Cytoskeletal Regulation Dominates Temperature-Sensitive Proteomic Changes of Hibernation in Forebrain of 13-Lined Ground Squirrels

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Abstract

13-lined ground squirrels, *Ictidomys tridecemlineatus*, are obligate hibernators that transition annually between summer homeothermy and winter heterothermy – wherein they exploit episodic torpor bouts. Despite cerebral ischemia during torpor and rapid reperfusion during arousal, hibernator brains resist damage and the animals emerge neurologically intact each spring. We hypothesized that protein changes in the brain underlie winter neuroprotection. To identify candidate proteins, we applied a sensitive 2D gel electrophoresis method to quantify protein differences among forebrain extracts prepared from ground squirrels in two summer, four winter and fall transition states. Proteins that differed among groups were identified using LC-MS/MS. Only 84 protein spots varied significantly among the defined states of hibernation. Protein changes in the forebrain proteome fell largely into two reciprocal patterns with a strong body temperature dependence. The importance of body temperature was tested in animals from the fall; these fall animals use torpor sporadically with body temperatures mirroring ambient temperatures between 4 and 21 °C as they navigate the transition between summer homeothermy and winter heterothermy. Unlike cold-torpid fall ground squirrels, warm-torpid individuals strongly resembled the homeotherms, indicating that the changes observed in torpid hibernators are defined by body temperature, not torpor per se. Metabolic enzymes were largely unchanged despite varied metabolic activity across annual and torpor-arousal cycles. Instead, the majority of the observed changes were cytoskeletal proteins and their regulators. While cytoskeletal structural proteins tended to differ seasonally, i.e., between summer homeothermy and winter heterothermy, their regulatory proteins were more strongly affected by body temperature. Changes in the abundance of various isoforms of the microtubule assembly and disassembly regulatory proteins dihydropyrimidinase-related protein and stathmin suggested mechanisms for rapid cytoskeletal reorganization on return to euthermia during torpor-arousal cycles.

Introduction

Hibernation includes some of the most dramatic physiological changes tolerated by mammals. In ground squirrels, this natural feature of life history involves transition from a summer homeothermic to a winter heterothermic state. In winter, animals are capable of 1–2 week torpor bouts that are rapidly and repeatedly reversed to inter-bout arousal (IBA) [1]. Body temperature (Tb) and vital signs drop precipitously during entrance into torpor, a period also associated with reduced metabolic rate and global ischemia [2,3,4]; hypoxemia is thought to be avoided during torpor as metabolic rate declines in concert with perfusion levels during entry into and maintenance of torpor to match oxygen delivery with demand. Small-bodied hibernators arouse from torpor at near-zero Tb to euthermic Tb via endogenous heat production mechanisms [1]. Rewarming episodes, in contrast to periods of torpor, represent periods of intense metabolic activity, high oxygen demand and restoration of perfusion to ischemic regions. The system is therefore challenged during rewarming when metabolic rates rise in advance of perfusion, leading to tissue hypoxemia and ischemia-reperfusion events [3,5]. Yet, these hibernators reversibly orchestrate torpor-arousal cycles throughout winter with no observable long-term tissue damage [5].

The brain is a central regulator of hibernation [6]. Yet, brain tissue experiences the temperature, oxygenation and metabolic rate declines associated with a reduced homeostatic set point during torpor [6,7,8], making damage-free recovery from the torpid state paramount. Indeed, evidence for neuroprotection in ground squirrel brain during hibernation, combined with rapid transitions between torpor and euthermia, suggest an innate adaptation to protect and maintain function in the brain during extreme physiological challenge [5]. Such defenses suggest that hibernator brains provide a unique model system for uncovering novel strategies of mitigating damage due to neurodegeneration and cerebral ischemia in humans [5,9].

Among the components of neuroprotection in the hibernator are: 1) depressed metabolic rate reduces oxygen demand to match the ~90% reduction of cerebral perfusion during torpor [7]; 2) a mechanism for cytoskeletal stabilization, including synapses,
during torpor allows rapid cytoskeletal reorganization on return to euthermy [5]; and 3) cellular homeostasis is maintained at low Tb [10]. Reduced Tb is associated with neural retraction across multiple brain regions in mammals: cell bodies, dendritic spines and synaptic profiles shrink [11], accompanied by general cytoskeletal breakdown as microtubules disassemble [12]. Yet, the hibernator’s brain is capable of rapid dendritic regrowth and synaptic rebuilding during each interbout arousal [13,14]. Entry into torpor is associated with 50–60% loss of synapses in golden-mantled ground squirrels (Callospermophilus lateralis), the result of decreased co-localization of pre- and post-synaptic protein markers, although their abundance does not change [15]. The disassembly of synapses occurring with each torpor bout likely complements channel arrest to reduce the potential for Ca2+ buildup at cold Tb and detrimental excitotoxicity [5]. Synaptic protein disassociation at low Tb may also create a pool from which to draw building blocks for synaptic regeneration upon arousal [15]. Although the phenomenon of dendritic retraction in hibernators is well described, the molecular components that enable rapid and reversible cytoskeletal and synaptic reorganization with each torpor-arousal cycle are unknown.

