**Stress alters hypothalamic gene expression in adolescent male hamsters**

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**Abstract**

In hamsters, a two-week exposure to chronic social stress in adolescence causes acceleration of agonistic behavior, enhanced adult aggression, impaired waiting impulsivity, and higher food intake, body fat, and long-term increased body weight. In adult rodents, stress is accompanied by widespread alterations in gene expression in the brain. The present research examined transcriptomic changes in the lateral, dorsomedial, and arcuate nucleus of the hypothalamus caused by adolescent stress using RNA Tag-sequencing. In each region, there were approximately 250 differentially upregulated and 250 downregulated genes. Many of the most significantly affected genes have been associated with metabolism and sex hormone function. For example, in the lateral hypothalamus, melanocortin 3 receptor, growth hormone releasing factor, both involved in metabolic processes, and neuropeptide VF precursor, involved in growth hormone inhibitory hormone production, were among the most upregulated in stressed subjects. In the dorsomedial hypothalamus, neuropeptide W, involved in feeding cessation, was significantly downregulated in stressed animals. Across both regions, G-protein coupled receptor 50, involved in thermoregulation, sleep, and sex-related mood disorders, was significantly altered, but in opposite directions. In the arcuate nucleus, a number of blood brain barrier- and inflammation-related genes were altered as well. Furthermore, there were consistent patterns of genetic ensembles identified through gene ontology analysis and weighted gene correlation network analysis that were altered across each region. Many of these involved roles in RNA processing, DNA methylation, myelination and synaptic organization. These findings reinforce prior behavioral, hormonal, and metabolic changes observed in this developmental model, and help guide future directions of research related to the negative consequences of early life stress.

Keywords: social stress, RNAseq, obesity, appetite, puberty, WGCNA

**1. Introduction**

Stress is an unavoidable circumstance to which species have developed various coping strategies, such as conditioned avoidance or learned helplessness [(Levine, 1956; Levine et al., 1956; Potegal et al., 1993; Seligman and Maier, 1967)](https://www.zotero.org/google-docs/?bFnneq). These behaviors are malleable and related to prior experience and time of exposure, and can be adaptive or maladaptive depending on following life circumstances. It is well known that prior stressful experiences during childhood and adolescence increase the likelihood of adult behavior, mood, and health disorders [(Hughes et al., 2017; Lipsky et al., 2021; Mancini et al., 2023; Pervanidou and Chrousos, 2012)](https://www.zotero.org/google-docs/?aNrj5f). Additionally, in many humans, chronic stress tends to cause increased consumption of high calorie foods, though in a smaller set of individuals, promotes anorexic behaviors [(Brambilla, 2001; Dallman et al., 2003; Tryon et al., 2013)](https://www.zotero.org/google-docs/?3tX5y8), which is also related to fetal or childhood experiences [(Baldwin et al., 2016; Lumey et al., 2007; Noll et al., 2007; Pervanidou and Chrousos, 2012)](https://www.zotero.org/google-docs/?mjqgxE).

One well-characterized example of this phenomenon in an animal model is the differential response to social defeat stress in adult and adolescent hamsters. In adult hamsters, a single loss causes conditioned defeat behaviors, in which losers/subordinate individuals show reduced aggression to conspecifics and likely lose any altercations in the future [(Huhman, 2006; Potegal et al., 1993)](https://www.zotero.org/google-docs/?FAl24X). Adult social defeat causes widespread *c-Fos* expression throughout the brain, including the hypothalamus [(Kollack-Walker et al., 1997)](https://www.zotero.org/google-docs/?WMRdWD), and persists through seven days of chronic defeat [(Kollack-Walker et al., 1999)](https://www.zotero.org/google-docs/?wPlo2Y). In adults, a single defeat also induces differential expression of over 500 gene transcripts in the basolateral amygdala [(McCann et al., 2019)](https://www.zotero.org/google-docs/?XCHVuA). Increasing the expression of genes in the medial prefrontal cortex via histone deacetylase inhibition also enhances social defeat behaviors in adults, while inhibition of acetylation reduces them [(McCann et al., 2017)](https://www.zotero.org/google-docs/?2801Pk).

Conversely, chronic exposure to social stress in adolescence leads to different behavioral outcomes [(Wommack et al., 2003)](https://www.zotero.org/google-docs/?F4Rl4o). Instead of presenting conditioned defeat, these individuals are more likely to attack smaller opponents once they reach adulthood [(Delville et al., 1998; Wommack et al., 2003)](https://www.zotero.org/google-docs/?5kCTJJ). This phenotype includes an acceleration of the maturation of agonistic behavior from play fighting to aggression [(Wommack et al., 2003)](https://www.zotero.org/google-docs/?8qWpgV), selective avoidance and enhanced risk assessment behaviors [(Bastida et al., 2009)](https://www.zotero.org/google-docs/?sHoKwy), altered motor impulsivity [(González Martínez et al., 2017; González-Martínez et al., 2020)](https://www.zotero.org/google-docs/?LapEuc), accelerated weight gain, higher food intake, enhanced body fat, long-term increased body weight and altered food-related preference behaviors [(Moran et al., 2025b, 2021)](https://www.zotero.org/google-docs/?uT1NxY). Additionally, adolescent stress alters innervation in a variety of systems involved in these behavioral and metabolic outcomes, including vasopressin, serotonin, and orexin [(Delville et al., 1998; Moran et al., 2025a)](https://www.zotero.org/google-docs/?9n1FuA). Clearly, a large number of behavioral and homeostatic-regulatory systems are impacted by adolescent stress.

In the present study, we used a global transcriptomic approach to identify potential gene targets in the hypothalamus that may be expressed due to chronic social stress exposure in adolescent male hamsters. We focused on three hypothalamic regions for analysis: lateral hypothalamus (LH), dorsomedial hypothalamus (DMH), and arcuate nucleus of the hypothalamus (ARC). We chose these regions as they are reciprocally connected and contribute to a variety of behaviors and homeostatic mechanisms that are altered in the adolescent stressed hamster model [(Cowley et al., 2001; Elias et al., 1998)](https://www.zotero.org/google-docs/?HoEzOc). The ARC and LH exert major influence over food intake and body weight, with the ARC monitoring internal energy states and sending signals to the LH, which then act downstream to influence appetitive and ingestive behaviors [(Anand and Brobeck, 1951; Keen-Rhinehart et al., 2013)](https://www.zotero.org/google-docs/?BGkUqG). The DMH is also involved in appetitive and homeostatic behaviors, as well as sex hormone production [(Bernardis, 1972; Kriegsfeld et al., 2018; Qi et al., 2009)](https://www.zotero.org/google-docs/?q7gfdd).

We predicted a number of genes or gene modules related to aggressive behavior, reward and decision making, metabolic processes, and development-related genes to be altered in stressed subjects. Specifically, given the phenotypic changes pertaining to food intake and metabolism that are observed in socially defeated adolescent hamsters, we hypothesized enhanced transcriptional changes to orexigenic processes and downregulation of anorexigenic processes.

**2. Methods**

*2.1 Animals and Housing*

Golden hamsters (*Mesocricetus auratus*) were bred in the laboratory from a colony initially obtained from Harlan Sprague–Dawley (Aura strain, Indianapolis, IN) and kept at the Animal Resource Center, an AAALAC-accredited facility. All procedures were consistent with NIH guidelines and approved by the IACUC from the University of Texas at Austin. Each litter was culled to six pups, including males and females, on postnatal day 7 (P7). On P25, all animals were weaned and single-housed in Plexiglas cages (19W x 43.2D x 26.5H cm) enriched with cotton pads for nesting materials and food piles, as this species hoards food in natural and laboratory environments [(Siegel, 1985; Waddell, 1951)](https://www.zotero.org/google-docs/?7ZXn6S).

