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Chronic Shift-Lag Alters the Circadian Clock of NK Cells and Promotes Lung Cancer Growth in Rats

Ryan W. Logan,* Changqing Zhang,* Sengottuvelan Murugan,* Stephanie O’Connell,* Dale Levitt,* Alan M. Rosenwasser,† and Dipak K. Sarkar*

Prolonged subjecting to unstable work or lighting schedules, particularly in rotating shift-workers, is associated with an increased risk of immune-related diseases, including several cancers. Consequences of chronic circadian disruption may also extend to the innate immune system to promote cancer growth, as NK cell function is modulated by circadian mechanisms and plays a key role in lysis of tumor cells. To determine if NK cell function is disrupted by a model of human shift-work and jet-lag, Fischer (344) rats were exposed to either a standard 12:12 light-dark cycle or a chronic shift-lag paradigm consisting of 10 repeated 6-h photic advances occurring every 2 d, followed by 5–7 d of constant darkness. This model resulted in considerable circadian disruption, as assessed by circadian running-wheel activity. NK cells were enriched from control and shifted animals, and gene, protein, and cytolytic activity assays were performed. Chronic shift-lag altered the circadian expression of clock genes, Per2 and Bmal1, and cytolytic factors, perforin and granzyme B, as well as the cytokine, IFN-γ. These alterations were correlated with suppressed circadian expression of NK cytolytic activity. Further, chronic shift-lag attenuated NK cell cytolytic activity under stimulated in vivo conditions, and promoted lung tumor growth following i.v. injection of MADB106 tumor cells. Together, these findings suggest chronic circadian disruption promotes tumor growth by altering the circadian rhythms of NK cell function.


Since the initial reports associating shift-work and cancer in humans, a majority of animal studies have primarily focused on the effects of disrupted molecular clocks on cellular proliferation pathways to promote tumor growth (15–19). Although these pathways are important for cancer development and progression, the relationship between circadian disruption and cancer may extend beyond perturbations of cell cycle processes. For example, the inflammatory response by macrophages is dysregulated by disruption to cellular clocks (20, 21). However, the underlying mechanism by which altered circadian immune function may promote the transition to particular disease states is unknown, especially in relation to cancer. There is little evidence implicating disruption to circadian mechanisms in impaired immune function, as well as tumor growth.

The role of NK cells as critical mediators of cancer immunosurveillance is well established (22, 23). In mice, depletion of NK cells is associated with increased tumor growth in spontaneous and induced tumor models (24, 25). Similarly, low NK cell activity is associated with increased risk for cancer in human populations (26, 27). To kill tumor cells, NK cells recruit pro-inflammatory cytokines, release cytolytic granules, and activate receptors on target cells (23, 28, 29). Of particular importance are the cytokolytic factors granzyme B and perforin and cytokines TNF and IFN-γ, which are critical factors involved in regulating NK cell-mediated killing of tumor cells (28–30). Decreases in any of these factors are associated with increased risk of developing infections and tumors (24, 25, 31–33).

The negative health consequences of circadian disruption are likely to involve alterations in NK cell function. Circadian mechanisms may optimize an immunological response by coordinating function between peripheral tissues and immunocompetent cells. Previously, we have reported that NK cell function, including cytokines, cytolytic factors, and cytolytic ability, is tightly controlled by the circadian system (34). Signals directly or indirectly derived from the circadian pacemaker, which is located

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Abbreviations used in this article: CT, circadian time; DD, total darkness; LD, light-dark (cycle); SCN, suprachiasmatic nucleus.

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in the suprachiasmatic nucleus (SCN) of the anterior hypothalamus, are transmitted to peripheral tissues by neural and endocrine pathways to synchronize and coordinate body physiology (34–36). Rhythmic sympathetic input to the spleen modulates NK cell function by regulating components of the molecular clock in these cells (37). Maximal cytotoxicity at specific circadian phases is postulated to be driven by molecular clock-coordinated expression of IFN-γ, TNF-α, perforin, and granzyme B (38, 39).

