals these associations were negative and located in the left prefrontal, the right parietal and the left occipital lobes. This is the first study to report that adiposity-related inflammation may reduce the integrity of some of the brain structures involved in reward and feeding behaviors.

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1. Introduction

Worldwide, a billion adults are overweight and at least 300 million are obese (World Health Organization, 2010). The prevalence of overweight people has increased dramatically in the US adult population and obesity-related diseases reached epidemic proportions in the last decade (Flegal et al., 2010). Excess body fat increases the risk of cardiovascular disease, hypertension, fatty liver, type 2 diabetes, and may contribute to a decline in cognitive abilities, and dementia (Ikeoka et al., 2010; Rizvi, 2010). Due to the meteoric increase in overweight and obese individuals, it is increasingly important to elucidate the cause and effects of excessive adipose deposition. All of the pathological conditions associated with excess weight may be related to adiposity-induced inflammation as obesity is characterized by a state of chronic low grade inflammation (Vachharajani and Granger, 2009). The adipose tissue of obese individuals is comprised of hypertrophied fat cells and infiltrating macrophages and lymphocytes. These immune cells secrete several pro-inflammatory cytokines and chemokines that may block insulin signaling and recruit other macrophages, which may propagate the low-grade inflammation and lead to further insulin resistance (Olefsky and Glass, 2010). High-fat diet has been found in experimental animal models to induce an inflammatory response in the hypothalamic areas that control feeding behavior and energy homeostasis by regulating downstream neurons (Velloso, 2009). Hypothalamic inflammation may result in the breakdown of the circuitry that maintains balance between energy intake and energy expenditure. Therefore, chronic low grade inflammation, in conjunction with a high calorie diet, is a possible mechanism that may contribute to the diseases associated with obesity (Hirai et al., 2010).

Several previous studies (Gunstad et al., 2008; Soreca et al., 2009; Ward et al., 2005) report that elevations in BMI are associated with the decreased volume in the frontal lobes and hippocampi (Raji et al., 2010), as well as the reductions in GM density in several regions involved in the taste and reward pathways (Pannacciulli et al., 2006). Additionally, Tataranni et al. (1999) suggest key roles in motivational feeding behavior for the paralimbic and limbic areas, such as the orbitofrontal cortex (OFC) and the hippocampal formation. Recent rodent and human studies also demonstrate that peripheral inflammation is associated with reductions in hippocampal gray matter volume (Marsland et al., 2008). Based on these observations, we hypothesized that the low grade inflammation present among overweight and obese individuals may be associated with structural/functional alterations to the orbitofrontal, limbic, and paralimbic brain structures that regulate feeding behavior. We used fibrinogen, a pleiotropic glycoprotein, as our marker of inflammation as it is associated with neuroinflammation (Ryu et al., 2009) and predicts weight gain in middle-aged adults (Duncan et al., 2000).

To test our hypotheses we utilized Diffusion Tensor Imaging (DTI) to generate brain apparent water diffusion coefficient (ADC) maps for lean and overweight groups. After the ADC maps were created we utilized a voxelwise correlational analysis, which is a fully automated whole-brain unbiased method, to ascertain regionally specific associations between fibrinogen and ADC value for each of the two groups. In addition, by utilizing structural images and operator-determined volumes, or using FreeSurfer, we derived the gray matter (GM) volume of medial and lateral OFC and hippocampus, structures that are involved in the circuitry of behavioral feeding control. We assessed group differences and their associations to inflammation in these specific regions of interest.

2. Results

The participants’ demographic and endocrine characteristics are summarized in Table 1. The groups were matched on

