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Neural Modulators of Temperament: a multivariate approach to personality trait identification in the horse

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ABSTRACT

A relationship between dopamine and temperament has previously been described in human cases of dopaminergic dysfunction. Adjustment in temperament prior to disease manifestation can enable the early identification of individuals at risk of such conditions, and scope exists to extend this application of temperament alterations to

cases of dopaminergic dysfunction in horses. A multivariate and mixed-methods approach utilising a questionnaire along with two inferred measurements of dopamine activity (Spontaneous Blink Rate [SBR] and Behavioral Initiation Rate [BIR]) were recorded from direct observation of animals (n=99) to identify the potential relationship between dopamine and temperament in horses. Principal components analysis (PCA) of 36 temperament variables revealed nine Principal Components, including 'Anxiety' and 'Docility', which accounted for 72.4% of the total variance. Component scores were calculated and correlated with SBR and BIR utilising Spearman Rank Correlation Coefficient analysis. The component 'Anxiety' was found to have a significant positive relationship with SBR, whereas 'Docility' was observed to have a significant negative relationship with SBR. These results indicate a relationship between dopamine and temperament within the horse that is certainly worthy of further study. Potential mechanisms involving neural dopaminergic and GABAergic systems are presented, in addition to how such alterations could be utilised to probe for equine dopamine dysfunction pending future research.

Highlights

- A significant positive correlation was observed between 'Anxiety' and dopamine levels
- A significant negative correlation was found between 'Docility' and dopamine levels
- Chronic dopamine adaptations may initially manifest as temperament alterations
- Potential exists to identify horses at risk of dopamine dysfunction development through analysis of temperament

Key words: Temperament; Dopamine; Equine; Behavior; Anxiety; Docility

1.0 Introduction

Equine temperament is defined as any characteristic of an individual which emerges at a young age and appears to be stable, over both time and situation (Lansade & Simon, 2010). However, this could neglect the distinct behavioral manifestation of temperament. Indeed, Stur (1987) suggested temperament is the sum of inherited and learned behavior patterns, whilst Kilgour (1975) highlights the distinct physical, hormonal and nervous characteristics of an animals' temperament. It is the combination of temperament and environmental conditions which are proposed to amalgamate as equine personality (see Randle, 2015). Temperament is of importance for the performance and leisure horse, for competition success as well as improving equine welfare and understanding (Visser *et al.*, 2001; Randle, 2015). It is perhaps for this reason that a plethora of temperament research does exist for the horse, utilising both questionnaire and behavior methodologies (Seaman *et al.*, 2002; Momozawa *et al.*, 2003; Visser *et al.*, 2008). Whilst a number of studies have investigated learning and memory functioning in horses in relation to temperament traits (Lansade & Simon, 2010; Lansade *et al.*, 2013; Valenchon *et al.*, 2013), the way in which this relates to neural functioning has not been investigated. This is perhaps surprising given Kilgour's (1975) early reference to 'nervous organisation' in his temperament definition, as well as the influence of prefrontal, striatal and hippocampal circuitry on such tasks (Izquierdo & Medina, 1997; Seger & Cincotta, 2005; Cartoni *et al.*, 2013).

Of particular interest to date is the relationship between the neurotransmitter dopamine (DA) and its role in temperament manifestation, with specific reference to human conditions of DA dysfunction, such as Parkinson's disease (PD), a hypodopaminergic condition (Burch

& Sheerin, 2005). An interesting aspect of PD onset is the emergence of a pre-morbid personality which can predate the emergence of motor symptoms, in some instances by decades (Todes & Lees, 1985). Indeed, PD patients often present with stoic, rigid and slow temperaments as a part of this characteristic parkinsonian personality (Dagher & Robbins, 2009), whilst also demonstrating low novelty seeking scores (Kaasinen *et al.*, 2001). This alteration in temperament has previously been linked to the hypodopaminergic characteristic of PD, though some conflicting evidence does remain (Kaasinen *et al.*, 2001).