Nevertheless, few data indicating potential relationships between the disruption of clocks in immune cells and the promotion of tumor growth are available. Therefore, the main goal of this study was to determine whether NK cells were susceptible to circadian disruption by exposing animals to a chronic shift-lag paradigm in which the daily light-dark (LD) cycle was repeatedly phase shifted to experimentally mimic the disruptive effects of human shift-work and jet-lag. Under the shift-lag schedule, circadian disruption was assessed by continuously monitoring voluntary running-wheel activity. We then examined circadian rhythms in NK cell function at the gene, protein, and physiological levels under both basal and stimulated conditions. Finally, we sought to determine whether alterations in NK cell function due to repeated shifting of the LD cycle were related to an increased risk of tumor development, using an induction model of NK-sensitive adenocarcinoma cells (MADB106) (40, 41). MADB106 cancer cells typically metastasize in the lungs following i.v. injection, and the method is well established for exploring the role of NK cell function in tumor growth (41–43). We hypothesized that chronic shifting would suppress NK cell function in response to tumor cells by altering the coordinated circadian expression of key immune factors. We also postulated that these changes in NK cell function would promote tumor growth.

Materials and Methods

Animals and apparatus

Male Fischer (F344) rats were obtained from Charles River Laboratories (Wilmington, MA) at ≈50 d of age. Upon arrival, animals were acclimated to the housing facility and maintained individually in running-wheel cages (wheel diameter: 34 cm; Mini Mitter, Bend, OR), placed within light- and sound-attenuating climate-controlled cubicles. Running-wheel activity was recorded by a computer and stored for later analysis of circadian parameters using the ClockLab software package (Actimetrics, Wilmette, IL). Animals were given free access to food and water for the duration of the experiment. All experimental procedures and animal treatment protocols were approved by Rutgers Animal Care and Facilities Committee and complied with National Institutes of Health policies.

Procedures

Animals were maintained initially under a standard 12:12 LD cycle with lights on at 6:00 h and lights off at 18:00 h for a 1- to 2-wk acclimation period to stabilize running-wheel activity, and then assigned randomly to either remain under the standard LD cycle (controls) or to undergo the chronic shift-lag protocol (n = 90 per lighting regimen used for various assays; see below). The shift-lag protocol consisted of 6-h LD phase advances repeated every 2 d for a total of 10 shifts (see Fig. 1A–C for a schematic representation of the control and shift-lag lighting regimens). For example, the first advance consisted of the lights coming on 6 h earlier than the previous day (lights on at 24:00 h), which resulted in a shortening of the dark period on the following day. Repeated phase advances were chosen owing to extensive prior data indicating that advances evoke greater circadian disruption than do phase delays. To determine whether the chronic shift-lag paradigm resulted in alterations of circadian expression of cytolytic factors, cytokines, and cytolytic capabilities in NK cells, a set of animals (n = 36 per lighting regimen) were sacrificed by decapitation at six time points (n = 6 per time point) 5–7 d into darkness (DD), which corresponded to circadian times (CTs) 3, 7, 11, 19, and 23. To avoid the potential masking effects of these circadian rhythms, animals were sacrificed 5–7 d into DD following the last advance of the LD cycle. Actograms were used to determine CTs by defining CT12 as the onset of running-wheel activity. Immediately upon sacrifice, tissues were collected and NK cells were separated from spleens for gene, protein, and cytotoxicity assays. To determine the effects of chronic shifting on NK cell cytotoxicity in response to tumor cells and the possible consequences on tumor growth, separate groups of animals (n = 30 per lighting regimen) were inoculated with MADB106 cells at CT19 in DD. Half of these ani-
mals (n = 10 per lighting regimen) were sacrificed 24 h after inoculation to assess immediate NK cell response to the presentation of tumor cells (40, 43). The remaining animals (n = 20 per lighting regimen) were placed for 8–14 wk under the same standard LD cycle as during the acclimation period, and then sacrificed for lung extraction and inspection of tumor metastases.

Enrichment of NK cells from spleen

Individual spleen tissue was processed as described previously (44). RBCs and granulocytes were removed from splenocyte suspensions by density centrifugation using Histopaque 1083 (Sigma-Aldrich). Splenocytes (~10 × 10^6 cells per spleen) were extracted from the middle layer, washed with RPMI-1640 (Invitrogen), and resuspended in buffer (PBS, 0.5% BSA). Splenocytes were incubated with primary Abs conjugated to FITC (BD Biosciences), anti-CD6 (OX52; T cells), anti-CD45RA (OX-33; mature B cells), and anti-RT1B (OX-6; dendritic cells and macrophages), followed by secondary incubation with anti-FITC microbeads, according to the manufacturer’s instructions (Miltenyi Biotec). NK cells were then enriched by magnetic separation (negative selection), using an AutoMACS Magnetic Separator (Miltenyi Biotec). The enriched fraction consistently yielded >5 × 10^6 cells per spleen with a purity of ~80–90%, which was assessed by flow cytometry. CD3^+ and CD8^+ cells in the enriched fraction were consistently below 3%, and the remaining 10–20% of cells were believed to be a mixture of premature hematopoietic and endothelial cells. Following enrichment, NK cells were divided and lysed in appropriate buffers for RNA and protein analyses or were used for cytotoxicity assays.

