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Brain putamen volume changes in newly-diagnosed patients with obstructive sleep apnea

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Abstract

Obstructive sleep apnea (OSA) is accompanied by cognitive, motor, autonomic, learning, and affective abnormalities. The putamen serves several of these functions, especially motor and autonomic behaviors, but whether global and specific sub-regions of that structure are damaged is unclear. We assessed global and regional putamen volumes in 43 recently-diagnosed, treatment-naïve OSA (age, 46.4 ± 8.8 years; 31 male) and 61 control subjects (47.6 ± 8.8 years; 39 male) using high-resolution T1-weighted images collected with a 3.0-Tesla MRI scanner. Global putamen volumes were calculated, and group differences evaluated with independent samples t-tests, as well as with analysis of covariance (covariates: age, gender, and total intracranial volume). Regional differences between groups were visualized with 3D surface morphometry-based group ratio maps. OSA subjects showed significantly higher global putamen volumes, relative to controls. Regional analyses showed putamen areas with increased and decreased tissue volumes in OSA relative to control subjects, including increases in caudal, medial, mid-ventral portions, and ventral regions, while areas with decreased volumes appeared in rostral, medial, caudal, and mid-ventral sites. Global putamen volumes were significantly higher in the OSA subjects, but local sites showed both higher and lower volumes. The appearance of localized volume alterations points to differential hypoxic or perfusion action on glia and other tissues within the structure, and may reflect a stage in progression of injury in these newly-diagnosed patients toward the overall volume loss found in patients with chronic OSA. The regional changes may underlie some of the specific deficits in motor, autonomic, and neuropsychologic functions in OSA.

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1. Introduction

Obstructive sleep apnea (OSA) patients show multiple neurobehavioral difficulties, including deficits in attention, psychomotor and executive functioning, memory, and affect (Beebe et al., 2003). However, the principal deficits in the syndrome involve the loss of motor control over the upper airway muscles, which fail to discharge in a timely fashion, when they should dilate the airway during diaphragmatic descent, and a loss of control over autonomic motor activity, with chronic, exaggerated sympathetic discharge (Somers et al., 1995), and inappropriate lagged or muted dynamic sympathetic responses to blood pressure or ventilatory challenges (Harper et al., 2003; Henderson et al., 2004; Macey et al., 2006). Those somatic and autonomic motor defects suggest central injury in motor and autonomic regulatory structures, and gross damage has been found in multiple sites, based on manual and voxel-based morphometry of high-resolution T1-weighted images (Kumar...
et al., 2008; Macey et al., 2002), diffusion tensor imaging (Kumar et al., 2012; Macey et al., 2008), and T2-relaxometry procedures (Cross et al., 2008; Kumar et al., 2009b). Several of the affected brain sites include structures that serve both somatomotor and autonomic motor roles; one such structure is the putamen, which shows overall injury, metabolic abnormalities, and functional deficits (Alkan et al., 2013; Emin Akkoyunlu et al., 2013; Kumar et al., 2012; Macey et al., 2006).

The evidence for putamen roles in autonomic regulation is well-known, especially from pathological conditions with dysautonomia, such as Multiple Systems Atrophy (Pastakia et al., 1986). Such a role is expected, considering the major projections received from the autonomically regulated insular cortices (Saper, 1982). The putamen somatic motor role appears to incorporate sensory and other signals to initiate effective motor action, with putamen stroke inducing conditions of “motor neglect” (Sapir et al., 2007). OSA represents a prime example of how signals fail to initiate and maintain upper airway muscle discharge, and the putamen may be playing a significant role in that upper airway muscle “neglect,” particularly considering its projections to brain areas regulating oral motor functions (Kunzle, 1978; Miyata and Sasaki, 1984).

The putamen also contributes to regulation of other neuropsychologic, cognitive, and learning functions which are deficient in OSA. These functions include memory, mood, language, and motivation, show differential topographic organization (Alexander and Crutcher, 1990; Badgaiyan et al., 2008; Booth et al., 2007; Bowman et al., 1996; Husain et al., 1991; Saper, 1982), and the sub-regions within that organization receive information from prefrontal, insular, and thalamic areas, and send dopaminergic projections to the ventral tegmentum, substantia nigra, and globus pallidus (Avendano et al., 2006; DeVito et al., 1980; Ferry et al., 2000; Gorbachwskaya, 1997; Saper, 1982; Schultz and Romo, 1987). Putamen volumes are reduced in depressed patients (Husain et al., 1991), and over half of OSA patients are depressed (Asghari et al., 2012); the majority of OSA subjects also show learning deficits (Wallace and Bucks, 2013).

