REM sleep’s unique associations with corticosterone regulation, apoptotic pathways, and behavior in chronic stress in mice

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One of sleep’s putative functions is mediation of adaptation to waking experiences. Chronic stress is a common waking experience; however, which specific aspect of sleep is most responsive, and how sleep changes relate to behavioral disturbances and molecular correlates remain unknown. We quantified sleep, physical, endocrine, and behavioral variables, as well as the brain and blood transcriptome in mice exposed to 9 weeks of unpredictable chronic mild stress (UCMS). Comparing 46 phenotypic variables revealed that rapid-eye-movement sleep (REMS), corticosterone regulation, and coat state were most responsive to UCMS. REMS theta oscillations were enhanced, whereas delta oscillations in non-REMS were unaffected. Transcripts affected by UCMS in the prefrontal cortex, hippocampus, hypothalamus, and blood were associated with inflammatory and immune responses. A machine-learning approach controlling for unspecific UCMS effects identified transcriptomic predictor sets for REMS parameters that were enriched in 193 pathways, including some involved in stem cells, immune response, and apoptosis and survival. Only three pathways were enriched in predictor sets for non-REMS. Transcriptomic predictor sets for variation in REMS continuity and theta activity shared many pathways with corticosterone regulation, in particular pathways implicated in apoptosis and survival, including mitochondrial apoptotic machinery. Predictor sets for REMS and anhedonia shared pathways involved in oxidative stress, cell proliferation, and apoptosis. These data identify REMS as a core and early element of the response to chronic stress, and identify apoptosis and survival pathways as a putative mechanism by which REMS may mediate the response to stressful waking experiences.

Significance

Sleep disturbances are common in stress-related disorders but the nature of these sleep disturbances and how they relate to changes in the stress hormone corticosterone and changes in gene expression remained unknown. Here we demonstrate that in response to chronic mild stress, rapid-eye-movement sleep (REMS), a sleep state involved in emotion regulation and fear conditioning, changed first and more so than any other measured sleep characteristic. Transcriptomic profiles related to REMS continuity and theta oscillations overlapped with those for corticosterone, as well as with predictors for anhedonia, and were enriched for apoptotic pathways. These data highlight the central role of REMS in response to stress and warrant further investigation into REMS’s involvement in stress-related mental health disorders.

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Data deposition: The data reported in this paper have been deposited in the Gene Expression Omnibus (GEO) database, https://www.ncbi.nlm.nih.gov/geo (accession no. GSE125441).

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leep is assumed to contribute to recovery from the wear and tear of wakefulness and to mediate adaptation to the waking experience, be it through memory consolidation or processing of emotional experiences, such as those associated with stressful events (1). Chronic stress is the most significant predictor of mood disorders (2) and major depressive disorder is anticipated to be the leading cause of disease burden by 2030 (3), while the true global burden of stress-related mental diseases might be largely underestimated (4). In animals, chronic stress leads to profound physiological changes, such as hypothalamic–pituitary–adrenal (HPA) axis regulation of corticosterone, neurogenesis, synaptic plasticity, and gene expression (5, 6). Chronic stress also leads to a plethora of behavioral disturbances, including depressive-like behavior, decreased responsiveness to rewards akin to anhedonia, a core symptom of depression, and sleep alterations (1, 7). The effects of chronic stress on sleep in rodents have been studied by applying physical, social, and environmental stressors. Several of these studies documented alterations in rapid-eye-movement sleep (REMS) and sleep continuity (8–11), while others reported changes in non-REMS (NREMS) and electroencephalogram (EEG) slow wave (delta) activity (12). In humans, chronic stress, alterations in the HPA axis regulating cortisol, and sleep disturbances have been associated with mood disorders (13, 14). However, the nature of sleep disturbances in major depression continues to be discussed, with some studies highlighting changes in NREMS (15, 16), and others REMS and sleep continuity (14, 17). Unresolved questions are how the various physiological and behavioral consequences of chronic stress interrelate and whether specific changes in sleep are early and core symptoms contributing to adaptation to chronic stress.

Stress triggers changes in gene expression in the brain and these transcriptome responses have been shown to be highly tissue/brain region-specific (6). Most studies have focused on the

depression | anhedonia | EEG theta power | machine learning | transcriptome
hippocampus and prefrontal cortex, identifying differential expression of genes related to inflammation, immune response, and neurogenesis (18–20). While transcriptomic changes underlyng neuroplastic adaptation to chronic stress have been extensively studied in the brain, very few animal studies have investigated the transcriptome response to stress in blood (21, 22). This is of interest in the context of translational studies because blood transcriptomic signatures of depression and treatment response have been identified in humans (23–25). Finally, the extent to which sleep and other behavioral and endocrine alterations in response to stress are related to changes in the transcriptome has not yet been comprehensively quantified.

Here, exposure to chronic stress was achieved using the well-validated unpredictable chronic mild stress (UCMS) paradigm in mice (7). UCMS elicits a broad range of physiological and ethological changes that are consistent with symptoms of major depressive disorder, and predicts the efficacy of antidepressant treatments (7, 26). This ethological “model” has been recognized for its high-translational potential in the context of stress-related disorders (26–28). The aims of the current study were: (i) to comprehensively characterize chronic stress-induced changes in REMS and NREMS, corticosterone, and behavioral variables, as well as the transcriptome in three stress- and sleep-related brain regions (hippocampus, prefrontal cortex, hypothalamus) and blood; and (ii) to investigate the interrelationship of these responses using machine learning and other robust statistical approaches.