RNA extraction, RT-PCR, and real-time RT-PCR

Total RNA was isolated from ~2 × 10^6 NK cells per spleen, using the RNasey Mini Kit (Qiagen). With the Superscript III First-Strand Synthesis SuperMix (Invitrogen) for RT-PCR, 100 ng total RNA was reverse-transcribed, and relative quantification of mRNA levels was performed by real-time RT-PCR (SYBR Green and TaqMan gene expression assays; Applied Bio-
systems), using an ABI Prism 7700 Sequence Detector. The following primer sequences were used. Per2F: 5'-GCGGACTTTCCATATTCTCTC-3'; R: 5'-
GCTTCCAGGCTTGATGAA-3'; Bmal1F: 5'-TCCGATAGCAGCTGA-
AACAC-3'; R: 5'-CTCGGTCACATCCTCAGACA-3'; perfdinF: 5'-GC
ATCGTGCTCCAAAGCCAGTG-3'; R: 5'-GCCAGCGGACCCCTGCT-
TCA-3'; and specifically designed probes were used for gram7nize-B and IFN-γ (Applied Biosystems). Analyses were performed using the standard curve method with GAPDH (Applied Biosystems) as the normalizing endogenous control. Relative mRNA expression levels were calculated as the percentage of the maximum value over the 24-h period (45). The sample with the maximum value was used to compute percentages for all other samples by dividing each ratio by the maximum and converting to a percentage. Values at each CT represent a mean and SE of these values.

Western blot

For protein analyses, enriched NK cells (~2 × 10^6 cells per spleen) were lysed with a buffer containing protease and phosphatase inhibitors (25 mM Tris-HCl, pH 7.4; 150 mM NaCl; 1% Nonidet P-40; 1 mM EDTA; and 5% glycerol with Pierce Halt Protease Inhibitor). Protein (50 μg) from total lysate was loaded and separated on a 4–20% SDS-PAGE and transferred onto polyvinylidene difluoride membranes. Membranes were incubated with primary Ab for 1 h at 4˚C in blocking buffer, 5% w/v nonfat dry milk in TBS and 0.1% Tween 20. Primary Abs were used as follows: anti-BMAL1 rabbit polyclonal (1:500; Millipore), anti-PER2 mouse monoclonal (1:100; BD Biosciences), anti-perforin rabbit polyclonal (1:250), anti-granzyme B mouse monoclonal (1:200), and anti–IFN-γ mouse monoclonal (1:250) Abs (Santa Cruz Biotechnology). Membranes were washed and incubated with horseradish peroxidase-conjugated secondary Abs (1:5000; Vector Laboratories) for 1 h at room temperature. Membranes were incubated with ECL Western blot chemiluminescence reagent (Pierce) and exposed to x-ray film, which was developed, and densitometry was performed using Image J analysis software (National Institutes of Health). Each protein was normalized to corresponding intensities for β-actin. Relative protein expression values were calculated as the per-
centage of the maximum value over the 24-h period (46). The sample with the maximum ratio was used to compute percentages for all other samples by dividing each ratio by the maximum and converting to a percentage. Values at each CT represent a mean and SE of these values.
NK cell cytotoxicity