In contrast, patients who exhibit schizophrenia, a hyperdopaminergic condition (Breier *et al.*, 1997) demonstrate an increased prevalence for anxiety disorders (Pallanti *et al.*, 2004; Achim *et al.*, 2009). Moreover, anxiety is thought to contribute as part of the prodromal stage of schizophrenia (Park *et al.*, 2016). When considered in the context of the proposed mechanism of anxiogenic drugs which ultimately elevates neural DA levels (Cancela *et al.*, 2001), this suggests that chronic DA reduction or elevation can manifest as a significantly altered temperament.

Horses can also present with hyper- and hypo-dopaminergic conditions, including stereotypic behaviours (SB; McBride & Hemmings, 2005; Roberts *et al.*, 2015) and pituitary pars intermedia dysfunction (PPID; McFarlane *et al.*, 2005), respectively. Interestingly, there is also an observation that horses with PPID present with significant alterations in temperament, including an increase in depression, lethargy and apathetic manifestations with disease progression (Bradaric *et al.*, 2013). However, this temperament alteration has not yet been linked with the reduction of DA which occurs within these animals. Furthermore, in the light of the pre-morbid personality associated with PD and anxiety within

the prodromal phase of schizophrenia, it is plausible that dopaminergic alterations could manifest prior to conventional diagnosis through analysis of temperament traits. This approach would however need to proceed mindful of the balance between genotypic determination of temperament and subsequent environmental modification of behavioural output. Whilst temperament and personality type research has been conducted in horses (for example see Momozawa *et al.*, 2003; Momozawa *et al.*, 2005a; Lloyd *et al.*, 2007, Nagy *et al.*, 2010; Ijichi *et al.*, 2013), the way in which the result of this relates to dopaminergic function has yet to be determined empirically. Interestingly however, Momozawa *et al.* (2005b) observed a significant relationship between *DRD4* (dopamine D4 receptor subtype) single nucleotide polymorphism, an A-G substitution causing an amino acid change from asparagine to aspartic acid (Hori *et al.*, 2013). Absence of the 'A' allele resulted in significantly increased 'Curiosity' and lower 'Vigilance' scores than those with the 'A' allele (Momozawa *et al.*, 2005b). Whilst persuasive of an influence of dopamine over temperament in the horse, the precise mechanisms as to how the *DRD4* gene directly influences temperament in any species is currently unknown and requires further research.

Previous research has demonstrated the successful utilisation of the proven DA correlates spontaneous blink rate (SBR) and behavioral initiation rate (BIR) to investigate dopaminergic function in oral and locomotor stereotypy in the horse (Roberts *et al.*, 2015), but as of yet no links have been made to any temperament data. It is for this reason that SBR and BIR were utilised here to examine the role of dopamine in the generation of temperament. A questionnaire was used to gather temperament data due to previous validation of this approach within the literature (Momozawa *et al.*, 2003; Momozawa *et al.*, 2005a; Lloyd *et al.*, 2007). Results of this study could provide a valuable basis for the early

identification of horses undergoing initial DA dysfunction thereby improving welfare through timely intervention.

2.0 Methods

2.1 Sample Population Characteristics and Management

A sample of 100 horses of varying backgrounds were sought, comprising 52 geldings, 44 mares and four stallions, aged 2-25 years (mean \pm SD 12.17 \pm 6.24 years) and included some stereotypy performing horses (n=3 crib-biters, n=15 weavers, n=9 box walkers) due to previously described linkage between SB, dopamine (Cabib & Bonaventura, 1997) and temperament traits (Dagher & Robbins, 2009; Park *et al.*, 2016). All animals were recruited through direct contact from private owners (n=40), and were kept at various locations in Gloucestershire and Wiltshire, in the South West of England during the study period (June 2014-January 2015). In addition, these animals differed in terms of management regime, breed (ranging from native breeds to sport horses) and use (from leisure to polo and competition disciplines such as polo and dressage) in order to source a representative cross-section of the UK horse population with the level of variance favoured for Principal Component Analysis (Ijichi *et al.*, 2013). As such, selection criteria required that the horse was free from acute clinical disease (excluding dopamine pathology) and able to be led from a head collar and lead rope (Ijichi *et al.*, 2013).