Results

Stress-Induced Physical, Neuroendocrine, and Behavioral Disturbances. We assessed the impact of the repeated exposure to an unpredictable stressful waking experience on a number of physiological and behavioral variables during the 9-wk protocol (Fig. 1A). Chronic mild stress significantly altered body weight and worsened coat state, an index of reduced grooming behavior (Fig. 1 B and C). Corticosterone regulation was compromised in the UCMS group, consistent with blunted HPA axis negative feedback (Fig. 1D). The dexamethasone (DEX)-induced corticosterone suppression results are not explained by handling and injection because the response to saline injection was not different between groups (P = 0.657) (SI Appendix, Fig. S1). Self-care behavior was reduced, as reflected by increased grooming latency and decreased grooming duration (Fig. 1 E and F). Quality of nest building, indicative of motivation, was also reduced in the UCMS group (Fig. 1G). Moreover, UCMS suppressed the progressive increase of consumption of a palatable stimulus, indicative of anhedonia (Fig. 1 J and K). Immobility during the forced swim test was increased (Fig. 1L), as was anxiety-like behavior (Fig. 1M).

Social disturbances were observed with increased aggressive behavior (i.e., decrease of attack latency and increased number of attacks) (Fig. 1N and Dataset S1), and decreased social preference for the novel congner (Fig. 1O). Exposure to UCMS reduced the weekly averaged locomotor activity during the dark (active) phase of the light-dark cycle, while activity remained unaffected during the light phase (Fig. 1 H and I and SI Appendix, Fig. S2). The lower locomotor activity of UCMS-subjected mice was also observed on stress-free days (i.e., “day”; P = 0.0099) (Fig. 2E). Expressed as a percentage of TST, REMS was increased, and these changes were observed both during the light and dark phases (SI Appendix, Fig. S3 C and D). Chronic mild stress also induced an increase in REMS continuity, with increased duration of REMS episodes (Fig. 2B) despite increased number of REMS episodes during both the light and dark phases (SI Appendix, Fig. S3 G, H, K, and L). In contrast, NREMS became more fragmented with an increased number of episodes of shorter duration (Fig. 2F and SI Appendix, Fig. S3 E, F, I, and J).

Quantitative EEG analysis, using baseline measurements as a covariate to control for individual differences in the EEG power spectra, showed that theta activity’s EEG hallmark of REMS was increased in the light (Fig. 2C) and dark phases (SI Appendix, Fig. S3 O and P). In contrast, NREMS delta activity was not affected by UCMS (Fig. 2G and SI Appendix, Fig. S3 M and N). Computation of relative EEG power spectra showed that changes in REMS were indeed mainly observed in the theta range, although some reduced activity in lower and higher frequencies was detected (Fig. 2D and SI Appendix, Fig. S4A). The increase in theta is not directly related to the duration of REMS bouts because power is a density measure that does not necessarily increase with bout duration. To further explore this issue, we compared theta power in long and short REMS bouts and nevertheless found that theta power is higher in long REMS bouts than in short REMS bouts in both UCMS and control groups (Dataset S1). We then compared theta power associated with long and short REMS bout lengths in light and dark periods before and after UCMS. We found that in both short and long REMS bouts, theta power was higher in the UCMS group, except for short REMS bouts in the light phase (Dataset S1).

In contrast to REMS, only minor changes were observed in the relative NREMS EEG power spectra (Fig. 2H and SI Appendix, Fig. S4B).

Temporal Associations of Phenotypic Alterations. The changes in 24-h REMS duration, and other measures of sleep duration across 24 h or during the light phase, were observed as early as day 3 of the UCMS protocol (Fig. 2A and SI Appendix, Figs. S3 A–D and S5). Degradation of coat state occurred from day 7, while differences in body weight, impairment of corticosterone regulation, self-centered behavior, and motivation appeared in weeks 3 and 4 (Fig. 1B and SI Appendix, Figs. S3 B and S4A). Locomotor activity in the dark period was reduced in the UCMS group during the last wk of the 9-wk protocol (SI Appendix, Fig. S2 B–E).

Effect Size and Stability of Chronic Stress Effects Across Phenotypes. The size of the effects of UCMS varied considerably across dependent measures, with the largest effect sizes observed for coat state, 24-h REMS duration, corticosterone regulation, and 24-h REMS expressed as percentage of TST (Fig. 3). Overall, most REMS and NREMS variances, including the number and length of sleep episodes, displayed a large (Cohen’s f^2 > 0.4) or medium effect size (Cohen’s f^2 > 0.25) (Fig. 3). Across behaviors, effect sizes of UCMS were large for despair behavior, aggression, self-centered behavior, social disturbances, anxiety-like behavior, and motivation. The impact of UCMS on 24-h TST, 24-h NREMS duration, and EEG delta power was small (Fig. 3). In addition, to assess to which extent UCMS-induced changes were stable within individuals, intraclass correlation (ICC) coefficients were computed for all dependent variables. ICCs ranged between 0.67 and 0.997 for body weight, locomotor activity, REMS EEG theta power, and NREMS delta power, suggesting that the response to UCMS is highly stable (i.e., ICC > 0.61 benchmarks defined by ref. 29) within individuals. Coat state, as well as REMS and NREMS expressed as a percentage of TST for 24-h, showed moderate trait stability (ICC = 0.5240 and 0.4671, respectively). Corticosterone regulation displayed a slight stability (ICC = 0.0066) (Dataset S1).

