Increased Neuronal Depolarization Evoked by Autoantibodies in Diabetic Obstructive Sleep Apnea: Role for Inflammatory Protease(s) in Generation of Neurotoxic Immunoglobulin Fragment

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1 Abstract

1.1 Aim
Obstructive sleep apnea increases in diabetes and morbid obesity. We tested a hypothesis that circulating autoantibodies in adult type 2 diabetes which increase in association with morbid obesity are capable of causing long-lasting neuronal depolarization and altered calcium release in mouse atrial cardiomyocytes.

1.2 Methods
Protein-A eluates from plasma of 14 diabetic obstructive sleep apnea patients and 17 age-matched diabetic patients without sleep apnea were tested for effects on depolarization and neurite outgrowth in N2a mouse neuroblastoma cells. The mechanism of autoantibody-mediated neurite outgrowth inhibition was investigated in co-incubation experiments of diabetic obstructive sleep apnea autoantibodies with specific antagonists of G-protein coupled receptors or the RhoA/Rho kinase signaling pathway. Following long-term storage of the protein-A eluates (to allow spontaneous proteolysis and IgG subunit dissociation), plasma autoantibodies from diabetic obstructive sleep apnea autoantibodies with specific antagonists of G-protein coupled receptors or the RhoA/Rho kinase signaling pathway. Following long-term storage of the protein-A eluates (to allow spontaneous proteolysis and IgG subunit dissociation), plasma autoantibodies from diabetic obstructive sleep apnea, cancer or control patients were compared for enhancement of inhibitory effects on endothelial cell survival. Size exclusion chromatography performed (in the presence or absence of a specific membrane type 1 matrix metalloproteinase inhibitor) was used to characterize the IgG autoantibody subunit(s) or fragments associated with peak neurotoxicity in diabetic obstructive sleep apnea.

1.3 Results
Diabetic obstructive sleep apnea (n = 14) autoantibodies caused a significant increase (P = 0.01) in membrane depolarization in N2a mouse neuroblastoma cells compared to control diabetic patients (n = 15) not suffering with obstructive sleep apnea. Process extension in N2A mouse neuroblastoma cells was significantly inhibited (P = 0.01) by diabetic obstructive sleep apnea (n = 9) autoantibodies compared to effects from identical 10 µg/mL concentrations of control diabetic autoantibodies in patients without obstructive sleep apnea. Ten micromolar concentrations of SCH-202676, a G-protein coupled receptor antagonist (n = 5) or ten micromolar concentration of Y27632, a selective Rho kinase inhibitor (n = 6), each significantly prevented (P < 0.001) neurite outgrowth inhibition by diabetic obstructive sleep apnea autoantibodies. Autoantibodies in representative patients with obstructive sleep apnea and either atrial fibrillation or left ventricular hypertrophy evoked acute large increases in intracellular Ca2+ in HL-1 mouse atrial cardiomyocytes. The magnitude of intracellular Ca2+ release was dose-dependently significantly correlated to the electrocardiographic Cornell voltage-duration product. Gel filtration of diabetic obstructive sleep apnea autoantibodies revealed peak neurotoxicity associated with MWs corresponding to IgG light chain dimer(s), monomers or half-light chains as well as a novel ~ 5.5 kD putative light chain fragment.

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1.4 Conclusions
These results suggest that diabetic obstructive sleep apnea autoantibodies may induce strong depolarization in neuronal cells and alter Ca^{2+} signaling in atrial cardiomyocytes consistent with a role in pathophysiology in subsets of diabetic obstructive sleep apnea having co-morbid atrial fibrillation or another clinically significant cardiac rhythm disturbance.

