



Histone Lysine Demethylase JARID1a Activates CLOCK-BMAL1 and Influences the Circadian Clock Luciano DiTacchio *et al. Science* **333**, 1881 (2011); DOI: 10.1126/science.1206022

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ways that are not feasible offline, there are also important limitations. First, unlike laboratory studies, we have little data on conditions that are known to influence mood, including demographic and occupational backgrounds that may influence when and how much people sleep, the level and timing of environmental stress, susceptibility to affective contagion, and access to social support. Second, lexical analysis measures the expression of affect, not the experience. Cultural norms may regulate the appropriateness of affective expression at different times of the day or week. Because these norms are unlikely to be universal, the robust patterns we observed across diverse cultures (as well as across days of the week) give us confidence that affective expression is a reliable indicator of diurnal individual-level variations in affective state.

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## Histone Lysine Demethylase JARID1a Activates CLOCK-BMAL1 and Influences the Circadian Clock

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In animals, circadian oscillators are based on a transcription-translation circuit that revolves around the transcription factors CLOCK and BMAL1. We found that the JumonjiC (JmjC) and ARID domain—containing histone lysine demethylase 1a (JARID1a) formed a complex with CLOCK-BMAL1, which was recruited to the *Per2* promoter. JARID1a increased histone acetylation by inhibiting histone deacetylase 1 function and enhanced transcription by CLOCK-BMAL1 in a demethylase-independent manner. Depletion of JARID1a in mammalian cells reduced *Per* promoter histone acetylation, dampened expression of canonical circadian genes, and shortened the period of circadian rhythms. *Drosophila* lines with reduced expression of the *Jarid1a* homolog, *lid*, had lowered *Per* expression and similarly altered circadian rhythms. JARID1a thus has a nonredundant role in circadian oscillator function.

o gain insight into the dynamics of chromatin modifications and the function of CLOCK-BMAL1 transcription factors in the circadian clock, we measured the state of two histone modifications that correlate with active transcription, acetylation of histone 3 (H3) lysine 9 (H3K9Ac), and trimethylation of H3 lysine 4 (H3K4me3) at the *Per2* promoter (*1*). In mouse liver, both modifications synchronously oscillated at the *Per2* gene promoter CLOCK-BMAL1 E2– binding site ("E-box") (*2*), with lowest amounts at circadian time (CT, the endogenous, free-running time) 3 hours after the onset of activity (CT3) and peak amounts at CT12 (Fig. 1A). The peaks of histone modification were followed by those of *Per2* mRNA abundance. We also found BMAL1 abundance rhythms at the E-box, which reached a maximum at CT9 (Fig. 1B). Histone acetyl-transferases (HATs) and histone deacetylases (HDACs) generate rhythms in histone acetylation and have important roles in circadian rhythms

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#### Supporting Online Material

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(3, 4). H3K4me3 modification at promoter regions correlates with transcriptional potential, which suggests that this mark helps maintain a transcriptionally poised state (5). Recently, the H3K4 methyltransferase MLL1 was shown to have a necessary role in CLOCK-BMAL1–dependent transcription (6).

We focused on a JumonjiC (JmjC) domaincontaining H3K4me3 demethylase family with four mammalian and one insect gene members (fig. S1). In murine liver chromatin, JARID1a was enriched at the Per2 E-box, and its profile at this site coincided with that of BMAL1 (Fig. 1B). Jarid1a expression in liver did not show robust oscillations (fig. S2), which suggested that recruitment of JARID1a to the Per2 promoter might be mediated by the circadian machinery. Indeed, JARID1a recruitment to the Per2 promoter E-box but not at a non-CLOCK-BMAL1 JARID1a target is reduced in  $Bmal1^{-/-}$  cells (Fig. 1C and fig. S17A). Consistently, immunoprecipitation of endogenous CLOCK or BMAL1 from nuclear extracts copurified with endogenous JARID1a (Fig. 1D). Similarly, CLOCK and BMAL1 associated with immunoprecipitated JARID1a (fig. S3). Overexpression of JARID1a enhanced CLOCK-BMAL1-mediated transcription from Per2 (Fig. 1E) and Per1 (fig. S4A) promoters in a dosedependent manner but failed to coactivate expression from an unrelated (E74-Luc) reporter (fig. S4D). Furthermore, coactivation of CLOCK-BMAL1 by JARID1a did not require its histone demethylase activity, as JARID1a mutants that carry a loss-of-function mutation (H483A, in which Ala replaces His<sup>483</sup>) (7) or that lack the JmjC domain enhanced CLOCK-BMAL1 activity, reversed

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HDAC1 repression, and increased acetylation amounts at the *Per2* E-box (Fig. 1F and figs. S4B, S4C, and S5).