2.2 Equine Temperament Questionnaire

Owners were asked to complete a two-part questionnaire. The first section of the questionnaire was designed to collect information regarding the management and feeding

regime of each individual, along with factors (i.e. breed) shown by other studies to impact upon temperament (Hausberger & Muller, 2002; Lloyd *et al.*, 2008). Section 2 was based on previous equine temperament questionnaires (Momozawa *et al.*, 2003; Momozawa *et al.*, 2005a; Lloyd *et al.*, 2007). A 1-9 Likert scale was used to assess 41 temperament descriptions. Each description was identified by a definition in order to ensure that participants had an agreed understanding of the temperament term being used. For example the temperament description '*Concentration*' was accompanied by the definition '*is trainable and undisturbed by the environment*'. All of the chosen temperament descriptions were selected from the aforementioned studies, though some were omitted due to repetition or irrelevance with reference to dopamine. One additional temperament description '*Impulsive*' with accompanying definition '*tends to act without forethought, regardless of the consequences*' was added, as impulsivity is thought to be characteristic of abnormal behaviors such as stereotypy (Garner & Mason, 2002) and has links back to dopamine transmission (Cools, 2008).

2.3 Spontaneous Blink Rate Observation

The method to assess SBR for all of the horses has previously been described in Roberts *et al.* (2015). To reduce stress for the animal during observations, all horses had visual contact with conspecifics. The purpose of this was for both ethical reasons i.e. not to isolate a social species, but also to avoid artificially inducing alterations in the horses SBR due to external factors. Each horse was stabled in their home stable and habituated to the researcher's presence by having the researcher stand calmly outside the stable with the horse in full view for 10 minutes prior to the observation period. Following the 10 minute habituation period, continuous method sampling was utilised where each full blink (defined

by Karson (1983) as bilateral paroxysmal brief repetitive eye closures occurring continuously) was recorded with a mechanical counter for 30 minutes. Considering the anatomy of the horse, it was difficult for a solo observer to record true bilateral eye closures, so only the left eye was observed for all horses. As such, the horses head collar remained on throughout to ensure the horse remained in position in such a way that the observer had full view of the horses left eye whilst outside the stable at all times. This procedure was repeated in the same manner over three consecutive days, allowing a mean SBR/30minutes to be calculated for each horse.

2.4 Behavioral Initiation Rate Observation

This procedure has previously been described for use in the horse by Roberts *et al.* (2015), based on BIR observations of bank voles (Garner & Mason, 2002). The horse was stabled within their home stable and habituated to the observers' presence for 10 minutes as described in Section 2.3. Following habituation, each behavioral initiation was recorded by continuous sampling utilising a mechanical counter for 30 minutes where the observer was positioned outside of the stable though with full view of the horse. All behaviors performed were defined by a pre-determined ethogram (McDonnell, 2003), and similar to Garner and Mason (2002) only the number, not the type, of behaviors was recorded. Furthermore, each bout of behavior was recorded as one initiation irrespective of the previous behavior, consequently the sequence 'Feeding – Grooming – Feeding – Drinking – Standing Rest' was recorded as four initiations (Garner & Mason, 2002; Roberts *et al.*, 2015). Thus, movement made as part of an identifiable behavior, for example lifting of the head whilst still undergoing mastication as part of feeding was accepted as part of the behavior and therefore not recorded as an initiation of a new behavior (Roberts *et al.*, 2015). Recording of

stereotypic behaviors was considered in terms of bouts of behavior, i.e. regardless of the number of weaves produced, each single bout of weaving was recorded as one behavioral initiation. BIR recording was repeated over three consecutive days in the same manner, allowing a mean BIR/30minutes to be calculated for each horse.