<table>
<thead>
<tr>
<th>Table 1 – Demographic and biological data.</th>
<th>Lean n=19</th>
<th>Overweight and obese n=44</th>
<th>T-Test</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male gender %</td>
<td>42%</td>
<td>52%</td>
<td>0.459</td>
<td>0.2</td>
</tr>
<tr>
<td>Age (years)</td>
<td>57.6±6.7</td>
<td>58.7±7.7</td>
<td>0.605</td>
<td>0.14</td>
</tr>
<tr>
<td>Education (years)</td>
<td>16.0±2.7</td>
<td>15.5±2.2</td>
<td>0.42</td>
<td>0.22</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>21.7±1.9</td>
<td>31.4±5.9</td>
<td>&lt;0.0001</td>
<td>1.92</td>
</tr>
<tr>
<td>Waist/hip ratio</td>
<td>0.94±0.08</td>
<td>0.99±0.06</td>
<td>0.003</td>
<td>0.99</td>
</tr>
<tr>
<td>Fibrinogen (mg/dl)</td>
<td>321±48.8</td>
<td>371±86.6</td>
<td>0.023</td>
<td>0.64</td>
</tr>
<tr>
<td>Hypertension %</td>
<td>42%</td>
<td>66%</td>
<td>0.078</td>
<td>0.49</td>
</tr>
<tr>
<td>Systolic BP (mm Hg)</td>
<td>114±16</td>
<td>123±19</td>
<td>0.84</td>
<td>0.48</td>
</tr>
<tr>
<td>Diastolic BP (mm Hg)</td>
<td>69±20</td>
<td>75±10</td>
<td>0.97</td>
<td>0.47</td>
</tr>
<tr>
<td>Glucose (mmol/l)</td>
<td>82±18</td>
<td>118±51</td>
<td>&lt;0.0001</td>
<td>0.81</td>
</tr>
<tr>
<td>Cholesterol</td>
<td>188±33</td>
<td>178±41</td>
<td>0.364</td>
<td>0.25</td>
</tr>
<tr>
<td>Triglycerides (mg/dl)</td>
<td>98±42</td>
<td>125±90</td>
<td>0.228</td>
<td>0.33</td>
</tr>
<tr>
<td>HbA1C (%)</td>
<td>5.83±1.5</td>
<td>6.52±1.80</td>
<td>0.021</td>
<td>0.65</td>
</tr>
<tr>
<td>Insulin (pmol/l)</td>
<td>6.0±4.6</td>
<td>13.8±10.5</td>
<td>&lt;0.0001</td>
<td>0.85</td>
</tr>
<tr>
<td>Periventricular WMH</td>
<td>0.40±0.89</td>
<td>0.70±0.70</td>
<td>0.40</td>
<td>0.39</td>
</tr>
<tr>
<td>Deep WMH</td>
<td>1.0±0.71</td>
<td>0.67±0.66</td>
<td>0.31</td>
<td>0.49</td>
</tr>
</tbody>
</table>

Data are given as mean±standard deviation except %data in percentage.

*Pearson chi-square test for data in percentage.

b Significant t-test (P value<0.05).

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gender, age, years of education, blood pressure (BP), hypertension, cholesterol, triglycerides, periventricular WMH and deep WMH. As expected, relative to lean subjects, overweight and obese individuals had significantly greater BMI ($t[58]=-9.69$, $P<0.0001$), as well as, larger waist/hip ratios and higher fibrinogen concentrations ($t[61]=-2.33$, $P=0.023$).

2.1. Association between gray matter volume and fibrinogen

Binary logistic regressions revealed that fibrinogen was associated with the residualized lateral OFC volumes which significantly classifies subjects with an excess of weight (76.2% of accuracy with a $P$ value of $<0.05$). However, the association of fibrinogen and the residualized hippocampal volumes (61.9% of accuracy with a $P$ value of $>0.1$), as well as the residualized medial OFC volumes (66.7% of accuracy with a $P$ value of $>0.08$), did not significantly predict a weight excess (see Table 2). The model involving fibrinogen and residualized lateral OFC is quite powerful at predicting overweight and obese individuals, with a sensitivity of 95.5% (see Fig. 1), but fails to predict lean individuals (specificity of 31.6%).

Given the sensitivity of the association between fibrinogen and residualized lateral OFC volumes to predict excess of weight, we wanted to ascertain whether fibrinogen predicted the lateral OFC volumes of the overweight and obese individuals. Correlational analyses, accounting for age and hypertension, showed that fibrinogen was significantly associated with the residualized-FreeSurfer-derived lateral OFC volume ($r=-0.37$, $P=0.016$) in the overweight and obese group but was only a statistical trend among lean participants ($r=-0.47$, $P=0.056$).