Regional structural changes within the putamen can thus alter information transfer between various brain sites. Manual volumetric procedures, using high-resolution T1-weighted images, can evaluate global putamen volume (Kumar et al., 2011), and 3D surface morphometry techniques can assess regional tissue changes, as well as extent of changes within the structures (Butters et al., 2009; Sabattoli et al., 2008; Shi et al., 2013; Thompson et al., 2004). Both procedures have been used to examine global and regional basal ganglia structures (Kumar et al., 2009a), including the putamen (Kumar et al., 2011). The extent of tissue changes within putamen sub-regions which serve specialized functions, or project to other brain areas serving unique roles, is unclear in OSA. Knowledge of localized changes is essential to disclose functional outcomes, since the functional topography of the putamen is not uniform.

Our aim was to examine global and regional putamen volumes in OSA relative to control subjects using high-resolution T1-weighted images with manual volumetric and 3D surface morphometry procedures, and to determine the nature of any volume changes. We selected newly-diagnosed, untreated OSA patients to avoid confounding measures with treatment interventions, and to determine tissue changes as early as possible in the syndrome development. Since OSA subjects show overall putamen damage, altered metabolites, and functional deficits (Alkan et al., 2013; Emin Akkoyunlu et al., 2013; Kumar et al., 2012; Macey et al., 2006) and severe autonomic, behavioral and motor deficits appear in the condition, we hypothesized that both left and right global and regional putamen integrity would be altered in OSA, compared to control subjects.

2. Materials and methods

2.1. Subjects

We studied 43 recently-diagnosed (based on overnight polysomnography; apnea–hypopnea-index ≥ 5) OSA subjects before any treatment, and 61 healthy control subjects. Both OSA and control subjects included here were subjects in previously-published manuscripts related to other issues in OSA (Cross et al., 2008; Kumar et al., 2008, 2009b, 2012; Macey et al., 2008, 2010, 2012, 2013). All OSA subjects were recruited from the Sleep Disorders Laboratory at UCLA, and control subjects were recruited from the UCLA campus and West Los Angeles area. OSA subjects were not taking any medications (β-blockers, α-agonists, angiotension-converting enzyme inhibitors, vasodilators, or serotonin reuptake inhibitors), and were without any diagnosed history of stroke, heart failure, or mental illness that might influence brain tissue composition. Control subjects were in good health, without any brain disorder which may induce brain tissue changes. We interviewed the control subjects, as well as their sleep partners, when available, to determine the potential for sleep disordered breathing, and subjects suspected of having such disturbed patterns underwent an overnight sleep study. OSA and control subjects with body weight more than 125 kg (scanner limitation), or with MRI incompatible metallic implants were excluded, as described at the MRI safety website (http://www.mrisafety.com/). All OSA and control subjects gave written and informed consent prior to the study, and the study protocol was approved by the Institutional Review Board at the University of California at Los Angeles.

2.2. Overnight polysomnography (PSG)

Overnight sleep studies were performed on all OSA subjects at the UCLA Sleep Disorders Center and Laboratory, and consisted of a 7–10 hour period (9:00 PM to 6:00 AM) monitoring of electroencephalogram (EEG — central and occipital), digastic electromyogram (EMG), electrocardiogram (EKG — lead II), right and left extra-ocular eye movement (EOG), thoracic and abdominal wall movement, air flow, O2 saturation, end-tidal CO2 levels, snore volume, bilateral leg movement, and sleep position. Acquired data were digitized, and stored on a computer for sleep evaluation.

2.3. Examination of daytime sleepiness and sleep quality

We used the Epworth Sleepiness Scale (ESS) and the Pittsburgh Sleep Quality Index (PSQI) questionnaires to examine daytime sleepiness and sleep quality, respectively, in both OSA and control subjects (Johns, 1991; Knutson et al., 2006). These commonly-used measures were self-administered, either before or after MRI data acquisition.

2.4. Assessment of depression and anxiety symptoms

We administered the Beck Depression Inventory (BDI)-II to examine depressive symptoms, and the Beck Anxiety Inventory to assess anxiety symptoms in OSA and control subjects (Beck et al., 1988, 1996), either before or after MRI data collection. Both inventories are self-administered questionnaires, with scores between 0 and 63, based on the severity of symptoms (Beck et al., 1988, 1996).