2.5 Statistical Analysis

A PCA was carried out with varimax rotation on all questionnaire items. Components with eigenvalues ≥ 1 were retained. Component loadings of ≥ 0.4 within each component were considered significant (McGrogan *et al.*, 2008) and checked for PCA suitability with the Kaiser-Meyer-Olkin (KMO) statistic. The Anderson-Rubin method for standardising component scores was applied to ensure that all component scores were directly orthogonal. Once component scores had been calculated for every horse for each temperament component, these were then correlated with the horses' mean SBR and BIR utilising a Spearman Rank Correlation Coefficient, similar to that conducted previously utilising both behavioral and questionnaire measures (Lloyd *et al.*, 2007). Mean SBR and BIR values were also calculated. A second PCA was conducted to investigate SBR and BIR as contributory towards component structure i.e. if for example SBR contributed towards a particular temperament component. To examine if sex, breed and usage had an influence on temperament and dopamine correlate data, Kruskal-Wallis with post-hoc pairwise comparisons and Bonferroni correction, or Mann-Whitney U tests where appropriate, were undertaken. To decipher whether median or mean-rank was presented, visual inspection of the box-plots produced via SPSS were utilised. Where distributions were similar median values were presented, though when distributions were dissimilar, median-ranks were utilised. Breeds were categorised into pony/cob types (14.2hh and under), sport horse types

(light breeds e.g. Thoroughbreds 14.3hh and over) and draught horse types (heavy breeds e.g. Irish Draught 14.3hh and over). Uses were categorised as leisure, so called traditional 'sports' disciplines (dressage, show jumping and cross-country jumping) or polo dependent on the horse's primary purpose. The significance level was set at $p < 0.05$. All statistical analyses were conducted in IBM SPSS version 22.

3.0 Results

3.1 PCA Data

A 99% return rate was achieved from the questionnaires, only animals with which temperament data were available were utilised for subsequent analysis, giving a total of $n=99$. Initially, all 41 temperament variables were utilised to perform the PCA. Five temperament traits (Timidity, Sociable, Protective, Subordinate and Permissive) were deemed unreliable following examination of the KMO statistic and were consequently removed from further PCA analysis. When the PCA was conducted with the remaining 36 temperament descriptions, nine components were extracted accounting for 72.4% of the total variance (see Table 1). The names of the components were derived from examination of the individual temperament definitions which contribute towards a component (Table 2). Previous research was also taken into account (Momozawa *et al.*, 2003; Momozawa *et al.*, 2005; Lloyd *et al.*, 2007; Nagy *et al.*, 2010; Ijichi *et al.*, 2013; See Appendix 1). As such, the components were named as follows, in descending order of percentage variance explained; 'Anxiety', 'Trainability', 'Excitability', 'Docility', 'Inquisitiveness', 'Irritability', 'Self-Reliance', 'Horse-Horse Interaction' and 'Horse-Human Interaction'.

Distributions of temperament traits and dopamine correlates were similar as assessed by visual inspection. Median 'Anxiety' score for mares (n=44; -0.54) versus geldings (n=55; 0.05) were significantly different ($U = 1534$, $z = 2.282$, $p = 0.023$). Furthermore, median 'Inquisitiveness' scores for mares (-0.23) was significantly lower than geldings (0.09) ($U = 1489$, $z = 1.965$, $p = 0.049$). Likewise, median 'Horse-Human Interaction' scores were significantly lower for mares (-0.13) than geldings (0.39) ($U = 1654$, $z = 3.127$, $p = 0.002$). No other significant differences were observed between mares and geldings for temperament components or dopamine correlates.

3.3 Effect of Breed

Distributions of temperament and dopamine correlates were not similar for pony/cob types (n=33), sport horse types (n=60) and draught types (n=6) as assessed by visual inspection of the box-plot, thus mean-ranks are provided. SBR ($\chi^2 (2) = 6.872$, $p = 0.032$), 'Anxiety' ($\chi^2 (2) = 8.665$, $p = 0.013$) and 'Excitability' ($\chi^2 (2) = 12.916$, $p = 0.002$) scores were significantly different between the three breed categories when corrected for ties. Pairwise comparisons with Bonferroni correction revealed significant differences in SBR between pony/cob types (39.35) against sports horse types (55.61; $p = 0.027$) though no difference was found between draught types (52.50) against either pony/cobs ($p = 0.907$) or sport horses ($p = 1.000$). Similarly, pony/cob types demonstrated a significantly lower mean rank 'Anxiety' score (39.36) than sport horse types (56.85; $p = 0.015$). No difference was observed between draught types (40.00) compared to pony/cob types ($p = 1.000$) or sport horse types ($p = 0.512$). A significantly lower 'Excitability' mean rank for pony/cob types (38.58) compared to sport horse types (58.23; $p = 0.005$) was also observed. There was no difference between draught types (30.50) with either pony/cob types ($p = 1.000$) or sports

horse types ($p = 0.072$) and no further differences were observed for BIR nor any of the remaining seven temperament components.