1.5 Key words
Diabetes mellitus; Obstructive sleep apnea; Neurotoxicity; Autoantibodies; Atrial fibrillation

2 Introduction
Obstructive sleep apnea (OSA) is characterized by sleep-disordered breathing and excessive daytime sleepiness [1]. Obstructive sleep apnea increases in adult type 2 diabetes (T2DM) and may affect as many as four in ten older men with T2DM [2]. Visceral obesity is an important underlying risk factor in OSA [3], although the mechanism for the association is not well understood. Airway patency is normally maintained by motor neuron outflow to pharyngeal dilator muscles during wakefulness and sleep [4]. Since synaptic input to pharyngeal dilator motor neurons normally decreases during sleep [4] a humoral factor which suppresses local neuronal excitability might contribute to apnea or hypopnea. In prior studies, we reported circulating autoantibodies which altered spontaneous activity in electrically-excitable cells. The autoantibodies were increased in plasma of diabetic patients having a cluster of complications including: painful neuropathy [5], atrial fibrillation [6], major depressive disorder [7]; and in cancer fatigue/depression [5].

In the present study we tested a hypothesis that plasma IgG autoantibodies in diabetic obstructive sleep apnea cause long-lasting neuronal depolarization compared to autoantibodies in diabetic patients without obstructive sleep apnea. The mechanism of diabetic OSA autoantibodies’ inhibitory effect on neurite outgrowth in N2A neuroblastoma cells was evaluating using specific antagonists of G-protein coupled receptors or the RhoA/ Rho kinase signaling pathway.

3 Subjects and Methods
3.1 Diabetic obstructive sleep apnea and control patients
Informed consent was obtained from all study patients prior to blood drawing. Patients were enrolled from the diabetes and endocrinology outpatient clinics at the Veterans Affairs New Jersey Healthcare System, East Orange, NJ. The baseline clinical characteristics in the patients whose autoantibodies were tested for ability to evoke neuronal depolarization are summarized in Table 1. Painful diabetic nephropathy is defined according to previously reported criteria [5]. Diabetic nephropathy is defined as urinary albumin excretion ≥ 300 milligrams/gram creatinine or urinary protein excretion ≥ 500 milligrams/gram creatinine. Diagnostic criteria and clinical evaluation in diabetic depression patients were previously reported [7]. A diagnosis of obstructive sleep apnea was based on chart review and included patients with concurrent use of a nasal or face-mask continuous positive airway pressure (CPAP) device at night with or without diagnostic results from overnight polysomnography testing. Moderate obstructive sleep apnea is herein defined as an apnea-hypopnea index (AHI) of 15-30 per hour; severe OSA as an AHI > 30 per hour. Morbid obesity is defined as body mass index 35-40 kg/m² or above with associated medical co-morbidities and ‘super obesity’ as BMI 50 kg/m² or higher.

<table>
<thead>
<tr>
<th>Risk factor</th>
<th>OSA (n=14)</th>
<th>No OSA (n=17)</th>
<th>P-value</th>
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</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>66.2 ± 13.3</td>
<td>64.6 ± 7.7</td>
<td>0.71</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>39.0 ± 9.3</td>
<td>30.5 ± 6.6</td>
<td>0.008</td>
</tr>
<tr>
<td>Depression (yes/no)</td>
<td>(7/7)</td>
<td>(9/8)</td>
<td>1.0</td>
</tr>
<tr>
<td>Nephropathy (yes/no)</td>
<td>(5/9)</td>
<td>(2/15)</td>
<td>0.2</td>
</tr>
<tr>
<td>Painful neuropathy (yes/no)</td>
<td>(6/8)</td>
<td>(6/11)</td>
<td>0.72</td>
</tr>
<tr>
<td>Atrial fibrillation (yes/no)</td>
<td>(7/7)</td>
<td>(2/15)</td>
<td>0.04</td>
</tr>
</tbody>
</table>

OSA- obstructive sleep apnea; BMI- body mass index
^P-value from T-test (continuous variables) or χ² test (dichotomous variables) comparing patients with or without OSA.