In order to ascertain the unique contribution of fibrinogen in explaining lateral OFC we ran hierarchical regression analyses among overweight and obese individuals and first controlled for factors that could also affect those volumes. In these analyses the normally distributed lateral OFC volumes were treated as the dependent variable to determine how much of their variance was explained by fibrinogen. After accounting for age and hypertension ($\Delta R^2=0.064$, $\Delta F P=0.285$), waist/hip ratio ($\Delta R^2=0.024$, $\Delta F P=0.330$), and lipid and sugar levels ($\Delta R^2=0.104$, $\Delta F P=0.242$), fibrinogen explained an additional and significant 9% of the variance ($\beta=-0.348$, $\Delta R^2=0.093$, $\Delta F P=0.046$) of the lateral OFC volume.

2.2. Association between GM ADC and fibrinogen

The voxelwise correlational analyses yielded different results for each of the two groups. Among lean individuals, and after adjusting for age and hypertension, we identified a total of 3 significant ($P<0.001$) cerebral clusters with an inverse correlation between GM ADC and fibrinogen. The largest cluster was located in the left prefrontal cortex and the other two were in the right parietal and left occipital lobes (see Table 3 and Fig. 2). In contrast, among overweight and obese individuals, after correcting for age and hypertension, we found 3 clusters of positive associations ($P<0.001$) between GM ADC and fibrinogen. These 3 significant clusters were located in the left and right amygdala, and the right parietal lobe (see Table 4 and Fig. 3).

3. Discussion

This study is the first to report an association between excess weight-associated inflammation and changes in some of the structures of the brain network thought to be involved in reward and eating behaviors. We demonstrated that among overweight and obese individuals, but not their lean counterparts, and after controlling for age, hypertension, waist/hip

<table>
<thead>
<tr>
<th>Table 2 – Binary logistic regression.</th>
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<tbody>
<tr>
<td><strong>Hippocampus</strong></td>
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<tr>
<td></td>
</tr>
<tr>
<td>Intercept</td>
</tr>
<tr>
<td>Fibrinogen</td>
</tr>
<tr>
<td>Volumes *</td>
</tr>
</tbody>
</table>

* Residualized volumes.
ratio and lipid and sugar levels, increased levels of fibrinogen were related to smaller lateral OFC volumes. Furthermore, we found a different anatomic pattern in the relationship between fibrinogen levels and microstructural integrity of GM regions (independent of age and hypertension) for the lean and overweight/obese individuals. Among lean individuals, higher fibrinogen levels were associated with lower ADC (less interstitial fluid) in the left prefrontal, right parietal and left occipital regions. However, among individuals with excess weight, elevations in fibrinogen concentration were associated with increased ADC (greater interstitial fluid) in both amygdala and the right parietal cortex. We did not expect to find an anatomical overlap in the areas detected by the different image analysis methods given the very different resolution and precision of the voxelwise and volumetric methods. To the best of our knowledge, this is the first report demonstrating that individuals with excess weight had reductions in microstructural integrity in some of the GM regions involved in eating behavior.

| Table 3 – Significant clusters of correlation between GM ADC and fibrinogen among lean individuals (P<0.001). |
|---|---|---|---|
| Talairach coordinates | Size (#voxels) |
| x | y | z |
| Left prefrontal | 56 | 8 | -3 | 146 |
| Right parietal | -30 | 97 | 12 | 145 |
| Left occipital | 30 | 113 | -19 | 132 |

* Represents the centroid of the cluster.

| Table 4 – Significant clusters of correlation between GM ADC and fibrinogen among overweight and obese individuals (P<0.001). |
|---|---|---|---|
| Talairach coordinates | Size (#voxels) |
| x | y | z |
| Left amygdala | 24 | 23 | -37 | 182 |
| Right amygdala | -23 | 24 | -37 | 155 |
| Right parietal | -52 | 54 | 19 | 124 |

* Represents the centroid of the cluster.