Here, we set out to define proteomic changes in the forebrain of 13-lined ground squirrels that occur in concert with physiological adjustments. The rationale behind this approach was that protein changes associated with neuroprotective elements supporting torpor-arousal cycles could be revealed by defining differences to alternative states. Brain proteomic variations in multiple stages of hibernation have not yet been documented. Six physiologically defined states encompassing summer, spring and the torpor-arousal cycle of winter were analyzed in addition to a group of fall animals transitioning from summer homeothermy to winter heterothermy. This experiment was intended to capture the broad-scale defense strategies employed across a variety of cell types to contend with Tb and metabolic rate declines during torpor. Two significant aspects of these data shed light on proteomic regulation in the forebrain of 13-lined ground squirrels. Body temperature segregates the abundance of many proteins, indicating distinct strategies for maintaining homeostasis in torpor compared to interbout arousals and summer homeothermy. There is also a dominant cytoskeletal signature among the proteins that change across the hibernator’s year, indicating its importance in neuroprotection.

Methods

Tissue Collection

Thirteen-lined ground squirrels purchased in July from the University of Wisconsin captive breeding program were housed at the University of Colorado individually at 18–21°C (14:10 L:D) with dry cat food, sunflower seeds and water provided ad libitum. Figure 1 describes sampled states. Summer active (SA) ground squirrels were euthanized Aug. 8. The remaining ground squirrels were maintained in standard housing until Oct. 5 when they were transferred to the hibernaculum (4°C, 0:24 L:D). To identify winter sampling states (Fig. 1) during hibernation, animals were each implanted with an intraperitoneal Tb recorder (iButton, Embedded Data Systems) and a telemeter (MiniMitter). Animals (IBA, Ent, LT, EAr) were collected during the specified Tb points or ranges (Fig. 1) within regular torpor-arousal cycles between 28-Dec and 26-Mar. Surgeries were performed under isoflurane anesthesia with post-operative analgesia with all efforts made to minimize suffering. Animals were transferred into the cold room only after a minimum 2-week post-surgical recovery. Food and water were removed once ground squirrels exhibited regular torpor in the cold room and returned to Spring animals (SpC) at least 10d before tissue collection. Samples were also collected from Fall Transition (FT) individuals between 21-Sept and 12-Oct to capture the physiological range of fall (FT animals were a mixture of individuals housed in standard laboratory housing or in the hibernaculum as or after the ambient temperature was lowered to 4°C and being torpid or not torpid at time of tissue collection [16]. Torpor in Fall Transition ground squirrels was evaluated from implanted Tb biologgers (iButton). Biometrics, housing and Tb data are presented in Table S1 for Fall individuals. This study was conducted according to the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. All animal care and procedures were approved by the University of Colorado School of Medicine Institutional Animal Care and Use Committee (protocol #44309).

For tissue collection, ground squirrels were anesthetized with isoflurane, then exsanguinated via cardiac blood draw and perfused with 6 volumes of ice-cold saline. After decapitation, brains were removed and placed on ice, then brainstem with cerebellum, olfactory bulbs and hypothalamus were dissected free. The remaining brain tissue, consisting of forebrain minus the hypotalamus was immediately frozen in N2(l), then stored at −80°C until processing. This sample was hemisected, and the right hemisphere was pulverized in N2(l) prior to bead-homogenization on ice (Bullet Blender, Next Advance) and centrifugation to retain the soluble, non-nuclear fraction as previously described [17]. Protein concentration of the homogenate was determined using a BCA assay (Pierce).

2D Gel Analysis and Protein Identification

The procedure for processing and labeling sample homogenates of equal protein content with Cy2, 3 or 5 fluoros (GE Healthcare) and their separation on 2D gels were as described [18], 24 gels, each containing 2 experimental samples (Cy3- and 5-labeled) and a pooled reference sample (Cy2-labeled) were imaged and imported along with Sypro Ruby stained pick gel images into DeCyder software (V7.0, GE Healthcare) for analysis [18]. Protein spots in all Cy2 reference images were matched automatically after manually landmarking ~400 spots. Each Cy3 and Cy5 spot from the 48 remaining gel images were matched and normalized for spot intensity corresponding to Cy2.