3.2 Obstructive sleep apnea and atrial fibrillation or dementia
Pt 1: 59-year-old male with super obesity (BMI 60 kg/m²), type 2 diabetes and obstructive sleep apnea. He suffers from paroxysmal atrial fibrillation, asthma, and pituitary-macroprolactinoma. Pt 2: 69-year-old male with morbid obesity, type 2 diabetes and severe obstructive sleep apnea. He developed atrial fibrillation with slow ventricular response requiring permanent pacemaker placement Pts 3-68-year-old male with super obesity (BMI 58 kg/m²), type 2 diabetes, obstructive sleep apnea and permanent atrial fibrillation. Results of overnight polysomnography testing included: apnea-hypopnea index (AHI) 34 per hour, minimum O₂ saturation 88%. Pt 4: A 68-year-old Caucasian male (with family history of Alzheimer’s dementia) who suffered with obstructive sleep apnea, moderate obesity, atrial fibrillation requiring pacemaker placement, hypertension, diabetic nephropathy, painful neuropathy, and depression. The patient died of unknown causes after two-years’ follow-up observation and treatment.

Pt 5- A 68-year-old Caucasian male with morbid obesity, severe obstructive sleep apnea, left ventricular hypertrophy, and hypertension. He experienced several episodes of hard syncope re-
sulting in multiple rib fractures. He was diagnosed with bradycardia secondary to sick sinus syndrome and was treated with a dual chamber cardiac pacemaker. He underwent overnight polysomnography: apnea/ hypopnea index (AHI) was 32.8 per hour, his lowest oxygen saturation (O₂ sat) was 72% and he experienced more than 200 episodes of bradycardia with duration of 392 minutes in 456 minutes total recorded time. He was doing well until age 73 years, when he died of unknown causes.

Pt 6 - A 73-year-old moderately- obese, African-American male with atrial fibrillation, OSA and glaucoma. Overnigh polysomnography parameters included: AHI 33 per hour, and minimum O₂ saturation 79%. Pt 7 - 73-year-old obese Caucasian male with seizure disorder, OSA, and diabetic depression who later developed dementia. Pt 8 - 53-year-old thin adult-onset type 1 diabetes with seizure disorder, schizophrenia, nephropathy, and dementia. Pt 9 - 74-year-old obese type 2 diabetes male with microalbuminuria, congestive heart failure and peripheral vascular disease.

3.3 Blood drawing
Baseline plasma samples were obtained from study participants prior to the initiation of study procedures.

3.4 Protein A chromatography
Protein- A chromatography was carried out as previously described [8]. The protein-A-eluate fractions consisted of total IgG isolated from plasma upon low pH elution from the protein-A column. The active protein-A eluate caused significant inhibitory activity in endothelial cell survival.

3.5 Endothelial cell survival assay
Bovine pulmonary artery endothelial cells (Clonetics, Inc. San Diego, CA) were grown in Medium 199 plus 10% fetal calf serum and endothelial cell growth medium (EGM, Clonetics, Inc., San Diego, CA). Endothelial cell number assays were carried out as previously reported [8]. After 48 hours' incubation in the presence of protein-A-eluate fractions, cells were washed with PBS and processed for the colorimetric estimation of cell number, i.e. cell associated acid phosphatase activity, as previously described [8]. Growth-promoting activity is expressed as a percentage of the control cell number for cells grown in the absence of protein-A-eluate fractions.

3.6 Mouse neuroblastoma cells
Mouse neuroblastoma N2A cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM), containing 10% fetal bovine serum, (FBS) (Invitrogen, Carlsbad, CA) and penicillin/streptomycin (100 U/mL and 100 mg/mL, respectively) at 37°C and 5% CO₂. Cells were plated in 96- well plates for 3 days prior to membrane depolarization experiments.

3.7 N2A cell/ neurite out growth
Cells were plated at low density in 35 mm dishes. Next recombinant human bFGF (10 ng/mL) in the presence or absence of human IgG test fractions were added to dishes in triplicate. Groups of 50–100 cells/dish were counted 2 days after the addition of test factors. Neurite outgrowth represents the percentage of N2A cells expressing more than one neurite. A neurite is defined as a cell process that is at least 2 cell diameters in length of the cell body. Results are expressed as % N2A cells expressing neurites (which represent the mean ±SD of triplicate determinations) compared to neurite expression in cells grown with 10 ng/mL human bFGF, but without added test protein-A-eluate fractions.