The cytotoxicity of enriched NK cells collected at each time point was determined by calcein AM assays using YAC-1 murine lymphoma cells (American Type Culture Collection) as target cells. YAC-1 cells were grown and maintained in RPMI 1640 without phenol red (Cellgro), containing 1% antibiotics (Sigma-Aldrich) and 10% FBS (Sigma-Aldrich). YAC-1 cells were washed and incubated with 5 mM calcein AM (Sigma-Aldrich) in serum-free RPMI-1640 for 10 min at 37°C. Labeled YAC-1 cells were washed and plated into U-bottom 96-well plates (Falcon) at a concentration of 5 × 10^4 cells per well. NK cells were added at various E:T ratios in triplicates. YAC-1 cells in RPMI alone were used to determine spontaneous calcein AM release, whereas maximal release was achieved by lysing target cells in buffer (0.1% Triton X-100). NK cells collected over the circadian timescale were preincubated for 12 h with IL-2 at 37°C (100 ng/ml; R&D Systems) prior to 4-h incubation with YAC-1 target cells. IL-2 was used to increase assay efficiency owing to the comparatively selective nature of this cytokine to increase cytoxicity in NK cells (47). NK cells collected from animals inoculated with MADB106 cells at CT19 were incubated at 37°C for 4 h with YAC-1 cells in the absence of IL-2. All assays were analyzed using a fluorescence plate reader (Tecan). The percent cytotoxicity for each sample was calculated as follows: % cytotoxicity = [experimental well – spontaneous well]/(max lysed well – spontaneous well)] × 100. The percentages at each E:T ratio were converted to LUs per 10^3 effector cells and based on 20% specific cell lysis (48).

Tumor cell inoculation and metastases

Following control and chronic shift-lag protocols, animals were placed under DD for 5–7 d and inoculated at CT19 with MADB106 tumor cells (1 × 10^5 cells/0.2 ml/rat) into the jugular vein, under isoflurane anesthesia (Henry Schein). As noted above, CT19 was determined by wheel-running data for each animal. This time was chosen based on previous studies and our pilot data showing a maximal response of NK cells during the dark phase of the LD cycle (45, 46). Isoflurane was used to minimize the amount of time the animal was anesthetized. Small incision sites (~1 cm) were closed with a surgical clip, and rats were placed back into their home cage. Surgical and inoculation procedures took ~5–7 min to complete per animal. Animals were sacrificed 6–8 wk after tumor cell inoculation, and each animal was inspected for the presence of visible tumors. As expected, tumors were located primarily in the lungs. Lungs were collected and placed for 24 h in Bouin’s solution (72% saturated picric acid solution, 23% formaldehyde 37% solution, and 5% glacial acetic acid). Lungs were washed with ethanol, and two individuals separately counted surface tumor metastases.

Plasma analysis of corticosterone

Blood samples (~80–100 μl) were collected at CT7 and CT19 in 15 μl EDTA and centrifuged for plasma. Plasma was analyzed for corticosterone levels by a competitive ELISA (IBL International) according to the manufacturer’s recommendations. All samples from both groups were run on a single 96-well plate to limit variability. Standard curves were used to determine corticosterone levels.

Data analysis

Two-way ANOVA with Bonferroni post hoc tests was used to determine differences between control and shift-lag groups at each circadian phase. The gene and protein expression data were tested for significant circadian rhythmicity, using CircWave v. 1.4 software (21, 49) to determine the best-fitting linear harmonic regression with an assumed period of 24 h and with a significance level of 0.05. The center of gravity of each best-fitting waveform, which comprises both sine and cosine components, in CircWave was used as the peak expression. The circadian acrophases of NK cell activity and protein and gene expression, tumor frequency data, and circadian periods. The Mann–Whitney U test was used to compare tumor frequencies between different defined chronotypes of the chronic shift-lag animals. Circadian period and robustness of running-wheel activity rhythms were derived using Lomb–Scargle periodograms during and following the control and shift-lag protocols (ClockLab).

Results

Circadian wheel-running activity rhythms

Control animals under the standard LD cycle displayed typical circadian entrainment patterns (Fig. 1A) with an approximate period of 24 h (Fig. 1D). Under the chronic shift-lag protocol, rats exhibited different patterns of locomotor activity rhythms. For most shift-lag animals, periodogram analysis revealed two distinct peaks, one corresponding to the mean period of the shift-lag schedule (i.e., ~21 h) and one corresponding to the typical, longer-than-24 h free-running period normally shown by rats under free-running conditions (Fig. 1E, 1F). However, individual variation was considerable among shift-lag animals in the relative prominence of these peaks and in a clear correspondence between the periodogram results and the actigraphic record within individuals (Fig. 1B, 1C). In DD following the shifting paradigm, rhythm amplitude was significantly reduced, with no observable change in period (Fig. 2A, 2B; p < 0.0001). Thus, the shift-lag protocol induced considerable circadian disruption and also resulted in light exposure during the animals’ subjective night.