The animals were kept in a reverse light cycle (14L-10D, lights off at 10:00 a.m.) and received food [Prolab RMH 1800 5LL2 rodent diet, Lab Supply, Dallas-Fort Worth, TX (3.15 kcal/gram; 21.2% protein, 13.7% fat, 65.1% carbohydrate)] and water *ad libitum*. For habituation prior and during behavioral tests, diet was supplemented with banana-flavored food pellets [Dustless Precision Pellets®, 45 mg, Primate Purified Diet, Banana flavor, Bio Serv, Flemington, NJ (3.45 kcal/gram; 20.2% protein, 6.3% fat, 52% carbohydrate)] as part of conditioning protocols. All behavioral procedures occurred between 11:00 am and 3:00 p.m. during the dark/active cycle. Body weights and food intake was monitored every two days. In this experiment we focus on male hamsters, as adolescent social stress exposure in females does not chronically alter cortisol levels or accelerate the development of agonistic behavior as it does in males, suggesting adolescent females habituate to repeated social stress (Taravosh-Lahn and Delville, 2004).

*2.2 Social Stress*

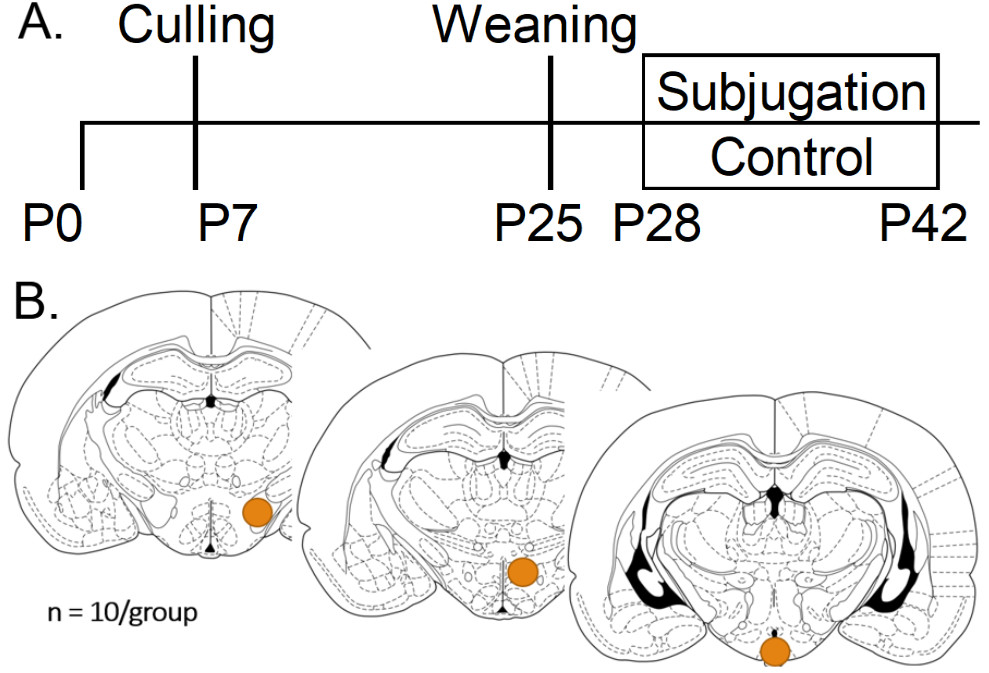
From P28 to P42, hamsters underwent either daily social stress or control handling procedures. This period corresponds to early puberty in this species [(Vomachka and Greenwald, 1979)](https://www.zotero.org/google-docs/?TLetE3). Experimental animals (Stressed) were placed in the home cage of an adult, experienced fighter male for 20 minutes [(Delville et al., 1998; Wommack et al., 2003)](https://www.zotero.org/google-docs/?59Wjb7). To control for handling and new-cage exposure, control subjects (Control) were placed into an empty, clean cage for 20 minutes (Figure 1A). While control animals eventually habituate to this process, the subjugated animals do not and continue to show a stress response at P42 [(Wommack and Delville, 2003)](https://www.zotero.org/google-docs/?SqOxKQ). All animals were checked for injuries after resident-intruder interactions, though none occurred as fights between golden hamsters rarely cause injuries [(Blanchard et al., 2003)](https://www.zotero.org/google-docs/?x5QQ4Q).

During stress, an observer recorded behaviors from the adult resident and the juvenile intruder. These recordings included the number of attacks by the resident, number of times the resident chased the intruder, number of times the intruder displayed his tail up (a mildly submissive posture), the number of time the intruder laid on his back without the resident’s physical influence (a highly submissive posture), and the number of times the resident flank marked his home cage.

*2.3 Tissue collection and RNA extraction*

20 subjects total were used in this analysis (10 Control, 10 Stressed). Subjects were rapidly decapitated 24 hours after final stress or control exposure and brains flash frozen on dry ice. Tissue was stored at -70°C until tissue punching. White adipose tissue deposits from mesenteric, epididymal, inguinal, and retroperitoneal areas were extracted and weighed, as described in previous studies [(Foster et al., 2006; Moran et al., 2021)](https://www.zotero.org/google-docs/?aWOUWi). The sum total of these is reported below alongside other recorded metabolic measures.

Regions were identified using a Golden Hamster stereotaxic brain atlas [(Morin and Wood, 2001)](https://www.zotero.org/google-docs/?9OtPKR). Tissue was extracted using Stoelting 0.75 mm tissue punch (Stoelting, Wood Dale, IL, Cat. No. 57401) centered on regions of interest: lateral hypothalamus (LH) (bregma -2.0 mm), dorsomedial hypothalamus (DMH) (bregma -2.3 mm), and arcuate nucleus of the hypothalamus (ARC) (bregma -2.9 mm) (Figure 1B). Samples were homogenized in 100 μl lysis buffer (Thermo Fisher Scientific, Waltham, MA; MagMax Total RNA isolation kit, Cat. No. A27828) with 0.7% beta-mercaptoethanol by vortexing at 3000 rpm speed for 15–20 s. Lysates were incubated at room temperature for 5 min and stored at − 80°C until RNA extraction. Once dissection of all individuals was completed, we proceed to RNA extraction using the KingFisher Flex (Thermo Fisher Scientific, Cat. No. 5400630l) with an additional DNase step to remove DNA contamination added according to the manufacturer's protocol. Samples from each brain region of all animals were extracted in the same batch. RNA quality was determined using RNA 6000 Nano Assay with BioAnalyzer (Agilent Technologies, Santa Clara, CA) and RNA concentration was determined with Quant-it RNA High Sensitivity assay kit (Thermo Fisher Scientific, Cat. No. Q33140). RNA samples were normalized to 100 ng/μl and stored at − 80°C before sequencing.