3.8 Membrane depolarization assays
After cell attachment, growth medium was removed and cells were washed and then incubated in modified Tyrode’s solution consisting of: 150 mM NaCl, 3 mM KCl and 30 mM HEPES, 10 mM D-glucose and 2 mM CaCl₂, pH 7.4. Test fractions (human IgG fractions) were added in the presence of 97 nM DiBAC₄ (Molecular Probes, Eugene, OR) - as previously reported [6]. Fluorescence was measured after 5 min or longer at room temp using a Fluoroskan Ascent FL (VWR, Inc., Franklin, MA); Ex = 485 nm, Em = 538 nm. Results are expressed as percent of change in gross fluorescence compared to cells to which no test protein-A-eluate fractions were added.

3.9 HL-1 cell culture
HL-1 atrial cardiomyocytes were developed (and generously provided) by Dr. William Claycomb (Louisiana State University Medical Center, New Orleans, LA). They were maintained in 5% CO₂/ 95% air at 37°C in Claycomb media (Sigma, St. Louis, MO) containing 10% FBS (Biocell, Rancho Dominguez, CA), 100 U/mL:100 ug/mL penicillin-streptomycin (Invitrogen, Carlsbad, CA), 0.1 mM norepinephrine (Sigma, St. Louis, MO), and 2 mM L-glutamine (Invitrogen, Carlsbad, CA).

3.10 Intracellular calcium measurement
HL-1 cells were grown in -βT3 dishes (Bioptechs, Inc., Butler, PA) and loaded with 2 µM Fura-2 acetoxy methyl ester (Invitrogen, Carlsbad, CA) for 30 min at 37°C in a balanced salt solution (BSS) (140 mM NaCl, 2.8 mM KCl, 2 mM CaCl₂, 2 mM MgCl₂, 10 mM HEPES, pH 7.2) as previously reported [6]. The cells were then left for a further 10-minute period in the bathing solution at room temperature to allow ester hydrolysis to go to completion. Using a dual-wavelength spectrofluorometer (Photon Technology International, Monmouth Junction, NJ) with excitation wavelengths at 340 and 380 nm and emission at 510 nm fluorescence measurements were performed at room temperature on the stage of an inverted fluorescence microscope (Nikon TE200). The release of intracellular Ca²⁺ was measured following exposure to 1μg/ml concentrations of the protein-A eluate(s).

3.11 Size exclusion (gel filtration) chromatography
G75 superfine Sephacryl (Pharmacia, Piscataway, NJ) equilibrated in 10 mM sodium phosphate, 0.15 M NaCl, pH 7.4 was packed into a 5 x 0.7 cm column (Sigma, St. Louis, MO) having a volume of 2.2 mL. An aliquot of the protein- A eluate (0.05 mL).
was added to the column and eluted in 10 mM sodium phosphate, 0.15 M NaCl, pH 7.4 at a flow rate of 0.22 mL/min, at 20°C. Each individual collected fraction (0.11 mL) was assayed for effects on N2A neurite outgrowth. For the experiments involving inhibition of endogenous MT1-MMP, protein-A eluates (100 µL) were incubated in the presence or absence of NSC 405020 (200 µM) in 50 mM HEPES, pH 6.8, containing 10 mM CaCl₂, at 37°C for 14 hours prior to size exclusion chromatography.

3.12 Chemicals

Protein-A agarose was obtained from Pierce Chemical Co., (Rockford, IL). N-(2,3-diphenyl-1,2,4-thiadiazol-5-(2H)-ylidene)methanaminehydrobromide (SCH-202676) and 3,4-dichloro-N-(1-methylbutyl)-benzamide, (NSC 405020) were from Sigma Chem. Co., Inc. (St Louis, MO). All other chemicals and reagents were analytical grade.

3.13 Protein determinations

Protein concentrations were determined by a bicinchoninic acid protein assay kit (Pierce Chemical Co., Rockford, IL).

3.14 Statistics

All data are the mean ±1 SE as indicated. Comparisons were made by Student’s t² test for a continuous variable, or by Chi-square (χ²) test for dichotomous variables with a significance level, P = 0.05.

