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Abstract

Introduction: Cranial electrotherapy stimulation (CES) is a noninvasive therapy that has been used for decades in the United States to treat anxiety, depression, and insomnia in the general population. The effectiveness of CES has been questioned by many and its use is considered controversial. In this study we are presenting data on one alcoholic patient using a newly engineered device we call Neuro-Electro-Adaptive Therapy 12™ [NEAT12]. This hybrid device utilizes TENS current characteristics yielding CES effects. This device has been found to primarily target the excitation of the Cingulate Gyrus region of the brain.

Case presentation: This is a 42 year old male who has been abstinent from alcohol for approximately two months. The data presented herein represents the pre to post qEEG differences of an alcoholic in protracted abstinence. This subject was evaluated both before and after using the NEAT-12 device. The pre to post comparisons suggest that the cortical potentials especially at the Cingulate Gyrus are up regulated after using the device. The absolute power changes obtained shows a decrease of more than 2 SD as noted in the delta wave spectrum. Also noted is an overall cortical increase in the alpha spectrum. The resting alert state of a neurotypical population is most prominently marked by a regulation of 7.5-11 Hz alpha throughout the cortex. The decreased in delta and theta suggests an up regulation of the prefrontal cortex and the anterior Cingulate Gyrus a site involved in substance use disorder (SUD).

Conclusion: A presence of dominant slow waves through the prefrontal cortex and the anterior Cingulate Gyrus is often associated with OCD, anxiety, impulsivity and cravings in addicted populations. It is conceivable that our initial finding of altered electrical activity of the brain using qEEG analysis suggests the NEAT-12 may induce a “normalization” of aberrant electrical activity of the cortical region of the brain known to occur during protracted abstinence of alcoholics. It may have utility as a putative anti-craving CES device and therefore warrants intensive investigation.

Introduction

The Cranial Electrical Stimulation (CES) technique appeared at the beginning of the 1960s and is aimed to act at the level of the central nervous system [1]. The current, composed of high frequency pulses interrupted with a repetitive low frequency, was delivered through three electrodes (a negative electrode placed between the eyebrows while two positive electrodes are located in the retro-mastoid region). We have recently introduced a new CES Device, the NEAT-12, in which shortcomings encountered with previous electrical stimulation techniques are avoided due to changes in the characteristics of the delivered current. The main property of CES is to potentiate some drug effects, especially opiates and neuroleptics, during anesthetic clinical procedures [2]. This potentiation effect permits a drastic reduction of pharmacological anesthetic agent and results in reduced post-operative complications. Despite numerous clinical and animal studies performed with this technique for several decades, CES mechanisms are not completely elucidated. Animal studies demonstrated that stimulation with CES releases 5-hydroxy-indol-acetic acid and enkephalins [2]. These results obtained without any undesirable outcome are encouraging signs. Continued investigation of this electrotherapeutic technique is warranted.

Brief History

CES received U.S. Food and Drug Administration (FDA) approval for the treatment of insomnia, depression, and anxiety in 1979 [3]. CES is the United States FDA term for the transcranial application of small amounts of electricity, usually less than 300-600 mA with a frequency of 100 Hz or lower. CES was imported into the United States from Europe and was originally introduced as electro sleep, possibly because it increases delta waves. CES and electronic medicine did not receive a particularly warm reception in American medicine until clinicians began to utilize the transcutanous electrical nerve stimulation (TENS) devices for pain. Electrotherapies have been used in psychiatry in the form of electroconvulsive therapy (ECT), which is still utilized in organic brain diseases such as Parkinson’s and organic-based depressions [4].

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Scrutiny of Table 1 indicates that abundant research has established that an electroencephalogram (EEG) recorded from a drug abuser has a predictable distributional electrical power (measured in microvolts squared), just as does the electrocardiogram (EKG). The predictable electrical signals recorded by the EEG, distinctive for each brain region, are regulated by the homeostasis of a complex neuroanatomical brain system that utilizes all known neurotransmitters. Just as the EKG can be used to assess heart dysfunctions, the EEG can assess a wide variety of brain dysfunctions related to developmental, neurological and psychiatric disorders, whether caused by structural or functional abnormalities.