Fig. 2 – Brain regions showing clusters of association between GM ADC and fibrinogen among lean individuals. Each column represents the 3 orthogonal orientations (axial, coronal, and sagittal) for the significant inverse correlation clusters (analysis controlling for age; minimum cluster size 100 voxels; p<0.001) overlaid on the T1 target image. The color bar represents the strength of the correlation.
We provided evidence that the fibrinogen concentrations differed significantly between lean subjects and individuals carrying excess weight. This is consistent with many previous studies that describe obesity as a state of chronic low grade inflammation. In obesity there is an infiltration of the adipose tissue by macrophages, leading to increases in inflammatory cytokine production (Vachharajani and Granger, 2009). Although the source of inflammation in obesity is known, the direction of causality of these associations requires further investigation. For instance, it is still unclear whether excess weight (increased adiposity) leads to inflammation in the brain and peripheral tissues, or whether genetic factors give rise to hypothalamic inflammation, which in turn leads to loss of control of eating behaviors and obesity (Wisse and Schwartz, 2009). However, a series of experimental studies proves that fat-rich diet induces an inflammatory status in the hypothalamus and triggers the resistance to anorexigenic signals (De Souza et al., 2005; Milanski et al., 2009) and inflammation, in combination with obesity, may result in the dysregulation of insulin signaling (Uysal et al., 1997). Furthermore, the impairment in signaling can have deleterious effects on neurons in the hypothalamus leading to disruptions in the normal eating behavior (Levin et al., 2004; Sherwin, 2008). Additionally, inflammation can directly damage brain tissue by impairing blood vessel integrity leading to an increase in inflammatory cells in the cerebrospinal fluid and perivascular spaces in the brain (Man et al., 2007).

The goal of this study was to ascertain whether excess weight and its associated inflammation affect some of the limbic and paralimbic structures, structures involved in feeding behavior. We determined that the association of fibrinogen and the lateral OFC volumes was a good model to predict an excess of weight. However, the fibrinogen and medial OFC volume and hippocampal volume were not strong predictors of excess weight. Moreover, there was an association between the increased inflammation and reductions in the volume of the lateral OFC. The lateral OFC plays a key role in the reward system by integrating information regarding reward outcome, and damage to this area may result in impaired decision making (Wallis, 2007). It is important to note that the OFC neurons receive gustatory and olfactory stimuli and respond to specific tastes and odors. In fact, the OFC contains the major cortical representation of tastes including sweet, salty, bitter and sour. These taste

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**Fig. 3** - Brain regions showing clusters of association between GM ADC and fibrinogen among overweight and obese individuals. Each column represents the 3 orthogonal orientations (axial, coronal, and sagittal) for the significant positive correlation clusters (analysis controlling for age; minimum cluster size 100 voxels; $p<0.001$) overlaid on the T1 target image. The color bar represents the strength of the correlation.

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representations can act as positive or negative reinforcers for the reward system (Rolls, 2000); therefore, the ability of the OFC to code rewards is likely to impact food selection (Zald, 2009). The support for the role of the OFC in food selection comes from an fMRI study reporting differential activation of this region during imagined intake of palatable versus unpalatable food (Slichter et al., 2010). It is possible that inflammation, by affecting the integrity of the OFC, impacts the reward system, and thus loosens the control on feeding behavior resulting in weight gain.

Contrary to our expectations, we found that age and hypertension did not significantly account for much variance in the OFC volume, which is in contrast with previous positive associations reported by Raz et al. (1997). They found a substantial age-related decline in the volume of the prefrontal gray matter in 148 healthy volunteers ranging in age from 18 to 77 years. Our narrower age-range and smaller sample size may be the reason why we did not observe an association.

Our DTI correlational analyses detected increased water diffusivity (an indication of increased water in GM possibly indicating loss of microstructural integrity) associated with increased fibrinogen concentration in the brains of overweight and obese individuals after accounting for age and hypertension. Increases in GM diffusion may be considered an increase of tissue water movement representing very subtle “atrophy” of the tissue. The mechanism(s) underlying this increase in diffusivity still needs to be clarified. ADC represents an average value of the diffusivity inside a voxel and we cannot determine if the ADC mainly reflects intracellular or extracellular diffusion. What is clear is that GM ADC, which increases in patients with the ADC mainly reflects intracellular or extracellular diffusion. Increases in GM diffusion may be considered an increase of tissue water movement representing very subtle “atrophy” of the tissue. The mechanism(s) underlying this increase in diffusivity still needs to be clarified. ADC represents an average value of the diffusivity inside a voxel and we cannot determine if the ADC mainly reflects intracellular or extracellular diffusion.

