Disorders of Sleep and Aggression Trait in Women Associated: What Underlies Staying “Woke”?

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Received: 09 Apr, 2019 | Accepted: 06 May, 2019 | Published: 13 May, 2019


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Abstract

Narcolepsy is a sleep disorder where the patient falls asleep unwillingly. It is thought to be related to hyper functioning central sleep centers in the brain. Sleep apnea is a disorder of breathing disruption during sleep. Genes of the dopamine system have been implicated with high dopamine: norepinephrine ratio. Since dopamine has also been associated with personality traits, the hypothesis we studied herein was that patients with narcolepsy and sleep apnea would score low in catecholamine settings causing aggression trait. We found that narcolepsy and sleep apnea diagnoses showed significantly lower aggression trait using an online test. The conclusion is that narcolepsy and sleep apnea patients are not aggressive in personality, and since aggressiveness is related to sympathetic nervous system activity, this would be predictable given the role of sympathetic nervous system in wakefulness.

This result sheds light on the connection of sleep and wakefulness and the autonomic nervous system as well as the central nervous system. It also enlightens on personality as a window to physiology in health and in illness.

Introduction

Personality is largely genetic with catecholamine genes showing associations with many traits [1-20], including dopamine receptors both D2 like receptors as well as D1 like receptors. Dopamine metabolizing enzymes such as monoamine oxidase type A as well as monoamine oxidase type B, catechol-O-methyltransferase has been found associated with many personality traits. Personality trait systems include the Cloninger traits of Novelty Seeking, Reward Dependence, Risk Avoidance and Persistence, and the Big Five traits of Openness to Experience, Conscientiousness, Extraversion, Agreeableness, and Neuroticism all of which have shown many genetic correlates. Still a consensus has not been reached about personality traits and genes just as little consensus exists about correlations between disease and personality traits.

A physiologic and genetic system showing promise of insights into health is one called NPA Personality Theory, a genetic theory based on sympathetic and parasympathetic nervous system correlates as causative of personality. Aggression as a correlate of sympathetic nervous system is one of three traits that are widespread. Narcissism correlates with the parasympathetic nervous system. Perfectionism affects the autonomic nervous system and thus modulates both Aggression and Narcissism in some individuals. Every individual putatively has one or more of these traits. Some individuals have modifiers of each trait some of which are associated with discernible unusual behavior patterns. A personality test as well as full explanation of the theory for these traits is available online [21-24].

Sleep is related to the central nervous system with dopamine as a prime factor in regulation of sleep and wakefulness. High dopamine activity seems related to sleep states, and high norepinephrine activity seems related to wakefulness [25-32]. Lower availability of DRD3 receptors was associated with daytime somnolence in Parkinson disease patients suggesting that high dopamine neurotransmission caused somnolence through activation at certain dopamine receptors [31]. Rodent studies have demonstrated also dopamine receptor agonists both centrally and peripherally [29,30] as productive of somnolence [26,28,32].

Since these neurotransmitters are also related to personality traits, there may be clinical correlation with personality traits and sleep patterns. Many genes and illnesses have shown correlation with various sleep related diagnoses. Indeed, if sleep has physiologic correlates with the autonomic nervous system, a personality survey based on autonomic nervous system function would be correlated to sleep diagnoses, and specifically lower aggression trait as a correlate of sympathetic nervous system with higher norepinephrine and epinephrine balance would be related to sleep apnea or narcolepsy incidence. Since sleep and aggression are correlated with sympathetic tone in opposite directions our hypothesis was that patients with diagnosis of sleep disorders such as sleep apnea and narcolepsy would show low prevalence of aggression trait.

Method

With a view of conducting a pilot study on personality and
Diagnoses, patients seen in the office over a year's time of one obstetrician-gynecologist were given opportunity to take the online NPA personality test. At the time of their office visit, all diagnoses as well as demographic factors and physician assessment of patient status as to these personality traits were recorded on an Excel spreadsheet with over 250 queries regarding a broad range of established diagnoses in general health as well as gynecologic health available for selection.

This study was not submitted to Institutional Review Board as the patient data was de-identified for the study and dealt with standard clinical data collected in taking history in an ob-gyn office.

After the data collection phase was completed, patients with diagnosis of narcolepsy or sleep apnea who had taken the online personality tests were to be reviewed for their results as to whether they had shown Aggression trait on their tests.