2.5. Magnetic resonance imaging

Brain studies were performed using a 3.0-Tesla MRI scanner (Magnetom Tim-Trio; Siemens, Erlangen, Germany), with a receive-only 8-channel phased-array head-coil, and a whole-body transmitter coil. Foam pads were used on both sides of the head to reduce head motion related artifacts, and subjects lay supine during data collection. We collected two high-resolution T1-weighted image scans using a magnetization prepared rapid acquisition gradient-echo (MPRAGE) pulse sequence [repetition time (TR) = 2200 ms; echo time (TE) = 2.2 ms; inversion time = 900 ms; flip angle (FA) = 9°; matrix size = 256 × 256; field of view (FOV) = 230 × 230 mm; slice thickness = 1.0 mm; slices = 176]. Proton density (PD) and T2-weighted images were collected using a dual-echo turbo spin-echo pulse sequence (TR = 10,000 ms; TE1,
2 = 17, 134 ms; FA = 130°; matrix size = 256 × 256; FOV = 230 × 230 mm; slice thickness = 4.0 mm; slices = 55; turbo factor = 5). Diffusion tensor imaging data were acquired using a single-shot echo-planar imaging with twice-refocused spin-echo pulse sequence (TR = 10,000 ms; TE = 87 ms; flip angle = 90°; band-width = 1346 Hz/pixel; matrix size = 128 × 128; FOV = 230 × 230 mm; slice thickness = 2.0 mm, b = 0 and 700 s/mm², diffusion directions = 12). Four separate DTI series were collected with the same imaging protocol for subsequent averaging. We used a parallel imaging technique, “generalized autocalibrating partially parallel acquisition (GRAPPA)”, with an acceleration factor of two for all scans.

2.6. Data processing and analysis

2.6.1. Overnight PSG data

All sleep recordings were interpreted by the UCLA sleep physicians. Sleep stages, sleep disorders breathing, and physiology were assessed first for total sleep time, and number and type (central, obstructive, or mixed — based on air flow and thoracic and abdominal strain gauges) of apneic episodes. Sleep records were scored by a registered PSG technician, and then each PSG was reviewed by the sleep physician. Using the digitized PSG signals, each one-min data epoch were scored as awake (W), rapid-eye movement sleep (R), or sleep stages N1–N3, based on recently-revised scoring criteria (Iber et al., 2007). The apnea–hypopnea-index (AHI), is a severity index of sleep disordered breathing, combining both apneas and hypopneas, and is calculated by dividing the number of apnea and hypopnea by the total sleep time. Subjects were classified as having OSA based on an AHI ≥ 5/h (1999). OSA was classified as “moderate” if AHI > 15, and “severe” if AHI > 30 (1999). The total number and duration of apnea, AHI, number of arousals, O₂ saturation variables, total sleep time, time in each state, and sleep efficiency variables were calculated.

2.6.2. MRI data

High-resolution T1-weighted, PD-, and T2-weighted scans were visually-examined to assess any major brain lesions (e.g., cystic lesions, major infarcts, or space occupying lesions). Both high-resolution T1-weighted scans were also examined for any imaging artifacts before processing. The high-resolution T1-weighted scans were resampled to 2.0 mm isotropic, and used to outline manually putamen structures, and determination of putamen volumes.

2.6.2.1. Realignment, averaging, and rigid-body transformation. We realigned both high-resolution T1-weighted scans to remove any potential variations between the two series, and averaged the images using SPM8 software. Averaged images of individual subjects, using unified segmentation, were bias-corrected for regional signal intensity differences, and image volumes were reoriented into Montreal Neurological Institute (MNI) common space with an affine transformation (Kumar et al., 2011). The affine-transformed images were resampled to a voxel size of 0.9 × 0.9 × 0.9 mm, and used to outline manually putamen structures (Kumar et al., 2011).

2.6.2.2. Gray matter, white matter, and cerebrospinal fluid and intracranial volume calculation. The averaged and reoriented images of individual subjects were partitioned into gray, white, and cerebrospinal fluid (CSF) probability maps using the unified segmentation method. Within the gray, white, and CSF probability maps, all voxels with a probability value > 0.5 were counted, and whole-brain gray, white, and CSF volumes were calculated. Whole-brain gray, white, and CSF volumes were added to determine total intracranial volume (TIV).