In contrast, JARID1b and JARID1c repressed CLOCK-BMAL1-mediated activation of *Per2* transcription (fig. S6, A and B). Accordingly, overexpression of JARID1b and JARID1c, but not JARID1a, reduced H3K4me3 modification at the *Per2* promoter (fig. S6C). Overall, these

Fig. 1. IARID1a coactivates CLOCK-BMAL1-mediated transcription of Per genes. (A) H3K4 trimethylation and H3K9 acetylation oscillations at the Per2 promoter E-box in mouse liver shown alongside Per2 mRNA expression (secondary y axis). Normalized average (+SEM, n = 3) of chromatin immunoprecipitations (ChIPs) performed with antibodies against H3K4me3 or H3K9Ac followed by QPCR analysis. Chromatin modifications shown as a percentage of total chromatin used (input) normalized to the corresponding lowest value (H3K4me3 = 0.27%, H3K9Ac = 0.25%) for ease of comparison. (B) Abundance of BMAL1 and JARID1a in mouse liver at the Per2 promoter E-box and flanking regions. (C) Absence of JARID1a from the Per2 promoter E-box in *Bmal1*<sup>-/-</sup> cells. (**D**) Association of endogenous JARID1a and CLOCK-BMAL1. Immunoprecipitates (IPs) obtained with the indicated antibodies from asynchronous U2OS (a human osteosarcoma cell line) nuclear extracts were immunoblotted (IB) with antibodies against JARID1a, CLOCK, BMAL1, or βACTIN. (E) Effect of JARID1a on CLOCK-BMAL1dependent transcription from a *Per2:Luc* reporter. Luciferase activity (light counts) from HEK293T cells transiently expressing full-length JARID1a cDNA and Per2:Luc reporter construct are shown. (F) Wild-type (WT) or demethylase-mutant JARID1a H483A (MUT) rescues CLOCK-BMAL1 activation of a Per2 reporter construct in Jarid1a<sup>-/-</sup> cells.

Fig. 2. JARID1a is required for normal circadian function. (A) Real-time bioluminescence from Jarid1a<sup>+/+</sup> (black) and Jarid1a<sup>-/-</sup> (red) mouse fibroblasts stably expressing a Per2:Luc construct. (Middle and right) Average period lengths (+SEM, n = 10) and amplitude estimates (\*\*\*P < 0.001, two-tailed Student's t test). (B) Per2, Cry1, Dbp, and SDF1 mRNA abundance in Jarid1 $a^{+/+}$  cells (black) and Jarid1 $a^{-/-}$ cells (red). Forty-eight hours after circadian synchronization, cells were collected at 3-hour intervals over the course of 24 hours, and the mRNA abundance was analyzed by reverse transcription (RT)-QPCR and normalized to that of Gapdh. Average normalized values ( $\pm$ SEM, n = 3) are plotted at various times after cell synchronization. (C) Representative real-time luminescence from U2OS cells stably expressing Bmal1:Luc and transfected with scrambled (S, black) or Jarid1a siRNA (red). Effects of different Jarid1a siRNA concentrations (10, 20, and 40 nM) on the endogenous Jarid1a mRNA abundance (average + SEM, n = 3) and period length (average + SEM, n = 5) relative to those of cells

results suggest that JARID1a enhances CLOCK-BMAL1 activation of the *Per2* promoter, whereas JARID1b and JARID1c might function to reverse its H3K4me3 levels during the repression phase of *Per* transcription.

To test whether JARID1a has a nonredundant role in the cell-autonomous oscillator, we stably transfected a *Per2:Luciferase (Per2:Luc)* reporter into fibroblasts derived from either *Jarid1a*<sup>+/+</sup>

(wild type; WT) or *Jarid1a<sup>-/-</sup>* (knockout, KO) mice (7) and monitored circadian rhythms in *Per2:Luc* luminescence. In cell populations (Fig. 2A and fig. S7A) and single cells (fig. S8), *Per2* transcriptional rhythms in a *Jarid1a<sup>-/-</sup>* genetic background showed a significantly shorter period than those of WT controls [WT = 23.7 ± 0.2 hours, KO = 22.4 ± 0.3 hours,  $P = 6.68 \times 10^{-8}$ , n = 10 each genotype, one-way analysis of variance



transfected with scrambled siRNA (40 nM). Representative results from at least three independent repetitions are shown.