Chronic shift-lag alters the circadian expression of clock genes in enriched NK cells

To determine whether chronic shifting affected the molecular clock in peripheral immune cells, we examined gene and protein expression of two canonical clock genes, Per2 and Bmal1 (52), from enriched NK cells collected during DD at 4-h intervals across the circadian cycle. In control animals, both Per2 and Bmal1 gene and protein expression displayed robust circadian rhythmicity (Fig. 3; all p < 0.01). Acrophase estimates (estimate ± SD) derived from fitted curves indicated Per2 and Bmal1 expression peaks were out of phase with one another at both the gene (Fig. 3A, 3C) and the protein (Fig. 3B, 3D) levels (Table I), as expected from previous studies (45). In addition, as hypothesized, chronic shift-lag altered the circadian expression of both clock genes and proteins; nevertheless, these displayed significant rhythmicity in shift-lag animals (Fig. 3; all p < 0.01). Comparison of acrophases for Per2 and Bmal1 revealed that expression rhythms were shifted in NK cells from shift-lagged animals compared with controls (Table I; Fig. 3A, 3C; p < 0.05), whereas Bmal1, but not PER2, phase of peak expression was affected by shift-lag (Table I; Fig. 3B, 3D). Moreover, two-way ANOVAs indicated significant group by time interactions (p < 0.05) for Per2 and Bmal1 gene and protein expression rhythms, which were further explored using Bonferroni post hoc tests to indicate significant changes in levels of expression at each CT. Levels of expression were significantly altered at particular CTs for Per2 and Bmal1 (Fig. 3), further indicating chronic shift-lag affected the rhythmic expression of core clock genes, with the most profound effects on Bmal1 expression.

Chronic shift-lag alters the circadian expression of the cytokine IFN-γ and cytolytic factors perforin and granzyme B in enriched NK cells

Alterations in circadian expression of core components of the molecular clock were paralleled by changes in expression rhythms of cytokines and cytolytic factors in NK cells. Consistent with previous results (37, 38), NK cells from control animals displayed robust circadian rhythms of gene and protein expression of perforin, granzyme B, and IFN-γ (Fig. 4; assumed period of 24 h, all p < 0.01), with expression peaks during the active phase (Table I). In line with our hypothesis, chronic shift-lag altered the rhythmic gene and protein expression of perforin, granzyme B, and IFN-γ (Fig. 4). Chronic shift-lag completely abolished the gene and protein rhythms of perforin in NK cells, as indicated by a lack of significant curve fitting and decreased gene expression across the entire circadian cycle (Fig. 4A, 4B; compared with controls, all CTs, p < 0.05). Although gene expression rhythms remained intact for granzyme B and IFN-γ in animals undergoing shift-lag (p < 0.01), these gene rhythms were shifted to earlier in the circadian cycle, as indicated...
by changes in acrophase (Table I) and changes in expression levels at particular phases (Fig. 4C, 4E). The protein expression of granzyme B and IFN-γ did not show a similar shift in peak expression (Table I; Fig. 4D, 4F). In general, alterations in rhythms of these factors due to shift-lag resulted in reductions of gene and protein levels near or at phases of peak expression in control animals (Fig. 4; p, 0.05). Therefore, chronic shift-lag appeared to differentially alter gene and protein expression of cytokines and cytolytic factors by either shifting the circadian rhythm to earlier in the cycle or attenuating levels at phases of peak expression.

Chronic shift-lag alters the circadian rhythmicity of NK cell cytolytic function

In an effort to investigate whether changes in circadian rhythms of cytokines and cytolytic factors due to chronic shift-lag had consequences for NK cell function, we completed a series of NK cell cytotoxicity assays across the circadian cycle. The cytotoxic function of NK cells displays a robust circadian rhythm with a peak during the subjective night (Table I; Fig. 5A; p, 0.001), which is in line with our previous results (46). In shifted animals, NK cell cytotoxicity remained rhythmic, with a shift in the rhythm by ∼16 h (Table I; Fig. 5A; p < 0.05). Importantly, it is evident that peaks of NK cell cytotoxicity observed in controls animals were dramatically reduced in the shifted group (Fig. 5A; p < 0.05). From these data, it is suggested that coordinated circadian peaks of expression of perforin, granzyme B, and IFN-γ are important for NK cell-mediated killing.