One of the most striking findings of this study was the strong positive correlation between inflammation and ADC in both the right and left amygdala among overweight and obese individuals. Both animal and human studies highlight key roles for the amygdala in feeding behavior and the reward system. Specifically, the amygdala is involved in the motivational control of appetite (Grundmann et al., 2005). Lesion studies in rats show that the amygdala and hypothalamus are part of the same pathway that regulates weight balance and feeding behavior (Hinton et al., 2004), and the activation of the amygdala by stress may be reduced by consuming high caloric foods (Dallman et al., 2003). In addition, the amygdala plays a role in food preference and selection; studies in primates demonstrate that animals with amygdala lesions are less discriminating and increase their selection of food that normal animals refuse. These data suggest that the amygdala performs a role in the avoidance of unpalatable foods (Machado and Bachevalier, 2007). Moreover, disconnecting the OFC from the amygdala in primates demonstrates a direct functional interaction between these two structures and suggests that they compose a neural system controlling adaptive response selection and decision making (Baxter et al., 2000). Furthermore, human data utilizing positron emission tomography provides evidence for the dissociable contribution of the amygdala and OFC in motivation and decision making (Arana et al., 2003). The amygdala is activated by high incentive information regardless of whether a choice is required, whereas, the medial OFC and the lateral OFC are both recruited during incentive judgment and goal selection.

Our results in overweight and obese individuals (but not lean) showed smaller OFC volumes and increased ADC in the amygdala was associated with elevations in the fibrinogen level, which may suggest structural and functional impairments in the circuitry controlling feeding behavior. The negative correlation between ADC and fibrinogen in some prefrontal, parietal and occipital brain regions of the lean group may be explained by the varying biological functions of fibrinogen depending on its levels. For instance, a high level of fibrinogen may be indicative of an inflammatory process, but at lower concentrations the fibrinogen in the blood contributes to cell proliferation, adhesion and migration and even myelination (Adams et al., 2004). In addition to the amygdala, we also found a positive correlation between fibrinogen and parietal and occipital areas in the overweight and obese participants. We postulate that these additional regions could be related to the chronic state of inflammation, which might impair the blood brain barrier more broadly, and that these were the regions that crossed statistical significance.

A significant strength of this report is that we used well-established automatic and unbiased methods for quantifying brain volumes. Further strengthening this report, we used operator determined volumes for the hippocampus, a structure with poor contrast differentiation from other adjoining gray matter regions such as the amygdala and the entorhinal cortex and for which automated methods such as FreeSurfer are not validated (Morey et al., 2009a, 2009b). Another strength of this report is that our participants were well matched on years of education, which is highly associated with socioeconomic status, thus removing this as a possible confound. Fibrinogen is an appropriate marker of neuroinflammation but we plan to assess other markers of systemic inflammation in future studies.

One of the potential weaknesses of this study is that even though sophisticated techniques were used to correct for the spatial distortions that occur in the echo-planar acquisition of the DTI data, we cannot completely rule out that these distortions did not influence our amygdala findings. However, this is highly unlikely given that both hemispheres were affected. Although direct measurement of the hypothalamus would be of great interest in a study such as this, currently there are no reliable and/or valid methods for obtaining hypothalamic volumes or even for the placement of regions of interest in this area on MRI images in humans. Better measurement methods need to be developed before this key structure is included in future studies.

Our study further affirms the suggestion that excess weight should be regarded as a disease that may have anatomical and physiological cerebral abnormalities associated with the phenotype.
4. Conclusion

This is the first report utilizing DTI-based MRI assessments of water diffusivity as well as structural volume measurements to ascertain the associations between obesity-mediated inflammation and abnormalities in the brain structure involved in the control of feeding behavior. This report provides evidence for a connection between inflammation, amygdala integrity and the volume of lateral OFC.

5. Experimental procedures

5.1. Participants

Sixty-three middle aged and elderly volunteers were recruited as part of a study of normal aging and metabolic dysregulation associated with obesity. The protocol was approved by the Institutional Research Board of the NYU School of Medicine. All participants signed written informed consent and were compensated for their time and inconvenience. Subjects were screened for cerebrovascular disease by inspection of the modified Fazekas Scale. A comprehensive panel of blood chemistries, glucose and insulin values. Significant head trauma, stroke, hydrocephalus, lacunar infarcts, seizures, mental retardation, or any neurological disorder excluded subjects from participation. All subjects were screened for cerebrovascular disease by inspection of white matter hyperintensity (WMH): score above 2 WMHs on the modified Fazekas Scale. A comprehensive panel of blood tests was performed after a 10–12 h overnight fast for the assessment of blood count, liver and lipid profiles, thyroid function, blood chemistries, glucose and insulin values.