Bivariate Associations Between Phenotypes. To assess the strength of associations between measured variables, we computed Kendall’s partial correlations between pairs of symptoms induced by chronic mild stress. We controlled for the effect of “group” (i.e., control vs. UCMS) to identify bivariate associations at the level of the individual independent of “unspecific” group effects. The increased percentage of REMS per TST (during the light phase and for 24 h) correlated negatively with DEX suppression; that is, more
REM was associated with the impairment of corticosterone regulation \( [\tau = 0.72, \text{nominal } P \text{ value } (P_{\text{nom}}) = 0.00034, \text{false-positive rate (FDR)-adjusted } P \text{ value (} P_{\text{adj}}) = 0.0252 \text{ and } \tau = 0.71, P_{\text{nom}} = 0.00037, P_{\text{adj}} = 0.0211, \text{respectively} ] \) (Fig. 4A and B). These associations were of a large effect size defined by \( \tau = 0.25 \) (30). Other large effect-size associations were observed; however, they reflected trivial relationships among dependent sleep variables (e.g., percentage of REMS and NREMS per TST).

**Effects of Chronic Stress on the Transcripome.** To gain insight into the molecular mechanisms underlying the phenotypes induced by UCMS, we performed RNA sequencing on three brain regions and whole-blood samples collected at the end of the UCMS paradigm.

**Differential gene expression and functional enrichment.** We first performed differential expression analysis between the UCMS and control groups. The number of differentially expressed genes (DEGs) was relatively small (range across the three brain regions and blood: 40–194) and the number of up-regulated genes was larger than the number of down-regulated genes in all tissues (Dataset S3). The fold-changes were relatively small (range of log_{2}-transformed fold-change: −1.65 to 1.18 (Dataset S3). The comparison of transcriptomic responses in the four tissues showed a robust overlap of DEGs between the prefrontal cortex and the hippocampus, while the commonalities between other tissues were weaker (Fig. 5A; for identity of these overlapping DEGs, see SI Appendix, Figs. S6 and S3). The three brain regions had only six common DEGs, encoding hemoglobin subunits (Hba-a1, Hba-a2, Hbb-b1, Hbb-ba), an erythroid-specific mitochondrially located enzyme (Alas2), as well as the noncoding RNA Rmr2. Only one DEG, the predicted gene Gm98221 (apolipoprotein L 7c pseudogene), was common to all four tissues and was among the most down-regulated DEGs in all tissues (Fig. 5A and Dataset S3). At the individual transcript level, a literature search revealed that numerous DEGs in all four tissues had been previously reported to be associated with sleep and circadian rhythms (prefrontal cortex: 35.1%; hippocampus: 18.7%; hypothalamus: 21.1%; blood: 17.1%), stress (prefrontal cortex: 40.5%; hippocampus: 35.2%; hypothalamus: 50.9%;
blood: 20%), neuropsychiatric symptoms (prefrontal cortex: 37.8%; hippocampus: 20.9%; hypothalamus: 29.8%: blood: 25.7%), mood disorders (prefrontal cortex: 16.2%; hippocampus: 8.8%; hypothalamus: 19.3%; blood: 2.9%), or neurodegenerative diseases, such as Alzheimer’s and Parkinson’s diseases (prefrontal cortex: 37.8%; hippocampus: 30.8%; hypothalamus: 36.8%; blood: 17.1%) (see SI Appendix, Fig. S6 and Dataset S3 for references). In addition, several DEGs in the prefrontal cortex (e.g., S100a8, S100a9, Lbp, Tgfβ2), hippocampus (e.g., Inava, Lbp, Rd12l2, Pla2g5, F3, Vegf; C4a2a, Tgip1, Cast, Lst1), and blood (e.g., Clec4n, Chil3, Reg3g, Bpif1) play a key role in the immune system, inflammation, and the hypothalamus (e.g., Glra3, Ptgdr, Pnicr, Oprkl, Kcnq2, Gpr6) and hypothalamus (Slc6a3, Slc5a7, Chat) showed several DEGs involved in neural transmission, including the down-regulation of neuropeptide genes encoded in adaptation to stress and social behavior in the hypothalamus (i.e., Ucn3, Avep, Otx, Vip). Some of the most up-regulated DEGs in blood are involved with DNA damage response (i.e., Mnd1, E2f7), while others have previously been associated with sleep deprivation or fragmentation (Fads3, Gm6166, Spig1, Hspa1a, Hspa1b, Sgcb3a1) (SI Appendix, Fig. S6a and Dataset S3).

To further characterize the effects of the 9-ω UCMS, we performed functional enrichment analysis using Gene Ontology (GO) processes and canonical pathway maps. The hypothalamus showed the largest number of enriched GO processes (n = 168) compared with the prefrontal cortex (n = 74), hippocampus (n = 37), and blood (n = 54). Ten processes were shared by the three brain regions (Fig. 5B). These included processes associated with the immune system (e.g., cytokocyte development and differentiation), circulatory system processes (e.g., regulation of blood pressure), and metabolic processes (e.g., oxygen transport, hydrogen peroxide metabolic process) (Fig. 5C; for detailed identity of the GO processes, see Dataset S4). In contrast, only two enriched GO processes were common to blood and brain regions. Response to stress was common to the blood, hypothalamus, and hippocampus, while regulation of receptor activity was shared by blood and hypothalamus (Fig. 5C and Dataset S4).