2.6.2.3. Putamen tracing and global volume quantification. We used the reoriented and resampled brain scans to manually-outline both left and right putamen structures with MRcron software. One investigator, who was blinded for subject group assignments, outlined putamen structures in all OSA and control subjects, based on a protocol described earlier (Kumar et al., 2011). Using these tracings, left and right global putamen volumes were calculated (Kumar et al., 2011).

2.6.2.4. Calculation of putamen mean diffusivity (MD) values. DTI data were processed as described previously (Kumar et al., 2012). Briefly, using diffusion weighted and non-diffusion weighted images, diffusion tensor matrices were calculated, matrices diagonalized, and principal eigenvalues were calculated. The principal eigenvalues were used to calculate MD maps, as described earlier (Kumar et al., 2012). MD maps, derived from each DTI series, were realigned, averaged to create one MD map per subject, and averaged MD maps were normalized to a common space. Similarly, non-diffusion images were realigned, averaged, and normalized to a common space. All normalized non-diffusion images were averaged to determine a background image, which was used to create left and right putamen regions-of-interest masks, guided by the previous findings (Kumar et al., 2012). Using the normalized MD maps and left and right putamen region-of-interest masks, average left and right MD values were derived.

2.6.2.5. Intra-tracer reliability. We examined intra-tracer reliability for outlining putamen structures by re-outlining those structures in 10 randomly-selected OSA and 10 control subjects by the same investigator who outlined structures in all OSA and control subjects. Total putamen volumes were used for intra-tracer reliability calculations. Intra-tracer reliability was high for putamen tracings (intra-class correlation = 0.91, p = 0.001), indicating consistent tracings across all subjects.

2.6.2.6. 3D Surface morphometry. We used 3D surface morphometry to visualize regional putamen volume variations between OSA and control subjects (Butters et al., 2009; Sabattoli et al., 2008; Shi et al., 2013; Thompson et al., 2004). Both left and right surface models were created from individual putamen tracings, and 3D meshes were generated in which each mesh consisted of a fixed number of equally-spaced points. A medial curve representing center of mass was determined in the anterior–posterior direction within each mesh, and using the medial curve, the thickness of the putamen (radial distance) was calculated at each surface point (Thompson et al., 2004). To visualize the local vol-

2.7. Statistical analysis

Data were evaluated using the IBM Statistical Package for the Social Sciences (IBM SPSS, V 20). We examined numerical demographic, biophysical, sleep, and neuropsychologic data (age, body mass index, ESS, PSQI, BDI-II, and BAI) with independent samples t-tests, and categorical demographic data (gender) with Chi-square tests. Global putamen volumes, putamen MD values, gray matter, white matter, CSF, and TIV values were examined between groups using independent samples t-tests. Global putamen volumes and putamen MD values were also examined with analysis of covariance, with age, gender, and TIV included as covariates for putamen volumes, and age and gender as covariates for MD values. Pearson’s correlation procedures were used to examine the association between global putamen volumes and BMI, sleep variables, neuropsychologic scores, putamen MD values, and O₂ saturation variables in OSA subjects. We used linear regression analysis to determine the impact of clinical, biophysical, and O₂ saturation values on global putamen volumes of OSA subjects. The dependent variables were left
and right putamen volumes, and independent variables were AHI, BMI, ESS, PSQI, BDI-II, BAI, and O2 saturation nadir values. We established intra-tracer reliability with an intra-class correlation procedure.

3. Results

3.1. Demographics

Demographic data are summarized in Table 1. Neither age nor gender differed significantly between OSA and control subjects (age, p = 0.56; gender, p = 0.38). Body mass indices were significantly higher in OSA compared to control subjects (p < 0.001).

3.2. Sleep, O2 saturation, and neuropsychologic variables

Sleep, O2 saturation, and neuropsychologic variables of OSA and control subjects are summarized in Table 1. Both ESS and PSQI scores significantly differed between OSA and control subjects (ESS, p < 0.001; PSQI, p < 0.001). BDI-II and BAI values were significantly higher in OSA, compared to control subjects (BDI-II, p < 0.001; BAI, p = 0.001).

