Orexinergic neuron numbers in three species of African mole rats with rhythmic and arrhythmic chronotypes.

Adhil Bhagwandin¹, Nadine Gravett¹, Jason Hemingway¹, Maritjie K. Oosthuizen², Nigel C. Bennett², Jerome M. Siegel³ & Paul R. Manger¹.

¹School of Anatomical Sciences, Faculty of Health Sciences, University of the Witwatersrand, 7 York Road, Parktown 2193, Johannesburg, Republic of South Africa.

²Mammal Research Institute, Department of Zoology and Entomology, University of Pretoria, Pretoria 0002, South Africa.

³Department of Psychiatry, UCLA School of Medicine, Brain Research Institute, Neurobiology Research, 151A3, Sepulveda VA Medical Centre, North Hills, CA, 91343.

Correspondence to: Paul Manger

School of Anatomical Sciences
Faculty of Health Sciences
University of the Witwatersrand
7 York Road, Parktown, 2193
Johannesburg, REPUBLIC OF SOUTH AFRICA.
Ph: +27 11 717 2497 Fax: +27 11 717 2422
Paul.Manger@wits.ac.za
Abstract

In the present study orexinergic cell bodies within the brains of rhythmic and arrhythmic circadian chronotypes from three species of African mole rat (Highveld mole rat – *Cryptomys hottentotus pretoriae*, Ansel’s mole rat – *Fukomys anselli* and the Damaraland mole rat – *Fukomys damarensis*) were identified using immunohistochemistry for orexin-A. Immunopositive orexinergic (Orx+) cell bodies were stereologically assessed and absolute numbers of orexinergic cell bodies were determined for the distinct circadian chronotypes of each species of mole rat examined. The aim of the study was to investigate whether the absolute numbers of identified orexinergic neurons differs between distinct circadian chronotypes with the hypothesis of elevated hypothalamic orexinergic neurons in the arrhythmic chronotypes compared to the rhythmic chronotypes. We found statistically significant differences between the circadian chronotypes of *F. anselli*, where the arrhythmic group had higher mean numbers of hypothalamic orexin neurons compared to the rhythmic group. These differences were observed when the raw data was compared and when the raw data was corrected for body mass ($M_b$) and brain mass ($M_{br}$). **For the two other species investigated no significant differences were noted between the chronotypes**, although a statistically significant difference was noted between all rhythmic and arrhythmic individuals of the current study when the counts of orexin neurons were corrected for $M_b$ – the arrhythmic individuals had larger numbers of orexin cells.

**Key words**: Orexin; hypocretin; circadian rhythmicity; sleep.
1. Introduction

It is well known that mole rats, subterranean, visually regressed rodents, have unusual patterns of circadian rhythmicity (Lovegrove and Papenfus, 1995; Lovegrove and Muir, 1996; Oosthuizen et al., 2003). Locomotor activity studies have shown that within a species of mole rat there are individuals that have a rhythmic chronotype, and others that are distinctly arrhythmic and this is seen in both social and solitary species of mole rat (Oosthuizen et al., 2003). A study of sleep patterns in the giant Zambian mole rat (Cryptomys mechowi) showed some differences in sleep patterns between rhythmic and arrhythmic chronotypes of this species, where arrhythmic individuals spend more time in waking with a longer average duration of a waking episode, and less time in nonREM (NREM) sleep with a shorter average duration of a NREM episode though with a greater NREM sleep intensity (Bhagwandin et al., 2011b).

The neurotransmitter orexin is known to promote wakefulness, particularly during motor activities (Estabrooke et al., 2001; Mileykovskiy et al., 2005; Lee et al., 2005; Saper et al., 2005). Orexinergic neurons, which are restricted to the hypothalamus, have ascending projections to the cerebral cortex and descending projections to the cholinergic (basal forebrain, lateral dorsal tegmental nucleus, pedunculopontine nucleus) and monoaminergic (locus coeruleus, ventral tegmental area, raphe) nuclear groups of arousal systems involved in the sleep-wake cycle (Peyron et al., 1998; Chemelli et al., 1999; Baumann and Bassetti, 2005). Loss of orexinergic neurons has been associated with narcolepsy (Chemelli et al., 1999; Lin et al., 1999; Siegel, 1999), narcolepsy with cataplexy in humans (Peyron et al., 2000; Thannickal et al., 2000), narcolepsy without cataplexy in humans (Thannickal et al., 2009) and the narcoleptic like symptoms in patients with Parkinson’s disease (Thannickal et al., 2007).

It is also known that orexins are involved in appetite regulation through activation of the leptin sensitive neurons of the arcuate nucleus and its subsequent feeding associated signaling to orexinergic neurons of the lateral hypothalamus (LH) and henceforth interaction between LH and the hypothalamic ventro-medial nucleus (VMH) (Stellar, 1954; Rodgers et al., 2002; Sakurai, 2003; Sakurai, 2005). Several studies have demonstrated increased orexinergic activity during states of vigilance and foraging and as a result have paired waking with hunger sensations and optimal feeding (Sakurai et al., 1998; Edwards et al., 1999; Haynes et al., 1999; Van Itallie 2006). Conversely anorexinergic activity has been associated
with sleep states, thereby pairing satiety with sleep (Rodgers et al., 2002; Nicolaidis, 2006, Van Itallie, 2006). These studies are indicative of a coordinated orexinergic control of appetite regulation with the sleep-wake cycle that is responsive to the SCN circadian clock and nutritional status (Sakurai, 2005; Van Itallie, 2006), plus, it has been suggested that disruption of the circadian cycle and sleep deprivation can affect energy balance resulting in changes in body composition (Taheri et al., 2004; Van Itallie, 2006).