4 Results

4.1 Baseline clinical characteristics in study groups

Body mass index was significantly increased in diabetes having obstructive sleep apnea (39.0 vs 30.3 kg/m², P = 0.011, Table 1) compared to age-matched diabetic patients without obstructive sleep apnea. Baseline atrial fibrillation was also significantly increased in diabetes having OSA vs. no OSA (50% vs 11.8%, P = 0.012, Table 1).

4.2 Depolarization induced by diabetic obstructive sleep apnea autoantibodies

Autoantibodies in diabetic obstructive sleep apnea (10 µg/mL) caused a near doubling in the mean level of N2A depolarization (129.1 vs 115.1 %, P = 0.012, Figure 1) compared to identical concentrations of autoantibodies in age-matched diabetic patients without OSA. These data are the first to suggest that circulating autoantibodies in diabetes having OSA cause increased neuronal depolarization.

4.3 Neurite inhibition by diabetic obstructive sleep apnea autoantibodies

Autoantibodies in diabetes having obstructive sleep apnea (10 µg/mL) caused significantly greater inhibition of neurite outgrowth in N2A cells (33% vs 50 %, P = 0.01, Figure 2A) compared to identical concentrations of autoantibodies in age-matched diabetic patients without OSA. Co-incubation of diabetic OSA autoantibodies (N = 5 patients) with (10µM) concentrations of the G protein- coupled receptor antagonist SCH-202676 significantly prevented (P < 0.001) N2A neurite inhibition from the autoantibodies (Figure 2B). Neurite outgrowth inhibition by diabetic OSA autoantibodies was also significantly prevented (P < 0.001) by co-incubating N2A cells with 10 µM concentrations of the selective Rho kinase inhibitor Y27632 (Figure 2C). Taken together, these results suggest involvement of one or more G-protein coupled receptor(s) and downstream RhoA/ Rho kinase signaling in the mechanism of autoantibody-mediated neurite retraction.

4.4 Long-term storage unmasks inhibitory endothelial cell activity in diabetic OSA protein-A eluates

In prior studies, we reported that peak EC inhibitory activity and neurotoxicity was unmasked following long-term storage (or furin treatment) in diabetic protein-A eluates from patients having a co-morbid neurodegenerative disorder, i.e. glaucoma or dementia [9]. In the present study, storage (9-60 months at 0-4°C) (n = 5) unmasked significant EC inhibitory activity preferentially in diabetic OSA protein-A eluates compared to the stored protein-A eluates of diabetes without OSA (n = 17) (Table 2). Patient subgroup having diabetes plus OSA included significantly more glaucoma suspects (4/5 vs. 4/17; P = 0.04; Fischer’s exact test) than the diabetes subgroup without OSA (not shown in Table 2). Potent EC- inhibitory activity was also unmasked following storage of a metastatic lung cancer protein-A eluate (n = 1), or in the freshly-isolated autoantibodies from breast cancer (n = 4), but not in older adults without cancer or diabetes (n = 3) (Table 2). These data suggest a shared role for specific proteases elaborated in certain cancers (or in OSA) in unmasking latent EC inhibitory autoantibody activity.

4.5 Peak neurite-inhibitory in diabetic OSA autoantibodies: apparent MWs

In prior studies, peak neurotoxicity in cancer fatigue/depression [5] and diabetic depression protein-A-eluates had apparent MWs of the G protein- coupled receptor antagonist SCH-202676 significantly prevented (P < 0.001) N2A neurite inhibition from the autoantibodies (Figure 2B). Neurite outgrowth inhibition by diabetic OSA autoantibodies (N = 6 patients) was also significantly prevented (P < 0.001) by co-incubating N2A cells with 10 µM concentrations of the selective Rho kinase inhibitor Y27632 (Figure 2C). Taken together, these results suggest involvement of one or more G-protein coupled receptor(s) and downstream RhoA/ Rho kinase signaling in the mechanism of autoantibody-mediated neurite retraction.