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| Figure 1. Experimental timeline and regions of interest. **A.** Diagram of experimental timeline. Postnatal day 0 (P0) represents birth. Litters were culled to 6 hamsters on P7 and weaned to individual housing on P25. Socially Stressed or Control procedure exposure occurred from P28 to P42. Animals were sacrificed on P43. **B.** Depiction of regions punched for Tag-seq. Left to right, lateral hypothalamus, dorsomedial hypothalamus, and arcuate nucleus of the hypothalamus. Punches were taken bilaterally. Brain drawings here are from the Paxinos and Watson Rat Brain Atlas [(Paxinos and Watson, 2007)](https://www.zotero.org/google-docs/?BzpaLR), though a hamster atlas [(Morin and Wood, 2001)](https://www.zotero.org/google-docs/?lqEJEt) was used to guide tissue collection. |

Samples were submitted to the Genome Sequence and Analysis Facility at the University of Texas at Austin for Tag-based RNA sequencing (Tag-seq). This method is an efficient and cost-effective approach specifically designed to measure abundances of polyadenylated transcripts yielding highly reliable data for differential gene expression analysis in well annotated genomes [(Lohman et al., 2016; Meyer et al., 2011)](https://www.zotero.org/google-docs/?4Ce5qx). It requires very few sequencing reads and is resilient to variation in sample integrity [(Stark et al., 2019)](https://www.zotero.org/google-docs/?7jbxsK). While it does not yield complete RNA sequences including splice variants (i.e. coding and non-coding RNA), it is suitable for identifying coding genes that are differentially expressed. With these transcriptome profiles, we identified differentially expressed genes (DEGs) between Stressed and Control hamsters. All samples were processed in the same batch throughout the process. Libraries were constructed with a protocol modified from [(Meyer et al., 2011)](https://www.zotero.org/google-docs/?xm0GAX) and [(Lohman et al., 2016)](https://www.zotero.org/google-docs/?6j376d). Reads were sequenced on the NovaSeq 6000 SR100 with minimum reads of 4 million and the target reads per sample of 5 million.

*2.4 Data analysis*

All statistical analyses were completed using R v4.3.2 [(R Core Team, 2014; Wickham et al., 2019)](https://www.zotero.org/google-docs/?qUj06k).

*2.4.1 Metabolic metrics analysis*

Welch’s t-tests were used to assess pairwise mean group differences in metabolic metrics. Welch’s t-tests reported as [mean ± standard deviation; t(df) t value with estimated degrees of freedom; p = p value with α = 0.05, two-tailed, d = effect size using Cohen’s d]. Trends (p < 0.1) are also reported and discussed further below.

*2.4.2 Bioinformatics analysis*

Raw RNA reads were processed to obtain gene count data by following the TagSeq data processing pipeline provided based on [(Meyer et al., 2011)](https://www.zotero.org/google-docs/?BXRXGs) and [(Lohman et al., 2016)](https://www.zotero.org/google-docs/?gkeHfz). Briefly, customized perl script utilizing FASTX-Toolkits and CUTADAPT 2.8 [(Martin, 2011)](https://www.zotero.org/google-docs/?2sW8NB) was used to remove reads with a homo-polymer run of “A” ≥ 8 bases and retain reads with minimum 20 bases and remove PCR duplicates. Processed reads were then mapped to the annotated BCM\_Maur\_2.0 *Mesocricetus auratus* genome using Bowtie2 and STAR [(Dobin et al., 2013; Harris et al., 2022)](https://www.zotero.org/google-docs/?1Y67U3). Differential gene expression analysis was conducted using Bioconductor package limma [(Ritchie et al., 2015)](https://www.zotero.org/google-docs/?zJdoqv). For each brain region, we conducted principal component analysis filtered gene counts data (filtered out genes with less than 10 counts for each sample) to visually inspect for outliers. No outliers were detected for any regions. Filtered read counts were then normalized to account for different library sizes among samples with a voom transform. Differentially expressed genes were identified between Control and Stressed subjects. We adjusted raw p-value via empirical false discovery rate (eFDR) [(Storey and Tibshirani, 2003)](https://www.zotero.org/google-docs/?RiTL5J). To estimate eFDR, we permuted sample IDs 5000 times and obtained a null distribution of p-values. The significance threshold for differentially expressed genes (DEGs) was set as 15% of change in the absolute values of log 2-fold change at the eFDR of 5%. We also performed Gene Ontology (GO) analysis to explore differences among identified DEGs within functional modules between groups using the clusterProfiler R package [(Wu et al., 2021)](https://www.zotero.org/google-docs/?c47Ejr).

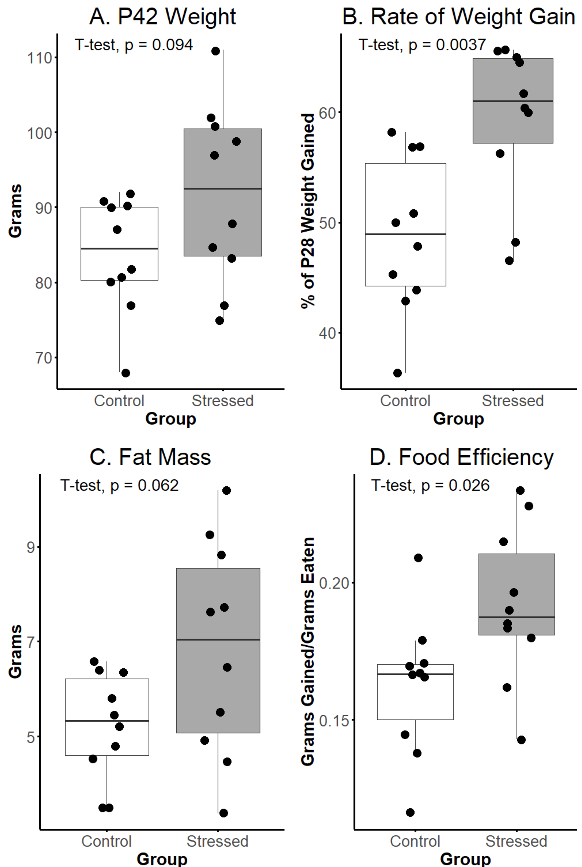
*2.4.3 Weighted gene co-expression network analysis (WGCNA)*

Weighted gene co-expression network analysis (WGCNA) was performed using the WGCNA R package [(Langfelder and Horvath, 2008)](https://www.zotero.org/google-docs/?lDNhTK). WGCNA constructs Pearson correlation matrices of RNA expression and clusters highly co-expressed genes into several modules. WGCNA then examines the association between given experimental factors and module eigengenes (MEs), assigned to colors, which are the first component from the principal component analysis (PCA) for each module. WGCNA further calculates module membership for each gene. Module membership (MM) is measured as the Pearson correlation between the gene expression level and the module eigengene, with an absolute value of module membership close to 1 indicating that the gene is highly connected to other genes in the module. Gene counts were normalized with the DESeq2 package and genes that had low expression counts (<15) in more than 75% of samples were filtered out prior to network construction. Each WGCNA model was constructed in signed hybrid mode. Power was selected by selecting a value with scale-free topology model fit above 0.8 with a minimal mean connectivity (12 for LH, 10 for DMH, and 6 for ARC). We tested whether each module eigengene was significantly different between groups using linear regression models, which is useful for identifying differential associations of functional modules and has been used in other WGCNA analysis [(Blighe, 2018; Lewis et al., 2019)](https://www.zotero.org/google-docs/?O54LxR) and report modules that are statistically significant between group (b = beta coefficient ± standard error;p = p, with α = 0.05, two-tailed). We performed GO analysis on these modules to identify functions of independently clustered modules. Finally, we extracted hub genes with very high functional module membership.