We used one-way ANOVA with false-discovery rate correction [19] to identify protein spots whose abundance differed significantly by hibernation state. Only spots present in ≥3 experimental samples from each state were included. Spots with ANOVA q<0.05 after FDR correction were examined on all gels and manually matched in additional samples where possible. 77% of protein spots ultimately identified from this assessment had abundance data for all sampled animals and a given sampled state was left with only 3 data points (3 of 6 possible) only once. Hibernation states were also classified with the machine-learning clustering tool, Random Forests [20,21]. The minimum required protein spots of known identity present on all gels that produced the lowest out-of-bag clustering error between the states were identified by Random Forests variable section algorithm (drop = 0.2, n = 100,000 iterations) [22]. All analyses were conducted on two datasets – the six base hibernation states with and without the Fall Transition — and were performed in R (v2.15.0, http://www.r-project.org) [23].

We used LC-MS/MS to attempt identification of protein spots that were significant by ANOVA after FDR or by Random Forests. Gel plugs (diameter = 1.4 mm, depth = 1 mm) containing
spots of interest and that were reproducible on the pick gels were picked and digested with an automated system as previously described [17]. We collected spectral data from tandem MS on the four highest peaks with 30s dynamic exclusion as previously described [17]. Tryptic fragments were analyzed with Spectrum Mill MS Proteomics Workbench (revision B.04.00, Agilent Technologies) against a protein database containing predicted sequences from the published 13-lined ground squirrel genome (Ensembl build 67, 20,000 entries, 1 missed residue allowed). Spots not identifiable using this database were subsequently searched against the National Center for Biotechnology Information mammal sequences from April 2012 and previously published actual sequences from arctic ground squirrels, Urocitellus parryii [24], permitting 2 missed residues. Including the 13-lined ground squirrel genomic information, this database contained 1,071,259 entries. Settings such as precursor and fragment ion mass tolerances, allowed missed cleavages, and amino acid modifications followed Hindle et al. [17].

We retained protein identifications (IDs) with at least two supporting peptides and a score ≥30. Protein spots which resolved into a unique protein ID with ≥3 supporting peptides and a score ≥30 were considered identified from a single pick gel. Spots that did not initially resolve into a unique ID were analyzed from 1–3 additional gels. The resulting spectral data were merged with an in-house program, ExtracTags (v4.1). The top hit from multiple IDs was retained only when it presented a 4-fold higher score, peptide recovery or spectral intensity than any other hit. Single best IDs for each analysis are presented in Tables S2&S3 and the best IDs for each analysis are presented in Tables S2&S3 and the complete file output including unresolved weak and multiple IDs is given in Table S4.

Additional Analyses & Data Visualization

Significant pairwise differences between individual hibernation states were determined by Tukeys HSD posthoc comparisons (Tables S5&S6). Data were visualized by heatmap and Random Forests MDS plots (n = 50,000 trees) based on all significant, identified protein spots and those determined to produce the lowest-error clusters, respectively. Protein spot clusters generated by heat map were submitted to the DAVID bioinformatics database (V6.7) to identify significantly enriched gene pathways [25], using settings as described [17]. The proximity of the Fall Transition proteome to the physiologically defined states was predicted with spot data from the six defined states as a training set in Random Forests [21] and tested using unsupervised classification.

We examined phosphoprotein staining as evidence of posttranslational protein modification (PTM) in forebrain samples from IBA and EAr animals (n = 3 per state). 150 µg of unlabeled protein from each sample were separated on 2D gels (DGE protocol), fixed overnight (50% MeOH, 10% acetic acid), then stained for phospho-proteins (ProQ Diamond, Molecular Probes) followed by total protein staining (SYPRO Ruby) as described [26]. Western blots confirmed total protein content in proteins of interest. 20 µg protein of IBA and LT ground squirrels (n = 3 per state) were separated with SDS-PAGE then transferred to PVDF membranes (Immobilon-FL, Millipore). Consistent protein loading and transfer were assessed with total protein stain (MemCode™ #24585, Pierce). After blocking (Odyssey blocking buffer, Li-Cor) blots were incubated overnight at 4°C with anti-DYSPL2 (1:1000) or anti-STMN1 rabbit pAbs (Cell Signaling), then for 1.5 h with a secondary antibody for anti-rabbit IgG (1:15,000 IRDye 800CW-conjugated, Li-Cor). Examination of serine phosphorylation was attempted by multiplexing immunoblots with mouse anti-pSer mAb (1:500, 3C171 Abcam) with secondary detection via anti-mouse IgG (1:15,000 IRDye 680RW-conjugated, Li-Cor). Blots were imaged with Odyssey near-infrared fluorescence detection (Li-Cor) and total protein densities were calculated in ImageJ (http://imagej.nih.gov/ij, National Institutes of Health, USA) and analyzed with ANOVA. 2D western blots following electrophoresis methods for phosphoprotein staining were also employed.
Results

Forebrain Proteomic Changes are Limited and Associated with Body Temperature

In six physiologically defined hibernation states of 13-lined ground squirrels (Fig. 1) we detected 84 protein spots from 3471 total spots analyzed with 2DGE that differed significantly among groups. 56 of these were successfully identified by LC-MS/MS, which represented 34 unique proteins due to multiple isoforms (Table S2). When the Fall Transition animals were included in the ANOVA, 88 protein spots significantly differed (54 identified; Table S3). These two ANOVAs combined to suggest 90 protein spots of interest.