To determine whether chronic shift-lag altered NK cell function in vivo, enriched NK cells were collected from animals injected with MADB106 tumor cells. On the premise that circadian activity of NK cell function is critical for immune response to cancer, tumor cells were injected at CT19—the phase of maximal cytolytic activity (Table I; Fig. 5A). As expected, NK cells from shifted animals collected 24 h following the i.v. introduction of tumor cells displayed significantly reduced cytolytic activity (Fig. 5B; p < 0.01). From these data, we interpret that shift-lag significantly altered the circadian rhythm of NK cell cytolytic function and the NK cells’ response to tumor challenge.
Decreased innate immune function in chronically shifted animals is associated with increases in lung tumor frequency and prevalence

Control and shift-lag animals injected at CT19 with MADB106 tumor cells were left for 6–8 wk to determine tumor frequency or prevalence. In line with our hypothesis, chronic shift-lag promoted tumor growth, as indicated by increased frequency and prevalence of tumors in the lungs (Fig. 6A, 6B; \( p < 0.001 \)). In addition, tumor load was compared between animals displaying two different chronotypes in response to the shifting protocol—animals that exhibited either free-running activity rhythms or re-entrainment (Fig. 1B, 1C, respectively). The shifted group that displayed repeated re-entrainment tended to have more tumors relative to the group that did not successfully synchronize to the shift-lag schedule, but this difference did not reach significance (Fig. 6C; \( p < 0.07 \)). Representative lungs from control and shifted (Fig. 6D, rows 1 and 2, respectively) animals are pictured. Two animals from the chronic shift-lag group were removed from the analysis because tumor growth was so invasive to the lungs and surrounding organs that each required sacrificing 3 wk following injection (not included in the analysis). No control animals required sacrificing prior to the planned 6–8 wk.

**Figure 3.** Chronic shift-lag alters the circadian expression of clock genes and proteins in enriched NK cells. Expression levels of Bmal1 and Per2 genes (A, C) and associated proteins (B, D) in NK cells separated from animals at different time points in DD following either control or shifted paradigms. Representative immunoblots are shown above respective protein plots across all CTs. Densitometric quantification of protein levels was done by Image J software. Sine wave fits using linear harmonic regression with an assumed period of 24 h for control and shifted groups (solid and dotted lines, respectively) are superimposed with group means ± SEMs for each CT. All curve fits are significant \( (p < 0.01) \). *\( p < 0.05 \), significant difference between groups at CT.

**Table I.** Chronic shift-lag alters the acrophases of gene, protein, and cytolytic activity rhythms in NK cells

<table>
<thead>
<tr>
<th>Gene</th>
<th>Protein</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Control</strong></td>
<td><strong>Shifted</strong></td>
</tr>
<tr>
<td>Per2</td>
<td>9.54 ± 3.26*</td>
</tr>
<tr>
<td>Bmal1</td>
<td>20.88 ± 2.21</td>
</tr>
<tr>
<td>Perf</td>
<td>22.24 ± 3.23</td>
</tr>
<tr>
<td>Gran B</td>
<td>20.66 ± 1.76</td>
</tr>
<tr>
<td>IFN-γ</td>
<td>18.97 ± 2.46</td>
</tr>
<tr>
<td><strong>Cytolytic Activity</strong></td>
<td></td>
</tr>
<tr>
<td>NK cells</td>
<td>17.35 ± 3.04</td>
</tr>
</tbody>
</table>

Acrophases were determined from curves fitted to gene and protein expression and NK cell activity levels over the circadian cycle. Acrophases (±SD) represent estimations of the circadian phase corresponding to the peak of the rhythm. One-way ANOVA with Newman–Keuls multiple comparison tests were used to compare acrophases between control and shifted groups \( (n = 6 \text{ per lighting regimen}) \) by gene and protein, or, in the case of NK cell activity \( (n = 4 \text{ per lighting regimen}) \), a Student \( t \) test was used.

*\( p < 0.05 \), significant differences between groups. **\( p < 0.05 \), significant differences between Per2 and Bmal1.