**3 Results**

*3.1 Metabolic metrics*

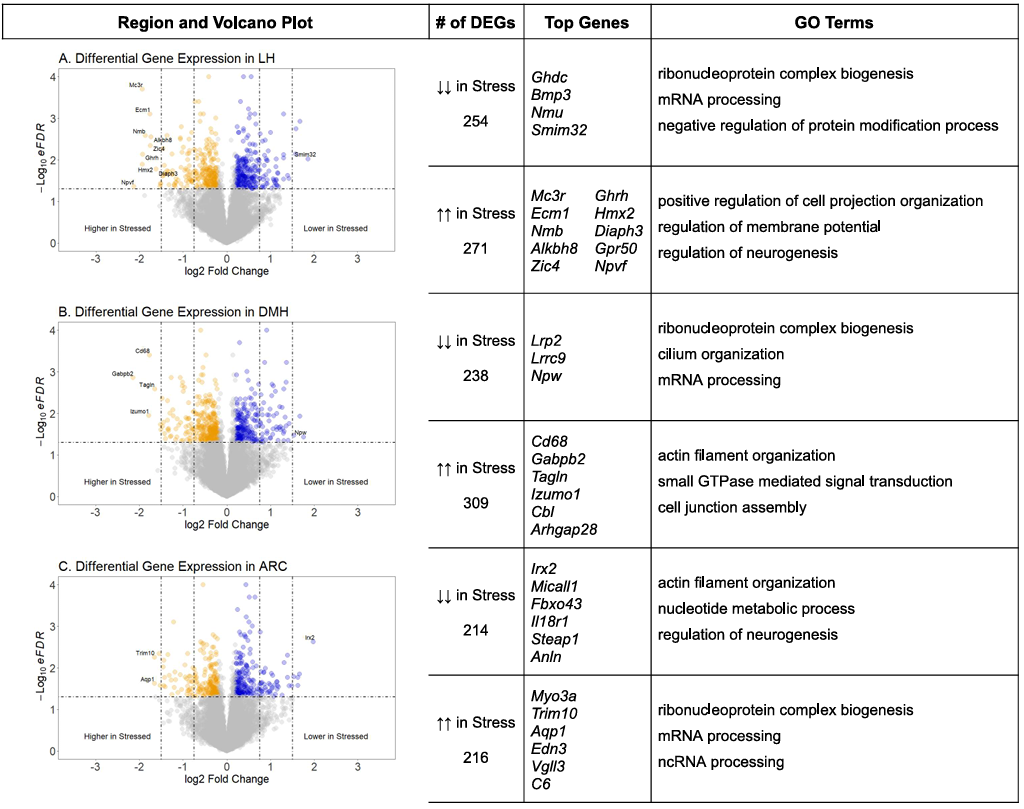
There were no group differences in body weight at P28, and Stressed subjected trended heavier at P42 [Control: 83.8 ± 7.66; Stressed: 91.8 ± 11.9; t(15.35) = 1.78, p < 0.094, d = 0.80], with Stressed subjects weighing about 10% more than controls. The rate of weight gain was significantly higher in Stressed subjects [Control: 48.89 ± 7.08; Stressed: 59.38 ± 7.01; t(18.00) = 3.33, p < 0.01, d = 1.49]. Body fat total at P42 trended higher in Stressed hamsters [Control: 5.21 ± 1.13; Stressed: 6.83 ± 2.25; t(13.27) = 2.04, p = 0.062, d = .91]. Despite no group difference in total food intake, Stressed subjects had enhanced food efficiency (gaining more weight per gram of food eaten) [Control: 0.162 ± 0.025; Stressed: 0.191 ± 0.028; t(17.75) = 2.42, p < 0.05, d = 1.09] (Figure 2).



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| Figure 2. Boxplots of significantly different metabolic measures between Stressed and Control subjects. A. Body weight at P42 in grams. B. Percent of weight at P28 gained by P42. C. Total fat mass of fat pads collected in grams. D. Total food efficiency over stress period as grams of weight gained divided by grams of food eaten. Boxplots are depicted with median line and first and third quartile ranges in boxes, with 95% range as whiskers. Points represent the y-axis value of each subject. Significance results of Welch two-sample, two-tailed t-tests are reported for each measure. |

*3.2 Stressed hamsters show differential gene expression in hypothalamic nuclei*

In the LH, out of 11,354 genes, 271 genes were upregulated in Stressed subjects and 254 genes were downregulated in Stressed subjects compared to Controls. In Figure 3A, volcano plots of gene expression and a selection of DEGs with the highest log 2-fold change and lowest eFDR are highlighted. Values of log2FC and eFDR for top genes across all regions are reported in Table 1. Gene Ontology (GO) terms are also reported in Figure 3, displaying functional modules associated with clusters of highly differentially expressed genes. In the DMH, out of 11,350 genes, 309 genes were upregulated by stress and 238 genes were downregulated by stress compared to Controls (Figure 3B). In the ARC, out of 11,444 genes, 216 genes were upregulated in the Stressed group and 214 genes were downregulated compared to Controls (Figure 3C). We also performed an overlap and conflict analysis on DEGs, compiling genes that were up- or downregulated in multiple regions in Table 2, and genes that showed differential up- or downregulation between multiple regions in Table 3. Only one gene, *Zfp36*, was upregulated across all 3 regions in Stressed subjects, and 33 other genes shared some overlap across multiple regions. 43 genes showed incongruent differential expression, being upregulated in one region and downregulated in another.



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| Figure 3. Results of differential gene expression (DEGs) in A. lateral (LH), B. dorsomedial (DMH) and C. arcuate nucleus (ARC) subregions of the hypothalamus. Genes were determined to be differentially expressed if the log2 Fold Change was >= |0.2| and the significant (eFDR) < 0.05. Top genes were identified as the greatest absolute value log2FC with an eFDR < 0.05. Functional modules identified by Gene Ontology enrichment analysis (GO terms) are on the right. |