Electrophysiological imbalances secondary to drug abuse have been well characterized (Table 1). These EEG abnormalities are similar in characteristic to the electrophysiological imbalances of various

<table>
<thead>
<tr>
<th>Drug of Abuse</th>
<th>Measure</th>
<th>Outcome</th>
<th>Comment</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alcohol</td>
<td>qEEG and LORETA mapping</td>
<td>Increase in absolute and relative beta power and a decrease in alpha and delta/theta power.</td>
<td>Detoxified patients compared to normal controls</td>
<td>Saletu et al. [5]</td>
</tr>
<tr>
<td>Alcohol</td>
<td>EEG</td>
<td>Subjects with family history have reduced relative and absolute alpha power in occipital and frontal regions and increased relative beta in both regions.</td>
<td>Family history of alcoholism compared no family history</td>
<td>Finn and Justus [6]</td>
</tr>
<tr>
<td>Alcohol</td>
<td>EEG</td>
<td>Alcoholics differ in resting EEG coherence having lower frontal alpha and slow –beta coherence in males and females.</td>
<td>Heavy drinkers compared to light drinkers</td>
<td>Kaplan et al. [7]</td>
</tr>
<tr>
<td>Alcohol</td>
<td>EEG</td>
<td>In alcohol-dependent subjects found higher central alpha and slow –beta coherence, but lower parietal alpha and slow –beta coherence in males.</td>
<td>Alcohol-dependence compared to controls</td>
<td>Michael et al. [8]</td>
</tr>
<tr>
<td>Alcohol</td>
<td>EEG</td>
<td>Higher left-temporal alpha and slow beta coherence and higher slow-beta coherence at right –temporal and frontal electrode pairs in alcoholic males and females.</td>
<td>Alcohol-dependence compared to controls</td>
<td>Winterer et al. [9]</td>
</tr>
<tr>
<td>Alcohol</td>
<td>EEG</td>
<td>Moderate to heavy drinking is associated with differences in synchronization of brain activity during rest and mental rehearsal. Heavy drinkers displayed a loss of hemispheric asymmetry of EEG synchronization in the alpha and slow –beta band. Moderately and heavy drinking males also showed lower fast-beta band synchronization.</td>
<td>Comparison of moderate – heavy to Heavy drinking</td>
<td>De Bruin et al. [10]</td>
</tr>
<tr>
<td>Marijuana</td>
<td>EEE</td>
<td>Acute THC exposure produced transient increases in either posterior alpha power, decreases in mean alpha frequency or increase in alpha synchrony and decrease in relative power of beta.</td>
<td>Acute effects of THC</td>
<td>Struve et al. [11]</td>
</tr>
<tr>
<td>Marijuana</td>
<td>qEEG</td>
<td>Significant association between chronic marijuana use and topographic qEEG patterns of persistent &quot;alpha hyperfrontality&quot; as well as reductions in alpha mean frequency. There was also elevated voltage of all non-alpha bands in chronic marijuana users. Finally there was a widespread decrease in the relative power of delta and beta activity over the frontal cortical regions in chronic marijuana users.</td>
<td>Chronic effects of THC exposure.</td>
<td>Struve et al. [12]</td>
</tr>
<tr>
<td>Heroin</td>
<td>qEEG</td>
<td>Qualitative changes were observed in more than 70% of heroin addicts in early abstinence and included low-voltage background activity with diminution of alpha rhythm, an increase in beta activity, and a large amount of low amplitude delta and theta waves in central regions. Also frequency shifts in fast alpha range at the frontal and central recording sites and a slowing of slow wave alpha mean frequency at the central, temporal, and occipital sites of recording heroin abusers who used heroin for at least 18 months.</td>
<td>Acute withdrawal</td>
<td>Polunia and Davydov et al. [13]</td>
</tr>
<tr>
<td>Heroin</td>
<td>qEEG</td>
<td>Abstinent alcoholics have an enhanced fast beta power compared to healthy controls.</td>
<td>Alcoholics compared to healthy controls</td>
<td>Franken et al. [14]</td>
</tr>
<tr>
<td>Heroin</td>
<td>EEE</td>
<td>Elevated synchrony within beta frequency during short term heroin withdrawal may reflect a state of CNS activation toward reward–seeking behavior, with this being a prerequisite to relapse among drug dependent patients.</td>
<td>Polydrug abusers with emphasis on heroin abuse.</td>
<td>Bauer et al. [15]</td>
</tr>
<tr>
<td>Cocaine</td>
<td>EGG</td>
<td>Acute effects of cocaine include increase in beta activity, increase in delta, increase in frontal alpha as well as an increase in alpha wave EEG associated with bursts of cocaine–induce euphoria.</td>
<td>Human studies</td>
<td>Prichep et al. [16]</td>
</tr>
<tr>
<td>Cocaine</td>
<td>qEEG</td>
<td>During protracted abstinence from cocaine qEEG effects include long-lasting increases in alpha and beta bands together with reduced activity in delta and theta bands.</td>
<td>Several studies reported similar effects on withdrawal.</td>
<td>Roemer et al. [18]</td>
</tr>
<tr>
<td>Cocaine</td>
<td>qEEG</td>
<td>Cocaine produced a rapid increase in absolute theta, alpha and beta power over the prefrontal cortex, up to 25 minutes after drug administration. The increase in theta power was correlated with a positive drug high, and the increase in alpha power was correlated with anxiety. Also an increase in delta coherence over the prefrontal cortex correlated with nervous energy.</td>
<td>qEEG profiles in cocaine-dependent patients in response to an acute, single-blind, self–administered dose of smoked cocaine base (50 mg) versus placebo.</td>
<td>Reid et al. [19]</td>
</tr>
<tr>
<td>Cocaine</td>
<td>qEEG</td>
<td>Changes occur 5-14 days after last reported crack cocaine use induced changes in brain function. These changes lasted up to six months.</td>
<td>Subjects with cocaine dependence have persistent changes in brain function</td>
<td>Vennemann et al. [20]</td>
</tr>
<tr>
<td>Cocaine</td>
<td>qEEG</td>
<td>qEEG techniques demonstrate an association between beta activity in the spontaneous EEG and relapse in cocaine abuse.</td>
<td>qEEG changes associated with relapse</td>
<td>Ceballos et al. [21]</td>
</tr>
</tbody>
</table>