Hierarchical clustering of spot intensities and visualization by heatmap revealed two dominant reciprocal patterns of protein abundance (Fig. 2A). In cluster 1, spot abundance was highest in homeothermic states (SA & SpC, Fig. 2B) and lowest in the cold-Tb states, LT and EAr. Warm- (IBA) or declining-Tb states (Ent) during winter resembled the homeothermic proteome or an intermediate abundance pattern, respectively. DAVID analysis of protein IDs belonging to cluster 2 (Fig. 2C) contained enrichments of proteins with roles in neural growth and differentiation and synaptic transmission (Table 1). Cluster 1 also revealed enrichment of the neurogenesis/differentiation pathway. Significantly, both clusters indicated differential regulation of the cytoskeleton (Table 1). The considerable protein overlap in the two clusters implies the importance of specific isoforms in regulating neural and cytoskeletal architecture during torpor-arousal cycles.

Random Forests clustering segregates the hibernation states by Tb (Fig. 3). The two cold-Tb groups, LT and EAr, are not separated under any conditions, even when the winter states are examined alone (Fig. 3B). Indeed, only three protein spots significantly differ between these two groups (Tukey posthoc, Table S5). The segregation among sampled states persisted when Random Forests was run in unsupervised mode (Fig. 4A). The mixed-physiology FT animals were used to test whether separation among hibernation states resulted from changes in Tb, or torpor status. We performed two assessments of Fall Transition (FT) individuals with unsupervised classification. Initially, 2D scaling plots were used to visualize the classification of FT individuals, which lie predominantly with the homeothermic states (Fig. 4A).

Subsequently, unsupervised clustering of FT quantified proximities of the mixed-physiology animals to each defined state (Fig. 4B). All but three FT individuals (which were cold-torpid) had a forebrain proteome most proximate to the homeotherms (Figs. 4A&B). Two additional individuals using torpor in standard laboratory housing shared the high proximities to homeothermic states observed in non-torpid animals (Fig. 4B). A significant positive correlation between Tb at time of tissue collection and proximities to SA, or to the euthermic states overall (SA+SpC+IBA; Fig. 4C) suggests that Tb rather than torpor per se, is related to the brain protein patterns observed in LT and EAr.

Cytoskeletal Proteins Dominate the Dataset

Random Forests clustering also revealed protein spots of interest. A combined 23 protein spots were key in the lowest-error clustering for comparisons among hibernation states (Figs. 3A&B). 18 (78%) of these spots overlapped between this analysis and the ANOVA. Approximately half of the spots important in separating the states by Random Forests were cytoskeletal components or binding proteins (represented by GENE SYMBOL, Table S4, below). Focusing on those with ≥1.5X maximum fold change (Fig. 5), the selected cytoskeletal proteins (Fig. 5A) generally increase from summer to winter. Two building blocks of nerve cell architecture, neurofilament heavy-chain (NEFH(1467,177) and α-tubulin isoform 1A (TUBA1A(1461) displayed a general winter elevation compared to summer. The remainder, DYPSL2(2412), CORO1A(101), STMN1(2906,2908), DPYSL2(1212), CORO1A(1401), STMN1(3983,2986), DPYSL2(1212), CORO1A(1401), STMN1(3983,2986) and sorcin (SRI2862). The abundance of synaptosomal-associated protein 25 (SNAP25(2512), involved in Ca2+-sensitive synaptic responsive peaks in SA and displays no significant pairwise differences among the winter states (Tables S5&S6). Peroxiredoxin 6 (PRDX6(1576) is least abundant in spring (Fig. 5B), and also does not significantly differ among the winter groups (Tables S5&S6). A limited complement of metabolic enzymes were included in these

Table 1. Significant pathway enrichments in squirrel forebrain for two primary hierarchical clusters.

| Annotation                          | Definition                             | Enrichment Score | Fold Enrichment | Genes                          |
|-------------------------------------|----------------------------------------|------------------|-----------------|--------------------------------|
| **Cluster 1**                       |                                        |                  |                 |                                |
| 1                                   | GTPase activity                        | 3.0              | 18.5            | RAB3A, DNM1L, TUBB2A, TUBA1A, DNM1, TUBA1C |
| 2                                   | Growth cone                           | 2.2              | 35.5            | DPYSL3, DPYSL2, SNAP25          |
| 3                                   | Neuron differentiation                | 1.7              | 7.3             | RAB3A, MAP2K1, TUBB2A, SNAP25   |
| 4                                   | Endocytosis                            | 1.7              | 10.9            | RAB3A, DLG4, DNM1               |
| 5                                   | Synaptic transmission                 | 1.6              | 13.4            | RAB3A, SYN2, DLG4, SNAP25, MBP   |
| 6                                   | Neurotransmitter secretion            | 1.6              | 70.2            | RAB3A, SYN2, SNAP25             |
| 7                                   | Cytoskeleton Regulation               | 1.3              | 5.5             | MAP2K1, MAP2K2, CFL1            |
| **Cluster 2**                       |                                        |                  |                 |                                |
| 1                                   | Neurogenesis/differentiation           | 1.7              | 26.3            | NDRG2, STMN1, DPYSL2           |
| 2                                   | Cytoskeleton                          | 1.6              | 4.0             | CORO1A, NEFH, NDRG2, STMN1, DPYSL2, TUBA1A |