(ANOVA)] (Fig. 2A). Amounts of endogenous Per2 transcript were also reduced and oscillated with a shorter period in Jarid1 $a^{-/-}$  cells (Fig. 2B). Similarly, other CLOCK-BMAL1 targets were significantly reduced in  $Jarid1a^{-/-}$  cells, and their rhythms were either absent or dampened (Fig. 2B and fig. S9). In contrast, abundance of SDF1 mRNA, a repressive target of the histone demethylase (KDM) activity of JARID1a (7), was increased in Jarid1a<sup>-/-</sup> fibroblasts. Acute small interfering RNA (siRNA)-mediated depletion of Jarid1a mRNA also shortened circadian period length in a dose-dependent manner (Fig. 2C and fig. S7B). Conversely, transient transfection reconstitution of either wild-type or demethylase mutant JARID1a into Jarid1a<sup>-/-</sup> cells resulted in a circadian period lengthening (fig. S10). These results indicate that JARID1a has a nonredundant role in maintaining the normal periodicity of the circadian oscillator.

The overall mechanism and several components of the circadian oscillator are conserved between mammals and insects. The *Drosophila* genome has only one *Jarid1* family gene, *little* 

Fig. 3. Circadian rhythm disruption in *lid*<sup>10424</sup> flies. (A) RT-QPCR of lid, per, cry, and tim mRNA in the heads of *y;ry* (black) and *lid*<sup>10424</sup>/CyO (red) flies collected every 4 hours after 72 hours in constant darkness (±SEM, n = 3). (B) Representative circadian double-plotted actograms of wild-type *lid*<sup>+/+</sup> (*y*;*ry*), *lid*<sup>10424</sup>/CyO, *lid*<sup>10424</sup>/*lid*<sup>10424</sup>;*gLid*, and lid<sup>10424</sup>/lid<sup>10424</sup>;gJmjC\* flies under LD or DD conditions. Arrowheads indicate transition from LD to DD. (C) (Left) Normalized average activity of *y;ry* (black) and *lid<sup>10424</sup>/CyO* flies (red) (n = 33) under LD and DD conditions. (Right) Summary of the circadian activity rhythms in various Drosophila strains tested. Histogram shows the percentage of flies with normal (R, rhythmic), weak (W) or no detectable (A, arrhythmic) circadian activity rhythms (+SEM). (D) Normalized average activity (+SEM; n = 16 to 75) in 30-min bins during the light:dark period show two major phases of activities in *v;ry* flies at morning and evening.

imaginal discs (lid) (fig. S1). Flies carrying the hypomorphic allele  $lid^{10424}$  (P-element insertion in the first intron) over the balancer chromosome CyO (lid<sup>10424</sup>/CyO) expressed less lid mRNA than did WT y;ry flies (Fig. 3A). Consistent with our observations in  $Jarid1a^{-/-}$  fibroblasts (Figs. 1 and 2), *lid<sup>10424</sup>/CyO* flies also expressed reduced amounts of *per*, *cry*, and *timeless* (*tim*) mRNA. These *lid*<sup>10424</sup>/*CyO* flies showed normal activity distribution under a 12 hours of light:12 hours of dark (LD) cycle (Fig. 3B), and their total activity  $(1030 \pm 44)$ , average beam breaks for 3 days in LD, SEM, n = 75) was equivalent (P > 0.13, oneway ANOVA) to that of WT  $y_{,ry}$  (831 ± 85, SEM, n = 16) or +/CyO (938 ± 60 SEM, n = 45) flies, which implied no deficit in locomotion or light:dark entrainment of the circadian clock.

However, under constant darkness (DD), the *lid*<sup>10424</sup>/CyO flies exhibited disrupted circadian activity rhythms. The majority of *lid*<sup>10424</sup>/CyO flies showed no or weak circadian activity rhythms (Fig. 3, B and C). Those *lid*<sup>10424</sup>/CyO flies with detectable rhythms showed significant period shortening (Fig. 3C) (*lid*<sup>10424</sup>/CyO = 22.91 ± 0.04 hours,

 $y;ry = 23.61 \pm 0.06$  hours; average  $\pm$  SEM, n = 16and 75; P < 0.001). Expression of a genomic copy of *lid* in *lid*<sup>10424</sup>/*lid*<sup>10424</sup> homozygous flies (lid<sup>10424</sup>/lid<sup>10424</sup>;gLid) restored Lid protein expression to near-WT amounts and also restored circadian activity rhythms (Fig. 3C and fig. S11). Furthermore, a demethylase-deficient Lid allele (8) partially rescued the circadian phenotype  $(lid^{10424}/lid^{10424};gJmjC^*)$ . Under LD and DD, the lid<sup>10424</sup> flies exhibited attenuated day:night differences in activity. Specifically, the lid<sup>10424</sup> flies exhibited increased activity during midday and reduced dark anticipation (Fig. 3D). Comparable circadian rhythm disruption and reduced expression of CLOCK-BMAL1 (or CYC) targets in both Drosophila and mammalian cells supports a conserved role of lid/Jarid1 in the insect and mammalian circadian oscillator.