**Suppressed NK cell function in animals undergoing chronic shift-lag may not be due to increases in plasma corticosterone**

Other studies have shown that repeated shifting of the LD cycle either shifts the rhythm or transiently increases concentration of plasma corticosterone (15, 53). To determine whether shifting increased plasma corticosterone, levels were measured at two time points, CT7 and CT19, at the time of sacrifice (5–7 d into DD; \( n = 4 \text{ per time point} \)). Corticosterone levels did not differ between groups at either time point (Fig. 7). At CT19, the time of maximal NK cytolytic activity in controls, corticosterone levels were not increased in shifted animals, suggesting that attenuation of NK function may not be due to activation of the stress axis.
Discussion
The circadian system regulates numerous cellular and physiological processes, including immune function (34). At the top of the circadian hierarchy lies the SCN, which coordinates timing of peripheral clocks by hormonal and autonomic nervous system signals (54). For example, immune tissues receive rhythmic sympathetic input that modulates clock gene expression and other circadian functions within immunomodulatory cells, including splenic NK cells (37, 55, 56). NK cell function seems to be passively regulated by clock mechanisms, as clock genes, cytokines, and cytolytic factors all oscillate in a circadian fashion (38, 45). The present study demonstrates that circadian coordination of the immune response may influence the degree to which the innate immune system initially reacts to circulating tumor cells. During a state of chronic circadian desynchrony, the cytolytic activity of NK cells is almost completely suppressed, quite possibly by disruption of the coordinated expression of multiple cytolytic factors. We postulate that this mechanism is responsible for the increased incidence of lung tumors in animals undergoing chronic shift-lag.

A number of epidemiological studies have linked disruptions in circadian rhythms to cancer susceptibility and clinical course, although the genetic and physiological mechanisms underlying these findings are poorly understood. Much attention has been focused on the role of the molecular clock in cell proliferation, DNA damage and repair, and apoptotic pathways within the developing tumor cells (16, 57–59). In contrast, the current study shows that the tumorigenic effects of circadian disruption depend on tumor surveillance mechanisms of NK cells. Specifically,
chronic shift-lag altered the expression of clock genes and the immunomodulators IFN-γ, perforin, and granzyme B in enriched NK cells at both the gene and the protein levels. Thus, the functional activity of NK cells was significantly decreased at normal phases of peak expression in shift-lag compared with control animals. Further, gene expression rhythms of granzyme B and IFN-γ displayed a shift by ~4 h to earlier in the circadian cycle, but similar shifts were not observed at the protein level. Perforin expression appeared the most affected by chronic shift-lag, as gene and protein rhythms were completely abolished in NK cells. It is plausible that these reductions of perforin hamper NK cell-mediated apoptosis and lysis of tumor cells in shifted animals (60–62), especially because granzyme B and IFN-γ were also reduced, which has been associated with increased tumor susceptibility (31).

Circadian coordination of NK cell function appears to be important for generating an efficient immune response. Consistent with our previous findings, circadian rhythms of perforin, granzyme B, and IFN-γ expression peak at similar phases during the subjective night, corresponding to the circadian peak of NK cell cytotoxicity (45, 46). In an effort to determine whether circadian coordination of NK cell function may have functional significance, we injected control and shifted animals with tumor cells at CT19 according to the hypothesis that chronic shift-lag would cause impairment of NK cell function, ultimately producing a shift in the rhythms of cytotoxicity. Somewhat surprisingly, however, chronic shift-lag dramatically attenuated NK cell function across the entire circadian cycle, suggesting that similar tumor frequencies might have been observed in shifted animals regardless of phase of tumor cell injection. The apparent discoordination of cytokines, cytolytic factors, and cytotoxicity, along with the attenuation of cytotoxicity following an in vivo tumor challenge, suggests that circadian disruption compromises the ability of NK cells to kill tumor cells. In our model, this compromise could lead to the promotion of tumor growth by increasing the probability of tumor cell retention following inoculation.

To assess circadian disruption at the whole-animal level, we monitored circadian activity rhythms throughout the experimental protocols. Because rhythmic locomotor activity is a reliable indicator of circadian pacemaker functioning, changes in amplitude and/or period due to chronic shift-lag can be inferred to reflect SCN activity. Inspection of actograms from our shifted animals yielded two different chronotypes. Although some shifted animals exhibited an apparent free-running period, essentially ignoring the changing light onsets, others displayed short circadian periods as a consequence of partial synchronization to the advancing shift-lag schedule. Using a similar shifting protocol with hamsters, the former has been observed (53). Nevertheless, despite shifting-induced alterations in rhythm amplitude and period, our animals remained completely rhythmic during the experiment and thus suggest the SCN maintains coherent rhythmicity even during rapidly changing environmental cues (53). Studies examining the effects of photic phase shifts on SCN clock gene expression have also demonstrated the SCN maintains rhythmicity with differing periods of re-entrainment, depending on frequency and direction of the shifts (63).