Patients were offered the online NPA personality test as a part of their office visit with the invitation stating that a study was being done to explore the relationships between personality and diagnoses. An Excel spread sheet was configured to list all diagnoses of each patient and was logged by the physician at the time of each patient's office visit.

Statistical analysis planned was chi square for difference between study group and entire population. The online test center returned details were published previously and detailed several diagnoses (275 of 1275 patients) differed significantly on personality diagnosis (688 of 1275 patients) and those with any psychiatric diagnosis (275 of 1275 patients) differed significantly on personality test to the physician who recorded the result and related the result to the patient with an invitation to answer any questions about the result. After the year of testing was done, correlations were to done between diagnoses and personality results in the entire group. Based on literature review on personality and illness, the expectation was that there would be correlations between low aggression trait and illness in a wide variety of types in our patient population of women.

**Results**

There were 4028 patients seen in the office during the study period of a year. 1235 patients took the online personality test. In looking at all patients, 50% showed aggression trait. Age showed minimal effect of lessening aggression trait. Eight patients had diagnosis of narcolepsy of them one took the online personality test while 18 patients had diagnosis of sleep apnea of whom ten took the online test. Of those patients with diagnosis of sleep apnea who took the test none had results showing aggression trait though of those ten patients one patient had ambiguous results on online test while the office assessment was non-aggressive personality. Because all study patients showed the expected result of not having aggression trait, there was no need for chi square test to compare study group with entire population 50 percent of whom had aggression trait result on the online test.

Results of this pilot study unrelated to sleep apnea or narcolepsy are described in previous article, but in summary, those with any medical diagnosis (688 of 1275 patients) and those with any psychiatric diagnosis (275 of 1275 patients) differed significantly on personality test from others. Details were published previously and detailed several differences in groups as well as exploring likely genetic correlates as causes of these differences.

This present paper explores possible meanings of score differences in a subset of patients in the above pilot study; patients with disorders of wakefulness.

**Discussion and Conclusion**

Sleep disorder may have implications for such diverse pathologies as sudden death in general, sudden infant death syndrome, fetal demise, neonatal respiratory distress, automobile and other transportation-related accidents, dementia, Alzheimer's disease and end-of-life process and many others[33-84]. The central commonality in these diverse pathologies could be catecholamine and other neurotransmitter processes such as the genetically controlled ratio of dopamine and norepinephrine activity emanating from the locus coeruleus in the midbrain and the processes of the sleep center in the ventral tegmental area [32,47]. Dopamine beta hydroxylase alleles which provide dopamine quantitative dominance fuels all downstream dopamine dominance, a situation producing more cognitive activity as opposed to more musculoskeletal action. In fact, somnolence as an extreme is found in individuals who evidence less trend to musculoskeletal activity, true to the model. Another example of an effect of this dopamine: norepinephrine dichotomy is the attention deficit hyperactivity diagnosis to include the carelessness allowing accidental injury and impulsive misbehaviors, where a shift toward norepinephrine as opposed to dopamine has presumably happened upstream. So pathologies of both extremes in the dopamine: norepinephrine ratio exists though more pathologies do appear to be related to the high dopamine: norepinephrine ratio. Further examples can be noted in the case of sudden death in adults, fetuses or children where high dopamine balance appears to be operative. In the example of Alzheimer's disease, other dementias and end of life events for the extremely elderly, the loss of norepinephrine seems at the heart of the sleep that overcomes wakefulness eventuating in the cessation of breathing and the end of life.

Just as genetic causes of illness are becoming better understood so are personality genetics becoming better elucidated. And they may be correlated [66,77].

Our study did not address genetic polymorphisms of patients to correlate with personality and illness but instead studied personality as a proxy for genetic polymorphism. Since 50% of personality is genetic, this approach has some validity though with some limitations given both the shared and non-shared environmental correlates of personality. Added to that limitation is the lack of consistencies of genetic studies of personality as many studies fail to yield replication in subsequent studies. Similarly, many studies of genetic causes of illness find a complex interweaving of genetic and environmental causes that seems to approximate the 50/50 division of nature vs nurture found in personality causation. So with some strong limitations in this type of endeavor, this pilot study was planned and carried out and yielded the expected outcomes in illness and personality associations and relationships in general and with many specific topics such as the one in focus in this paper, that of sleep apnea and narcolepsy.