Given this scenario on the action of the orexinergic neurons, and the clearly distinct circadian chronotypes of the African mole rats, along with differences in the neuropeptide population of the suprachiasmatic nucleus of the mole rats (Negroni et al., 2003), it was thought that an investigation of orexinergic neuronal numbers may yield results of interest in relation to function, especially so in the mole rats. In the current study, orexinergic neuronal numbers of three species of African mole rat, the Highveld mole rat (Cryptomys hottentotus pretoriae), Ansel’s mole rat (Fukomys anselli) and the Damaraland mole rat (Fukomys damarensis) were quantified. It should be noted that Ansel’s and Damaraland mole rats were previously considered to be members of the genus Cryptomys. All three species of mole rat have a reduced visual system, live a subterranean lifestyle and have unusual patterns of circadian rhythmicity (Lovegrove and Papenfus, 1995; Lovegrove and Muir, 1996; Oelschlager et al., 2000; Cernuda-Cernuda et al., 2003; Negroni et al., 2003; Oosthuizen et al., 2003; Gutjahr et al., 2004; Nemec et al., 2004; Nemec et al., 2008). Brains from circadian distinct rhythmic and arrhythmic individuals of each species were examined. The unusual circadian patterns of individuals within a species provide interesting models to examine whether the absolute numbers of identified orexinergic neurons differs between circadian chronotypes amongst the species examined.
2. Materials and Methods

A total of 18 brains (15 males and 3 females – see table 1) from three species of wild caught African mole rat, the Highveld mole rat (*Cryptomys hottentotus pretoriae*) (CHP) (average body mass: 127.9g; average brain mass: 1.5g), Ansel’s mole rat (*Fukomys anselli*) (FA) (average body mass: 84.5g; average brain mass: 1.2g), and the Damaraland mole rat (*Fukomys damarensis*) (FD) (average body mass: 107.8g; average brain mass: 1.8g) were utilized in this study. For the purposes of the current study, three rhythmic and three arrhythmic individuals per species were used. A single mole rat was housed in an enclosure (30 x 25 cm) and wood shavings were used to line the floor. The mole rats used in the present study were caught from wild populations in South Africa under permission and supervision from the appropriate wildlife directorates. All animals were treated and used according to the guidelines of the University of Pretoria Animal Ethics Committee, which parallel those of the NIH for the care and use of animals in scientific experimentation.

2.1 Determination of Rhythmicity patterns

Infrared detectors were used to detect activity patterns in the mole rats. The mole rats were exposed to differing light conditions that included, light (L) and dark (D) cycles, DD cycles and LL cycles. Initially the experimental protocol started with a 12L:12D cycle and once entrainment was achieved the light cycle was switched to constant darkness for 1 month. Recorded behavioral activity data was analyzed using Actiview and Clocklab software which were also used to generate actograms. The sensitivity of the y-axis remained the same during analysis of each mole rat. For the purposes of the current study it was of importance to determine whether the mole rats were able to re-entrain to light cycles and switch their activity according to the light. If a mole rat was able to re-entrain it was deemed ‘rhythmic’ and the converse were applied to ‘arrhythmic’ mole rats. For details regarding the specifics of circadian rhythm determination see Oosthuizen et al. (2003).

2.2 Orexin Immunohistochemistry

Individuals of each species of mole rat were perfused between 17:00 and 20:00 on three consecutive days, i.e. a day per species. The mole rats were euthanized (Euthanaze,
200mg sodium pentobarbital/kg, i.p.) and then perfused intracardially. The perfusion was initially done with a rinse of 0.9% saline solution at 4°C, followed by a solution of 4% paraformaldehyde in 0.1M phosphate buffer (PB) (approximately 1 l/kg of each solution). Brains were removed from the skull and post-fixed overnight in 4% paraformaldehyde in 0.1M PB, and then allowed to equilibrate in 30% sucrose in PB. The brains were blocked and the diencephalon frozen and sectioned coronally (50 µm section thickness). Tissue from rhythmic and arrhythmic chronotypes of all three species of mole rat in the present study was processed concurrently. A one in two series of stains was made for Nissl and orexin-A immunohistochemistry. Sections used for the Nissl series were mounted on 0.5% gelatine coated glass slides, cleared in a solution of 1:1 chloroform and absolute alcohol, then stained with 1% cresyl violet.