Figure 1: Diabetic obstructive sleep apnea (OSA) autoantibodies caused significantly greater mean depolarization in N2A neuroblastoma cells compared to control diabetic autoantibodies (No OSA). Results are mean ±/- SE; P = 0.01 for the difference; N = number of participants in each patient subgroup.
Figure 2: Diabetic obstructive sleep apnea (OSA) auto antibodies A) caused significantly greater mean inhibition of neurite outgrowth in N2A neuroblastoma cell compared to control diabetic auto antibodies (No OSA). B) Inhibition of neurite outgrowth by diabetic OSA auto antibodies (N= 5 pts) was completely blocked by co-incubation with (10 µM) SCH-202676. C) Diabetic OSA auto antibodies’ neurite inhibition (N=6) was also significantly antagonized by co-incubated N2A cell with (10 µM) Y27632, a selective Rho kinase inhibitor. Results are mean+/- SE.

corresponding to IgG light chains (23 kD) or half-light chains (11.5 kD) [6]. In the present study, peak neurotoxicity in diabetic OSA/atrial fibrillation protein-A-eluates, (i.e. Pts 1, 2) had apparent MWs of ∼22 kD, and ∼5.5 kD (Figure 3A). An additional neurotoxic peak in a third diabetic OSA/AF patient had apparent MW ∼11 kD (Pt 3 eluate; Figure 3B).

4.6 Diabetic OSA autoantibodies induce Ca2+ release in HL-1 atrial cardiomyocytes

Protein A eluates (1 µg/ mL) from two representative diabetic patients with OSA and either atrial fibrillation (Pt 4) or symptomatic bradycardia requiring permanent pacemaker placement (Pt 5) caused large increases in intracellular Ca2+ (Figure 4A-B) in atrial cardiomyocytes associated with interruption in spontaneous Ca2+ oscillations (Figure 4A-B). Autoantibody-induced intracellular Ca2+ release (in HL-1 cardiomyocytes) was present in a significantly higher proportion of diabetic patients with co-morbid OSA plus atrial fibrillation or another symptomatic cardiac dysrhythmia (6/7 vs. 1/6; P= 0.03; Fischer’s exact test) compared to type 2 DM without OSA, AF or symptomatic cardiac dysrhythmia (not shown in Figure 4). In an age-matched subset of type 2 DM having AF or left ventricular hypertrophy, peak amplitude of Ca2+ release (in HL-1 cells) was significantly correlated with the Cornell voltage-duration product (r = 0.708, P = 0.007)(Figure 4C). The Cornell product is an estimate of the time-voltage integral of the electrocardiographic QRS complex which is useful in the ECG detection of left ventricular hypertrophy [10].

4.7 Possible involvement of MT1-MMP in unmasking neurotoxicity in diabetic OSA autoantibodies

Membrane type 1 matrix metalloproteinase (MT1-MMP) is a broad spectrum proteinase implicated in prostate, breast and lung cancer invasion through its ability to degrade collagen and non-collagenous proteins in the extracellular matrix [11, 12]. MT1-MMP expression also increases in adipogenesis (e.g. morbid obesity) [13], under atrial stretch conditions [14] and via effects of pro-inflammatory cytokines in vascular cells [15]. MT1-MMP has catalytic and pexin domains which bind and unfold triple helical collagen [16] prior to collagenolysis. NSC 405020 is a novel pexin domain-specific MT1-MMP inhibitor [17]. To test for involvement of MT1-MMP in the generation of neurotoxic autoantibody fragments, we incubated diabetic OSA/glaucoma (or/dementia) protein A eluates (Pt 6, 7) in the presence or absence of half-maximal inhibitory concentrations of NSC 405020 followed by gel permeation chromatography. Protein-A- eluates not exposed to NSC 405020 displayed peak neurotoxicity having an apparent MW of ∼5.5 kD (Fig 5A). Protein- A- eluates exposed to NSC405020 displayed a shift in peak neurotoxicity toward a higher MW species (∼43 kd) (Figure 5B).