A guiding hypothesis seems to be that neurons in the sleep centers are affected by genes of the dopamine system such that high dopamine activity promotes sleep and high norepinephrine activity promotes wakefulness. And in personality genetics higher dopamine seems to promote non-aggressiveness and higher norepinephrine activity seems to promote aggressiveness. This scheme seems to play out in the central nervous system as well as in the autonomic nervous system with the principle that sympathetic nervous system seems more active in fight or flight aggressive actions while parasympathetic nervous system seems more active in resting functions such as gastrointestinal functions.

Because of the central nervous system and the peripheral nervous system control of all physiology, disorders associated with sleep disorders also include schizophrenia and autism. Dopamine has been found central to pathophysiology of both schizophrenia and autism, and the dopamine: norepinephrine circuitry especially the locus coeruleus as the prime site of action of dopamine beta hydroxylase, the enzyme determining the dopamine: norepinephrine ratio is...
thought central [33]. Dopamine Beta Hydroxylase gene (DBH) has linkage with other genes such as the ABO blood group (ABO), and ABO blood group maternal-fetal incompatibility as been found to be associated with lower hippocampal volume in schizophrenia [33,34]. Schizophrenia has been noted through basic physiologic variants to be associated with a wide range of health issues starting with cardiovascular and renal illness and including metabolic issues such as osteoporosis [55]. Vitamin D deficiency was found in a Dutch study to be 4.7 times more common among those with bipolar disorder, schizophrenia or schizoaffective disorder [58]. Osteoporosis may be related through Vitamin D genetics having as it does strong interactions with midbrain dopamine: norepinephrine ratio. Thus osteoporosis is associated with schizophrenia and other psychoses [44-45,56-60,64,74,75]. And similarly autism has been found to have associations with osteoporosis as well [62-64]. Environmental causes of osteoporosis have not been found to explain this, and the finding holds for all ethnicities and genders studied. So instead of being related to environmental or socioeconomic variation, these linkages of such diverse pathologies as osteoporosis and schizophrenia would logically have a major commonality genetically.

These associations have been thought to be a manifestation of evolutionary concepts demonstrating the concept of tradeoffs [61] that is, illnesses occurring in clusters have continued genetically because of some tradeoff advantages that are linked to the illnesses by virtue of their common biochemistry and in fact brain chemistry. Thus an evolutionarily wrought larger volume in cerebral brain centers allows more neurotransmission with some accompanying pathologic chemistries producing collateral damage. The relatively rapid attainment of larger neo-cortex would logically carry deficiencies since the ancestral organism had survived untold generations without the larger neo-cortex, and given that mutations are statistically speaking detrimental for the organism, a bad outcome would be assumed. In the case of the larger neo-cortex and accompanying increase in neurotransmission, damages include that of sleep apnea, narcolepsy, osteoporosis, and putatively such far-reaching illnesses as schizophrenia, autism, Alzheimer’s disease, sudden death in and fetuses and infants, sudden death, and various anomalies.

The allele for dopamine beta hydroxylase that produces high dopamine: norepinephrine ratio is not the ancestral allele and appears to be on the ascendency in human populations. So since any evolutionary event has to be explained by changes in the production of progeny that survive to produce progeny, the issue of high dopamine: norepinephrine ratio producing individuals with seemingly more pathologies may demonstrate that fitness has its drawbacks or tradeoffs that have to be faced. And downstream dopamine related genes seem to fine tune such that the benefits of high dopamine neurotransmission can be retained while the risks ameliorated. One example of that would be the metabolizing enzyme COMT appearing to be manifesting an evolutionary path where the lower metabolizer is not the ancestral state but is instead the evolutionary winner in human populations. Additionally, dopamine receptors with polymorphisms producing less activity make up for the profound suppression of adenylyl cyclase caused by the D2 type receptor activation. So understanding one neurotransmitter role is fraught with the complexity of checks and balances upstream and downstream. The production of surviving progeny would appear to always be the goal of evolution, and even a very high burden of disease from a genetic background that allows surviving progeny may survive even with high disease burden. Such may be the case of the high dopamine: norepinephrine ratio in the midbrain.

Progeny that survive may live shorter lives, but reproduction is the prime feature. Diseases such as schizophrenia and autism shorter lives and decrease progeny. Why does schizophrenia continue to exist? Why is autism in fact on the rise? The answer seems to be in the trade-offs whereby these pathologies are hard-wired by the large neo-cortex that was selected in early man. A set of environmental challenges existed. Man’s life was short and brutal, and a larger neo-cortex with higher dopamine: norepinephrine ratio for memory and analysis of environment survived.

