Oxytocin and Vasopressin Receptors in the Pouched Rat

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5 Abstract

This is the abstract. It consists of two paragraphs.

6 Introduction

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- The neuropeptides, oxytocin (OT) and vasopressin (VP), have receptor distributions in the brain that can modulate a variety of social behaviors such as parental care, affiliation, and aggression, among other behaviors (Caldwell and Albers, 2015). The densities of their associated receptors, oxytocin (OTR) and vasopressin (V1aR and V1bR) receptors, are often species- and sex-dependent.
- What do we know about the distribution and relative densities of receptors (i.e. what can it tell us, what's been done on behavior?)
 - -How might life history differences play into patterning of central distribution of these receptors?
- -What tends to be conserved?
- -How does comparative analysis help? (Kelly and Ophir, 2015)
 - -Why did we do this study? What were we examining?
- -We wanted to describe the presence and relative density of OTR and V1aR in pouched rat brains in males and females, to see if there were differences in presence and density between sexes
- -We wanted to explore how the patterning of these receptors might differ from other rodents and see if
 it further supports the ideas found in the recent metaanalysis (Freeman et al., 2020)(see where pouched
 rats fall in this framework)

22 Methods

All work with animals was approved under the U.S. Army Medical Research and Materiel Command (USAMRMC) Animal Care and Use Review Office (ACURO) and the Cornell Institutional Animal Care and Use
Committee (IACUC 2014-0043). All tissues were collected from wild-caught animals from Morogoro, Tanzania
(6°49'49"S, 37°40'14"E). Prior to collection, animals were housed individually in standard rabbit enclosures and

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maintained on a 12:12 h light:dark light cycle, at 21°C and 45% humidity. Males and females were kept in separate rooms. Animals were fed a standard rodent diet supplemented with dog kibble and fresh fruit and vegetable treats.

Chewing bones, a metal 'stovepipe' hutch, and dog puzzle toys were given as behavioral enrichment. Newspaper was given for nesting material.

Animals were euthanized via CO₂ inhalation and brains were swiftly removed and frozen using liquid nitrogen or powdered dry ice and stored at -80°C prior to sectioning. Ten male and ten female brains were used for this study. Brains were blocked coronally by removing the cerebellum, then split sagittally next to the midline into two hemispheres, and one hemisphere (preferably the left if unblemished) was coronally sectioned at 20µm thick using a Leica cryostat (CM1950, Leica Biosystems, Nussloch, Germany) set at -20°C. Due to the large size of the pouched rat brains, we mounted every 3rd section and kept 6 serial sets. Sections were collected from the olfactory bulbs to the start of the cerebellum, and mounted on Superfrost Plus Microscope sides (Fisher Scientific, Pittsburg, PA USA).

Microscope slides were stored at -80°C until the autoradiography procedure.

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On two of the sets of slides, we used ¹²⁵I radioligands to label oxytocin receptor (ornithine vasopressin analog, ¹²⁵I-OVTA); NEX 254, PerkinElmer; Waltham, MA) or vasopressin 1a receptor (vasopressin (Linear), V-1A
antagonist (Phenylacetyl1, 0-Me-D-Tyr2 [¹²⁵I-Arg6]-); NEX 310, PerkinElmer), as described by Ophir and colleagues (Ophir et al., 2013). Following processing and air-drying, we exposed radiolabeled tissue to film (Kodak
Carestream Biomax MS) for 6 days for OTR and 2 days for V1aR to account for differing degrees of decay at the
time of use. In each film cassette, we included two ¹²⁵I microscales (American Radiolabeled Chemicals; St Louis,
MO), to allow for the conversion of optical density to receptor density. We inferred that receptor density relates
to optical density of exposed film to use optical measurements as a proxy for receptor density. We digitized films
on a Microtek ArtixScan M1 (Microtek, Santa Fe Springs, CA) and measured optical densities using NIH ImageJ
Software. We calculated receptor density by first converting optical density to disintegrations per minute (dpm),
adjusted for tissue equivalence (TE; for 1 mg in the rat brain), by fitting curves generated by radiographic standards
and extrapolating based on these standard curves for each film.