GO biological processes in the hypothalamus were involved in developmental processes (e.g., cell fate commitment), nervous system processes (e.g., regulation of sensory perception), immune system (e.g., regulation of C-C chemokine binding, myeloid cell homeostasis), cell communication (e.g., G protein-coupled receptor signaling pathway), and behavior (grooming and aggressive behaviors) (Fig. 5C and Dataset S4). One enriched pathway involved in protein folding and maturation (i.e., posttranslational processing of neuroendocrine peptides) was observed (SI Appendix, Fig. S7 and Dataset S4). In the brain regional cortex and hippocampus, functional enrichment analysis identified 37 processes associated with inflammatory and immune response (some of which were shared; e.g., response to IFN-β; leukocyte migration involved in inflammatory

![Fig. 2. Time course of UCMS-induced alterations on sleep and the EEG.](image-url)

Fig. 2. Time course of UCMS-induced alterations on sleep and the EEG. (A) Duration of REMS per 24 h. (B) Duration of REMS episodes per 24 h. (C) EEG theta power density (6–9 Hz) in REMS during the 12-h light phase expressed as the percentage of theta power in baseline. (D) Relative EEG power spectra in REMS during the 12-h light phase (averaged spectra of all EEG recording sessions during the 9-ω UCMS protocol). (E) Duration of NREMS per 24 h. (F) Duration of NREMS episodes per 24 h. (G) EEG delta power density (1–4.5 Hz) during the 12-h light phase expressed as the percentage of delta power in baseline. (H) Relative EEG power spectra in NREMS (averaged spectra of all EEG recording sessions during the 9-ω UCMS protocol). Data are L5means ± 95% CI (controls: gray; UCMS: red; n = 8 per group). *P < 0.05, **P < 0.01, ***P < 0.001 (post hoc comparisons for significant “treatment” × “day” interaction, except for Δ and Δ: effect of “treatment” in general linear mixed model). For detailed statistics, see Dataset S1.

![Fig. 3. Effect size of UCMS-induced physical, behavioral, neuroendocrine, and sleep alterations.](image-url)

Fig. 3. Effect size of UCMS-induced physical, behavioral, neuroendocrine, and sleep alterations. Effect sizes of repeated (Cohen’s d) and nonrepeated measures (Cohen’s d) were converted to Cohen’s f using the following formula: Cohen’s f = 2 × Cohen’s d, with large effect size: >0.40; medium: 0.25–0.40; small: 0.10–0.25. Bar colors correspond to those displayed in Fig. 1A and SI Appendix, Fig. S5 for all measured phenotypes. For values, see Dataset S1.
response) among others (Fig. 5C and Dataset S4). Enriched pathways evoked by chronic stress were involved in transcription and development; however, none were significant in the hippocampus after FDR adjustment (SI Appendix, Fig. S7 and Dataset S4). In blood, functional enrichment also identified biological processes involved in immune and inflammatory response (e.g., regulation of cytokine production) and signaling pathways (e.g., nitric oxide-mediated signal transduction, TNF-mediated signaling pathways). In addition, 10 processes were associated with RNA cleavage and the unfolded protein response (Fig. 5C and Dataset S4).

**Bivariate associations for physical, behavioral, neuroendocrine, and sleep alterations.** (A) Kendall’s partial correlation between pairs of phenotypes (i.e., after removing the effect of the experimental groups). The phenotypes (the averaged last three measurements were used for repeated measures) were ordered according to their phenotypic categories. Correlations were considered significant at an FDR < 0.05 (P_{adj>0} symbolized by black-framed square), computed with the Benjamini-Hochberg procedure for multiple testing correction. For detailed statistics, see Dataset S2. (B) Example of a correlation from (A) illustrated for percentage of REMS per TST during the light (L) phase and impairment of the corticosterone regulation (τ = 0.72, \( P_{adj} = 0.00034, P_{adj>0} = 0.00197; n = 8 \) animals per group; gray: control mice, red: UCMS subjected animals). DEX supp., dexamethasone suppression.

**Fig. 4.** Bivariate associations for physical, behavioral, neuroendocrine, and sleep alterations. (A) Kendall’s partial correlation between pairs of phenotypes (i.e., after removing the effect of the experimental groups). The phenotypes (the averaged last three measurements were used for repeated measures) were ordered according to their phenotypic categories. Correlations were considered significant at an FDR < 0.05 (\( P_{adj>0} \) symbolized by black-framed square), computed with the Benjamini-Hochberg procedure for multiple testing correction. For detailed statistics, see Dataset S2. (B) Example of a correlation from (A) illustrated for percentage of REMS per TST during the light (L) phase and impairment of the corticosterone regulation (\( τ = 0.72 \), \( P_{adj} = 0.00034, P_{adj>0} = 0.00197; n = 8 \) animals per group; gray: control mice, red: UCMS subjected animals). DEX supp., dexamethasone suppression.

**Fig. 5.** Characterization and functional enrichment of genes differentially expressed following chronic mild stress. Overlap of (A) DEGs and (B) significantly enriched GO biological processes for DEGs in the prefrontal cortex, hippocampus, hypothalamus, and whole blood. (C) GO biological processes associated with DEGs. Outer track: tissue; second track: overarching themes associated with GO processes; third track: tissues in which GO processes were found; inner track: overlap of processes, colors corresponding to overarching theme. \( n = 8 \) per group for brain regions; \( n = 7 \) controls vs. \( n = 9 \) UCMS group for blood. Enrichment analyses were performed using MetaCore and significance was set at \( P_{adj} < 0.05 \). Information is available in tabular format (Datasets S3 and S4).

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suffer from the multiplicity problem. In addition, they are not necessarily best suited to identify sets of transcripts that predict specific complex phenotypes. Thus, we applied elastic-net learning, a multivariate approach based on a generalized linear model using penalized regression, to identify sets of features predicting specific phenotypes. We performed this analysis using all transcripts identified by RNA sequencing, that is, not just the DEGs, focusing on sleep variables and some variables associated with stress and mood disorders. We aimed to identify transcriptomic features that were specifically associated with sleep and behavioral variables both within the control and UCMS group (i.e., at the level of the individual). To accomplish this, “unspecific group” effects (i.e., control vs. UCMS) on these variables need to be removed from the analysis. We therefore applied normalization procedures to control for group effects (Materials and Methods). The features that associate with behavioral variables, as identified by elastic-net after application of the normalization procedures, indeed contained very few transcripts (30 of 1,595) identified by the group-level analysis (DEGs; see previous section). This demonstrates that this approach yields information that is different from the DEG approach. The number of features in the various identified predictor sets was overall small and varied between variables and across tissues (range: 1–333) (Dataset S6). To gain insights into molecular mechanisms associated with a given sleep or behavioral variable and to control for behavioral correlates of the sleep and other variables, we then performed functional enrichment analysis of predictor sets focusing on pathway maps.