Source: Miller et al. Post Graduate Medicine (in press) with permission.

Table 1: qEEG, EEG changes in Substance Abuse.
psychiatric diseases such as generalized anxiety, ADHD, depression, OCD and other impulsive disorders.

Abnormal behaviors involving dopaminergic gene polymorphisms often reflect an insufficiency of usual feelings of satisfaction, or Reward Deficiency Syndrome (RDS). RDS results from a dysfunction in the “brain reward cascade,” a complex interaction among neurotransmitters (primarily dopaminergic and opioidergic). Individuals with a family history of alcoholism or other addictions may be born with a deficiency in the ability to produce or use these neurotransmitters.

A number of earlier studies have examined the effect of CES on pain, headaches, fibromyalgia, smoking cessation, closed head injuries, and opiate withdrawal [22-24]. There have been a number of studies including some randomized double-blind, controlled experiments [25-31] on generalized anxiety and related symptoms using CES.

Substance use Disorder and CES

The use of CES as a potential modality for drug abuse was reviewed many years ago by Blum’s laboratory [32]. At that time we showed that in high risk drug abusers the P300 amplitude increased significantly (n=14, P<0.03) when the pre and post qEEGs were compared. The CES device at a frequency of 100 pulses/sec, 1.0 mA, 20% duty cycle, in a square wave form, was worn on the forehead and wrist for 40 minutes. Moreover, fifteen drug abusing individuals were evaluated in an eyes closed and eyes open computerized power spectral analysis for delta, theta, alpha, beta activity before and after 40 minutes of CES. Significant electrophysiological changes occurred in all of the subjects with abnormal baselines. When the pre and post-CES brain electrical activity mapping were compared, significant increases in delta, alpha and beta activity occurred, with a significant decrease in theta activity. We suggested that CES is a therapy that may beneficially alter the abnormal electrophysiology associated with drug abuse and it may normalize patterns of electrotherapy. Others have also studied the effects of CES and drug abuse [33-36]. Specifically, in a double blind placebo controlled investigation [35] it was shown that CES, comprising of the combination of a constant current with a pulse square of square impulses of 70-80 Hz, is an effective method to correct affective disturbances (anxiety, depression) in alcoholic patients. REF Krupitsky et al. [35] in Russia found that the medical effects of CES are accompanied by changes in the metabolism of GABA and monoamines, but not of beta-endorphin. These changes were accompanied by a decrease in the latency of alpha-rhythm appearance eyes closed qEEG.