Three sequential sections were measured for density by encircling the regions of interest using NIH ImageJ software. The software program calculated mean optical density values and area for regions of interest (ROI). We measured background labelling by measuring optical density from an area of cortex in the same section for each region of interest. To correctly identify ROI, we Nissl-stained a third set of tissue to use as a reference, in conjunction with anatomical landmarks identified using a Rat brain atlas. The three measurements for each individual's ROIs and background were averaged separately, and background was subtracted from the ROI to yield receptor density.

These final measurements were used for all statistical tests, tables, and figures.

OTR density was measured in the olfactory bulb (OB), anterior olfactory nucleus (AON), prefrontal cortex (PFC), piriform cortex (Pir), nucleus accumbens (NAcc), lateral septum (LS), endopiriform (Den), claustrum

(VCL), lateral bed nucleus of the stria terminalis (BSTI), medial bed nucleus of the stria terminalis (BSTm), ventral pallidum (VPall), medial preoptic area (MPOA), anterior hypothalamus (AH), paraventricular thalamus (PVT), suprachiasmatic nucleus (SCN), paraventricular nucleus (PVN), magnocellular hypothalamic nucleus (MCPO), medial habenula (MHb), central amygdala (CeA), medial amygdala (MeA), basolateral amygdala (BLA), ventromedial hypothalamus (VMH), zona incerta (ZIR), lateral hypothalamus (PrFLH), hippocampal CA1, hippocampal CA2, hippocampal CA3, dentate gyrus (DG), premammilary ventral nucleus (PMV), ventral tegmental area (VTA), periaqueductal gray (PAG), medial geniculate (MG), superior colliculus (SuG), and the ventral CA3. V1aR was measured in the same regions except for the MCPO and the MHb.

To calculate 'relative binding' on a 4-point scale, we used the following definitions: mean OTR < 35 dpm/mg: absent/-, 35 to 490: present/+, 490 to 945: moderate/++, 945 to 1400: dense/+++; mean V1aR < 100 dpm/mg: absent/-, 100-1367: present/+, 1367 to 2634: moderate/++, 2634 to 3800: dense/+++. To compare receptor densities between sexes, we conducted t-tests for each region. We used a bonferroni correction to adjust for multiple comparisons. Test statistics were considered significant when p < 0.05.

We used the framework from Freeman et al. (2020) to compare pouched rat overall binding patterns to those in other rodents (Freeman et al., 2020). This framework uses overall OTR and V1aR binding patterns to examine similarities among species, genera, and family groups. Briefly, relative binding patterns within a species are converted to a 4-point scale, using wording and data from previously published studies. These data are then used in a principal components analysis, and species are plotted along the PC1 and PC2 components, with vectors in the plot serving as weights of each variable, and the direction indicating loading on PC1 and PC2.

We mapped pouched rat relative binding data onto this plot based on this framework and the data presented in this study. In addition to superimposing the pouched rat onto the PCA biplot, we recreated the comparative permutational anova (Adonis function) to examine whether genus or family groups predicted similarities among species' relative binding patterns. All analyses were conducted in R 4.0.2, with the vegan package for the 'adonis' function, and stats package for t tests and principal components analysis (R Development Core Team, 2016). PCA biplots were made using the ggbiplot function in the ggbiplot package with some aesthetic changes.

Results

After comparing male and female densities in the measured regions, most regions showed no differences between sexes. The superior colliculus had higher densities of OTR in females compared to males (Table 1, Female mean: 199.35, Male mean 51.46, $t_{(13.58)} = 2.79$, p = 0.01), however, this was no longer significant after corrections for multiple comparisons (adjusted $\alpha = 0.001$).