REMS and NREMS. The size of the predictor sets for REMS and NREMS parameters was similar for sleep duration and continuity (493 and 464, respectively), but few significantly overlapped (n = 73) (Dataset S7). Common predictors were seen in the prefrontal cortex, primarily between REMS continuity and NREMS duration (n = 29) and continuity (n = 39), as well as in the hippocampus between REMS bout count and NREMS duration (n = 3) and bout count (n = 2) (Dataset S7). For REMS variables, several included transcripts involved in neural transmission, sleep, and circadian rhythms (Dataset S8). Six predictors for REMS variables play a key role in the regulation of NF-kB signaling, while 26 predictors of NREMS were associated with mitochondrial function (Dataset S8).

Whereas REMS predictor sets were significantly enriched for many canonical pathways (n = 193), only three pathways were identified in the NREMS predictor sets (Fig. 6 A–C and Dataset S9). Enriched pathways associated with predictors of REMS duration and REMS theta power were most strongly observed in the hippocampus (n = 35 and 42, respectively), while pathways associated with predictors of REMS bout length (n = 90) and REMS bout count (n = 26) were primarily enriched in the prefrontal cortex (n = 30), hippocampus (n = 43), and hypothalamus (n = 17), and in the hippocampus (n = 9) and blood (n = 14), respectively (Fig. 6C). REMS duration upregulated several enriched pathways associated with stem cells (n = 14) and development (n = 6) (Fig. 6 and Dataset S9). Across REMS theta power, REMS continuity and tissues, 24 pathways were involved in the immune response, including various IL, IFN, and Toll-like receptor signaling pathways. Remarkably, 11 enriched pathways were common to REMS theta power and REMS bout length, with eight of them associated with apoptosis and cell survival, such as the TNF receptor (TNFR)-1 signaling pathway, the role of inhibitor of apoptosis proteins (IAP), the endoplasmic reticulum (ER) stress response pathway (five of these pathways are listed under the “generic theme”) (Dataset S9), stem cells (n = 5), immune response (n = 3), development (n = 3), as well as several generic metabolic and signaling pathways (SI Appendix, Fig. S8). Predictor sets of anhedonia were enriched in several pathways involved in development (n = 12) and stem cell processes (n = 4) and generic pathways (SI Appendix, Fig. S8). Nine enriched pathways were found in predictor sets for despair behavior, primarily in the hippocampus (n = 7) and included circadian rhythm process (SI Appendix, Fig. S8 and Dataset S9).

Pathways shared between REMS, corticosterone regulation, and anhedonia. More than one-third of the 40 pathways associated with corticosterone regulation (37.5%) overlapped with pathways for REMS bout length or for EEG theta activity in the prefrontal cortex and hippocampus, respectively (SI Appendix, Fig. S8). Thirteen of the 15 common pathways were associated with apoptosis and cell survival (Table 1). They included apoptotic pathways involved in the extrinsic death receptor pathway (e.g., TNFR-1 signaling, FAS signaling cascade, and apoptotic TNF-family pathways) and the intrinsic mitochondrial pathway (role of IAPs in apoptosis; regulation of apoptosis by mitochondrial proteins) (Table 1 and SI Appendix, Fig. S8). In addition, REMS variables, and in particular REMS bout length and EEG theta activity, also shared seven pathways with anhedonia. These included pathways involved in apoptosis and survival and response to oxidative stress (Table 1). Finally, no overlap was observed between pathways associated with NREMS variables, corticosterone regulation, and any of the investigated behavioral variables (SI Appendix, Fig. S9).

Discussion

REMS Enhancement, a Core Response to Chronic Mild Stress. The UCMS paradigm induces changes in physical, behavioral, and neuroendocrine variables in accordance with previous reports (7, 20, 28). The simultaneous and longitudinal assessment of a wide range of variables allowed for a comparison of the magnitude of changes and the temporal emergence of these physical, behavioral, and neuroendocrine alterations. This approach demonstrated that increase in REMS variables (i.e., 24-h duration, bout length, bout count, EEG theta oscillations) exhibited not only large effect sizes but were also among the earliest responses induced by stress. The longitudinal assessment of sleep also showed that REMS and NREMS respond differently to chronic stress. The increased continuity (i.e., bout length, bout count, and duration of REMS) and the increase in EEG theta activity during REMS, primarily reflecting hippocampal theta activity, imply that REMS is affected by UCMS in a positive manner. Of particular interest is the increase in both REMS bout duration and theta power, because in the rat it has been reported that theta power decreases in the course of a REM bout (31). In contrast, NREMS continuity was decreased and EEG delta power in NREMS was not affected. The changes observed in sleep and their effect sizes agree well with metaanalyses performed in clinical depressive populations, according to which effect sizes for REMS are larger than those for NREMS and sleep continuity (14, 17).