3.4 Effect of Use

Distributions of temperament and dopamine correlates were similar for leisure ($n=56$), traditional sports discipline ($n=27$) and polo ($n=19$) horses as assessed by visual inspection of the box-plot, thus median values are provided. 'Anxiety' ($\chi^2 (2) = 9.418, p = 0.009$), 'Excitability' ($\chi^2 (2) = 8.138, p = 0.017$) and 'Inquisitiveness' ($\chi^2 (2) = 6.002, p = 0.05$) were significantly different between the three use categories when corrected for ties. Pairwise comparison with Bonferroni correction highlighted that the leisure animals demonstrated a lower median 'Anxiety' score (-0.21) compared to sport discipline horses (0.50; $p < 0.009$). No significant difference was observed for median 'Anxiety' score between polo horses (-0.01) and leisure ($p = 0.335$) nor traditional sports discipline animals ($p = 1.000$). Leisure animals demonstrated a lower median 'Excitability' score (-0.36) compared to traditional sports discipline horses (0.49; $p = 0.019$). Polo animals median score (-0.30) showed no significant difference between either leisure ($p = 1.000$) or sports discipline horses ($p = 0.112$). Furthermore, leisure animals demonstrated an increased median 'Inquisitiveness' score (0.06) compared to polo horses (-0.51; $p = 0.049$), though no difference was apparent between traditional sports discipline horses (0.09) and leisure ($p = 1.000$) or polo animals ($p = 0.130$). No other temperament traits, SBR or BIR were found to demonstrate statistical significance between uses.

4.0 Discussion

The combined PCA and correlational analysis supported the hypothesis that one known measure of inferred DA activity (SBR) is correlated with the two temperament components 'Anxiety' and 'Docility', a finding not yet reported in the equine literature, although the range of SBR data gathered is roughly in-line with our previous work (Roberts *et al.*, 2015). SBR is correlated with striatal DA levels, with increases indicating higher, and decreases lower, levels of production and release of DA (Kaminer *et al.*, 2011). Therefore, our data suggest that SBR may not only be a potentially useful predictor of central dopamine function in relation to behavioral output and as a risk factor for onset of pathology, but also as a proximate predictor tool. Furthermore, given that SBR does not appear to contribute to temperament traits themselves, this could indicate the suitability of SBR as a potential measure of underlying temperament components as opposed to being a direct causal factor. Temperament and personality have previously been cited as highly influential for horse purchase and breeding, and a need to incorporate an objective temperament assessment identified by Graf *et al.* (2013). Thus the use of SBR as part of pre-purchase veterinary examination may allude to the temperament of the animal and be beneficial to the horse owning population. Indeed, improved understanding of equine temperament prior to purchase could benefit human safety, given that unanticipated horse behavior was highlighted as a contributory factor in 61% and 39% of injuries in children under 15 and adults above 15 respectively (Northey, 2003).

4.1 'Anxiety' and Dopamine

Due to the causal links between stress and DA (Cabib *et al.*, 1998), it is possible that anxious horses are more sensitive to environmental stressors, such as restricted feeding or social isolation; common stressors faced by stabled horses (McAfee *et al.*, 2002; Ninomiya

et al., 2007). Underlying this increased responsiveness to stress, the anxious individuals may have elevated striatal DA in comparison to less anxious animals kept under the same environmental conditions. This could allow the initiation of active coping in an attempt to gain control over the environment, similar to the elevated dopamine levels observed in the active coping DBA mouse strain (Cabib & Bonaventura, 1997; Cabib & Puglisi-Allegra, 2012). A similar process could be occurring with the anxious horses, as evidenced by the elevation in SBR in these individuals.