4.8 Diurnal variation in plasma EC autoantibody growth activity

The 25-75% ammonium sulfate pellet fraction of adult microalbuminuric diabetes plasma includes the IgG fraction and was reported to contain FGF-like, EC growth stimulatory activity [18]. Plasma EC stimulatory activity displayed a diurnal variation: peak activity (e.g. Pt 8, 9) occurred in the morning (0600 – 1000 hours) with much less, if any, stimulatory activity in the evening.

Table 2: Unmasking of EC inhibitory activity in protein A eluates after long-term storage

<table>
<thead>
<tr>
<th>Co morbidity</th>
<th>Before storage</th>
<th>After storage</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diabetes &amp; OSA (n=5)</td>
<td>98 ± 4%</td>
<td>80±15%</td>
<td>0.03</td>
</tr>
<tr>
<td>Diabetes without OSA(n=17)</td>
<td>92 ± 15%</td>
<td>93 ± 26%</td>
<td>0.84</td>
</tr>
<tr>
<td>Diabetes &amp; Lung cancer (n=1)</td>
<td>85 ± 7%</td>
<td>28 ± 10%</td>
<td>0.01</td>
</tr>
<tr>
<td>Breast cancer (n=4)</td>
<td>66 + 15%</td>
<td>NT</td>
<td></td>
</tr>
<tr>
<td>Non-diabetic control (n=3)</td>
<td>102 + 3%</td>
<td>NT</td>
<td></td>
</tr>
</tbody>
</table>

^T-test: comparing activity before and after storage
^ ^P = 0.02: compared to non-diabetic controls

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Increased Neuronal Depolarization Evoked by Autoantibodies in Diabetic Obstructive Sleep Apnea: Role for Inflammatory myocytes which activates MT1-MMP expression in vascular endothelial cells or static stretch in atrial myocytes. They cause cyclic stretch in vascular endothelial cells or static stretch in atrial myocytes, respectively which activates MT1-MMP expression.

2

dicative of a role in altered Ca$^{2+}$ release in atrial cardiomyocytes in-... release [21] or cell depolarization [22]. Our finding of a dose-dependent association between Ca$^{2+}$ release in atrial cardiomyocytes and increased left ventricular mass (QRS voltage-duration) suggests the autoantibodies may be causally-related to cardiac hypertrophy. Diabetic AF/OSA autoantibodies caused quite large intracellular Ca$^{2+}$ release in atrial cardiomyocytes indicative of a role in altered Ca$^{2+}$ signaling underlying clinically-significant cardiac rhythm disturbances.

Hypertension and left ventricular hypertrophy both increase substantially in obstructive sleep apnea [23]. They cause cyclic stretch in vascular endothelial cells [24] or static stretch in atrial myocytes [25], respectively which activates MT1-MMP expression required for extracellular matrix remodeling. Integrin ($\beta$1) responds to mechanical force, e.g. stretch, by activating G12/13 family heterotrimeric G-proteins capable of activating downstream RhoA/ROCK signaling [26]. In a prior report, diabetic EC inhibitory autoantibodies caused loss of endothelial cell adhesion in association with stress fiber activation and RhoA/Rho kinase signaling [27]. Although the receptor is unknown, the autoantibodies may bind to site(s) in the extracellular matrix affecting HSPG-integrin interaction(s) perhaps triggering intracellular signaling events.

Glaucomatous optic atrophy is characterized by remodeling of the optic nerve head associated with increased local expression of MT1-MMP [28]. Our finding of a significant association between latent neurite inhibitory activity in diabetic OSA autoantibodies and glaucomatous cupping (i.e. glaucoma suspect) is novel and perhaps consistent with a reported association between glaucoma and OSA [29]. It suggests that tissue-specific proteolytic remodeling may contribute to optic disc neurite loss in part by unmasking (latent) neurotoxicity in a subset of circulating autoantibodies. Endothelial cell inhibitory activity in the stored or freshly-isolated protein-A eluates of plasma in metastatic lung and breast cancer (Table 2) may be consistent with elaboration of proteolytic activity, (e.g. MT1-MMP or closely-related MMPs) reported in these cancers [11,12].