Transcripts and Associated Processes Affected by Chronic Mild Stress. Transcriptome changes, assessed by differential expression and thus primarily reflecting effects of stress at the group level, were relatively small and most changes were observed in the hippocampus, which is consistent with previous reports (20, 21, 32). One tentative conclusion from these data are that effects of stress on sleep and changes in gene expression converge on the hippocampus (33), and an emerging question is whether REMS-related phenomena, such as EEG theta power, reflect or direct these hippocampal changes. On the other hand, our data also highlight that the cortex and hypothalamus are responsive to...
stress. In fact, the hypothalamus showed the largest number of enriched GO processes. Furthermore, many enriched biological processes were shared across brain regions. These include processes associated with inflammatory and immune responses, and parainflammation thus appears to be a common mechanism in the three brain regions investigated. This is in line with a recent framework emphasizing that inflammatory signals contribute to restore homeostasis (34) and agrees well with the emerging view that chronic stress and stress-related diseases, such as major depression, share inflammation as a common mediator (35, 36). Given the changes in DEX-induced corticosterone suppression, it may seem surprising that we did not observe changes in transcripts related to glucocorticoid or mineralocorticoid receptors in either the hippocampus or other brain regions. Our findings are, however, consistent with previous UCMS studies in which no change in their gene expression was observed (18, 20, 37).

Transcriptomic Predictors of Phenotypic Variation Identified Using Machine Learning. Transcriptomic predictor sets were overall relatively small in accordance with a previous study (38). Hippocampal transcriptomic predictors of 24-h REMS duration were associated with pathways involving stem cells differentiation and hedgehog signaling. Inhibition of hedgehog signaling by glucocorticoid treatment has been shown to decrease hippocampal cell differentiation (39). In addition, several enriched pathways in apoptosis and cell survival were among the molecular signatures characterizing REMS continuity variable (bout length) and EEG theta power, in the cortex and hippocampus, respectively. The overlap between pathways for theta power and REMS bout length may point to common mechanisms underlying theta and bout length regulation. The identified predictors and related pathways specifically associated with REMS in the cortex, a brain region not necessarily implicated in the generation of REMS, may reflect an effector system by which REMS exerts its adaptive response to chronic stress. The ER stress-response pathway was common to REMS continuity and theta power. ER stress, which may lead to apoptosis (40), has been recently shown to be induced during social isolation in Drosophila (41). A number of circadian-related transcripts were identified as predictors of NREMS variables, as well as despair and anhedonia-like behaviors. This is consistent with a recent study correlating UCMS-induced depressive-like behavior with circadian rhythm alterations in brain tissues (42), and the growing recognition that circadian rhythmicity may play a role in mood regulation (43, 44). It should be noted that while we observed changes in sleep, at the behavioral level circadian rhythmicity was not much affected, although the reduced activity during the dark period may be interpreted as a reduction in circadian amplitude. NREMS and REMS shared very few predictors, further emphasizing the contrast between these two sleep states observed at the electrophysiological level in this study. Furthermore, no overlap was observed between pathways associated with predictors for NREMS variables, corticosterone regulation, or behavioral phenotypes.

One aim of the present analyses was to investigate to what extent transcriptomic changes in the brain are reflected in the blood transcriptome. The results demonstrate that, at the level of individual DEGs or associated processes/pathways, there were no significant overlaps between brain and blood transcriptic changes. However, even though the blood transcriptome may not be directly informative about changes in the brain, the elastic-net approach indicated that whole blood contains predictors of behavior (anhedonia) and REMS (bout count), which ultimately may be useful for biomarker development.

Close Associations Between REMS, Corticosterone Regulation, and Apoptotic Pathways. One major theme emerging from the multilevel analyses is the robust effects of stress on REMS and the close link between changes in REMS and dysregulation of corticosterone. These data may be interpreted as evidence for a shared role of REMS and corticosterone within the context of “adaptation” to the waking experience. REMS has been proposed to play a central role in emotional processing and memory consolidation (45–48). A causal role for EEG theta activity during REMS was demonstrated in contextual and extinction memory consolidation in rodents and humans (49, 50). Furthermore, REMS is suppressed by most antidepressants (51) and some antidepressants interfere with the homeostatic control of REMS (52).

While REMS enhancement and alterations in the HPA axis negative-feedback regulation of corticosterone have been previously reported in preclinical studies of chronic stress or stress-vulnerable rodents (8, 9, 28), the present data demonstrate the close association between REMS% and corticosterone suppression (Fig. 4). This result, as well as previous findings in humans (53, 54), suggest that these phenomena share common causal mechanisms. Hypothalamic neuropeptides, such as vasoactive intestinal peptide (VIP), arginine vasopressin (AVP), and melanin-concentrating hormone (MCH), whose encoding genes were downregulated in the present study, are potential candidates for orchestrating this association because they have been implicated in the regulation of REMS (55–58) and the HPA axis (59, 60). In humans, increased REMS (61) and HPA axis dysregulation (13) have been shown to correlate with remission and recovery in major depressive disorder.

Further evidence for the close association between REMS and corticosterone regulation emerged from the transcriptomic analyses. We identified 15 overlapping pathways between the corticosterone regulation and REMS continuity variables and theta power. These pathways were primarily involved in apoptosis...
Limitations of this study include a relatively small sample size and the choice of setting the statistical significance at FDR-adjusted \( P < 0.05 \), which may have led to an underreporting of significant effects. Another limitation relates to experimental constraints, which precluded an assessment of the temporal association between behavioral phenotypes and transcriptomic changes. Nevertheless, the high intraindividual stability of many of the phenotypes indicates that the observed transcriptomic changes at the end of the experiment are relevant to the phenotypes throughout the UCMS. This study was only conducted in males. Hypotheses based on the present data may need to be tested in females.