3.3. Global gray, white, and CSF volumes and TIV

Global gray matter, white matter, CSF volumes, and TIV values of OSA and control group are summarized in Table 1. Gray matter, white matter, and CSF volumes, as well as TIV values, did not differ significantly between OSA and control subjects (gray matter, p = 0.40; white matter, p = 0.27; CSF, p = 0.62; TIV, p = 0.82).

3.4. Global putamen volumes and MD value differences

Both left and right global putamen volumes of OSA and control subjects are summarized in Table 1, and scatter plots are shown in Fig. 1. The left and right global putamen volumes were significantly higher in OSA compared to control subjects (independent samples t-tests; left, p = 0.041; right, p = 0.025); values were significantly higher in OSA, even after controlling for age, gender, and TIV (ANCOVA; left, p = 0.043; right, p = 0.027).

Average putamen MD values of OSA and control subjects are summarized in Table 1. Both left and right putamen MD values were significantly lower in OSA, compared to control subjects (independent samples t-tests; left, p = 0.017; right, p = 0.001); values were

### Table 1

Characteristics of OSA and control subjects.

| Variable type | Variables | OSA (Mean ± SD) | Controls (Mean ± SD) | [A] vs [B] p values |
|---------------|-----------|-----------------|----------------------|-------------------|
| Demographic   | Age (years) | 46.4 ± 8.8 (n = 43) | 47.4 ± 8.9 (n = 61) | 0.56 |
|               | Gender (Male-Female) | 31:12 | 39:22 | 0.38 |
| Biophysical   | BMI (kg/m²) | 30.1 ± 5.2 (n = 43) | 24.8 ± 3.7 (n = 61) | < 0.001* |
| Sleep         | AHI (events/h) | 31.4 ± 19.3 (n = 43) | – | – |
|               | ESS | 10.3 ± 4.9 (n = 43) | 5.5 ± 3.5 (n = 61) | < 0.001* |
|               | PSQI | 9.3 ± 4.0 (n = 43) | 3.8 ± 2.5 (n = 61) | < 0.001* |
| O2 saturation | Baseline SaO2 (%) | 96.3 ± 2 (n = 40) | – | – |
|               | SaO2 nadir (%) | 77.9 ± 10.3 (n = 36) | – | – |
|               | SaO2 nadir change (%) | 18.3 ± 10.6 (n = 36) | – | – |
|               | Obstructive length of nadir (s) | 40.6 ± 21.4 (n = 30) | – | – |
| Neuropsychological | BDI-II | 9.3 ± 8.3 (n = 43) | 3.9 ± 4.4 (n = 61) | < 0.001* |
|               | BAI | 10.7 ± 12.4 (n = 43) | 3.9 ± 5.0 (n = 61) | 0.001* |
| Brain (volume) | Gray matter (mm³) | 700,469.7 ± 73,557.7 (n = 43) | 713,303.3 ± 77,941.7 (n = 61) | 0.40 |
|               | White matter (mm³) | 445,347.5 ± 56,252.4 (n = 43) | 458,578.8 ± 63,000.5 (n = 61) | 0.27 |
|               | CSF (mm³) | 315,275.5 ± 167,583.0 (n = 43) | 258,249.2 ± 175,544.6 (n = 61) | 0.62 |
|               | TIV (mm³) | 1,461,092.6 ± 206,276.6 (n = 43) | 1,470,131.3 ± 199,550.1 (n = 61) | 0.82 |
| Putamen (volume) | Left putamen (mm³) | 4146.6 ± 660.0 (n = 43) | 3909.1 ± 502.4 (n = 61) | 0.041 |
|               | Right putamen (mm³) | 4152.8 ± 654.5 (n = 43) | 3897.6 ± 487.8 (n = 61) | 0.025 |
| Putamen (MD values) | Left putamen (×10⁻⁶ mm²/s) | 803.17 ± 84.52 (n = 43) | 870.50 ± 84.48 (n = 61) | 0.017 |
|               | Right putamen (×10⁻⁶ mm²/s) | 784.75 ± 67.94 (n = 43) | 889.12 ± 112.18 (n = 61) | 0.001** |