Table 1: OTR densities by region and sex

Region	Sex	Density mean \pm SE (dpm/mg)	N
Olfactory Bulb	F	126.27 ± 39.56	9
Olfactory Bulb	M	142.74 ± 39.34	9
Accessory Olfactory Nucleus	\mathbf{F}	186.38 ± 40.35	9
Accessory Olfactory Nucleus	\mathbf{M}	144.1 ± 30.98	9
mPFC	\mathbf{F}	117.96 ± 36.91	9
mPFC	\mathbf{M}	94.51 ± 32.08	9
Infralimbic Cortex	\mathbf{F}	138.22 ± 43.25	9
Infralimbic Cortex	M	72.54 ± 20.32	9
Nac Core	\mathbf{F}	46.18 ± 14.48	9
Nac Core	M	37.34 ± 12.45	9
Nac Shell	\mathbf{F}	81.44 ± 17.96	9
Nac Shell	M	51.75 ± 13.37	9
Caudate Putamen	\mathbf{F}	30.8 ± 13.39	9
Caudate Putamen	M	23.93 ± 8.42	9
Piriform Cortex	\mathbf{F}	222.05 ± 39.71	9
Piriform Cortex	M	192.1 ± 35.8	9
Lateral Septum I	\mathbf{F}	55.24 ± 12.25	11
Lateral Septum I	M	57.94 ± 13.57	9
Lateral Septum D	\mathbf{F}	48.7 ± 7.59	11
Lateral Septum D	M	56.22 ± 12.33	9
Lateral Septum V	\mathbf{F}	71.16 ± 16.63	11
Lateral Septum V	M	103.53 ± 35.63	9
Endorpiriform Cortex	\mathbf{F}	147.07 ± 22.53	10
Endorpiriform Cortex	M	111.82 ± 19.71	9
Claustrum	\mathbf{F}	192.58 ± 47.69	10
Claustrum	M	193.17 ± 28.09	9
BSTm	F	585.91 ± 117.64	10
BSTm	M	532.58 ± 91.23	9
BSTi	\mathbf{F}	687.03 ± 132.48	10
BSTi	\mathbf{M}	425.66 ± 79.05	9
BSTv	F	101.57 ± 21.35	10