Note. Hierarchical Cluster 1, with seven significantly enriched functional annotation clusters, contains proteins decreased in cold Tb, whereas hierarchical Cluster 2, with two significant functional annotation clusters, contains proteins increased at cold Tb and decreased during euthermia (see Fig. 2). Enrichment score gives the mean p-value among members of each annotation cluster (negative log). Enrichment data were calculated by DAVID Bioinformatics Resources v6.7 [25].

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Posttranslational Modifications (PTMs) Support Abundance Shifts During Heterothermy and Torpor

The nature of intra-winter protein abundance cycles may suggest mechanisms of regulation in heterothermy. Increased gene expression by protein synthesis in the homeothermic season, or during euthermia between torpor bouts is the likely explanation for spot abundance increases during warm-Tb states such as SA, SpC, IBA and Ent. However, changes in subcellular sequestration or PTM are additional explanations for an increased relative protein spot abundance. A modification such as phosphorylation or acetylation creates isoelectric point shifts for protein isoforms that would result in a train of spots on 2D gels. If steady-state levels of a protein remain constant, we expect that PTMs conferring a change in protein activity should manifest as spots bearing the same ID but having reciprocal abundance patterns within a train. This dataset contains several examples of same-identity spots appearing in trains and bearing reciprocal patterns.

γ-tubulin isoform 1A (TUBA1A) may appear winter-depleted (spots #1399, #1449, Table S7) or winter-elevated (spots #1400 and #1461, Fig. 5A, Table S7). Given that transcription and protein synthesis effectively cease at the cold-Tbs of torpor [27,28], spot abundance increases in the cold-Tbs of LT and EAr, exhibiting the highest maximum fold change of four metabolic proteins, being 1.35-fold higher in EAr compared to SA (Table S7). Normalized abundance data for all identified spots are presented in Table S8.

Figure 2. Forebrain proteome heat map. The forebrain proteome of cold-Tbs ground squirrels demonstrated a reciprocal abundance pattern to euthermic animals. (A) Heat map y-axis: Individual ground squirrels grouped by state. Heat map x-axis (bottom): Protein spots (Spot Number.GeneID) that differed significantly by hibernation state (ANOVA q) and identified by LC-MS/MS were hierarchically clustered (X-axis top: dendrogram). Blank squares in the heat map represent samples in which a given protein spot was not recovered. (B & C) Means with 95% confidence intervals are plotted for the two reciprocal clusters.

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Figure 3. Sampled state separation by Random Forests. Data were clustered by Random Forests (50,000 trees) after variable selection to determine the minimum input protein spots required for the lowest classification error among groups. This was accomplished with (A) 22 protein spots for the base states and (B) 11 for winter states only (see Fig. 5).

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The forebrain proteome in hibernating ground squirrels appears in several ways to be an exception among tissues studied to date. The magnitude and diversity of proteins recovered on 2D gels was the highest yet (3417 protein spots on the master gel), however, the degree of proteomic response across the hibernation cycle was minimal (only 84 protein spots or ~2.5% differed significantly among the six physiologically defined states). Also, in contrast to previous proteomic studies of skeletal muscle, heart and kidney following the methods used here [17,29,30], there is a notable paucity of metabolic enzymes contained in the proportion of the brain proteome that varies with hibernation. Only four metabolic enzymes differed significantly among the hibernation
transaldolase had the largest fold change (1.35X, Table S7). This enzyme of the pentose phosphate pathway aids in connecting the non-oxidative portion of this energy-producing pathway with glycolysis. The pentose phosphate pathway provides cells with metabolites essential for nucleic acid synthesis such as ribose-5-phosphate and reducing cofactors such as NADPH [31]. Transaldolase abundance peaks early in the rewarming period, a time consistent with a need for elevated NADPH production to maintain the reduced glutathione pools that protect against oxidative damage [32].