In Jarid1a<sup>-/-</sup> mouse fibroblasts, the Per2 promoter region showed an increased abundance of H3K4me3 (fig. S12A), but a significantly reduced amount of acetylated H3K9 relative to those of WT controls (Fig. 4A). Conversely, overexpression of JARID1a in human embryonic





**Fig. 4.** Dynamic interaction between HDAC and JARID1a correlates with normal histone acetylation at *Per* promoter. (**A**) H3K9Ac detected by ChIP-QPCR in *Jarid1a<sup>-/-</sup>* or WT cells (average +SEM, n = 3; \*\*P < 0.05). (**B**) H3K9 acetylation at the endogenous *Per2* promoter after overexpression of JARID1a or HDAC1 (\*\*P < 0.001, \*\*\*P < 0.0001, n = 3 per group, one-way ANOVA). (**C**) ChIP-QPCR quantification of HDAC1 at E-box of *Per2* promoter in mouse liver. For

(A) to (C), chromatin modifications are presented as percent of total chromatin used for immunoprecipitation (% input). (**D**) Effect of various concentrations of TSA on *Per2:Luc* in *Jarid1a<sup>-/-</sup>* cells. Representative real-time traces are shown. (**E**) Acetylated histone H3 (lane 1) was incubated with 100 ng HDAC1 (lane 2), along with purified JARID1a (lane 3) or JMJD6 (lane 4), and analyzed by immunoblot.

kidney–293 cells (HEK293T cells) increased accumulation of H3K9Ac at the *Per2* promoter (Fig. 4B), but not at a control promoter (fig. S12C). This suggests that coordinated increases in both H3K4me3 and histone acetylation are required for sufficient *Per* induction. Recruitment of JARID1a to *Per* promoters might promote H3K9 acetylation levels during *Per* transcription.

Lid inhibits histone deacetylase Rpd3 to activate transcription at specific loci (9). Such HDAC inhibition by Lid, which is independent of its KDM activity, is consistent with the observed activation of CLOCK and BMAL1 by JARID1a (Fig. 1F and figs. S4 and S5). JARID1a also coimmunoprecipitated HDAC1 in mammalian cells (fig. S13). Chromatin immunoprecipitation followed by quantitative PCR (ChIP-QPCR) analysis from mouse liver chromatin showed a nearly constant HDAC1 occupancy at the Per2 E-box (Fig. 4C). Further, its presence at the Per2 promoter was unchanged in Jarid1a<sup>-/-</sup> cells (fig. S12B). Although HDAC1 likely mediates repression of Per2 transcription (3) (Fig. 4C and fig. S14), its presence during the activation phase is consistent with enrichment of HDACs at promoters of highly expressed and inducible genes (5). The high-amplitude H3K9 acetylation rhythms in the presence of relatively constant amounts of HDAC1 at the Per2 promoter could be partly explained by transient inhibition of HDAC activity. Loss of JARID1a might lead to higher HDAC activity during activation phase, low H3K9 acetylation, and reduced Per transcriptional activation by CLOCK-BMAL1. Indeed, pharmacological inhibition of HDAC by trichostatin A (TSA) led to increased histone acetylation at the Per2 promoter, rescued the period-length defect, and improved the amplitude of Per2:Luc rhythm in Jarid1a<sup>-/-</sup> cells (Fig. 4D). Similar TSA treatment slightly increased the period length of WT cells, but dampened Per2:Luc oscillations (fig. S15). Affinity-purified JARID1a impaired the ability of HDAC1 to deacetylate acetylated histone H3 in vitro, whereas another JmiC domaincontaining protein, JMJD6, could not (Fig. 4E and fig. S16).