For immunohistochemical staining the sections were first treated for 30 min with an endogenous peroxidase inhibitor (49.2% methanol: 49.2% of 0.1PB: 1.6% of 30% H2O2) followed by three 10 min rinses in 0.1M PB. This was followed by a 2 h pre-incubation, at room temperature, in a solution (blocking buffer) containing 3% normal goat serum (NGS, Chemicon), 2% bovine serum albumin (BSA, Sigma), and 0.25% Triton X-100 (Merck) in 0.1M PB. The sections were then incubated in a primary antibody solution containing the appropriately diluted antibody in blocking buffer for 48 h at 4°C under gentle agitation. To reveal orexinergic neurons, we used the anti-orexin-A antibody AB3704 (Chemicon, raised in rabbit) at a dilution of 1:1500. This step was followed by three 10 min rinses in 0.1M PB, after which the sections were incubated in a secondary antibody solution for 2 h. The secondary antibody solution contained a 1:1000 dilution of biotinylated anti-rabbit IgG (BA-1000, Vector Labs) in 3% NGS and 2% BSA in 0.1M PB. After three 10 min rinses in 0.1M PB, the sections were incubated for 1 h in AB solution (Vector Labs) and again rinsed three times. The sections were then treated in a solution of 0.05% diaminobenzidine in 0.1M PB for 5 min, following which 3µl of 30% H2O2 was added to the 1 ml of solution in which each section was immersed. Staining development was monitored visually and checked under a low power stereomicroscope. Development was allowed to continue until the immunopositive neurons became readily identifiable from the background stain. Development was arrested by placing the sections in 0.1M PB, and then rinsed twice more in the same solution. Sections were mounted on glass slides coated with 0.5% gelatine and left to dry overnight. They were then dehydrated in a graded series of alcohols, cleared in xylene, and coverslipped with Depex. Two controls were employed in the immunohistochemistry, including the omission of
the primary antibody and the omission of the secondary antibody in selected sections, which eliminated staining.

2.3 Quantitative analysis

The number of orexinergic (Orx+) positive cells was determined with stereological techniques on a one in two series of sections through the complete hypothalamus in all 18 brains. All analyses were done blind to the rhythmicity status of the individual. A Nikon E600 microscope with three axis motorized stage, video camera, Neurolucida interface and Stereo-Investigator software (MicroBrightfield Corp., Colchester, Vermont) was used for the stereological counts. In an attempt to achieve the most accurate estimation of Orx+ neurons, a pilot study was first conducted in an individual of each species. The pilot study determined the best counting frame size and grid size and these parameters were then used for all individuals of each species investigated. A 275 x 185 µm counting frame and a 300 x 400 µm sampling grid were employed in each individual. Only orexinergic neurons with clearly visible nuclei were marked in the sampling grids (Table 1). For the calculation of total neurons numbers, we measured section thickness in a random sample of 50 sections from each species in the regions where orexinergic neurons were present and used these measurements to calculate the species average mounted thickness. The ‘optical fractionator probe’ function of the software computationally determined the total number of Orx+ cell in the hypothalamus of each individual using the following formula:

\[ N = Q/(SSF \times ASF \times TSF) \]

Where: N – was the total estimated neuronal number, Q – was the number of neurons counted, SSF – was the section sampling fraction (in the current study this was 0.5), ASF – is the area sub fraction (this was the ratio of the size of the counting frame to the size of the sampling grid), and TSF – was the thickness sub fraction (this was the ratio of the dissector height relative to 50 µm). In order to determine TSF we used the average mounted section thickness calculated for each species (as described above, Table 1), subtracted the total vertical guard zones (10 µm) to give dissector height and used the ratio of dissector height to cut section thickness (50 µm) to provide TSF for each species and applied this to all individuals within a species.

A function in the stereology programme called the “nucleator probe” facilitated the
estimation of the mean cross-sectional area, volume and length of the orexinergic positive
cells. Only neurons with a distinct nucleus were chosen for analysis. The “nucleator probe”
was employed in conjunction with the optical fractionator and stereology procedures for
systematic random sampling to identify cells (Gunderson, 1988). In total, seven probes were
used in the current study namely: the optical fractionators, optical fractionator using number
weighted section thickness, physical dissector, physical fractionator, Schmitz nearest
neighbour, Cavalieri estimator for area and volume, and combination of planes with the
optical fractionator for absolute length.

2.4 Statistical analysis

In addition the quantitative values generated using stereological techniques; data for
body mass ($M_b$), brain mass ($M_{br}$) and encephalisation quotient ($EQ$) were compiled for
analysis. $EQ$ in each individual was determined using the formula (from Manger, 2006):

$$EQ = M_{br}/0.069M_b^{0.718}$$

Comparisons were conducted between species (CHP vs. FA, CHP vs. FD and FA vs.
FD) and within species (CHP rhythmic vs. CHP arrhythmic, FA rhythmic vs. FA arrhythmic,
and FD rhythmic vs. FD arrhythmic). These comparisons were made for $M_b$, $M_{br}$, $EQ$, number
of Orx+ cell bodies (counts), estimated volume of an Orx+ cell body, estimated cross-
sectional area of an Orx+ cell body and estimated length of an Orx+ cell body. The values
obtained for the counts, volume, area and length of Orx+ cell bodies in each mole rat were
divided by the $Mb$ of each individual and the $M_{br}$ of each individual thereby creating an index
that corrected for $M_b$ and $M_{br}$. These indices were statistically analyzed between species and
within species for the counts, volume, area and length. The indices also facilitated
comparisons between all rhythmic individuals and all arrhythmic individuals. All data below
are presented as the mean with its standard error. All statistical tests were performed using
PAST (PAleontological Statistics, v2.02; Hammer et al., 2001). Initially the Kruskal-Wallis
test was used for the aforementioned comparisons. When significance was observed ($p<0.05$),
multiple Bonferroni-corrected Mann-Whitney U tests were used as a post-hoc to the Kruskal-
Wallis for the detection of specific differences. p-values for statistically significant
differences (again where $p<0.05$) are provided in the results
3. Results