While there have been a number of studies some of which are negative on the use of CES for the treatment of anxiety, insomnia and depression less is known about the use of CES in substance use disorder [37-39]. In fact there have been no studies related to Substance Use Disorder (SUD) in humans that we can find by doing a PubMed search (7-21-10) after 2000.

Method

Subject

The male was 42 years old with a two month period of sobriety. The subject was recruited from G & G Holistic Addition Center alumni group. He was diagnosed using DSM –IV criteria for SUD with emphasis on alcoholism. He was currently employed but complained about anxiety and a continuation of craving, “white knuckle” sobriety.

Study design

The study was approved by the PATH Foundation NY Institutional Review Board as part of an approved research project related to genetic testing and novel approaches to SUD treatment (RB NIH registration is # IRB00002334). The patient signed a consent statement prior to the experiment. The patient was not taking any psychoactive medication at the time of the evaluation. He was drug free as assessed by urine testing. A qEEG analysis obtained one-day prior to NEAT-12 treatments was considered as the pretest. At the exact same time only one day later the patient utilized the NEAT 12 for a one-hour period. Immediately after this treatment the subject was re-evaluated by qEEG analysis.

The NEAT 12 has twelve independent programs associated to patient’s condition whether it is acute, sub-acute or chronic. The program used in this case study was for post chronic conditions. The current characteristics were 25 uA, at a frequency of 0.4 Hz, polarity change 2.25 seconds every 17 pulses with a modulated pulse. The electric parameters for the NEAT-12 device are shown in Table 1 and Figure 1.

The memory of the NEAT-12 device stores data for 12 different therapeutic waveforms and pulse sequences, each of which is described in Table 2.

In a current search of the existing devices approved for market by the FDA for Cranial Electrical Stimulation no device was found to have any similar waveform characteristics. Many claimed to have waveform harmonics which are trade secrets but from our research the duration, polarity and modulation of our waveform has very little resemblance of any other device on the market today.

qEEG procedure

Nineteen electrodes using an electro-cap consistent with the International 10/20 systems were placed. Routine EEG was recorded on a Cadwell Easy II using a linked ear montage and with electrodes.

Figure 1: Description of NEAT Device and treatment protocol
digitally referenced to the Cz electrode allowing for retrospective montage analysis of all data. Using data gathered under technical conditions as listed above, 59.99 seconds of EEG were selected and subjected to quantitative analysis of absolute power, relative power, power asymmetry and coherence. These measurements are logarithmically transformed and referenced to age-adjusted population norms.

## Results

Frequency band magnitude (uV) topographies of the EC condition are presented below (Figure 2).

The absolute power changes represented in the above images shows a decrease of more than 2 SD as noted in the delta wave spectrum. Also noted is an overall cortical increase in the alpha spectrum. The resting alert state of a neuro typical population is most prominently marked by a regulation of 7.5-11 Hz alpha throughout the cortex. The decrease in delta and theta suggests an up regulation of the idling frequencies of the prefrontal cortex and the anterior Cingulate Gyrus. A presence of dominant slow waves through the prefrontal cortex and the anterior Cingulate Gyrus is often associated with OCD, Anxiety, and impulsivity.

To validate these findings a percentage change in microvolts squared was performed as presented by the below data (Figure 3). The percentage of relative power change is congruent with the previous findings noting a 25% decrease in prefrontal delta and an increase