Ventral Pallidum F 107.32 ± 15.18 10 Ventral Pallidum M 78.04 ± 23.15 9 MPOA F 95.77 ± 19.61 10 MPOA M 88.22 ± 17.61 9 Anterior Hypothalamus F 90.27 ± 24.22 10 Anterior Hypothalamus M 74.19 ± 17.25 7 PVT F 50.91 ± 16.77 10 PVT F 50.91 ± 16.77 10 PVT M 64.52 ± 27.91 7 SCN F 80.48 ± 23.44 10 SCN F 14.44 ± 16.18 9 PVN F 14.44 ± 16.18 9 PVN B 118.68 ± 37.66 9 Medial Hypothalamic Nucleus M <th>Region</th> <th>Sex</th> <th>Density mean \pm SE (dpm/mg)</th> <th>N</th>	Region	Sex	Density mean \pm SE (dpm/mg)	N
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Medial Amygdala M 380.99 ± 82.29 9 Basolateral Amygdala F 627.27 ± 120.17 9 Basolateral Amygdala M 712.9 ± 155.24 9 VMH F 1242.79 ± 245.41 9 VMH M 1224.42 ± 165.26 9 Zona Incerta F 408.51 ± 67.31 10 Zona Incerta M 415.41 ± 58.07 9 Lateral Hypothalamus F 176.03 ± 32.85 10 Lateral Hypothalamus M 171.5 ± 41.95 9 CA1 F 121.81 ± 25.41 11 CA1 M 107.19 ± 15.49 9	Central Amygdala	M	734.04 ± 92.77	9
Basolateral Amygdala F 627.27 ± 120.17 9 Basolateral Amygdala M 712.9 ± 155.24 9 VMH F 1242.79 ± 245.41 9 VMH M 1224.42 ± 165.26 9 Zona Incerta F 408.51 ± 67.31 10 Zona Incerta M 415.41 ± 58.07 9 Lateral Hypothalamus F 176.03 ± 32.85 10 Lateral Hypothalamus M 171.5 ± 41.95 9 CA1 F 121.81 ± 25.41 11 CA1 M 107.19 ± 15.49 9	Medial Amygdala	F	328.84 ± 51.14	9
Basolateral Amygdala M $712.9 \pm 155.24 = 9$ VMH F $1242.79 \pm 245.41 = 9$ VMH M $1224.42 \pm 165.26 = 9$ Zona Incerta F $408.51 \pm 67.31 = 10$ Zona Incerta M $415.41 \pm 58.07 = 9$ Lateral Hypothalamus F $176.03 \pm 32.85 = 10$ Lateral Hypothalamus M $171.5 \pm 41.95 = 9$ CA1 F $121.81 \pm 25.41 = 11$ CA1 M $107.19 \pm 15.49 = 9$	Medial Amygdala	M	380.99 ± 82.29	9
VMH F 1242.79 ± 245.41 9 VMH M 1224.42 ± 165.26 9 Zona Incerta F 408.51 ± 67.31 10 Zona Incerta M 415.41 ± 58.07 9 Lateral Hypothalamus F 176.03 ± 32.85 10 Lateral Hypothalamus M 171.5 ± 41.95 9 CA1 F 121.81 ± 25.41 11 CA1 M 107.19 ± 15.49 9	Basolateral Amygdala	F	627.27 ± 120.17	9
VMH M 1224.42 ± 165.26 9 Zona Incerta F 408.51 ± 67.31 10 Zona Incerta M 415.41 ± 58.07 9 Lateral Hypothalamus F 176.03 ± 32.85 10 Lateral Hypothalamus M 171.5 ± 41.95 9 CA1 F 121.81 ± 25.41 11 CA1 M 107.19 ± 15.49 9	Basolateral Amygdala	M	712.9 ± 155.24	9
Zona Incerta F 408.51 ± 67.31 10 Zona Incerta M 415.41 ± 58.07 9 Lateral Hypothalamus F 176.03 ± 32.85 10 Lateral Hypothalamus M 171.5 ± 41.95 9 CA1 F 121.81 ± 25.41 11 CA1 M 107.19 ± 15.49 9	VMH	F	1242.79 ± 245.41	9
Zona Incerta M 415.41 ± 58.07 9 Lateral Hypothalamus F 176.03 ± 32.85 10 Lateral Hypothalamus M 171.5 ± 41.95 9 CA1 F 121.81 ± 25.41 11 CA1 M 107.19 ± 15.49 9	VMH	M	1224.42 ± 165.26	9
Lateral Hypothalamus F 176.03 ± 32.85 10 Lateral Hypothalamus M 171.5 ± 41.95 9 CA1 F 121.81 ± 25.41 11 CA1 M 107.19 ± 15.49 9	Zona Incerta	F	408.51 ± 67.31	10
Lateral Hypothalamus M 171.5 ± 41.95 9 CA1 F 121.81 ± 25.41 11 CA1 M 107.19 ± 15.49 9	Zona Incerta	M	415.41 ± 58.07	9
CA1 F 121.81 ± 25.41 11 CA1 M 107.19 ± 15.49 9	Lateral Hypothalamus	F	176.03 ± 32.85	10
CA1 M 107.19 ± 15.49 9	Lateral Hypothalamus	M	171.5 ± 41.95	9
	CA1	F	121.81 ± 25.41	11
CA2 F 106.13 ± 11.57 11	CA1	M	107.19 ± 15.49	9
	CA2	F	106.13 ± 11.57	11