The absence of sweeping changes in metabolic enzymes across the hibernation cycle as seen previously in tissues including liver [33], heart [29], skeletal muscle [17] and even brainstem [18] leads to the supposition that passive temperature-regulation of enzyme activity is important in the forebrain. In other words, cerebral metabolism is not actively suppressed in advance of torpor, but rather enzymatic processes are slowed via the Q10 effect as Tb declines [34]. Our results do not support the expected seasonal change in metabolic strategy in this tissue [3], perhaps highlighting the importance of circulating substrate delivery to brain, rather than tissue-level production [35]. This lack of evidence for substrate switching to accompany the winter fast is in contrast to seasonal mRNA expression in cerebral cortex consistent with ketone use and glucose conservation [36]. This difference could reflect post-transcriptional gene regulation or the difference in tissue complexity between the two studies.

Fall Transition individuals shared the majority of their forebrain proteome with summer and spring (Fig. 4). Prior experience with torpor, and in fact the collection of tissues during torpor bouts in standard laboratory housing when Tb was 22°C did not affect this similarity to the homeotherms. Rather, Tb or a physiological correlate seems to be a more significant seasonal mechanism of cell regulation in forebrain as opposed to a coordinated fall reprogramming to generate a proteome prepared for winter heterothermy, with subsequently minor changes observed across torpor-arousal cycles. This is evidenced by the proteomic similarity of fall cold- but not warm-torpid animals to those in winter torpor (LT) or rewarming (EAr) (Fig. 4). Overall, a shift between cold and warm Tb dominated the proteomic changes observed in 13-lined ground squirrel forebrain (Figs. 2&4). Again, this response is markedly different from multi-state proteomic examinations of other tissues (skeletal muscle, kidney, heart) conducted in our lab, in which a summer-winter shift comprised the bulk of protein changes [17,29,30].

While control of metabolic enzyme activity during torpor may derive largely from passive effects of low Tb, the strong signature of temperature oscillation in other proteins suggests that distinct homeostatic regulatory events do occur. Sorcin (soluble resistance-related calcium binding protein) may be one such example. The abundance of sorcin (SRI2962) is decreased during cold-Tb states (Fig. 5B). Ca²⁺ binding induces a conformational change in sorcin that promotes translocation from a soluble into a membrane-associated state. Its low abundance at cold-Tb can be explained by a general transition to a membrane-associated state where it would not be collected in the soluble fraction of the homogenate analyzed here. In rat caudate-putamen nucleus (CPN) sorcin is localized with smooth endoplasmic reticulum (SER) and mitochondria, two Ca²⁺-storing organelles [37]. In heart, sorcin interacts with

Figure 4. The fall proteome tracks body temperature. Fall transition individuals were held in warm (Tb = 18–20°C, #41–46) or cold rooms (Tb = 4°C, #47–52). (A) Unsupervised classification of all states using the most important predictors determined in Fig. 3A maintains the segregation of states by Tb. The mixed physiology of Fall animals can test the effect of Tb. (B) Unsupervised classification quantified proximity of FT individuals based on the defined groups as training data. In all but three individuals, more than 50% of the FT proteome was similar to summer or spring euthermic ground squirrels (x-axis: animal ID). The three exceptions were cold torpid (asterisk denotes torpid), and instead had proteomes more proximate to LT/EAr. The absence of a similar pattern in the two warm-housed but torpid individuals (Tb = 18–21°C), supports the hypothesis that the LT/EAr proteome is defined by Tb, and not torpor per se. (C) Proximity to the euthermic groups (SA+SpC+IBA) correlates significantly with Tb.

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Cell Structure across the Hibernation Season

Cytoskeletal signals dominate this dataset. The cytoskeleton or its regulation was functionally enriched (Table 1) among proteins following the two most commonly observed abundance patterns (Fig. 2). Further, more than half of the proteins with \( \geq 1.5 \) fold change and identified as important for separating the states by Random Forest were cytoskeletal components or binding proteins (Fig. 5A). Changes in neural cell architecture across torpor-arousal cycles are a well-documented aspect of hibernation biology [11,13,14]. Neural retraction occurs during every torpor bout [11] with re-establishment of neural profiles and function during each interbout euthermic period [11,13,14]. The ER likely plays a role in this process as well, as it controls local \( \text{Ca}^{2+} \) levels in growth cones and synaptic compartments [42]. Ca\(^{2+}\) is a key second messenger in growth cones, and decreased Ca\(^{2+}\) ER-outflow (via dantrolene) reduces axonal outgrowth [42]. The inverse, a relative increase in ER-Ca\(^{2+}\) release during euthermia would support the neural recovery observed after each torpor cycle and is consistent with the patterns noted for the Ca\(^{2+}\)-regulating protein sorcin. This overrepresentation of cytoskeletal proteins that change seasonally and more importantly within torpor-arousal cycles supports the idea that synaptic remodeling is a significant neuroprotective feature in this portion of brain. Neuroprotection in this manner may derive from or promote the quiescence observed in this region during torpor and explain the absence of similarly abundant cytoskeletal proteins in a dataset from the more active brainstem [18]. mRNAs involved in cytoskeletal remodeling are also overrepresented in cerebral cortex, but not in the constitutively active hypothalamus [36]. Interestingly, no overlap exists between observed changes in protein (this study) and mRNA levels [36], however, many mRNAs corresponded to extracelluar matrix proteins, a cell component excluded from our soluble protein preparation.