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| **Table 1. Summary of Top Differentially Expressed Genes** | | | | | | | | |
| **LH** | | | **DMH** | | | **ARC** | | |
| **gene** | **log2FC** | **eFDR** | **gene** | **log2FC** | **eFDR** | **gene** | **log2FC** | **eFDR** |
| *Npvf* | -2.13038 | 0.0438 | *Gabpb2* | -2.14412 | 0.0014 | *Irx2* | 1.978426 | 0.0024 |
| *Hmx2* | -1.93924 | 0.013 | *Izumo1* | -1.78528 | 0.0114 | *Micall1* | 1.660793 | 0.0142 |
| *Mc3r* | -1.93781 | 0.0004 | *Cd68* | -1.76592 | 0.0004 | *Aqp1* | -1.65694 | 0.024 |
| *Ghrh* | -1.93195 | 0.0074 | *Npw* | 1.758465 | 0.037 | *Trim10* | -1.65321 | 0.0056 |
| *Nmb* | -1.86515 | 0.0026 | *Lrp2* | 1.668749 | 0.0118 | *Steap1* | 1.627907 | 0.0266 |
| *Smim32* | 1.864586 | 0.0096 | *Tagln* | -1.64991 | 0.0026 | *Il18r1* | 1.626486 | 0.0172 |
| *Ecm1* | -1.75846 | 0.0008 | *Lrrc9* | 1.529791 | 0.034 | *Myo3a* | -1.54983 | 0.0046 |
| *Zic4* | -1.75188 | 0.0046 | *Cbl* | -1.51526 | 0.0186 | *Fbxo43* | 1.53378 | 0.0164 |
| *Alkbh8* | -1.73714 | 0.0028 | *Arhgap28* | -1.50624 | 0.0246 | *Edn3* | -1.52287 | 0.0268 |
| *Ghdc* | 1.679592 | 0.0012 | *Smad6* | -1.48152 | 0.0044 | *Rgs22* | 1.487865 | 0.0272 |
| *Diaph3* | -1.62275 | 0.0168 | *Exoc3l2* | -1.47295 | 0.0204 | *Ncf2* | -1.45757 | 0.0096 |
| *Bmp3* | 1.585725 | 0.0018 | *Ctxnd1* | -1.45043 | 0.0164 | *Etfbkmt* | -1.44306 | 0.0066 |
| *Nmu* | 1.575957 | 0.0072 | *Gpr50* | 1.429156 | 0.0176 | *Trmt2b* | -1.43991 | 0.0272 |
| *Acta1* | -1.53146 | 0.044 | *Col1a1* | -1.42618 | 0.0312 | *Morn5* | 1.427706 | 0.027 |
| *Gpr50* | -1.51998 | 0.0384 | *Evpl* | -1.40366 | 0.0408 | *Siglec1* | -1.42272 | 0.017 |
| *Clrn1* | -1.51716 | 0.0364 | *Gas2l2* | 1.403227 | 0.045 | *Rac2* | -1.4084 | 0.0288 |
| *Krt77* | -1.46919 | 0.0246 | *Atp7b* | 1.367329 | 0.0006 | *Eln* | 1.391259 | 0.005 |
| *Irs4* | -1.45883 | 0.0054 | *Crtam* | 1.357569 | 0.0202 | *Jmjd4* | 1.387232 | 0.017 |
| *Isl1* | -1.43804 | 0.0234 | *Krtcap3* | 1.356831 | 0.0018 | *Ptgis* | -1.34655 | 0.0048 |
| *Slc7a14* | -1.43257 | 0.0136 | *Dock6* | -1.35372 | 0.0148 | *Atp8b1* | -1.3178 | 0.0154 |

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| Table 1. Summary of top highly differentially expressed genes across all regions analyzed. LH: lateral hypothalamus, DMH: dorsomedial hypothalamus, ARC: arcuate nucleus of the hypothalamus. log2FC: log2 fold change in expression between groups. Positive values indicate lower levels of transcript in Stressed subjects; negative values indicate higher levels in Stressed subjects. eFDR: enhanced false discovery rate, or permuted p-value. |

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| **Table 2. Differential Gene Expression Region Overlaps** | | | |
| **Gene** | **Direction** | **Regions** | **Description** |
| *Zfp36* | Up | LH, DMH, ARC | zinc finger protein 36 |
| *Apcdd1* | Up | LH, DMH | adenomatosis polyposis coli down-regulated 1 |
| *Armh3* | Up | LH, ARC | armadillo-like helical domain containing 3 |
| *Clec2l* | Up | LH, DMH | C-type lectin domain family 2, member L |
| *Cpne1* | Up | LH, ARC | copine I |
| *Ebpl* | Up | LH, ARC | emopamil binding protein-like |
| *Ecpas* | Up | LH, ARC | Ecm29 proteasome adaptor and scaffold |
| *Entpd3* | Up | LH, DMH | ectonucleoside triphosphate diphosphohydrolase 3 |
| *Gpc1* | Up | LH, DMH | glypican 1 |
| *Ltbr* | Up | DMH, ARC | lymphotoxin B receptor |
| *Mrpl3* | Up | DMH, ARC | mitochondrial ribosomal protein L3 |
| *Mtmr10* | Up | LH, DMH | myotubularin related protein 10 |
| *Nek8* | Up | LH, ARC | NIMA (never in mitosis gene a)-related expressed kinase 8 |
| *Npr1* | Up | LH, DMH | natriuretic peptide receptor 1 |
| *Sprn* | Up | LH, DMH | shadow of prion protein |
| *Tchh* | Up | LH, ARC | trichohyalin |
| *Angptl4* | Down | LH, ARC | angiopoietin-like 4 |
| *Aopep* | Down | DMH, ARC | aminopeptidase O |
| *Ccdc18* | Down | LH, DMH | coiled-coil domain containing 18 |
| *Clk1* | Down | LH, DMH | CDC-like kinase 1 |
| *Fos* | Down | LH, ARC | FBJ osteosarcoma oncogene |
| *Hmgn5* | Down | LH, DMH | high-mobility group nucleosome binding domain 5 |
| *Icam2* | Down | LH, ARC | intercellular adhesion molecule 2 |
| *Med9* | Down | DMH, ARC | mediator complex subunit 9 |
| *Nek3* | Down | DMH, ARC | NIMA (never in mitosis gene a)-related expressed kinase 3 |
| *Nt5c2* | Down | LH, DMH | 5'-nucleotidase, cytosolic II |
| *Parp12* | Down | LH, ARC | poly (ADP-ribose) polymerase family, member 12 |
| *Pex12* | Down | LH, DMH | peroxisomal biogenesis factor 12 |
| *Rad23a* | Down | LH, DMH | RAD23 homolog A, nucleotide excision repair protein |
| *Rspry1* | Down | DMH, ARC | ring finger and SPRY domain containing 1 |
| *Smim15* | Down | LH, DMH | small integral membrane protein 15 |
| *Steap1* | Down | DMH, ARC | six transmembrane epithelial antigen of the prostate 1 |
| *Tent5a* | Down | DMH, ARC | terminal nucleotidyltransferase 5A |
| *Tut1* | Down | DMH, ARC | terminal uridylyl transferase 1, U6 snRNA-specific |

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| Table 2. Differential gene expression overlap analysis. Direction indicates if genes were up- or downregulated in socially stressed hamsters and in which regions. |