<table>
<thead>
<tr>
<th>Program</th>
<th>Name</th>
<th>MicroAmperage</th>
<th>Frequency</th>
<th>*Pulse Width</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Low Freq Steady</td>
<td>25</td>
<td>0.44 Hz</td>
<td>1.12s</td>
<td>This program is for those individuals that have a simple diagnosis of stress, anxiety or insomnia in aftercare treatment.</td>
</tr>
<tr>
<td>2</td>
<td>Med Freq Steady</td>
<td>25</td>
<td>2.5 Hz</td>
<td>0.02s</td>
<td>This program is for those individuals that have a moderate diagnosis of stress, anxiety or insomnia in aftercare treatment.</td>
</tr>
<tr>
<td>3</td>
<td>High Freq Steady</td>
<td>25</td>
<td>7 Hz</td>
<td>0.714s</td>
<td>This program is for those individuals that have a severe diagnosis of stress, anxiety or insomnia in aftercare treatment.</td>
</tr>
<tr>
<td>4</td>
<td>Low Freq modulated</td>
<td>25</td>
<td>0.44 Hz</td>
<td>High 1.85s</td>
<td>This program is for those individuals that have a simple diagnosis of stress, anxiety or drug addiction in aftercare treatment.</td>
</tr>
<tr>
<td>5</td>
<td>Med Freq modulated</td>
<td>25</td>
<td>2.5 Hz</td>
<td>High 0.32s</td>
<td>This program is for those individuals that have a moderate diagnosis of stress, anxiety or drug addiction in aftercare treatment.</td>
</tr>
<tr>
<td>6</td>
<td>High Freq modulated</td>
<td>25</td>
<td>7 Hz</td>
<td>High 0.114s</td>
<td>This program is for those individuals that have a severe diagnosis of stress, anxiety or drug addiction in aftercare treatment.</td>
</tr>
<tr>
<td>7</td>
<td>Low Freq Steady</td>
<td>100</td>
<td>0.44 Hz</td>
<td>1.12s</td>
<td>This program is for those individuals that have a simple diagnosis of stress, anxiety or insomnia that are in acute treatment.</td>
</tr>
<tr>
<td>8</td>
<td>Med Freq Steady</td>
<td>100</td>
<td>2.5 Hz</td>
<td>0.02s</td>
<td>This program is for those individuals that have a moderate diagnosis of stress, anxiety or insomnia that are in acute treatment.</td>
</tr>
<tr>
<td>9</td>
<td>High Freq Steady</td>
<td>100</td>
<td>7 Hz</td>
<td>0.714s</td>
<td>This program is for those individuals that have a severe diagnosis of stress, anxiety or insomnia that are in acute treatment.</td>
</tr>
<tr>
<td>10</td>
<td>Low Freq modulated</td>
<td>100</td>
<td>0.44 Hz</td>
<td>High 1.85s</td>
<td>This program is for those individuals that have a simple diagnosis of stress, anxiety or drug addiction that are in acute treatment.</td>
</tr>
<tr>
<td>11</td>
<td>Med Freq modulated</td>
<td>100</td>
<td>2.5 Hz</td>
<td>High 0.32s</td>
<td>This program is for those individuals that have a moderate diagnosis of stress, anxiety or drug addiction that are in acute treatment.</td>
</tr>
<tr>
<td>12</td>
<td>High Freq modulated</td>
<td>100</td>
<td>7 Hz</td>
<td>High 0.114s</td>
<td>This program is for those individuals that have a severe diagnosis of stress, anxiety or drug addiction that are in acute treatment.</td>
</tr>
</tbody>
</table>

*The measurement of the pulse width reported is without a load. Also, in a non-modulated program the pulse width measurement of individual pulses, the pulse duration range is 1 microsecond to 1000 microseconds with a 1000 ohm load.

**Table 2: NEAT-12 programs.**

![Maps of EEG power spectra for band ranges](image_url)
in cortical potential marked by the prefrontal increase in activity throughout the theta and alpha spectrum. The overall alpha increase of 15-25% suggests an up regulation in post synaptic potentials noting that the brain has a more power to self-regulate.

Discussion

In comparison of pre to post CES treatment the aforementioned absolute power ranges in these findings support the use of the CES technology in ameliorating the electrophysiological correlates of obsessive compulsion, anxiety, impulsivity, and cravings in addicted individuals [3-4,22-45].

The exact mechanism of NEAT-12 is unclear. However several preliminary studies have shown that CES alters various neurotransmitters or hormone levels in the brain [43]. One study demonstrated increased catecholamine levels in men and women and increased thyroxin production in men following log-term therapy with CES [36]. Others reported that platelet monoamine oxidase –B (MAO-B) activity and plasma concentration of Gamma –amino butyric acid (GABA) increased following CES, in conjunction with clinical improvement in anxiety and depression [31]. Shealy [44] has repeatedly shown that in both normal volunteers and in volunteers with treatment resistant depression, that CES was associated with significant elevations of MAO-B activity and plasma concentration of Gamma –amino butyric acid (GABA).