These results support a model in which JARID1a mediates transition from repression to robust activation of *Per* transcription. During the activation phase, both H3K4me3 and acetylated histones at CLOCK-BMAL1 target sites accumulate in parallel with *Per* transcription. JARID1a appears to be recruited to the *Per* promoter in complex with CLOCK-BMAL1, where it inhibits HDAC1 activity to enhance histone acetylation and transcriptional activation. Although purified JARID1a

or Lid inhibits HDAC activity in vitro, this mode of regulation is likely gene- and context-dependent in vivo. *Jarid1a<sup>-/-</sup>* cells have increased amounts of H3K4me3 at the *Per* promoter (fig. S12), and overexpression of the JmjC domain of JARID1a slightly repressed *Per* transcription (fig. S5), which suggest that, in vivo, JARID1a may mediate demethylation or that its presence in the transcriptional complex of target genes is required for proper recruitment of other proteins that carry out this function. However, whereas the lysine demethylase activity of JARID1a is dispensable for normal circadian rhythms in mammalian cells, its HDAC-inhibiting function appears to be necessary for proper oscillator function.

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Supporting Online Material www.sciencemag.org/cgi/content/full/333/6051/1881/DC1 Materials and Methods Figs. S1 to S17 References (10–17)

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# Superfast Muscles Set Maximum Call Rate in Echolocating Bats

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As an echolocating bat closes in on a flying insect, it increases call emission to rates beyond 160 calls per second. This high call rate phase, dubbed the terminal buzz, has proven enigmatic because it is unknown how bats are able to produce calls so quickly. We found that previously unknown and highly specialized superfast muscles power rapid call rates in the terminal buzz. Additionally, we show that laryngeal motor performance, not overlap between call production and the arrival of echoes at the bat's ears, limits maximum call rate. Superfast muscles are rare in vertebrates and always associated with extraordinary motor demands on acoustic communication. We propose that the advantages of rapid auditory updates on prey movement selected for superfast laryngeal muscle in echolocating bats.

aryngeal echolocation and insectivory characterize about 70% of present-day bat species (1-3). Over the course of an attack on a flying insect, bats increase their echolocation call emission rates as they progress from prey detection, through approach, to the terminal buzz (1, 2, 4) (Fig. 1A). Increasing call emission rates

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means more information updates per unit time from returning echoes on the relative position of the target. All aerial hawking bats studied to date produce the buzz, which is sometimes subdivided into "buzz I" and "buzz II" phase calls, the former occurring at rates of ~100 to 160 calls/s, and the latter  $\geq$ 160 calls/s (1, 5) (Fig. 1B). Bats do not call at rates exceeding those reached during this final stage of aerial hawking attack (2, 4), and we hypothesize that call production, echo processing, or both limit maximum echolocation call rate.

Laryngeal nerve-cut experiments reveal that each call a bat emits is under active neuromus-

cular control (6, 7). Consequently, muscle performance might place an upper limit on the rate at which bats produce calls. Alternatively, if prey echoes overlap with or return after the next call is emitted, accuracy in target ranging may suffer as a result of ambiguity in matching echoes to calls (1, 8). While hunting, most species avoid potential ambiguity by not producing the next call until target echoes reach the bat's ears (1), potentially limiting maximum call rates during the buzz. To investigate these hypotheses, we first measured sound production during aerial attack sequences in free-flying Daubenton's bats (Mvotis daubentonii, Vespertilionidae) using a 12-microphone array (Fig. 1C) (9) and determined when the start and the end of each prey echo (Fig. 2, A to C) would impinge upon the bat's ears relative to both the source call and the next call emitted (10). Our data show that during a buzz, echoes from individual calls terminate before the start of the next call (Fig. 2C and fig. S1), suggesting no ambiguity in callecho matching. In fact, for buzz II calls, the repetition rate could theoretically exceed 400 calls/s without any such ambiguity (Fig. 2C and fig. S1), a rate twice as high as the 190 calls/s observed in our study (Fig. 1). Our results also demonstrate that, because call duration decreases during the buzz, there is no overlap between a call and its echo until the bat is less than 5 cm from its target (Fig. 2B), corroborating previous estimates for





**Fig. 1.** Echolocation and flight kinematics during aerial attack sequences in *Myotis daubentonii*. (**A**) Spectrogram and oscillogram of emitted echolocation calls as recorded on the center microphone of the array. (**B**) Instantaneous call repetition rates from sequence in (A). The start of buzz I is defined as call repetition rate >100 calls/s (cyan circles, light gray zone). In buzz II, the repetition rate is >160 calls/s (red circles, medium gray zone); here, peak fundamental frequency of the calls drops from 45 to 25 kHz. (**C**) Four reconstructed flight paths from a single bat using a 12-microphone array (*10*). Red arrows indicate flight direction.