In all three species of mole rat, orexinergic immunopositive (Orx+) cell bodies were located within the hypothalamus, as previously reported in all mammals studied to date, including mole rats (Bhagwandin et al., 2011a). Stereological counts of Orx+ cell bodies were found to be statistically significantly different only between the distinct circadian chronotypes of Ansel’s mole rat (Fukomys anselli) and between all rhythmic and arrhythmic individuals of the current study when the counts of Orx+ cell bodies were corrected for body mass ($M_b$). Estimation of cell volume, cross-sectional area and length showed statistically significant differences between species without correction for $M_b$ and brain mass ($M_{br}$) and statistically significant differences between species when these variables were corrected for $M_b$ and $M_{br}$ though no statistically significant differences were reached for these parameters within species when similar comparisons were made.

3.1 Orexinergic cell body distribution

All three species of mole rat expressed Orx+ neurons only within the hypothalamus and were observed as sharing common neuronal locality within the lateral hypothalamic area (LHA), perifornical region (PFR) and the lateral ventral hypothalamic supraoptic area (LVHA), similar to those reported in two other species of mole rat (Bhagwandin et al., 2011a). Thus, there appears to be three distinct clusters of Orx+ neurons in the hypothalamus of the mole rats studied, a large homogeneous cluster spanning the lateral and perifornical regions (the main cluster), a distinct cluster extending into the region of the zona incerta (the zona incerta cluster), and a final cluster in the ventral lateral hypothalamus adjacent to the diminutive optic tracts (the optic tract cluster). The mole rats, from both chronotypes and all three species, exhibited neuronal cell bodies that were morphologically homogenous in all three clusters, that were ovoid in shape, a varying mixture of bi- and multi-polar types and that showed no specific dendritic orientation (Figs. 2, 3).

3.2 Comparisons of body mass ($M_b$), brain mass ($M_{br}$) and encephalisation quotient (EQ)
Mean $M_b$ in *C. hottentotus pretoriae* was 132.8 ± 22.4 g for the rhythmic group and 123 ± 17.6 g for the arrhythmic group. *F. anselli* had a mean $M_b$ of 90.6 ± 2.8 g for the rhythmic group and 78.5 ± 10.9 g for the arrhythmic group (Table 2). *F. damarensis* had a mean $M_b$ measurement of 121.9 ± 22.7 g for the rhythmic group and 93.7 ± 9.6 g for the arrhythmic group. $M_b$ comparisons between species were only statistically significant (Kruskal-Wallis p-value: 0.0064; Mann-Whitney p-value: 0.008) between *C. hottentotus pretoriae* and *F. anselli* where it was observed that *C. hottentotus pretoriae* had a higher mean $M_b$ (Table 3).

Mean $M_{br}$ in *C. hottentotus pretoriae* measured 1.4 ± 0.09 g for the rhythmic group and 1.5 ± 0.10 g for the arrhythmic group. *F. anselli* had a mean $M_{br}$ measurement of 1.2 ± 0.03 g for the rhythmic group and 1.2 ± 0.03 g for the arrhythmic group. *F. damarensis* had a mean $M_{br}$ measurement of 1.8 ± 0.11 g for the rhythmic group and 1.8 ± 0.07 g for the arrhythmic group (Table 2). Statistically significant differences were noted between the *C. hottentotus pretoriae* and *F. anselli* (where the *C. hottentotus pretoriae* had a higher mean $M_{br}$, Kruskal-Wallis p-value: 0.0039; Mann-Whitney p-value: 0.004) and *F. anselli* and *F. damarensis* (where *F. damarensis* had a higher mean $M_{br}$, Kruskal-Wallis p-value: 0.0039; Mann-Whitney p-value: 0.004) (Table 3).

Mean $EQ$ in *C. hottentotus pretoriae* was calculated as 0.6 ± 0.06 for the rhythmic group and 0.7 ± 0.10 for the arrhythmic group. *F. anselli* had a mean $EQ$ calculation of 0.7 ± 0.01 for the rhythmic group and 0.8 ± 0.09 for the arrhythmic group. *F. damarensis* had a mean $EQ$ calculation of 0.9 ± 0.09 for the rhythmic group and 1 ± 0.04 for the arrhythmic group (Table 2). Statistically significant differences were obtained when *F. damarensis* was compared to *C. hottentotus pretoriae* (where *F. damarensis* had a higher mean $EQ$, Kruskal-Wallis p-value: 0.02; Mann-Whitney p-value: 0.02) and when *F. damarensis* was compared to *F. anselli* (where the *F. damarensis* showed a higher mean $EQ$, Kruskal-Wallis p-value: 0.02; Mann-Whitney p-value: 0.02) (Table 3).