From a mechanistic standpoint, when DA agonists are administered, there is a partnership between the elevation of DA levels and the emergence of behaviors characteristic of anxiety (McCullough & Salamone, 1992; Cancela *et al.*, 2001). This DA elevation appears to result from GABA disinhibition at the level of the midbrain i.e. the ventral tegmental area (VTA) (Biggio *et al.*, 1990; Nikulina *et al.*, 2005). A similar progression could also be apparent in environmentally induced anxiety. For example in rodents chronic stressors such as restricted feeding, social isolation and restricted locomotion are known to induce mu-opioid proliferation (Nikulina *et al.*, 2005), and therefore resulting in GABA disinhibition, in addition to sensitisation of the dopaminergic pathways (Cabib *et al.*, 1998). Combined, this could indicate a relationship between mu-opioid receptor proliferation and GABAergic disinhibition, perhaps giving rise to elevated DA. These chronic stressors are strikingly similar to those faced by the stabled horse, and induce similar neural adaptations. For example, dopaminergic pathways are suggested to be sensitised within stress-linked conditions of the horse including oral (McBride & Hemmings, 2005) and locomotor stereotypy (Roberts *et al.*, 2015). Further research indicates mu-opioid receptor density is significantly elevated within horses displaying oral stereotypy at the VTA, as well as dorsal and ventral striatal regions (Hemmings *et al.*, 2006). Thus, it is possible that in the more

anxious animals, mu-opioid receptor sensitisation and GABAergic disinhibition giving rise to DA elevation has occurred, or is well underway. This could well be a perpetuating issue given that DA elevation within the amygdala, a structure which also receives dopaminergic input from the VTA and is characteristic for its fear gating and anxiety inducing functions, potentiates anxiety in both rodent and human models (Kienast *et al.*, 2008). Additionally, this elevation in dopamine effectively removes inhibitory control of the medial prefrontal cortex over amygdala function (de la Mora *et al.*, 2010) giving rise to prolonged 'Anxiety' type responses. This could therefore provide a mechanism by which elevated 'Anxiety' is observed in this population of horses presenting with raised SBR.

Importantly in rodents, persistent DA elevation is the final step to instigate active coping, leading to stereotypy manifestation in stressed, captive animals (Cabib & Bonaventura, 1997). The emergence of an elevated 'Anxiety' type temperament in the horse could similarly be indicative of significant underlying neural adaptations which pre-date stereotypy development. Timely removal of key stressors could prevent progression to the stereotypic behavioral end-point in this regard. Indeed, as neural sensitisation of DA appears to be permanent (or at least recalcitrant) where rodent species are concerned, prophylaxis rather than remediation is recommended (Cabib *et al.*, 1998). This latter point is particularly important given the proposed habitual mechanisms of crib-biting behavior (Hemmings *et al.*, 2007; Parker *et al.*, 2009; Roberts *et al.*, 2015), and also with the highly motivated (albeit non-habitual) phenotype thought to contribute to weaving behavior (Roberts *et al.*, 2015). Also of interest with regard to environmentally induced stereotypy is that under improved management regimes, which reduce chronic stress and potentially the normalisation of DA levels within the neural circuitry, the high 'Anxiety' horses may then become more biddable in the context of training and management. Indeed, the performance of increasingly anxious

type behavior is a part of the prodromal phase of schizophrenia (Park *et al.*, 2016) a hyper-dopaminergic phenomenon. As such, the use of increasing anxiety is currently under investigation as a potential predictor of an oncoming schizophrenic episode (Park *et al.*, 2016). In light of this evidence, it is plausible then that the more anxious horses are presenting as such due to consistently elevated levels of the neurotransmitter DA.

However, when investigating the work completed by Nagy *et al.* (2010), an interesting question arises. It was reported that crib-biting horses demonstrated a significantly lower 'Anxiety' score when compared to the control animals (Nagy *et al.*, 2010). Whilst this initially appears to contradict what is being suggested here, the data reported by Nagy *et al.* (2010) may in fact lend support. Crib-biting animals present as hypodopaminergic, as evidenced by a significantly decreased SBR and significant reduction in caudate dopamine receptor sensitivity (McBride & Hemmings, 2005; Roberts *et al.*, 2015). This is despite post-mortem evidence of increased receptor sensitivity to dopamine within the nucleus accumbens (McBride & Hemmings, 2005). This neural work supports the notion that SBR is a reflection of midbrain DA transmission into the dorsal striatum (caudate nucleus) rather than the ventrally sited nucleus accumbens (Taylor *et al.*, 1999). However, elevations in nucleus accumbens DA are of significant behavioral relevance, and have been considered crucial to the putative self-stimulatory stress reducing aspects of crib-biting. It may be that following the onset of a stress coping function i.e. crib-biting performance, leads to the diminishing anxiety in crib-biting animals as reported by Nagy *et al.* (2010). Indeed, recent evidence appears to support the stress coping function of crib-biting (Freymond *et al.*, 2015). Overall then, it is plausible that initial elevations in DA could manifest as increased 'Anxiety', though potentially following the development of stereotypy as a part of the behavioral repertoire