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| **Table 3. Differential Gene Expression Region Incongruence** | | | |
| **Gene** | **Region Up** | **Region Down** | **Description** |
| *Tchh* | LH, ARC | DMH | trichohyalin |
| *Abhd14a* | LH | ARC | abhydrolase domain containing 14A |
| *Btg1* | LH | DMH | BTG anti-proliferation factor 1 |
| *Cckar* | LH | DMH | cholecystokinin A receptor |
| *Efl1* | LH | DMH | elongation factor like GTPase 1 |
| *Gjc1* | LH | ARC | gap junction protein, gamma 1 |
| *Gpr50* | LH | DMH | G-protein-coupled receptor 50 |
| *Gtf2h3* | LH | ARC | general transcription factor IIH, polypeptide 3 |
| *Hdac4* | LH | DMH | histone deacetylase 4 |
| *Htr1b* | LH | DMH | 5-hydroxytryptamine (serotonin) receptor 1B |
| *Ift88* | LH | DMH | intraflagellar transport 88 |
| *Irs4* | LH | DMH | insulin receptor substrate 4 |
| *Med24* | LH | ARC | mediator complex subunit 24 |
| *Mustn1* | LH | ARC | musculoskeletal, embryonic nuclear protein 1 |
| *Nol12* | LH | DMH | nucleolar protein 12 |
| *Nsun7* | LH | DMH | NOL1/NOP2/Sun domain family, member 7 |
| *Pdyn* | LH | DMH | prodynorphin |
| *Rcc1l* | LH | DMH | reculator of chromosome condensation 1 like |
| *Riox1* | LH | ARC | ribosomal oxygenase 1 |
| *Slc16a11* | LH | ARC | solute carrier family 16 (monocarboxylic acid transporters), member 11 |
| *Slc7a14* | LH | DMH | solute carrier family 7 (cationic amino acid transporter, y+ system), member 14 |
| *Vps18* | LH | ARC | VPS18 CORVET/HOPS core subunit |
| *Desi1* | DMH | LH | desumoylating isopeptidase 1 |
| *Dut* | DMH | LH | deoxyuridine triphosphatase |
| *Fkbp15* | DMH | LH | FK506 binding protein 15 |
| *Gpr146* | DMH | ARC | G protein-coupled receptor 146 |
| *Irx2* | DMH | ARC | Iroquois homeobox 2 |
| *Mid1ip1* | DMH | ARC | Mid1 interacting protein 1 (gastrulation specific G12-like (zebrafish)) |
| *Mss51* | DMH | LH | MSS51 mitochondrial translational activator |
| *Mtg2* | DMH | LH | mitochondrial ribosome associated GTPase 2 |
| *Pex7* | DMH | ARC | peroxisomal biogenesis factor 7 |
| *Sidt1* | DMH | LH | SID1 transmembrane family, member 1 |
| *Skida1* | DMH | LH | SKI/DACH domain containing 1 |
| *Slc9a3r2* | DMH | LH | solute carrier family 9 (sodium/hydrogen exchanger), member 3 regulator 2 |
| *Smad6* | DMH | ARC | SMAD family member 6 |
| *Tmc7* | DMH | LH | transmembrane channel-like gene family 7 |
| *Tph2* | DMH | ARC | tryptophan hydroxylase 2 |
| *Bmp3* | ARC | LH | bone morphogenetic protein 3 |
| *Fnbp4* | ARC | DMH | formin binding protein 4 |
| *Iws1* | ARC | DMH | IWS1, SUPT6 interacting protein |
| *Snrpb2* | ARC | DMH | U2 small nuclear ribonucleoprotein B |
| *Stim2* | ARC | DMH | stromal interaction molecule 2 |
| *Zcchc8* | ARC | DMH | zinc finger, CCHC domain containing 8 |

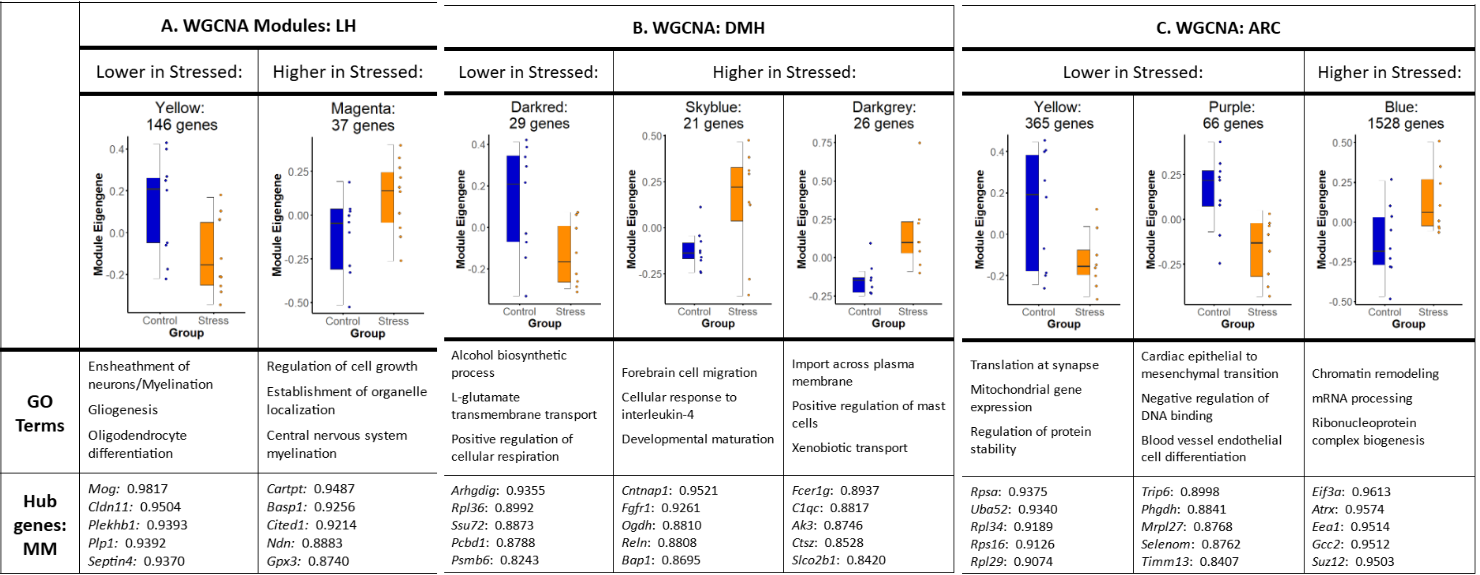
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| Table 3. Differential gene expression incongruence analysis. First organized by which regions incongruent genes were upregulated in stressed subjects, then which regions they were downregulated in. |

*3.3 WGCNA reveals co-expressed gene modules uniquely associated with social stress*

Supplemental Figure 1 summarizes all WGCNA modules identified. Modules differentially expressed between groups are highlighted in Figure 4, including the GO terms most associated with differentially expressed modules. Genes with high module membership (MM - genes that are most highly correlated with other genes in the module) are also listed. In the LH, WGCNA identified 14 modules of highly correlated genes. Two of these (yellow and magenta) exhibited significant differences in ME expression between groups (Figure 4A). The yellow module (b = -0.224 ± 0.097, p < 0.05) was most strongly associated with myelination and had ME scores that were significantly downregulated in the LH. The magenta module (b = 0.212 ± 0.099, p < 0.05) was also associated with myelination processes along with cell growth and organization and was upregulated in stressed individuals.

In the DMH, WGCNA identified 31 modules. Three of these (darkred, skyblue, and darkgrey) were differentially expressed between groups (Figure 4B). The darkred module (b = -0.251 ± 0.107, p < 0.05) was associated with alcohol related bioprocesses and glutamate transport and was downregulated with stress. The skyblue module (b = 0.254 ± 0.107, p < 0.05) was associated with neuronal development and immune function and was upregulated in stress. The darkgrey module (b = 0.315 ± 0.095, p < 0.01) was also upregulated in stress and was also associated with membrane transport and immune function.

In the ARC, WGCNA identified 11 modules. Three of these (yellow, purple, and blue) were differentially expressed between groups (Figure 4C). The yellow module (b = -0.235 ± 0.110, p < 0.05) was associated with synaptic function and cellular energy use and was lower in Stressed subjects. The purple module (b = -0.321± 0.095, p < 0.01) was also lower with Stressed animals and was associated with DNA processing and blood vessel function. The blue module (b = 0.262 ± 0.106, p < 0.05) was upregulated in Stressed subjects and was associated with mRNA processing and epigenetic functions.