### 3.3 Stereological counts of Orx+ cell bodies

Stereological counts of Orx+ cell bodies in the *C. hottentotus pretoriae* revealed a mean of 9047 ± 2200 for the rhythmic group and 11547 ± 777 for the arrhythmic group. Counts from *F. anselli* revealed a mean of 5786 ± 80 for the rhythmic group and 9299 ± 724
for the arrhythmic group. *F. damarensis* revealed a mean of 9850 ± 1454 for the rhythmic group and 9275 ± 1518 for the arrhythmic group (Fig. 4) (Table 2).

Comparisons of Orx+ cell body counts between rhythmic and arrhythmic groups in each species revealed a statistically significant difference only between the circadian chronotypes of *F. anselli* (where the mean number of Orx+ neurons in the hypothalamus of the arrhythmic group was higher than the rhythmic group, Kruskal-Wallis p-value: 0.008; Mann-Whitney p-value: 0.009). Even though statistical significance was not reached in the other two species investigated, the mean number of Orx+ neurons in the hypothalamus of *C. hottentotus pretoriae* was higher for the arrhythmic group than the rhythmic group. For *F. damarensis* stereological counts of Orx+ cell bodies yielded similar totals for both the rhythmic and arrhythmic groups. When the counts of Orx+ cell bodies were corrected for *M*₇ and *M*₂₉, statistically significant differences were noted only between the circadian chronotypes of *F. anselli*. In both cases the arrhythmic groups had a higher mean numbers of Orx+ neurons in the hypothalamus than the rhythmic groups (Kruskal-Wallis p-value: 0.0028; Mann-Whitney p-value: 0.003). Additionally, when all rhythmic individuals of all species studied here were compared to all arrhythmic individuals, a statistically significant difference was only reached in counts that had been corrected for *M*₇ (where the arrhythmic group had a higher mean number of Orx+ neurons in the hypothalamus than the rhythmic group, Kruskal-Wallis p-value: 0.02; Mann-Whitney p-value: 0.03) (Table 3).

### 3.4 Stereological estimation of volume, area and length of Orx+ cell bodies

Stereological estimation of the volume of an Orx+ cell body in *C. hottentotus pretoriae* presented a weighted mean of 2068 ± 98 µm³ for the rhythmic group and 2281.2 ± 116.7 µm³ for the arrhythmic group. *F. anselli* presented a weighted mean of 3228.6 ± 172.2 µm³ for the rhythmic group and 3337.3 ± 128.7 µm³ for the arrhythmic group. *F. damarensis* presented a weighted mean of 2429.5 ± 116.2 µm³ for the rhythmic group and 2163.2 ± 88.9 µm³ for the arrhythmic group (Fig. 4) (Table 2). Statistically significant differences were noted: (1) between species (where *F. anselli* showed a higher mean volume of an Orx+ cell body than the other two species, Kruskal-Wallis p-value: 0.003; Mann-Whitney p-value: 0.0005); (2) when volume was corrected for *M*₇ between species (where *F. anselli* had a higher mean volume of an Orx+ cell body than *C. hottentotus pretoriae*, Kruskal-Wallis p-value: 0.004; Mann-Whitney p-value: 0.0008), and where *F. anselli* had a higher mean
volume of an Orx+ cell body than *F. damarensis* (Kruskal-Wallis p-value: 0.004; Mann-Whitney p-value: 0.008); and (3) when volume was corrected for *M*, between species (where *F. anelli* had a higher mean volume of an Orx+ cell body than the other two species, Kruskal-Wallis p-value: 0.003; Mann-Whitney p-value: 0.005, and where *C. hottentotus pretoriae* had a higher mean volume of an Orx+ cell body than *F. damarensis*, Kruskal-Wallis p-value: 0.03; Mann-Whitney p-value: 0.03) (Table 3).

Stereological estimation of the cross-sectional area of an Orx+ cell body in the *C. hottentotus pretoriae* revealed a weighted mean of 180.9 ± 4.8 µm² for the rhythmic group and 193.3 ± 5.9 µm² for the arrhythmic group. *F. anelli* revealed a weighted mean of 247.7 ± 8.1 µm² for the rhythmic group and 250.7 ± 5.8 µm² for the arrhythmic group. *F. damarensis* revealed a weighted mean of 202.4 ± 5.4 µm² for the rhythmic group and 189.3 ± 4.9 µm² for the arrhythmic group (Fig.4) (Table 2). Statistically significant differences were noted: (1) between species (where *F. anelli* showed a higher cross-sectional area of an Orx+ cell body than the other two species, Kruskal-Wallis p-value: 0.003; Mann-Whitney p-value: 0.005); (2) when cross-sectional area was corrected for *M*, between species (where *F. anelli* had a higher cross-sectional area of an Orx+ cell body than *C. hottentotus pretoriae*, Kruskal-Wallis p-value: 0.004; Mann-Whitney p-value: 0.005, and where *F. anelli* had a higher cross-sectional area of an Orx+ cell body than *F. damarensis*, Kruskal-Wallis p-value: 0.01; Mann-Whitney p-value: 0.01); and (3) when area was corrected for *M*, between species (where *F. anelli* had a higher cross-sectional area of an Orx+ cell body than the other two species, Kruskal-Wallis p-value: 0.003; Mann-Whitney p-value: 0.005, and where *C. hottentotus pretoriae* had a higher cross-sectional area of an Orx+ cell body than *F. damarensis*, Kruskal-Wallis p-value: 0.04; Mann-Whitney p-value: 0.03) (Table 3).