SD, standard deviation; BMI, body mass index; AHI, apnea-hypopnea-index; ESS, Epworth Sleepiness Scale; PSQI, Pittsburgh Sleep Quality Index; SaO2, O2 saturation; BDI-II, Beck Depression Inventory II; BAI, Beck Anxiety Inventory; CSF, cerebrospinal fluid; TIV, total intracranial volume; *, equal variances not assumed; **, p value adjusted for age, gender, and TIV; ##, p value controlled for age and gender.
3.5. Relationships between global putamen volumes and BMI, sleep, O2 saturation, and neuropsychologic variables

coefficients of correlation and significance levels among putamen volumes, BMI, sleep, O2 saturation, left and right putamen MD values, and neuropsychologic variables are tabulated in Table 2. significant correlations appeared between global putamen volumes, BMI (left r = −0.35 p = 0.022; right r = −0.32 p = 0.036), and putamen MD values (left r = −0.48 p = 0.035; right r = −0.46 p = 0.047) in OSA subjects. However, no significant correlations emerged between putamen volumes and AHI, PSQI, or ESS scores. the BAI scores significantly correlated with putamen volumes (left r = −0.39 p = 0.01; right r = −0.40 p = 0.008); however, BDI-II scores did not show any correlations. the O2 saturation nadir showed a significant association with left putamen volume (r = −0.34 p = 0.045); but, other O2 saturation variables did not show any relationships.

3.6. Regional putamen volume changes

multiple regions within the putamen showed increased and decreased ratios of radial distance of surface points from the medial curve, indicating increased or decreased regional volume in OSA, compared to control subjects. Regions with decreased volumes within the putamen in OSA subjects included bilateral rostral (Fig. 2a, b, f, n, and q), mid-dorsal (Fig. 2c, g, and h), medial-caudal (Fig. 2d, j, and p), and mid-ventral sites (Fig. 2k, p), and increased volumes included the left caudal-dorsal (Fig. 2e), right mid-dorsal (Fig. 2i), left mid-ventral portions (Fig. 2l, m), and right caudal-ventral area (Fig. 2o).

3.7. Putamen volume and linear regression analyses

Regression analyses findings are summarized in Table 3. Oxygen saturation nadir (p = 0.045, B = −20.46) was an independent predictor for left global putamen volume in OSA subjects. However, no variable independently predicted right global putamen volume in these subjects.
Injury to ventral putamen sub-regions served by dopamine-based neurotransmitters would have significant effects on affective behaviors dependent on reward (Seeman et al., 2006); those behaviors are heavily affected in OSA (Asghari et al., 2012). A primary characteristic of OSA is impaired autonomic nervous system regulation, reflected especially in excessive sympathetic activity and impaired timing to autonomic challenges (Harper et al., 2003; Henderson et al., 2002; Somers et al., 1995). Injury to both the insular cortices and dorsal putamen likely contributes to that deficit, since both regions serve autonomic roles; the putamen receives projections from the insular cortices (Saper, 1982), and both structures show overall injury and functional deficits during autonomic and respiratory challenges (Henderson et al., 2004; Kumar et al., 2012; Macey et al., 2006). Since a close relationship exists between transient elevation in blood pressure with both diaphragmatic (Trelease et al., 1985) and upper airway action (Marks and Harper, 1987), and upper airway muscle suppression is more sensitive to blood pressure elevation than the diaphragm, distorted regulation of dynamic sympathetic patterns has the potential to contribute to the “motor neglect” of upper airway muscles found in OSA.

### 4.4. Tissue volume changes: processes

Several mechanisms may underlie the global and regional putamen tissue changes. OSA subjects are intermittently exposed to hypoxia from the repeated, successive cessations of airflow during sleep, which is followed by rapid re-oxygenation at apnea termination. Moreover, exaggerated blood pressure swings occur over the course of each apneic event, with substantial changes in perfusion accompanying the vascular alterations (Yan et al., 2009). Hypoxic and ischemic conditions induce energy pump failure, altering sodium and potassium ion distribution, and affecting intracellular and extracellular water, resulting in neuronal and axonal swelling (Hossmann, 1971; Lowry et al., 1964; Mintorovitch et al., 1994; Oehmichen et al., 2001). The axonal and neuronal swelling increases volumes of affected brain structures. However, chronic stages of hypoxia/ischemia would be accompanied by a reduction in axons, neurons, and glia, leaving a net loss of tissue volume over control values. Since OSA subjects showed higher global putamen volumes, relative to control subjects, we believe that the tissue changes are representative of acute changes. These changes likely contribute to manifestations of characteristics in OSA.