The fetal pathologies have their genesis in the earliest of embryonic events. And it is this timing of the creation of pathologies that pinpoints brain chemical and anatomic embryology as the cause, specifically the roots of the autonomic nervous system in the midbrain and medulla oblongata. The formation of the neural tube is the first discernible event in embryology and therefore the genesis of all body forming events that follow. So downstream phenomenon reflect early embryonal events.

Sudden infant death syndrome of unknown cause as well as fetal demise has been thought to result from central nervous system midbrain malfunction [49,50,52]. Central sleep centers through a high dopamine: norepinephrine ratio connects third trimester fetal demise with sudden infant death syndrome [50,84]. A prime genetic cause of this is low activity Dopamine Beta Hydroxylase (DBH) as it codes for an enzyme that determines that ratio [48,53]. And because of linkage disequilibrium of DBH with ABO blood group gene [83], correlation is found not just with previously mentioned pathologies such as schizophrenia, autism and osteoporosis but also with ABO and sudden infant death and fetal demise [34-36,38-40,42-44] as well as with ABO gene and fetal anomalies such as congenital heart disease [35,36] and spina bifida [38] as well as cleft lip and cleft palate [42-44]. ABO gene has also been associated with osteoporosis [38-40]. Most studies have trended toward finding that non-O ABO groups are more likely to be affected when maternal ABO O is involved, this related to the higher antibody formation of ABO O maternal vs ABO non-O fetuses. And those fetuses with dopamine predominance have more suppression effects as they have more dopamine neurotransmission at the D2 like receptors. This may explain the lack of neuro-resilience eventuating in more pathology at all fetal stages, certainly supporting miscarriage and fetal anomalies as well as more schizophrenia and autism in surviving fetuses.

Because personality trait testing is genetically associated with brain factors, this can be used as a clinical inroad into behavior and broader medical factors [33-83].

Our finding that sleep disorders of sleep apnea and narcolepsy are associated with lack of aggressiveness in personality is consistent with this theoretical framework used to explore the complex function of sleep and wakefulness.

The study was extremely limited as only one patient of the 8 patients with diagnosis of narcolepsy took the online test and only 10 of 18 patients with diagnosis of sleep apnea took the online test. Furthermore, the test used to test for aggression trait in personality is invalidated. Aggression trait is defined many different ways by different researchers and has subcategories such as stage aggression, reactive aggression, and many others. In some studies it is related to such concepts as impulsivity, hyperactivity, and other many traits. The NPA system used in our study aimed to elicit fight or flight traits built on traditional theories about the sympathetic nervous system and behavior.
This study was a pilot study to generate hypotheses, and these hypotheses will need much further study before any conclusions can be reached on such topics. Results of analysis of this pilot study suggested significant difference in personality in patients with diagnoses versus those without diagnoses both for medical diagnoses and for psychiatric diagnoses [80]. Our study showed a trend toward low aggression trait consistent with research supporting association of several diagnoses with each other and with causative catecholamine activities in osteoporosis, in third trimester fetal demise, fetal and neonatal disease/anomalies [35-37,41-43,49,52,56], schizophrenia [34,35,46-48,51,54-61,79] and autism [33,51,62,63] as well as Vitamin D deficiency with ABO genes [45,46,77], Alzheimer’s [76] bone fracture [62], and autoimmune diseases [61].

Schizophrenia studies are extensive given the profound morbidity of that individuals, families and societies accrue related to these individuals. Some authorities have thought of autism as a childhood form of schizophrenia, just to highlight the profound disturbances noted. But schizophrenia typically has on set in late puberty. Medical advances are significant in ability to control symptoms be they negative (lack of motivation) or positive (distortions of reality) and were heralded by the finding that antihistaminic preparations offered some control of symptoms. This led to development of dopamine blocking medications. The lack of specificity in blockade has made these treatments problematic. So a clear central cause of schizophrenia and autism is most to be sought. Starting at the beginning of the organism, in embryology seems logical. The development, antenatal, viral or other causes can be placed in their right perspective.

And in the case of higher aggression trait, this genetic set related to dopamine: norepinephrine ratio includes ADD or ADHD [65] and Parkinson’s disease [46] as well as some types of autoimmune diseases [61], and alertness deficits with resultant accidental injury [62].

This group of diagnoses may share several genetic causes from pleiotropy so that the study of these issues as a genetic anlage has promise to be rewarding [61].

References