Despite the dominance of \( T_b \) or correlated physiological regulation observed in the forebrain proteome of 13-lined ground squirrels, we also uncovered limited examples of seasonal regulation. For instance, all detected isoforms of neurofilament heavy-chain (NEFH) displayed a seasonal winter increase (Fig. 5A). Neurofilaments are the most prominent cytoskeletal component of neurons in adults and occur as heteropolymers containing light- medium- and heavy-chain polypeptides. They are a major structural component of neurons, specifically implicated in maintaining their diameters and therefore nerve conduction velocities [43]. Neurofilament accumulation in non-hibernators is generally pathological and occurs in association with several conditions, including amyotrophic lateral sclerosis (ALS). We postulate that the winter-elevation of NEFH is a general signal of seasonal cytoskeletal plasticity and limited protein synthesis during euthermic periods to support neuronal remodeling.

Microtubules are another component of the neural cytoskeleton that are regulated across hibernation. Along with the actin cytoskeleton, controlled transport, development and extension of microtubule arrays are important for axonal growth [44], dendritic spine development and therefore synaptic plasticity [45,46]. Microtubules are dynamic polymers of \( \alpha \) - and \( \beta \)-tubulins dimers. Several isoforms of both \( \alpha \) - and \( \beta \)-tubulins displayed

Figure 5. Abundance of protein spots important for separating clusters. The relative abundance of (A) cytoskeletal elements and binding proteins and (B) non-cytoskeletal proteins highlighted by Random Forests as important inputs for hibernation state clustering (Fig. 3). Mean values in each base state are plotted for spots that also had \( \geq 1.5X \) maximum fold change. A broken vertical line divides the sequential homeothermic from heterothermic states.

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overall summer-winter abundance shifts, but remained consistent throughout winter (e.g., TUBA1A1461, Fig. 5A). Two forms of β-tubulin (TUBB2A1641, TUBB4B1527) peaked in summer and were consistently winter-depressed. Only a single TUBA1C1396 isoform was identified, which was also consistently lowest in winter. However, several isoforms of α-tubulin-1A chain (TUBA1A) were identified (Table S2). These isoforms occurred in two reciprocal patterns, having intra-seasonal consistency, but an abundance change between the homeothermic and heterothermic periods. This observation suggests that the PTM status of this protein changes seasonally, with the natural dynamic instability of microtubules in homeothermy adjusted to handle the additional requirements of heterothermic plasticity. Our phosphoprotein staining experiment does indicate that several of the detected α-tubulin isoforms are phosphorylated, however this protein presents on 2D gels as several large spot trains and the location of phosphorylated isoforms within the trains suggests that this is not the sole regulatory PTM. Indeed, many PTM are known for tubulins [47,48] and phosphorylation is not the most common form [48]. Our initial finding suggests that tubulins, and α-tubulin in particular may be an avenue for further study of winter neuroprotection.

Torpor-arousal Cycle Regulation of Microtubules

The retraction of dendritic spines and synaptic structures at cold temperatures, such as during torpor is well documented in hibernators and non-hibernators alike [49,50] and is likely driven by cold-induced disassembly of microtubules [12]. Dendritic spines are capable of regeneration upon re-warming in non-hibernators [49] but this process occurs more rapidly in hibernators [11]. In fact, the rate of dendrite extension during arousal from torpor occurs more quickly than rates observed in development or in response to environmental enrichment in non-hibernators [11]. If cytoskeletal components themselves display
within-winter consistency, neuroprotection may instead exist in the form of regulatory proteins that are activated at specific points across heterothermy.

One contender is the cytoskeletal assembly regulator stathmin 1 (STMN1). Two stathmin isoforms (#2893, #2896) were the two most significantly changing proteins across the hibernation cycle (Table S4) and among the most important predictors of group classification by Random Forests. The abundance of both stathmin isoforms is high in cold-Tb states and consistently low otherwise (Fig 5A). Microtubule polymerization is subject to dynamic instability, or rapid cycling between growth (‘rescue’) and disassembly (‘catastrophe’). The overall assembly or disassembly of microtubule arrays results from an imbalance in dynamic instability favoring either catastrophe or rescue. Polymerization is in part determined by the cytoplasmic availability of free tubulin [51]. Stathmin binds two tubulin dimers and thus removes them from the free soluble pool [52,53], promoting microtubule disassembly by increasing the frequency of catastrophe. Additionally, stathmin can bind directly to microtubules [54,55], acting primarily on the minus end but capable of increasing the frequency of catastrophe at both microtubule ends [55]. Stathmin’s tubulin-binding ability is regulated by PTM, specifically it is inhibited by phosphorylation at up to four serine sites [56]. Efforts to unequivocally determine the phosphorylation status of the stathmin isoforms identified here were unsuccessful. Neither 1D nor 2D western immunoblotting was sufficient to quantify total stathmin. The isoforms appeared dephosphorylated when evaluated by phosphoprotein stain and by pSer antibody labeling in 1D or 2D western blots, however this result is difficult to validate in the absence of definitive stathmin antibody labeling, likely an effect of the very low abundance of this protein in our forebrain sample. Thus the regulatory role of stathmin remains unclear. However, if dephosphorylated stathmin is most abundant at cold-Tb, its absence due to proteolysis or inhibition by unclear. However, if dephosphorylated stathmin is most abundant at cold-Tb, its absence due to proteolysis or inhibition by