Body mass index (BMI), computed as the weight in kilograms divided by height in meters squared, was used to identify overweight and obese individuals. Another measure of adiposity, waist/hip ratio, was used to account for differences in muscle mass. Participants were considered lean if they had a BMI between 18.0 and 24.9 kg/m², those with BMI between 25.0 and 29.9 kg/m² were considered overweight and those with BMI ≥ 30 kg/m² were considered obese individuals.

Subjects were considered hypertensive if they received an antihypertensive medication or had a systolic blood pressure of 140 mm Hg or higher, or a diastolic blood pressure of 90 mm Hg or higher. Subjects were considered hypertensive if they received antihypertensive medication or had a systolic blood pressure of 140 mm Hg or higher, or a diastolic blood pressure of 90 mm Hg or higher. Blood pressure was measured in triplicate for each subject.

Plasma fibrinogen concentration was measured by the prothrombin-time derived method with reference to the Clauss fibrinogen assay using ACL TOP 500 CTS coagulation analyzer with closed tube sampling (Instrumentation Laboratory, Beckman Coulter Inc.). The prothrombin time was calculated as the difference between the patient's sample and the reference.
et al. (2007) using the Multimodal Image Data Analysis System (MIDAS).

The MPRAGE scans were also used to determine the intracranial vault size, which is obtained by manually outlining the supratentorial compartment. This was done by underlining the margins of the dura and the tentorium on the sagittal images as described in detail elsewhere (Gold et al., 2007). To account for the individual variability in brain size, both the operator-derived and the FreeSurfer estimates of the GM volumes were residualized to intracranial volume. All the operator-derived volumes were conducted blind to the identity and BMI of participants.

5.3 Statistical analysis

We checked the normality of our data using the Shapiro–Wilk test. We performed two-tailed independent sample t-tests and chi-square tests to examine the group differences in demographic, endocrine and MRI brain volumetric data. The effect sizes are reported as Cohen’s coefficient. Data were analyzed using SPSS for Windows version 17.0 (SPSS, Inc., Chicago, IL). We used Matlab® version 2008a to carry out the binary logistic regressions that predict the BMI, defined as a categorical dependent variable, from the continuous predictor variables, fibrinogen and residualized volumes. Given the association between fibrinogen and residualized volumes, we predicted whether a particular subject belongs to the lean or the overweight and obese groups. Our model was constructed using maximum likelihood estimation by iteratively reweighted least squares and the goodness of the fit was assessed by the likelihood ratio test. Furthermore, using SPSS, we assessed the strength of the significant association using two-tailed partial correlation and accounted for age and hypertension and a hierarchical regression approach and accounted for potential confounders, such as age (step 1), and waist/hip ratio (step 2). In step 3, we controlled for hypertension, lipid levels and blood sugar, all of which may be associated with microvascular disease, especially in an older population. Fibrinogen acted as a surrogate marker for inflammation; therefore, fibrinogen was included as the last step in the regression to demonstrate the additional power of inflammation on structural changes and to determine if obesity-induced inflammation (fibrinogen values) were associated with the integrity of brain areas involved in the control of feeding behavior. To ascertain the possible associations between fibrinogen levels and the GM tissue density measured on the DTI images (ADC), we computed the correlations between ADC and fibrinogen using voxelwise correlational analysis (Hoptman et al., 2004). To reduce the risk of the escalation of type 1 error due to multiple comparisons, only clusters of at least 100 contiguous significant voxels (0.1 cc) were defined as a significant. We chose a false discovery rate (FDR) less than 0.01 according to the original Benjamini and Hochberg (1995) procedure. We selected a significance (p-value) threshold of 0.001 to ensure that the FDR would be kept below 0.01. Because of the wide age distribution (aged 42 to 74 years) we controlled for age and hypertension to ensure that the correlations were not driven by either age or hypertension-related changes in the brain structure. The correlation map was registered to the standard Montreal Neurological Institute (MNI) T1 MRI template and visualized with AFNI (Analysis of Functional NeuroImages).

Acknowledgments

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