'Anxiety' could decrease, at least in crib-biting horses. Further research would be required to investigate such an effect in weaving animals.

4.2 'Docility' and Dopamine

'Docility' was negatively correlated with SBR, and thus inferred levels of DA activity. A comparative phenomenon is observed in the inbred mouse strain C57, whereby in response to an inescapable stressor, a net reduction in DA transmission leads to reduced anxiety linked behaviors such as locomotion (Cabib, 2006; Cabib & Puglisi-Allegra, 2012). Rather than the often quoted pathological condition of learned helplessness, the depressed activity in the C57 strain is thought to signal a passive form of coping, which promotes energy thriftiness in the face of insurmountable stress (Cabib & Puglisi-Allegra, 2012). The reduced SBR which accompanies 'Docility' in the data reported herein, could well indicate a similar strategy is adopted by the horse in response to persistent low level stressors such as restricted locomotion and social isolation.

From a pathology prediction standpoint, in stark contrast to those with conditions characterised by elevated DA such as schizophrenia and drug addiction, human patients with PD typically present with personalities which are categorised as stoic, rigid and slow-tempered (Dagher & Robbins, 2009). Novelty-seeking type behaviors are also reduced in PD, with twin studies also demonstrating that individuals who later develop Parkinson's are more self-controlled than their non-affected twin (Menza, 2000). It is thought that the emergence of such traits are directly related to underlying degeneration of DA containing neurons (Kaasinen *et al.*, 2001). When considering the individual temperament variables

'slow' and 'patient' which contribute to the 'Docility' temperament trait in this cohort of horses (Table 2), it is reasonable to propose that the more 'docile' horses are also more slow-tempered and self-controlled. Parallels could be drawn between Parkinson's disease personality and conditions of the horse which feature reduced DA such as PPID. Typically, PPID is suspected when overt symptoms, such as hirsutism and hyperhidrosis are observed in combination with advancing age, at which point it is likely that the influence of elevated proopiomelanocortin (POMC) peptide levels are already having a significant detrimental effect on health (McFarlane, 2011). Indeed, ACTH is characteristically elevated in PPID horses prior to pharmacological treatment, with this posing long term health issues leading to reduced welfare status of the animal (Durham *et al.*, 2014). Early intervention with regards to PPID development could be essential for prolonged quality of life and improved welfare (McFarlane *et al.*, 2011). Despite this, there is a lack of peer reviewed publications specifying the early indicators of PPID, which when treated earlier could significantly improve prognosis and prevent life threatening consequences of the disease (McFarlane *et al.*, 2005). Therefore, our finding that a decrease in SBR is correlated with higher 'Docility' could provide an important early indicator for those at risk of PPID development. This is of significance given that depression, lethargy and an apathetic outlook are observed following PPID diagnoses (Bradaric *et al.*, 2013), and could therefore link towards a more 'docile' temperament. Thus, should the horse demonstrate an uncharacteristic alteration in temperament by increasing in 'Docility', this may signify that there are alterations with regards to DA physiology that could indicate the development of PPID. This is highly noteworthy, as previously McFarlane (2011) suggested that observed docility was due to increased beta-endorphin circulation. In light of the findings here however, it seems probable that increased 'Docility' is a reflection instead of DA reduction. Taken together, should the owner notice elevated 'Docility' even in the absence of PPID

symptomology, an in-depth endocrine work up should certainly be considered, but also proposes an interesting area for further research.