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| Figure 4. Results of weighted gene coexpression network analysis (WGCNA) in A. lateral (LH), B. dorsomedial (DMH) and C. arcuate nucleus (ARC) subregions of the hypothalamus. All noted modules are significantly differentially expressed between groups, with boxplots of module eigengene value shown per region. Below are Gene Ontology (GO) terms associated with differentially expressed modules, and hub genes with highest module membership (MM). |

**4. Discussion**

Stressed hamsters gained weight at an increased rate, had higher food efficiency, and tended to have more body fat than controls, supporting prior work in this model [(Moran et al., 2021)](https://www.zotero.org/google-docs/?cyaD99). Overall, approximately 500 genes were differentially expressed between groups in each region sampled. In the present analysis, we focused on hypothalamic regions functionally associated with appetite and other metabolic processes. We observed changes in expression of a number of orexigenic transcripts across regions. Due to prior research in the lab, we had a particular interest in LH orexin/hypocretin, expecting its expression to be elevated in stressed subjects that gain more weight than controls. Interestingly, there was only a trend that LH orexin expression (*hcrt*) was upregulated in stressed hamsters. This was not necessarily surprising, given prior results from our lab that suggests stress-induced differences in orexin are not at the innervation level, but more likely caused by downstream receptor changes or other firing mechanisms [(Moran et al., 2025a)](https://www.zotero.org/google-docs/?znVemS). However, growth hormone releasing hormone (*ghrh*) transcript was upregulated in the LH of stressed subjects. It was recently reported that activation of growth hormone receptors in LH orexin neurons promotes their cellular and related behavioral functions [(Tavares et al., 2024)](https://www.zotero.org/google-docs/?sOFPxm), which could be related to a change in orexin firing, if not orexin expression or innervation. Additionally, the LH WGCNA magenta module that was upregulated in stressed subjects was also related to cellular growth. Neuromedin U was also downregulated in the LH. In rats, this neuropeptide is typically expressed in the arcuate nucleus, and has an anorexic influence [(Howard et al., 2000)](https://www.zotero.org/google-docs/?q1cIaT), making its downregulation in our enhanced eaters unsurprising. NMU downregulation may be a compensatory mechanism, as NMU administration induces stress-response behaviors such as self-grooming in rats and mice [(Hanada et al., 2001)](https://www.zotero.org/google-docs/?tlyQHz). Similarly, neuropeptide W (NPW) was downregulated in the DMH. NPW activity inhibits food intake and stimulates thermogenesis [(Mondal et al., 2003)](https://www.zotero.org/google-docs/?zEDisF). Interestingly, NPW is widely involved in other stress, growth, and sex hormone related processes, making it an excellent candidate for further study in this model [(Baker et al., 2003; Niimi and Murao, 2005)](https://www.zotero.org/google-docs/?YHpqNt).

There was also upregulation of Neuropeptide VF precursor (*Npvf*) in the LH in stressed hamsters. *Npvf* is a precursor peptide for RFRP-3, which makes the upregulation in the LH and absence of change in the DMH interesting. In hamsters, occurrence of RFRP-3 expressing cell bodies can extend to perifornical areas [(Kriegsfeld et al., 2010)](https://www.zotero.org/google-docs/?Q9HGBA). While the majority of DMH *Npvf*-expressing cells did not differ between groups, a subset of these more laterally located neurons are upregulated by stress. Nevertheless, possible upregulation of RFRP-3, the formerly-putative gonadotropin inhibitory hormone [(Kriegsfeld et al., 2018, 2010; Yano et al., 2003)](https://www.zotero.org/google-docs/?ptn94E), aligns with prior findings of short-term, 50% reductions in testosterone in stressed male hamsters [(Wommack et al., 2004)](https://www.zotero.org/google-docs/?q9M52T). Stressed adolescent male hamsters also tend to hoard more food [(Moran et al., 2025b, 2021)](https://www.zotero.org/google-docs/?uiB927), and RFRP-3 is involved in food hoarding behavior in hamsters, though more so in low estrogen-state females [(Benton et al., 2018)](https://www.zotero.org/google-docs/?vekwgD). Additionally, in mice, whole hypothalamic *Npvf* expression is decreased in cold temperatures alongside enhanced thermogenesis, regardless of diet, leptin, or body fat [(Jaroslawska et al., 2015)](https://www.zotero.org/google-docs/?HpLZ3c), suggesting it may have a metabolic role independent of RFRP-3 that yet functionally coordinates with its gonadotropin inhibitory role. *Npvf* upregulation may also alter sleep patterns, as *Nvpf* expression is necessary for sleep in zebrafish [(Lee et al., 2017)](https://www.zotero.org/google-docs/?pjhAae). Here, *Nvpf* may be playing competing roles regarding metabolic processes that require further study.

In the LH, the melanocortin 3 receptor was significantly upregulated in stressed subjects. *Mc3r* codes for a receptor of melanocyte-stimulating hormone, which has anorexigenic downstream effects [(Maniam and Morris, 2012; Roselli-Rehfuss et al., 1993)](https://www.zotero.org/google-docs/?PXGgTa). Since the LH is also the home of α-MSH containing neurons [(Legrand et al., 2015; Olszewski et al., 2001)](https://www.zotero.org/google-docs/?c0UdP7), MC3R upregulation suggests a local action altered by stress, potentially in competing homeostatic processes.

We also observed differential up- and down-regulation of the orphan receptor G protein-coupled receptor 50 (*Gpr50*) between the LH and DMH. *Gpr50* is involved in cellular leptin sensitivity and leptin-induced thermogenesis in mice [(Bechtold et al., 2012; Khan et al., 2016; Sidibe et al., 2010)](https://www.zotero.org/google-docs/?UQ46i6). These processes are likely altered by stress in hamsters, both in relation to body fat gain and a number of other differentially expressed genes. We observed no *Gpr50* expression in the ARC (which very likely included tissue from the median eminence due to their proximity) despite consistent expression in the median eminence in other rodents and humans [(Sidibe et al., 2010)](https://www.zotero.org/google-docs/?o2k1eX). *Gpr50* is also necessary for cellular stress-induced mitochondrial autophagy responses [(Liu et al., 2024)](https://www.zotero.org/google-docs/?nkOjw3), deepening its metabolic role. Along this line, the DMH WGCNA darkred module, which is involved in cellular respiration, was also downregulated in stressed subjects. The five most central hub genes in this module have few functions classically associated with stress exposure, emphasizing the importance of analyzing transcriptomic datasets from more than just a DEG perspective, and highlighting these genes as potential focal points of future studies. Additionally, *selenom*, a hub gene in the downregulated ARC WGCNA purple module, has also been previously associated with the development of insulin resistance, obesity, and neuroinflammation when it is exogenously inhibited or knocked-out [(Cai et al., 2024; Lin et al., 2023)](https://www.zotero.org/google-docs/?W9CDpt). While others have investigated its role in the hippocampus [(Lin et al., 2023)](https://www.zotero.org/google-docs/?gCZ5Gu) and adipose tissue [(Cai et al., 2024)](https://www.zotero.org/google-docs/?iyarso), its role in the hypothalamus is a potential target of future research.

Other genes in the downregulated ARC WGCNA purple module and the DMH darkred module are associated with ribosomal and mitochondrial ribosomal proteins (*Rps*, *Rpl*, and *Mrpl*). However, Gene Ontology terms derived from differential gene expression suggest opposing patterns. In the ARC, one highly upregulated GO term was ribonucleoprotein complex biogenesis, while the same GO term was highly downregulated in the LH and DMH in stressed subjects. Others have found that chronic social stress upregulates many of these ribosomal genes in whole-hypothalamus RNAseq in C57BL/6J mice [(Smagin et al., 2016)](https://www.zotero.org/google-docs/?U7mhPl). Our findings clearly show there is more nuance to these mechanisms, requiring further study.