Stereological estimation of the length of an Orx+ cell body in *C. hottentotus pretoriae* revealed a weighted mean of 7.3 ± 0.09 µm for the rhythmic group and 7.5 ± 0.11 µm for the arrhythmic group. *F. anelli* revealed a weighted mean of 8.6 ± 0.14 µm for both rhythmic and arrhythmic groups. *F. damarensis* revealed a weighted mean of 7.8 ± 0.09 µm for the rhythmic group and 7.5 ± 0.10 µm for the arrhythmic group (Fig. 4) (Table 2). Statistically significant differences were noted: (1) between species (where *F. anelli* showed a longer length of an Orx+ cell body than the other two species, Kruskal-Wallis p-value: 0.003; Mann-Whitney p-value: 0.005); (2) when length was corrected for *M*, between species (where *F. anelli* had a longer length of an Orx+ cell body than *C. hottentotus pretoriae*, Kruskal-Wallis p-value: 0.004; Mann-Whitney p-value: 0.005); and (3) when volume was corrected for *M*.
between species (where *F. anselli* had longer length of an Orx+ cell body than the other two species, Kruskal-Wallis p-value: 0.001; Mann-Whitney p-value: 0.005, and where *C. hottentotus pretoriae* had a longer length of an Orx+ cell body *F. damarensis*, Kruskal-Wallis p-value: 0.02; Mann-Whitney p-value: 0.03) (Table 3).

4. Discussion

The initial aim of the current study was to determine whether rhythmic and arrhythmic chronotypes had different numbers of hypothalamic orexin immunopositive neurons with the hypothesis of elevated hypothalamic orexinergic neurons in the arrhythmic chronotypes as a result of the involvement of orexins in the sleep-wake cycle. The results of the current study indicated statistically significant differences between the circadian chronotypes of *F. anselli*, where the arrhythmic group had higher mean numbers of hypothalamic orexin neurons compared to the rhythmic group. No statistically significant differences in orexinergic neuronal numbers were observed in the other two species studied. A statistically significant difference was noted between all rhythmic and arrhythmic individuals of the current study when the counts of orexin neurons were corrected for *M*.*

4.1 Orexins, sleep-wake cycle and circadian rhythmicity

The results of the present study indicated statistically significant differences between circadian chronotypes of *F. anselli* for the counts of hypothalamic Orx+ neurons when the raw data was compared. In this case the arrhythmic chronotype consistently displayed higher mean counts of hypothalamic Orx+ neurons than the rhythmic chronotype. A similar tendency, though not statistically significantly, was observed when the raw data was compared between the chronotypes of *C. hottentotus pretoriae*; however, a similar comparison between chronotypes of *F. damarensis* showed that the mean count of Orx+ neurons in the rhythmic group was higher than the rhythmic group, but statistical significance was not reached. The analysis of *F. anselli* supports the hypothesis of elevated Orx+ neurons in the hypothalamus. Given the previous suggestions for the role of orexins in the sleep-wake cycle (Siegel, 2004; Baumann and Bassetti, 2005), the results of the current study indicate that the arrhythmic chronotypes of *F. anselli* and possibly *C. hottentotus pretoriae* would theoretically have an enhanced maintenance of waking (despite the reduced motor activity of
the arrhythmic chronotype compared to the rhythmic, as indicated by their actograms, with a net result that possibly enhances vigilance during quiet waking) and a reduced amount of time spent in NREM relative to the rhythmic chronotypes. The study of sleep in rhythmic and arrhythmic chronotypes of the Giant Zambian mole rat (*Fukomys mechowii*) (Bhagwandin et al., 2011b) has shown that the arrhythmic group statistically spent more time awake and spent less time in NREM compared to the rhythmic group.