**Table 1. Enriched pathways shared by REM sleep and corticosterone regulation, or REM sleep and anhedonia**

| REMS variables     | Overlapping pathways                                                                 | Apoptosis and survival related pathway |
|--------------------|-------------------------------------------------------------------------------------|---------------------------------------|
| **Corticosterone regulation** |                                                                                       |                                       |
| Bout length       | TNFRI1 signaling pathway* (PFC)                                                      |                                       |
| Bout length       | Role of IAP-proteins in apoptosis* (PFC)                                              |                                       |
| Bout length       | Granzyme B signaling* (PFC)                                                          |                                       |
| Bout length       | Cytoplasmic/mitochondrial transport of proapoptotic proteins Bid, Bmf, and Bim* (PFC) |                                       |
| Bout length       | Apoptotic TNF-family pathways* (PFC)                                                  |                                       |
| Bout length       | FAS signaling cascades* (PFC)                                                        |                                       |
| Bout length       | Endothelial differentiation during embryonic development* (HYP)                      |                                       |
| Bout length       | Role of Apo-2L(TNFSF10) in Prostate Cancer cell apoptosis* (PFC)                     |                                       |
| Bout length       | Resistance of melanoma cells to Apo-2L(TNFSF10)-induced apoptosis* (PFC)             |                                       |
| Bout length       | Inhibition of apoptosis in gastric cancer* (PFC)                                      |                                       |
| Bout length       | Neutrophil resistance to apoptosis in chronic obstructive pulmonary disease and proresolving impact of lipid mediators* (PFC) |
| Bout length       | Apo-2L(TNFSF10)-induced apoptosis in melanoma* (PFC)                                  |                                       |
| Bout length       | Apoptotic pathways and resistance to apoptosis in lung cancer cells* (PFC)           |                                       |
| Duration          | Endothelial differentiation during embryonic development* (HIP)                     |                                       |
| Theta power       | Role of IAP-proteins in apoptosis* (HIP)                                              |                                       |
| Theta power       | TNFR-1 signaling pathway* (HIP)                                                      |                                       |
| Theta power       | IL-5 signaling via JAK/STAT* (HIP)                                                   |                                       |
| Theta power       | Role of Apo-2L(TNFSF10) in prostate cancer cell apoptosis* (HIP)                     |                                       |
| Theta power       | Inhibition of apoptosis in gastric cancer* (HIP)                                      |                                       |
| Theta power       | Apoptotic pathways and resistance to apoptosis in lung cancer cells* (HIP)           |                                       |

| **Anhedonia**      |                                                                                       |                                       |
| Bout count         | Metabolism in pancreatic cancer cells* (BLO)                                          |                                       |
| Bout length        | Cytoplasmic/mitochondrial transport of proapoptotic proteins Bid, Bmf, and Bim* (PFC) |                                       |
| Bout length        | Glucocorticoids-mediated inhibition of proconstrictory and proinflammatory signaling in airway smooth muscle cells* (HIP) |
| Bout length        | Memory CD8+ T cells in allergic contact dermatitis* (HYP)                             |                                       |
| Duration           | EGF-induced proliferation of Type C cells in secondary proliferative zone of adult brain* (HIP) |
| Theta power        | Memory CD8+ T cells in allergic contact dermatitis* (HIP)                             |                                       |
| Theta power        | NRF2 regulation of oxidative stress response* (HIP)                                   |                                       |

Tissue in which enriched pathways associated with predictors of REMS variables is indicated in parenthesis (i.e., BLO, blood; HIP, hippocampus; HYP, hypothalamus; PFC, prefrontal cortex). The column “Apoptosis and survival related pathway” indicates whether a pathway was related to apoptosis and survival as some pathways listed under a generic theme in MetaCore are also implicated in apoptotic and antiapoptotic pathways. For each pathway, the functional theme to which it belongs to is listed according to the specified footnote.

*Apoptosis and survival.
†Stem cells.
$^{\dagger}$Generic (according to MetaCore).
$^{\ddagger}$Immune response.

and cell survival and included several members of TNF signaling, which triggers a broad spectrum of actions at the cellular level, including processes involved in the mitochondrial intrinsic pathway. Involvement of apoptotic pathways in depression and stress has been reported in recent human and animal studies. Blood transcriptomic studies in humans show that major depressive disorder and antidepressant response are associated with enrichment in apoptosis signaling processes and pathways (62, 63). Repetitive transcranial magnetic stimulation to treat depression counteracts hippocampal neuronal apoptosis and HPA axis disturbances induced by UCMS in rats (64). In humans, chronic insufficient sleep, characterized by REMS alterations (65), is associated with apoptosis-related blood mRNA biomarkers (66), and sleep restriction in mice alters apoptotic pathway signaling (67).

We also identify shared molecular pathways underlying the interindividual variation in anhedonia, a core symptom of depression, with REMS continuity and theta power in response to chronic stress. They included pathways involved in oxidative stress and apoptosis with a pathway involved in the transport of proapoptotic proteins linking the Jun amino-terminal kinases (JNK) signal transduction pathway and the mitochondrial apoptotic machinery (68). A causal role for mitochondrial genes was recently proposed as part of the processes in the striatum, linking REMS and a stress-induced anxiety-like phenotype (38), and the contribution of mitochondrial dysfunction in major depression is emerging (69).

Considering the link between REMS, cell proliferation, and apoptosis (70, 71), the severe alterations of hippocampal neuronal plasticity and HPA axis functioning in mood disorders (72), and the strong responsiveness of the hippocampus to stress hormones (73), our results linking REMS continuity, theta oscillations, corticosterone regulation, and cell apoptosis as well as the shared pathways between REMS and anhedonia, shed a new light on the pathological framework of stress-related conditions.