We observed a number of transcriptomic changes supporting our hypothesis that stress induces alterations to development-related gene expression. Across regions, a variety of both up- and down-regulated DEGs, GO terms, and WGCNA modules were related to RNA processing, development, neurogenesis, and maturation. As these data are from adolescent subjects, changes in development-related processes is not surprising. In fact, these changes could reflect longer-term alterations in gene expression. For example, in mice, similar developmental modules, such as those involved in multicellular organism development, nervous system development, and cell differentiation, are altered in adult animals that were exposed to early life stress, [(Peña et al., 2019)](https://www.zotero.org/google-docs/?QrPrKO), suggesting that these changes can be quite long-lasting. Nevertheless, a primary GO term related to highly DEGs, mRNA processing, was downregulated in stressed hamsters in the LH and DMH, but upregulated in the ARC. This somewhat aligns with WGCNA-derived modules in these regions. The ARC WGCNA blue module was upregulated and contributes to RNA processing. Many of the hub genes in this module are directly related to chromatin remodeling, which is well known to be altered by stress [(Smith and Workman, 2012)](https://www.zotero.org/google-docs/?33IQRu), and suggests potentially long-term epigenetic changes in this model. Despite these changes, it is interesting that we observed significant upregulation of growth hormone releasing hormone transcript (*ghrh*) in the LH, but not other hypothalamic regions.

WGCNA highlighted modules related to myelination both up- (LH magenta) and downregulated (LH yellow) by stress in the LH. Others have shown links between social dominance behaviors and myelination. In mice exposed to a shift in social environment that retained a prior dominant status in a previous colony, social reorganization was associated with myelination and oligodendrocyte development genes and modules in the amygdala [(Milewski et al., 2024)](https://www.zotero.org/google-docs/?4pMbPc). Direct chronic social defeat in mice also induces downregulation of myelination related gene expression, followed by reduction in oligodendrocyte number and myelin protein levels and thickness in the medial prefrontal cortex, hippocampus, and nucleus accumbens [(Bonnefil et al., 2019; Lehmann et al., 2017)](https://www.zotero.org/google-docs/?LiEtGo), though there is some variability in the prefrontal cortex and basolateral amygdala [(Cathomas et al., 2019; Poggi et al., 2022)](https://www.zotero.org/google-docs/?cf63ZO). Our results are interesting in conjunction with the fact that adolescent-stressed hamsters become more socially aggressive, suggesting that myelinating processes may play a broader role in response to stress and social behaviors. Furthermore, rats that experienced chronic social stressors from mid- to late-adolescence have altered hippocampal white matter structure, especially paired with obesogenic dietary conditions [(Ontiveros-Ángel et al., 2024)](https://www.zotero.org/google-docs/?2BPoV2). In humans that experienced early life abuse, these traits are correlated, such that in abused individuals, increased BMI is associated with decreased forebrain white matter connectivity, particularly with the lateral hypothalamus [(Luo et al., 2020)](https://www.zotero.org/google-docs/?mkRyiV). Hub genes associated with these myelination-related modules included *mog*, myelin oligodendrocyte glycoprotein, and *cartpt*, cocaine and amphetamine regulated transcript (CART) preprotranscript. While the role of *mog* in response to stress is less understood, other studies have found that *cartpt* expression is positively correlated with stress susceptibility in the hippocampus [(Long et al., 2023)](https://www.zotero.org/google-docs/?AWQ7MI), but positively associated with stress resilience in the anterior cingulate [(Funayama et al., 2023)](https://www.zotero.org/google-docs/?9FS8MA). The fact that we observed no differential expression of *cartpt* transcript despite its role in a functionally altered WGCNA module suggests increased study of CART in the LH is warranted. Thus, the effects of stress on myelinating processes and their interaction with metabolic function is important for continued research.

Gene expression in our ARC samples was not as expected in some aspects. While others have found that adolescent stress in rats resulted in reduced *Pomc* and *Cartpt* expression in the ARC alongside reduction in body weight [(Krolick et al., 2022)](https://www.zotero.org/google-docs/?48Hz1X). Though it is interesting that we observed no changes to the typical energy-sensing systems in the ARC, this species difference in gene expression may align with species typical metabolic outcomes [(Moran and Delville, 2024)](https://www.zotero.org/google-docs/?OKDFaF). While we did observe differential expression in over 400 genes in the ARC, many of them, such as *Aqp1*, are involved in ventricle function [(Venero et al., 2001)](https://www.zotero.org/google-docs/?75TeKu), or immune function, such as *Ncf2* or *Crlf1* [(Crisponi et al., 2022; Fernandez-Twinn et al., 2012)](https://www.zotero.org/google-docs/?DAZPJF). WGCNA also highlighted the purple module in the ARC, related to blood vessel development, which was downregulated in stressed subjects. These processes may be related to stress and obesity through the neuroinflammatory effects of both of these conditions [(Ontiveros-Ángel et al., 2024)](https://www.zotero.org/google-docs/?4PYCIt), thus causing changes in information received by metabolic-sensory arcuate neurons instead of gross changes in their expression. Follow up studies are needed to investigate this further. One of the downregulated GO terms derived from highly DEGs in the ARC was related to actin filament organization, further suggesting impairment of mechanical cellular sensory mechanisms.

More broadly, a plasma membrane permeability related module in the DMH (darkgrey) was upregulated in stressed subjects. One hub gene in this module, *c1qc*, encodes the complement C1q C chain. *C1qc* is a component of the complement cascade, relevant for proper immune function. Interestingly, upregulation of *c1qc*-related proteins in individuals that experienced trauma at age 12 was positively correlated with likelihood of psychotic experiences at age 18 [(Föcking et al., 2021)](https://www.zotero.org/google-docs/?bwDWY1). Furthermore, *c1qc* is upregulated in atherosclerotic arteries [(Cui et al., 2023)](https://www.zotero.org/google-docs/?2eb1cS). Therefore both behaviorally and physiologically, elements of the complement cascade warrant further study in the stressed hamster model. The DMH WGCNA skyblue module was also related to immune function, specifically cellular response to interleukin-4. Hub genes here included *fgfr1*, fibroblast growth factor receptor. *Fgfr1* has been more classically associated with obesogenic states in adipocytes [(Scioli et al., 2014; Ye et al., 2016)](https://www.zotero.org/google-docs/?dVNoEC), and other fibroblast growth factor mechanisms are involved in the neural response to stress hormones [(Molteni et al., 2001)](https://www.zotero.org/google-docs/?fxmtyP). Finally, the upregulation of *zfp36* across all regions sampled in stressed subjects is notable. *Zfp36*, zinc finger protein 36, is also involved in immune-inflammatory responses and could be a critical hypothalamic target of future study [(Makita et al., 2021)](https://www.zotero.org/google-docs/?cwwWMv).

Overall, the study outcome supported our initial hypotheses. While our focus was on appetite- and obesity-related genes, it is clear that the experience causes profound alterations in multiple systems. The present data highlight systems related to myelination and immune function in the brain from multiple analytical perspectives as potential drivers of stress-induced changes. Future studies will likely further relate our findings to the behavioral and metabolic consequences of social stress during adolescence, whether they result in short- or long-term effects.

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