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Our results provide mechanistic insight into three aspects of neuroprotection in the forebrain of 13-lined ground squirrels; metabolic rate, cytoskeletal dynamics and Ca^{2+} regulation. First, although it is acknowledged that metabolic rate depression is necessary to balance the ~90% reduction in cerebral perfusion during torpor, we documented that few changes occur among metabolic enzymes in this brain region between summer and winter, or across winter heterothermy. We therefore postulate that the ground squirrel forebrain exploits passive temperature reduction via the Q_{10} effect to depress metabolic rates and cell functions during torpor. Subsequent elements of neuroprotection are likely necessary to both stabilize the cell at low Tb, and to shepherd damage-free recovery from the torpid state.

Second, our data provide insight into the molecular mechanisms that both stabilize cells in the cold and protect them during arousal. The majority of proteins that vary with hibernation physiology were associated with the cytoskeleton, highlighting its importance in defining the torpid and aroused states. While cytoskeletal proteins themselves appear to differ from summer across the entire heterothermic period, their regulatory factors track changes in Tb. Both cytoskeletal regulators highlighted here are affected by PTM [56,60], specifically phosphorylation; we have demonstrated a phosphorylation shift associated with isoforms of opposing abundance patterns in one of these, DPYSL2. It is not surprising that phospho-modifications could be temperature-dependent regulatory mechanisms, as this process is rapid, reversible and unaffected by the dramatically decreased transcription and translation in place at low Tb [27,28]. Indeed, phosphorylation also regulates the function of tau, another microtubule binding protein implicated in cytoskeletal remodeling during hibernation [61,62]. It is noteworthy that both regulatory proteins discussed here (STMN1 & DPYSL2) differ from other microtubule-associated proteins including tau in that they more efficiently bind tubulin heterodimers as opposed to microtubules.

Third, these data also highlight Ca^{2+} homeostasis as an element of cell stabilization during torpor-arousal cycles. Ca^{2+} signaling has wide-reaching effects in regulating general cell functions. At least one protein implicated in Ca^{2+} homeostasis [SRI, sorcin] differed in the soluble fraction of 13-lined ground squirrel forebrain in response to hibernation, and specifically Tb. This reveals an element of the strategy in place to regulate intracellular Ca^{2+} handling during torpor.

It is clear that despite relatively few proteomic differences in forebrain across the hibernator’s annual cycle, key proteins do change. Body temperature is the most relevant factor in defining protein abundance changes, as evidenced by the temperature segregation of the mixed-physiology Fall animals. This suggests that the forebrain manages homeostasis during its quiescent torpor periods distinctly from interbout arousals. Components of neuroprotection in hibernation revealed by their response to Tb may be useful therapeutic targets for human neuropathies.

**Supporting Information**

**Table S1** Biometrics and housing information for Fall Transition 13-lined ground squirrels used in this study. (XLS)

**Table S2** Summary of all unequivocally identified proteins that differed among six base physiological states of hibernation in 13-lined ground squirrels.


**Table S3** Statistical summary for all unequivocally identified proteins (n = 56), which differed significantly among defined physiological states after multiple test correction.

**Table S4** Statistical summary for all unequivocally identified proteins (n = 62), which differed significantly for either one-way analysis or multiple ANOVA test correction.

**Table S5** Statistical summary for all unequivocally identified proteins (n = 54), which differed among the defined physiological states and the Fall Transition states.

**Table S7** Data summary for all unequivocally identified proteins (n = 62), which differed significantly for either one-way analysis or multiple ANOVA test correction.

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**Table S8** Normalized protein spot abundance data from all sampled 13-lined ground squirrels. Group identifies sampled state abbreviation and ID reflects individual animal identifier. Remaining column headings denote spot number on the master gel and official gene symbol. Data are presented for all identified spots, selected by significance in any statistical procedure or in select cases as non-significant but robust control spots.

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**Author Contributions**

Conceived and designed the experiments: AGH SLM. Performed the experiments: AGH. Analyzed the data: AGH. Contributed reagents/materials/analysis tools: SLM. Wrote the paper: AGH SLM.
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