Region	Sex	Density mean \pm SE (dpm/mg)	N
CA2	M	127.54 ± 25.3	9
CA3	F	39.48 ± 10.47	11
CA3	${\bf M}$	45.77 ± 10.29	9
Premammillary Nucleus	F	535.99 ± 110.84	10
Premammillary Nucleus	${\bf M}$	499.3 ± 106.84	9
VTA	F	523.97 ± 74.13	9
VTA	${\bf M}$	628.2 ± 174.1	9
PAG	F	191.21 ± 35.76	10
PAG	${\bf M}$	106.35 ± 28.22	7
Superior Colliculus	\mathbf{F}	199.35 ± 46.21	10
Superior Colliculus	\mathbf{M}	51.46 ± 26.08	7

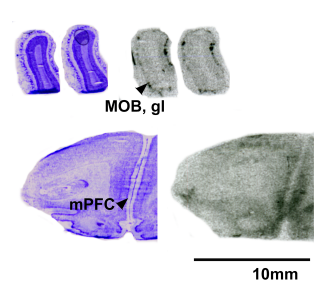


Figure 1: OTR in the forebrain of the pouched rat. Images on left are Nissl stain, associated autoradiograms on right show OTR density. MOB = Main olfactory bulb, mPFC = medial prefrontal cortex.

- Pouched rats had very dense binding in the VMH, with moderate binding in the BST, CeA and VTA. There
- was only a low level of binding in the OB and mPFC (Table 2, Figure 1).

Table 2: OTR relative densities in select regions

Region	Relative.Binding
Olfactory Bulb	+
Nucleus Accumbens	+
mPFC	+
Ventral Pallidum	-
Lateral Septum	+
BST	++
CeA	++
MeA	+
Hippocampus	+
VMH	+++
VTA	++

Table 3: V1aR densities by region and sex

Region	Sex	Density mean \pm SE (dpm/mg)	N
Olfactory Bulb	F	1456.47 ± 244.89	9
Olfactory Bulb	M	1841.49 ± 278.59	8
Accessory Olfactory Nucleus	F	1287.3 ± 173.38	9
Accessory Olfactory Nucleus	M	1596.88 ± 235.76	8
mPFC	F	588.16 ± 73.53	11
mPFC	M	604.41 ± 115.62	9
Infralimbic Cortex	F	388.15 ± 52.78	11
Infralimbic Cortex	M	426.68 ± 111.74	9
Nac Core	F	1319.1 ± 113.88	11
Nac Core	M	1748.56 ± 266.36	8
Nac Shell	F	1389.26 ± 169.07	11
Nac Shell	M	1687.58 ± 362.54	9
Caudate Putamen	F	1725.67 ± 148.69	11
Caudate Putamen	M	2329.94 ± 322	9
Piriform Cortex	F	604.16 ± 87.76	11
Piriform Cortex	M	649.05 ± 144.85	9
Lateral Septum I	F	3602.8 ± 355.47	11
Lateral Septum I	M	2885 ± 369.04	9
Lateral Septum D	F	3451.24 ± 276.7	11
Lateral Septum D	M	3613.44 ± 207.77	9
Lateral Septrum V	F	2395.85 ± 311.31	11
Lateral Septrum V	M	2351.65 ± 325.22	9
Endopiriform Cortex	F	296.6 ± 58.56	11
Endopiriform Cortex	M	260.42 ± 57.86	9
Claustrum	F	446.08 ± 74.68	11
Claustrum	M	334.95 ± 58.2	9
BSTm	F	1552.05 ± 161.04	11
BSTm	M	1452.1 ± 229.77	9
BSTi	F	1297.66 ± 80.59	11
BSTi	M	1056.38 ± 188.24	9
BSTv	F	885.07 ± 108.44	11