Peak neurotoxicity having apparent MWs of ~11 kD, ~23 kD and ~43 kD are characteristic of half-light chains, light chains and LC dimers, respectively. Yet more study is needed to determine whether a novel ~5.5kD diabetic OSA autoantibody fragment is derived from specific light chains. A preliminary search

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of database(s) of light chain variable genes revealed that lambda gene family (II or III) members (i.e. 2e2.2 or DPL 23) contain an unusual dibasic K-R amino acid pair at positions 52,53 in complementarity- determining region 2 (CDR2) [30]. DPL23 is a λ3 light chain variable gene which also contains a potential MT1-MMP consensus recognition sequence, PX-G-I [31], at amino acid positions 55-58 immediately adjacent to CDR2. Cleavage near dibasic residues within CDR2 might generate a light chain fragment having the observed MW of 5-6 kD.

DPL23 gene usage was reported in anti-neurotrophic viral glycoprotein antibodies such as anti-rabies [31] and anti- HIV-1 antibodies [30]. The crystallographic structure of a DPL 23-derived light chain component in anti-HIV-1 envelope antibodies was solved by W.D. Tolbert, X. Wu, M. Pazgier and is reported "(4fze-id,)") in the Protein Database, European Pazgier and Molecular Biology Laboratory-EBI, (Welcome Genome Campus, Hinxton, Cambridge shire, UK). The light chain has a secondary structure which is rich in hydrogen-bonded, β turns spanning the CDR2 region, i.e. amino acid residues 48-58 (Protein Database, EMBL-EBI). A postulated function of the pexin domain in MT1-MMP is to disrupt hydrogen- bonding in triple helical collagen causing unfolding which permits access to the MT1-MMP catalytic domain [32]. DPL23 is among a restricted group of κ or λ LC genes reported to encode anti-SS-A/Ro autoantibodies, such as were isolated from salivary gland- infiltrating B cells in primary Sjögren's syndrome [33]. Evidence for antigenic selection in anti-SS-A/Ro autoantibody LCs includes a somatic mutation in the Vk gene L6 which replaces asparagine (N) at position 51 with lysine (K) resulting in a dibasic KR amino acid pair in CDR2, analogous to the dibasic KR pair in 'unmutated' DPL23 gene-encoded LCs. It is of interest that a subset of anti-SS-A/ Ro auto antibodies in lupus are disease-causative of neonatal congenital heart blocks [33]. Evidence for antigenic selection as a driving force in certain pathogenic autoantibody LC fragments comes from "hypermutation" in the DPL 23-derived, anti-HIV-1 envelope antibody LC ("4fze") Among other amino acid replacements, the 4fze LC CDR2 region contains an unusual aspartic acid tandem repeat (DD) aa (positions 50-51) adjacent to the conserved KR dibasic pair. Another λ3 light chain variable gene product harboring an unusually large number of acidic and charged residues within CDR2 was reported as pathogenic in human membranoproliferative glomerulonephritis through its ability to bind and prevent complement binding to factor H [34]. In lupus anti-DNA antibodies, negatively-charged residues, i.e. aspartic acid, present in the κ LC CDR1 or CDR2 regions cause reduced binding to DNA [35] which promotes self-tolerance and escape from clonal deletion. Taken together, these findings suggest that high level of inflammation (in HIV disease, cancer, systemic autoimmunity, morbid obesity) may drive immunity to self-antigens leading to the selection of a restricted subset of λ light chain gene products prone to cleavage (by inflammatory proteases) resulting in formation of a neurotoxic fragment or fragments.

Two-fold increased neuronal depolarization evoked by diabetic OSA autoantibodies is significant. It suggests a novel humoral immune-mediated mechanism of disordered neural regulation of breathing in obstructive sleep apnea affecting older morbidly-obese type 2 diabetes. Our preliminary data showing peak plasma neurotoxicity at nighttime is of interest. One possibility is enhanced release of a protease which has a normal nocturnal pattern of expression. One such candidate is MMP2, activated by MT1-MMP, which showed increased expression at nighttime in orchestrating physiologic desquamation of surface corneal epithe-
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8 References