It is known that the bifurcated ascending arousal system incorporates firstly, a pathway to the thalamus which excites thalamic relay neurons that transmit information to the cerebral cortex and secondly, a branch that bypasses the thalamus to activate neurons in the lateral hypothalamic area, basal forebrain and cerebral cortex (Saper, 1985; Hallanger et al., 1987; Saper et al., 2001, 2005; Jones, 2003). Orexinergic neurons, with cell bodies restricted to the hypothalamus, have ascending projections to the cerebral cortex and descending projections to the cholinergic (basal forebrain, lateral dorsal tegmental nucleus, pedunculopontine nucleus) and monoaminergic (locus coeruleus, ventral tegmentum area, raphe) nuclear groups of arousal systems (Peyron et al., 1998; Chemelli et al., 1999; Baumann and Bassetti, 2005; Bhagwandin et al., 2011a). The exact role of orexins in REM is unclear as some studies suggest increased orexinergic activity during REM while others report the contrary (Less et al., 2005; Mileykovskiy et al., 2005); however, a recent study has postulated that orexin neurons mediate the circadian timing of REM by suppressing REM during the active period (Kantor et al., 2009). In light of the results of the current study it would appear that the increased numbers of hypothalamic orexinergic neurons observed within *F. anselli* arrhythmic chronotypes, and potentially *C. hottentotus pretoriae*, through possible increased activity and the resulting effect on wakefulness and motor activities, may facilitate a higher level of vigilance allowing for extended foraging and feeding activity.

Baumann and Bassetti (2005) proposed that orexins are under control of the suprachiasmatic nucleus (SCN) and that orexins do not significantly affect circadian rhythmicity. Saper (2005) described a 3 stage circuitry integrator for circadian rhythms that involve a pathway from the SCN to the subparaventricular zone (SPZ) and finally to dorsomedial nucleus of the hypothalamus (DMH). The DMH communicates directly with the lateral hypothalamic area, the region with the highest population of orexinergic neurons, and the ventrolateral preoptic nucleus (VLPO), the area primarily concerned with the promotion of slow wave sleep (SWS). The rationale behind the 3 stage pathway lies in the observation that in both diurnal and nocturnal animals, the SCN is primarily active during the light period...
and the VLPO during the dark period (Sherin et al., 1996; Gaus et al., 2002). Therefore, in animals that are nocturnal vs. diurnal, there should be an intervening circuitry that allows the circadian cycle to be set at opposite phases (Saper, 2005). It is well known that mole rats have unusual patterns of circadian rhythmicity (Oosthuizen et al., 2003; Lovegrove and Muir, 1996; Lovegrove and Papenfus, 1995) and that mole rats have different neuropeptide populations in the SCN compared to other rodents (Negroni, 2003). Therefore, the increased counts of Orx+ neurons in the hypothalamus within F. anselli arrhythmic chronotypes, and potentially C. hottentotus pretoriae, may alter the circadian control of wakefulness in these individuals.

4.2 Orexins and appetite regulation

The results of the present study indicated statistically significant differences between circadian chronotypes of F. anselli when the raw data for counts of Orx+ neurons were corrected for $\textit{M}_{b}$ and $\textit{M}_{br}$, where the arrhythmic groups showed higher mean counts than the rhythmic groups. This can be interpreted as the arrhythmic chronotype having a higher number of Orx+ neurons per gram of $\textit{M}_{b}$ or $\textit{M}_{br}$ compared to the rhythmic chronotype. Since the initial demonstration of orexin induced hyperphagia in rats (Sakurai et al., 1998), orexins have become synonymous with mechanisms of appetite regulation and increased food intake that is driven by orexin involvement in motivated behavior (Rodgers et al., 2002). Cai et al. (2001) have reported activation of the lateral hypothalamic orexin neurons in response to low blood glucose levels and an empty stomach, signals that indicate replenishment, and concluded that this context-dependency explains the effects associated with orexin activation – increased food intake with delayed onset of satiety and increased wakefulness with vigilance and behavioral activity. It is also known that orexins are involved in appetite regulation through activation of the leptin sensitive neurons of the arcuate nucleus and its subsequent feeding associated signaling to orexinergic neurons of the lateral hypothalamus (LH) and henceforth interaction between LH and the hypothalamic ventro-medial nucleus (VMH) (Stellar, 1954; Rodgers et al., 2002; Sakurai, 2003, 2005). Therefore it would seem fair to infer that the elevated counts of hypothalamic orexinergic neurons in the arrhythmic chronotype of F. anselli may yield a greater drive for motivated behavior and theoretically a higher food intake compared to the rhythmic chronotype.
Interestingly though, the results of the present study showed a statistically significant difference for the counts of Orx+ neurons that were corrected for $M_b$ when all rhythmic individuals were compared to all arrhythmic individuals from all three species of mole rats examined. This indicated that the arrhythmic individuals have a higher number of Orx+ neurons per gram of $M_b$. Given the effect of orexins on motivated behavior and food intake, this may be indicative of a higher body mass in the arrhythmic individuals compared to the arrhythmic individuals. This was not the case as it was observed in the present study that the mean body mass of the arrhythmic chronotype in each species investigated was lower than that of the rhythmic chronotype. A possible explanation for this could lie in the assessment of the impact of chronic administration of orexin-A on food intake on rats (Haynes et al., 1999; Yamanaka et al., 1999). Both these studies indicated an increased food intake during the light period and a compensatory reduction of food intake during the dark period, and both agree that chronic administration of orexin-A does not alter 24 h food intake or body mass gain. In addition, Lubkin and Stricker-Krongrad (1998) postulated that orexins may increase overall metabolic rate to support increased motor activity; however, Nicolaidis (2006) suggests that overall metabolic rate be divided into resting metabolism and locomotion-related metabolism and based on resting metabolism, a hypometabolic but physically active animal has a higher (total) metabolism than a hypermetabolic animal that remains quiet. This concept supports the observations made in the mole rats assessed in the current study as it has been previously reported that subterranean mole rats have lower resting metabolic rates than surface dwelling mammals (Bennett and Faulkes, 2000; Zelova et al., 2007, 2009). Furthermore, it has been suggested that sleep duration tracks the metabolism of nutrients in a meal and the size of a meal, with the effect that a higher nutritional content and larger meal size produces a longer sleep duration (Danguir and Nicolaidis, 1980a,b). Despite this, brain metabolism is conserved and independent from that of the periphery, the ventromedial nucleus, dorsomedial nucleus and paraventricular nucleus, metabolic strategic areas, are capable of reflecting metabolic changes resulting from bodily depletion and repletion (Nicolaidis, 2006). Therefore, it may be possible that a higher peripheral metabolism may result in a lower body mass despite the elevated Orx+ neurons observed in the arrhythmic chronotype compared to the rhythmic chronotype.