Region	Sex	Density mean \pm SE (dpm/mg)	N
BSTv	M	835.57 ± 116.42	9
Ventral Pallidum	F	616.94 ± 104.42	11
Ventral Pallidum	M	470.4 ± 142.24	9
MPOA	F	626.46 ± 84.5	11
MPOA	M	522.26 ± 75.01	9
Anterior Hypothalamus	F	364.98 ± 97.49	9
Anterior Hypothalamus	M	601.09 ± 288.18	8
PVT	F	63.75 ± 58.22	9
PVT	M	245.11 ± 131.63	8
SCN	F	207.91 ± 109.3	9
SCN	M	85.55 ± 71.97	8
PVN	F	514.58 ± 146.84	8
PVN	M	520.91 ± 161.33	7
Central Amygdala	F	1710.61 ± 186.92	10
Central Amygdala	M	1439.79 ± 188.75	9
Medial Amygdala	F	841.82 ± 76.12	10
Medial Amygdala	M	928.64 ± 160.08	9
Basolateral Amygdala	F	120.35 ± 66.19	10
Basolateral Amygdala	M	139.62 ± 74.36	9
VMH	F	390.55 ± 130.5	10
VMH	M	576.42 ± 141.68	9
Zona Incerta	F	705.31 ± 78.81	10
Zona Incerta	M	738.07 ± 94.39	9
Lateral Hypothalamus	F	745.93 ± 91.88	10
Lateral Hypothalamus	M	915.03 ± 143.4	9
CA2	F	119.25 ± 56.35	11
CA2	M	63.4 ± 81.27	9
Dentate Gyrus	F	281.61 ± 107.99	11
Dentate Gyrus	M	347.83 ± 172.25	9
Premammillary Nucleus	F	1458.58 ± 333.09	11
Premammillary Nucleus	M	2206.81 ± 616.26	9
VTA	F	1132.57 ± 186.43	11

Region	Sex	Density mean \pm SE (dpm/mg)	N
VTA	M	1278.42 ± 203.04	9
PAG	F	734.99 ± 97.71	11
PAG	M	947.68 ± 293.67	7
Medial Geniculate	\mathbf{F}	93.87 ± 50.87	11
Medial Geniculate	\mathbf{M}	147.82 ± 117.32	7
Superior Colliculus	F	819.1 ± 146.2	11
Superior Colliculus	\mathbf{M}	744.4 ± 229.16	7
Ventral CA3	F	67.1 ± 78.53	5
Ventral CA3	M	119.62 ± 148.52	2

Table 4: V1aR relative densities in select regions

Region	Relative.Binding
Olfactory Bulb	++
Nucleus Accumbens	++
mPFC	+
Ventral Pallidum	+
Lateral Septum	+++
BST	++
CeA	++
MeA	+
PVN	+
Hippocampus	-
Dentate Gyrus	+
Premammillary Nucleus	++
VMH	+
VTA	+

- -Where did we see binding for OTR
- 95 -Where did we see binding for V1aR
- -Sex differences
- 97 -Predict with PCA



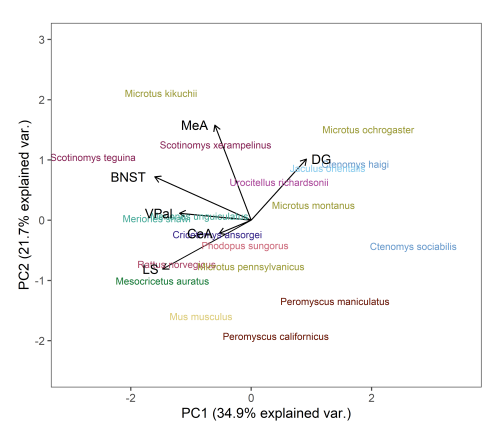


Figure 2: a PCA, V1aR with species maximized

a Cricetomys a Meriones a Mus a Phodopus a Scotinomys a Jaculus a Microtus a Peromyscus a Rattus a Urocitellus