Both increased and decreased localized volume changes appeared in OSA subjects within the putamen over controls, suggesting that the changes result from both early and long-standing tissue alterations in these newly-diagnosed, treatment-naïve subjects, or that portions of the putamen may be more vulnerable to hypoxic or perfusion changes through the nature of tissue composition or extent of vascularization. Overall, the predominant changes appear to be in acute stages, since global putamen volumes were found to be significantly increased in OSA over controls. These findings suggest that a series of changes occurs...
within tissue before axonal and neuronal loss, with early, acute stages of pathology associated with increased tissue volume, while chronic stages of pathology are accompanied by decreased volumes. Tissue changes in both acute and chronic stages of pathology likely contribute to functional deficits served by subareas within the structure.

It is possible, although speculative, that the OSA condition may result from putamen tissue changes, i.e., that the failure in upper airway action results from initial putamen dysfunction from injury which may progress, cumulating with the exaggerated autonomic and other manifestations found in OSA. Initial putamen injury, perhaps by carbon monoxide poisoning, accidental hypoxic exposure, or reaction to toxic substances acting on the multiple neurotransmitters in the structure, has the potential to interfere with normal putamen functioning, disrupting normal regulation of the upper airway. This initial dysfunction may escalate to a condition that self-perpetuates or worsens, with all of the autonomic and somatomotor outcomes reported for OSA. Determination of the feasibility of such an outcome would require basic studies of insults to the putamen in an animal model that results in interference with upper airway muscle action paralleling the coordination loss found in human OSA.

### 4.5. Limitations

One limitation of this study is the absence of data on the prediagnosis duration of OSA subjects. However, we are reasonably confident that the predominant brain tissue changes in the OSA subjects are acute in nature. MRI-based mean diffusivity (MD) procedures, which measure water diffusion within tissue, and are influenced by the presence of tissue barriers and extracellular/extra-axonal fluid, show reduced MD values in acute disease stages of ischemic/hypoxic pathology, but show increased values in chronic stages of such pathology (Ahlhelm et al., 2002; Chan et al., 2009; Hossmann et al., 1994; Loubinoux et al., 1997; Matsumoto et al., 1995; Shereen et al., 2011; Warach et al., 1996). Since these same OSA subjects showed reduced MD values across various brain sites (Kumar et al., 2012), we believe that the predominant tissue pathology in this sample derives from damage in acute stages.

### 5. Conclusions

Newly-diagnosed OSA subjects without treatment show significantly increased global putamen volumes over control subjects, with sub-regions within the structure showing both increased and decreased volumes. Regional sites with decreased volumes emerged in rostral, mid-dorsal, medial-caudal, and mid-ventral areas, and increased volumes appeared in caudal-dorsal, mid-dorsal, mid-ventral, and caudal-ventral portions. These areas contribute to regulation of neuropsychologic, autonomic motor, and somatomotor functions deficient in OSA, and especially to oral motor functioning, a special concern in the syndrome due to airway collapse in obstructive apnea. The pathological processes contributing to tissue changes may include intermittent hypoxia and impaired perfusion, both of which can affect glial tissue necessary for neuronal support; such tissue is extremely sensitive to hypoxia and ischemia. The finding of regional increased and decreased volume variation indicates that final volume loss earlier found in long-term OSA occurs with a progression of changes in the tissue.

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**Table 3**

Comparison of multivariate p values and B values of AHI, BMI, sleep, neuropsychologic, left and right putamen MD values, and O₂ saturation variables for the linear regression analyses in OSA subjects.

| Side | Covariates | Multivariate p values | B  |
|------|------------|-----------------------|----|
| Left putamen volume | AHI | 0.573 | – |
| | BMI | 0.175 | – |
| | ESS | 0.892 | – |
| | PSQI | 0.950 | – |
| | BDI-II | 0.897 | – |
| | BAI | 0.567 | – |
| | O₂ saturation nadir | 0.045 | –20.46 |
| Right putamen volume | AHI | – | – |
| | BMI | – | – |
| | ESS | – | – |
| | PSQI | – | – |
| | BDI-II | – | – |
| | BAI | – | – |
| | O₂ saturation nadir | – | – |

AHI, apnea-hypopnea-index; BMI, body mass index; ESS, Epworth Sleepiness Scale; PSQI, Pittsburgh Sleep Quality Index; BDI-II, Beck Depression Inventory II; BAI, Beck Anxiety Inventory; MD, mean diffusivity.
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