Our group Blum et al. [46]. have now completed a number of new studies using qEEG and functional magnetic resonance imaging (fMRI) showing that NAAT (specifically, Synaptose Complex Variant [KB220-Z™]) normalizes the electro-activity of the pre-frontal and cingulated gyrus region of the brain similar to the NEAT 12 devise in protracted poly-drug abusers with Reward Deficiency Syndrome (RDS) [47,48]. Use of this natural non-addicting, safe putative D2 agonist may find its place in recovery from (RDS) including addiction behavioral research involving cue induced reward site activation using fMRI studies in China indicate that an acute dose of KB220-Z compared to placebo in a double blinded study at rest activated the dopaminergic NAc as well as smoothing out abnormalities observed in the Putamen of heroin addicts. Additional addiction behavioral research involving cue induced reward site activation using fMRI is ongoing in China. These results will provide important information that could ultimately lead to significant improvement of recovery for victims of RDS having dopamine deficiency.

Positive outcomes demonstrated by qEEG imaging in a randomized Triple-blind placebo controlled study involving oral KB220-Z, showed an increase of Alpha and low Beta activity [48]. Moreover, preliminary unpublished evidence derived from our fMRI studies in China indicate that cortisol levels in the NAc are increased after KB220-Z treatment compared to placebo. This effect was observed in both fMRI and PET scanning to determine chronic effects of KB220-Z on numbers of D2 receptors and direct interaction at NAc.

To maintain homeostasis of the complex neuroanatomical brain system that utilizes neurotransmitters, poly-drug abusers who are known to be deficient in the neurotransmitters, may require continuous supplemental Neuroadaptogen Amino-Acid Therapy (NAAT) for neurotransmitter augmentation [46].

It is noteworthy, that alcohol may produce addiction by normalizing abnormal baseline states such as irritability, hyperexcitability,
dysphoria, impulsiveness, or anxiety [50]. The results of animal studies have revealed that alcohol can induce sensitivity of neuronal membranes, chemically altered proteins, and adaptations to ion channels and factors related to the binding and release of neurotransmitters and neuromodulators including serotonin, pro-opiomelanocortins, enkephalins, gamma aminobutyric acid, glutamate receptors, and dopamine, norepinephrine and acetylcholine [51]. These effects could manifest via dopamine abducts with acetaldehyde leading to opioid-like actions and potential opioid receptor binding [52-54]. It is quite possible that modulation by NEAT-12 could offset impulsiveness, for example, as only one important target requiring intensive investigation.

While NEAT-12 are not strictly a CES device but a hybrid whereby we utilized current characteristics of TENS, we are encouraged that in larger cohort double-blinded studies it will produce similar CES effects as observed in this case study. A limitation to the use of this device is that its effects are only transient and must be coupled with natural Dopamine D2 agonistic therapy such as NAAT or newly developed variants for further longer term D2 activation in both the posterior VTA and Cingulate gyrus.

Conclusion

The results of this pilot study are encouraging and we are proposing that NEAT-12 could provide adjunctive benefits to the addicted individual by reducing break-through aberrant cravings especially in conjunction with NAAT.

We propose following validation in large controlled studies, the systematic coupling of, RDS Inventory scale, genetic testing, personalized nutrigenomic (NAAT) solutions and NEAT 12, and Comprehensive Analysis of Reported Drugs™ (a medical monitoring personalized nutrigenomic (NAAT) solutions and NEAT 12, and systematic coupling of, RDS Inventory scale, genetic testing, personalized nutrigenomic (NAAT) solutions and NEAT 12, and Comprehensive Analysis of Reported Drugs™ (a medical monitoring personalized nutrigenomic (NAAT) solutions and NEAT 12, and comprehensive approach to the resolution of the ongoing difficulty we could revolutionize the RDS recovery field and ultimately provide an important contribution to the resolution of the ongoing difficulty we face in treating addiction and relapse [55,56].

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Conflict of Interest

Roger L. Waite, and Kenneth Blum own stock in Bio Clarity LLC., worldwide distributors of the NEAT 12 device based on US and Foreign Patents pending. 

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