4.3 Effect of Breed, Use and Sex

Pony/cob types were found to have significantly reduced SBR, 'Anxiety' and 'Excitability' scores than sports horses. In other studies (see Cabib & Puglisi-Allegra, 2012) utilising rodents, decreased dopamine release is associated with a passive, less active stress coping style. Decreased SBR, along with lower 'Anxiety' and 'Excitability' as observed here constitutes evidence for a similar phenomenon in the horse. On the other hand, our data is somewhat at odds with the work of Lloyd *et al.* (2008) who observed similar 'Anxiousness' and 'Excitability' in both welsh ponies and cobs compared to Thoroughbreds and Arabs. However, Lloyd *et al.* (2008) formed sample groups based on breed rather than the more generalised type designation employed in this study, which may account for the discrepancy.

Interestingly, geldings scored significantly higher than mares for 'Anxiety', 'Inquisitiveness' and 'Horse-Horse Interaction'. Duberstein and Gilkeson (2010) observed that mares were more 'Anxious' than geldings, a finding directly opposing to the results presented here. Whereas Le Scolan *et al.* (1997) previously found no significant difference between mares and geldings with their study of equine temperament. It should be noted however, that both of these previous studies utilised lower numbers of horses (n=18 and n=72 respectively) which may account for the lack of inter-study agreement. Given the gender imbalance often

observed in some equine sporting disciplines (i.e. mares are generally favoured as polo ponies) this area is certainly worthy of further investigation.

Finally, leisure animals were found to have significantly reduced 'Anxiety' and 'Excitability' compared to those engaged in traditional sports disciplines, though had significantly elevated 'Inquisitiveness' scores than polo ponies. It could reasonably be postulated that polo ponies and sport horses are more likely to be managed under environmental conditions (i.e. housing and feeding regimen) that promote chronic stress potentially giving rise to the elevated 'Anxiety' and 'Excitability' along with the reduced 'Inquisitiveness' scores observed here.

5.0 Conclusion

This research is the first demonstration of a relationship between temperament and inferred neural DA levels within the horse. Whilst this study cannot elucidate the precise underlying mechanisms governing the DA and temperament relationship, extrapolation from previous research provides a sound basis upon which to develop future work. The ability to utilise the presentation of either increased 'Anxiety' or 'Docility' to indicate altered neural DA function at an early stage could prove valuable for the welfare of at risk animals. The potential ramifications of this work, and the group differences observed, therefore highlights this is an area that warrants further research.

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Appendix 1

Table 3. Component names attributed to the extracted components. Note – superscripts sharing a number of the same row are indicative of the source of the variable. *highlights that these are not exact matches, though indicate similarities in findings

Component Number	Component Name	Matching Variables	Reference/s
1	Anxiety	Nervousness ^{1,3} Excitability ¹ Panic ^{1,3} Inconsistent ¹ Skittishness ^{1,3} Suspicious ² Tense ² Apprehensive ² Fearful ²	Momozawa <i>et al.</i> (2003) ¹ Momozawa <i>et al.</i> (2005a) ¹ Lloyd <i>et al.</i> (2007) ² Nagy <i>et al.</i> (2010) ³
2	Trainability	Concentration ^{1,2} Trainability ^{1,2} Memory ^{1,2} Patience ¹ Cooperation ²	Momozawa <i>et al.</i> (2003) ¹ Momozawa <i>et al.</i> (2005a) ¹ Nagy <i>et al.</i> (2010) ²
3	Excitability	Active ¹ Excitable ¹	Lloyd <i>et al.</i> (2007) ¹
4	Docility		None Similar
5	Inquisitiveness	Curious ¹ Opportunistic ¹	Lloyd <i>et al.</i> (2007) ¹
6	Irritability	Inconsistent ¹ Stubborn ^{1,2} Irritable ²	Momozawa <i>et al.</i> (2005a) ^{1*} Lloyd <i>et al.</i> (2007) ^{2*}
7	Self-Reliance		None Similar
8	Horse-Horse Interaction	Friendliness_Horse ¹ Popular ²	Momozawa <i>et al.</i> (2005a) ^{1*} Lloyd <i>et al.</i> (2007) ^{2*}
9	Horse-Human Interaction		Ijichi <i>et al.</i> (2013)*