References


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Table Legends

Table 1: Stereological parameters used for rhythmic and arrhythmic individuals of *C. hottentotus pretoriae*, *F. anselli* and *F. damarensis*.

Table 2: Table presenting data for rhythmic and arrhythmic individuals of *C. hottentotus pretoriae*, *F. anselli* and *F. damarensis* and the respective weighted means for: $M_{br}$, $M_b$, $EQ$ estimated cell counts, average estimated volume, average estimated cell area, and average estimated cell length.

Table 3: Table showing results of statistical comparisons between various data sets. (*) denotes statistical significance as determined using the post hoc Mann-Whitney U and Bonferroni correction factor when $p<0.05$ (p-values included in text). Letters in subscript indicate the species or chronotype of mole rat where the higher mean was observed. CHP = *C. hottentotus pretoriae*, FA = *F. anselli*, FD = *F. damarensis*, and AR = arrhythmic.

Figure Legends

Figure 1: Actigrams illustrating circadian patterns of locomotor activity in rhythmic and arrhythmic individuals from *C. hottentotus pretoriae*, *F. anselli* and *F. damarensis*. These actigrams indicate that the rhythmic mole rats have a predictable period of activity during the light period whereas locomotor activity in the arrhythmic animals is irregular.

Figure 2: Photomicrographs showing the orexin-A immunoreactive neurons within the hypothalamus of rhythmic and arrhythmic chronotypes of *C. hottentotus pretoriae* (A, B), *F. anselli* (C, D) and *F. damarensis* (E, F). Scale = 1 mm and applies to all.

Figure 3: High power photomicrographs showing the morphology of orexin-A immunoreactive neurons within the hypothalamus of rhythmic and arrhythmic chronotypes of *C. hottentotus pretoriae* (A, B), *F. anselli* (C, D) and *F. damarensis* (E, F). Scale = 100 µm and applies to all.

Figure 4: Graphs showing the various parameters of stereological data for arrhythmic and arrhythmic individuals (grey bars) of *C. hottentotus pretoriae*, *F. anselli* and *F. damarensis* and the respective weighted means (black bars). The graphs indicate average estimated values for somal number, volume, area and length.
Figure 1

Cryptomys hottentotus  Fukomys anselli  Fukomys damarensis

Rhythmic  Arrhythmic
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<th>Sampling grid size (µm)</th>
<th>Cut thickness (µm)</th>
<th>Average species mounted thickness (µm)</th>
<th>Vertical guard zones (top and bottom, µm)</th>
<th>Section interval</th>
<th>Number of sampling sites</th>
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|                      |           |            |    |        |        |      |        |
| Between Species Corrected for Body Mass | Counts | Volume | Area | Length |
| CHP vs.. FA          | -        | *_FA     | *_FA| *_FA   |
| CHP vs.. FD          | -        | -        | -   | -      |
| FA vs.. FD           | -        | *_FA     | *_FA| -      |
| Within species Corrected for Body Mass | CHP R vs.. NR | - | - | - | - |
| FA R vs.. NR         | *_AR     | -        | -   | -      |
| FD R vs.. NR         | -        | -        | -   | -      |

|                      |           |            |    |        |        |      |        |
| Between species Corrected for Brain Mass | Counts | Volume | Area | Length |
| CHP vs.. FA          | -        | *_FA     | *_FA| *_FA   |
| CHP vs.. FD          | -        | *_CHP    | *_CHP| *_CHP  |
| FA vs.. FD           | -        | *_FA     | *_FA| *_FA   |
| Within species Corrected for Brain Mass | CHP R vs.. NR | - | - | - | - |
| FA R vs.. NR         | *_AR     | -        | -   | -      |
| FD R vs.. NR         | -        | -        | -   | -      |

|                      |           |            |    |        |        |      |        |
| All R vs. NR Corrected for Body Mass | EQ | Counts | Volume | Area | Length |
| All R vs. NR         | -        | *_AR     | -    | -     | -     |

|                      |           |            |    |        |        |      |        |
| All R vs. NR Corrected for Brain Mass | EQ | Counts | Volume | Area | Length |
| All R vs. NR         | -        | -        | -    | -     | -     |