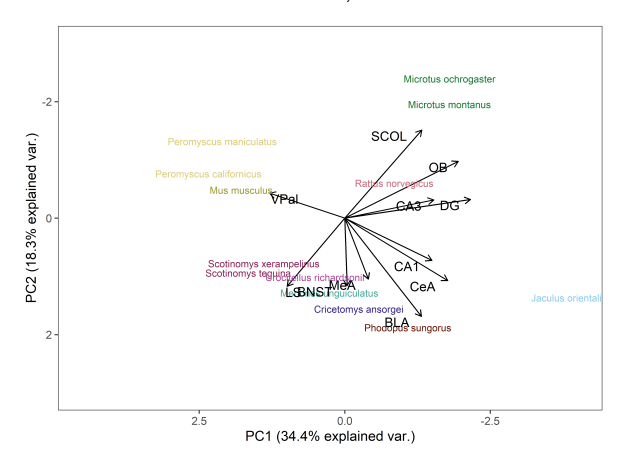


Figure 3: a PCA, V1aR with regions maximized

a Cavia a Georychus a Mesocricetus a Octodon a Rattus

a Cricetomys a Heterocephalus a Microtus a Otomys a Scotinomys

a Ctenomys a Meriones a Mus a Peromyscus a Urocitellus

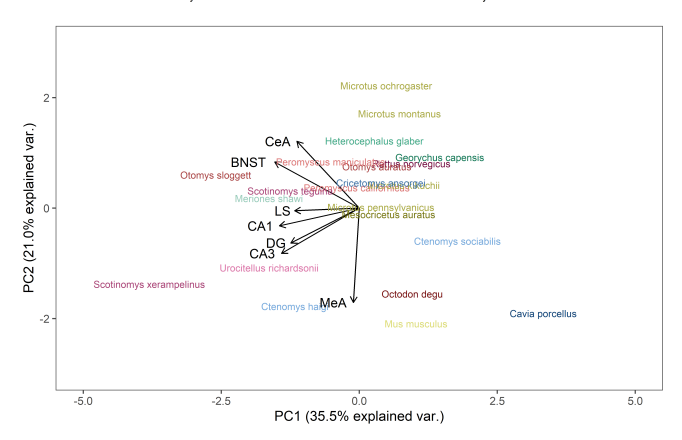


Figure 4: a PCA, OTR with species maximized

a Cavia
 a Heterocephalus
 a Mus
 a Rattus
 a Cricetomys
 a Meriones
 a Octodon
 a Scotinomys
 a Ctenomys
 a Microtus
 a Otomys
 a Urocitellus

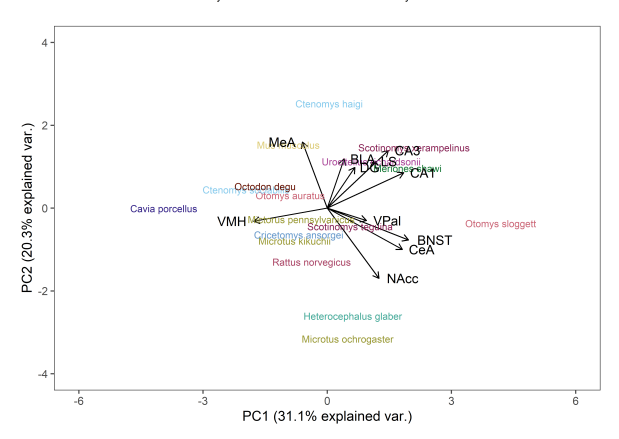


Figure 5: a PCA, OTR with regions maximized

98 Discussion

-We found OTR in ... V1aR in....

-Sex differences in densities/presence and absence

-We found that overall patterns were similar to.....

-Caveats

-Unknown age

-Unknown reproductive status

-Different experiences possible

-What this means, similarities to other species

-Relevance for behavior or life history

What still needs to be known?

109 References

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