**Limitations.** Limitations of this study include a relatively small sample size and the choice of setting the statistical significance at FDR-adjusted \( P < 0.05 \), which may have led to an underreporting of significant effects. Another limitation relates to experimental constraints, which precluded an assessment of the temporal association between behavioral phenotypes and transcriptomic changes. Nevertheless, the high intraindividual stability of many of the phenotypes indicates that the observed transcriptomic changes at the end of the experiment are relevant to the phenotypes throughout the UCMS. This study was only conducted in males. Hypotheses based on the present data may need to be tested in females.
be tested in future studies in which sex differences in sleep disturbances and their underlying molecular mechanisms in the context of chronic stress could be investigated (74).

Whether mice recover from the depressive-like phenotype after cessation of the UCMS was not assessed in this study. Other studies reported persistence of alterations at the transcriptomic, metabolic, and behavioral levels for several days or weeks after the end of UCMS (75–78). How this pattern of recovery relates to changes in REMS has not yet been studied in detail.

**Conclusion**

This study in mice provides a comprehensive characterization of sleep changes induced by chronic stress, with REMS increase being the earliest marker of a stress response. Our data show that interindividual variation in REMS continuity and theta oscillations during REMS, and apoptosis processes including mitochondrial pathways, changes in corticosterone regulation, and anhedonia are interrelated. Alteration in corticosterone regulation and REMS have both been implicated in the response to emotional experiences. Given the prominence of REMS alterations in mood disorders and the herein-identified correlates of REMS, further study of the function of REMS parameters—such as its duration, continuity, and the theta oscillations during REMS—in the response to stress is warranted.

**Materials and Methods**

**Animals.** Male BALB/c mice (n = 18; B&K Universal Ltd) underwent EEG/EMG surgery, as previously described (79) (see SI Appendix for details). After recovery, mice were randomly assigned to the control or UCMS group. Baseline data collection was performed, after which the 9-wk UCMS protocol started. Mice were daily subjected during the dark period to various socio-environmental low-intensity stressors according to an unpredictable schedule (27) (Fig. 1A and SI Appendix, Table 1). Experimental procedures were approved by the University of Surrey Animal Welfare and Ethical Review Board and were carried out in accordance with the United Kingdom Animals (Scientific Procedures) Act 1986.

**Physical, Behavioral, and Corticosterone Regulation Assessments.** Body weight, coat state, self-centered behavior (grooming test), motivation (nest building test), anhedonia (reward-driven exploratory test), social preference (social novelty preference test), aggressiveness (resident-intruder test), anxious (novelty-suppressed feeding test), and despair behavior (forced swim test) were assessed as previously described (27, 28, 80, 81) (see SI Appendix for details). The DEX suppression test was used to evaluate the HPA axis negative-feedback-regulated corticosterone (28) (see SI Appendix for details).

**Sleep and Locomotor Activity.** No stressor was applied during the sleep recordings. The data analyzed consisted of 24-h recordings starting at dark onset. EEG power spectra were computed for consecutive 10-s epochs by a fast Fourier transform (see SI Appendix for details). Locomotor activity was measured as previously described (79). Averaged daily activity for the 12-h light and dark periods were analyzed per week.

**Transcriptome Analysis.** Tissues (prefrontal cortex, hippocampus, hypothalamus, and whole blood) were collected 14–16 h after the last stress exposure. For details of RNA sequencing, see SI Appendix. Differential expression analysis (control vs. UCMS) was performed with the nonparametric Rank Product statistical method that is independent of interclass variability (82), using the R Bioconductor package RankProd. Significance was set at a p-value < 0.05. To robustly select relevant transcriptomic predictors, a form of penalized regression, referred to as elastic net, was performed using the R package glmnet (83). Analyses were focused on sleep variables, three behavioral variables, and corticosterone regulation, the values of which were z-scored within group to identify associations with phenotypes independent of any "unspecific" group effect. We only report predictor sets for variables achieving a positive Pearson correlation r between observed and cross-validated prediction values > 0.31. For functional annotation, lists of genes associated with a variable (e.g., REMS duration) were subjected to GO enrichment analyses (GO processes or Pathway maps) using MetaCore (Clarivate Analytics; https://portal.genego.com/; updated June 2018). Functional analyses were performed using the respective tissue-specific transcriptome, as identified by RNA sequencing, as background. Significant enrichment was defined by Pnom < 0.05 and Padj < 0.05.

**Statistics.** Unless otherwise stated, data were analyzed with SAS 9.2 (SAS Institute). For repeated measures, data were analyzed as dependent variables in a general linear mixed model using PROC MIXED for ANOVA with group (treatment: UCMS vs. control) and time (day, treated as repeated measure with spatial power anisotropic variance-covariance matrix) as categorical explanatory variables with baseline as a covariate (no group effect was found at baseline for all measures). Post hoc multiple pair-wise comparisons (UCMS vs. control) were assessed using the Bonferroni adjustment. Output data are expressed as least-squares means (LSmeans) with 95% confidence intervals (CIs). For nonrepeated measures, PROC TTEST for pair-wise comparisons was used with Padj < 0.05. To robustly select relevant transcriptomic predictors, a form of penalized regression, referred to as elastic net, was performed using the R package glmnet (83). Analyses were focused on sleep variables, three behavioral variables, and corticosterone regulation, the values of which were z-scored within group to identify associations with phenotypes independent of any "unspecific" group effect. We only report predictor sets for variables achieving a positive Pearson correlation r between observed and cross-validated prediction values > 0.31. For functional annotation, lists of genes associated with a variable (e.g., REMS duration) were subjected to GO enrichment analyses (GO processes or Pathway maps) using MetaCore (Clarivate Analytics; https://portal.genego.com/; updated June 2018). Functional analyses were performed using the respective tissue-specific transcriptome, as identified by RNA sequencing, as background. Significant enrichment was defined by Pnom < 0.05 and Padj < 0.05.

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