Furthermore, examination of circadian activity rhythms during the exposure to the shift-lag schedule revealed that repeated re-entrainment may have led to increased tumorigenesis relative to animals expressing stable free-running rhythms under the same conditions. Because re-entrainment is associated with internal circadian desynchrony at both the molecular and the physiological levels (64, 65), these preliminary observations support the hypothesis that dissociation among multiple circadian clocks may indeed underlie the pathophysiological effects of shift-lag exposure.

Nevertheless, although rhythmicity is maintained in the SCN, major SCN outputs that relay temporal information to other brain areas and peripheral tissues are compromised under shift-lag conditions (63, 64). Clocks in peripheral tissues re-entrain at a slower rate, compared with the SCN, following a single advance or delay, including those in the spleen (63, 65). Sympathetic input to the spleen may be a major output pathway by which the SCN coordinates NK cell function (37). It is interesting to speculate that...
chronic shift-lag impedes temporal coordination of the immune system by attenuating SCN-derived output signals to “preautonomic” neurons (66). Indeed, homeostasis of the sympathetic nervous system is disrupted in chronically shifted animals (17).

The molecular clock in NK cells was altered by chronic shift-lag, which was evident by advances in the rhythms and reductions of Bmal1 gene and protein expression, as well as a delay in the expression rhythms of Per2. Bmal1 rhythms were slightly attenuated but remained rhythmic. The delay of the Per2 gene was much more dramatic (~6 h) than the pattern observed in the protein rhythm (~1 h). Although shifting-induced reductions in NK cell function were correlated with changes in clock gene expression, these data can only suggest the involvement of molecular clocks in NK cells in regulating cytolytic abilities. Similarly, in macrophages, disruptions to clock genes due to repeated shifting has been associated with dysregulation of inflammatory responses (20). To determine the extent to which molecular clocks within NK cells regulate cell-specific functions, further studies are necessary to tease apart the interactions between clock and immune-related pathways. At the molecular level, a potential intersection could be clock modulation of NF-kB, a key transcription factor involved in cellular immune response (67). When activated, NF-kB can promote gene transcription of IFN-γ (68), perforin (69), and granzyme B (70) by direct binding to enhancer elements.

Other factors could be involved in mediating the effects of chronic shift-lag on NK cell function beyond involvement of circadian mechanisms. Circulating glucocorticoids and melatonin oscillate in antiphase, peaking during the early and late circadian cycle, respectively, and are known modulators of NK cell function (71, 72). However, several points of evidence suggest glucocorticoids and melatonin may not necessarily be critical for maintaining immune rhythms. In macrophages and NK cells, functional rhythms persist despite the absence of glucocorticoid rhythms (73, 74) or lack of melatonin production (73, 75). Furthermore, consistent with a previous result (20), our study suggests dysregulation of NK cell function induced by chronic shift-lag is not due to increased circulating corticosterone, especially during times of peak functioning. Corticosterone increases induced by chronic shifting paradigms may be transient and play a role in resynchronization of the circadian system (49, 53). Nevertheless, at the time points of NK cell collection, corticosterone was not increased in our shifted animals.

In conclusion, we have demonstrated that chronic states of circadian desynchronization severely attenuated NK cell function, possibly by altering the circadian rhythms of cytokines, cytolytic factors, and cytotoxicity. In addition, suppressed NK cell function was observed during an in vivo immune-stimulated condition in shifted animals, suggesting circadian disruption has substantial effects on the tumor response of NK cells. Importantly, loss of rhythms in NK cell cytolytic activity was associated with an increased frequency of tumors. More detailed studies are required to further understand the integration of physiological and molecular changes in immune function caused by disruption to the circadian system. For example, whether the chronic shift-lag model used in this study alters noradrenergic signaling of the sympathetic nervous system to impact NK cell function has yet to be determined (17). Molecular studies, such as large-scale gene expression arrays in primary and secondary immune tissues, of wild-type and clock mutant mice may also provide a wealth of information regarding the molecular links between circadian and immune systems. Understanding these intersections could be invaluable to developing novel therapeutics to restore circadian functioning in immune cells, which could aid in preventing disease progression (59, 76, 77).

Disclosures
The authors have no financial conflicts of interest.

References