ANNUAL REVIEWS

Annual Review of Neuroscience

Navigating Through Time: A Spatial Navigation Perspective on How the Brain May Encode Time

John B. Issa,^{1,*} Gilad Tocker,^{1,*} Michael E. Hasselmo,² James G. Heys,³ and Daniel A. Dombeck¹

¹Department of Neurobiology, Northwestern University, Evanston, Illinois 60208, USA; email: d-dombeck@northwestern.edu

²Center for Systems Neuroscience, Boston University, Boston, Massachusetts 02215, USA ³Department of Neurobiology and Anatomy, University of Utah, Salt Lake City, Utah 84112, USA

Annu. Rev. Neurosci. 2020. 43:73-93

First published as a Review in Advance on January 21, 2020

The Annual Review of Neuroscience is online at neuro.annualreviews.org

https://doi.org/10.1146/annurev-neuro-101419-011117

Copyright © 2020 by Annual Reviews. All rights reserved

*These authors contributed equally to this article



www.annualreviews.org

- Download figures
- Navigate cited references
- Keyword search
- Explore related articles
- Share via email or social media

Keywords

timing, animal behavior, neural circuits, interval timing, spatial navigation, episodic memory

Abstract

Interval timing, which operates on timescales of seconds to minutes, is distributed across multiple brain regions and may use distinct circuit mechanisms as compared to millisecond timing and circadian rhythms. However, its study has proven difficult, as timing on this scale is deeply entangled with other behaviors. Several circuit and cellular mechanisms could generate sequential or ramping activity patterns that carry timing information. Here we propose that a productive approach is to draw parallels between interval timing and spatial navigation, where direct analogies can be made between the variables of interest and the mathematical operations necessitated. Along with designing experiments that isolate or disambiguate timing behavior from other variables, new techniques will facilitate studies that directly address the neural mechanisms that are responsible for interval timing.

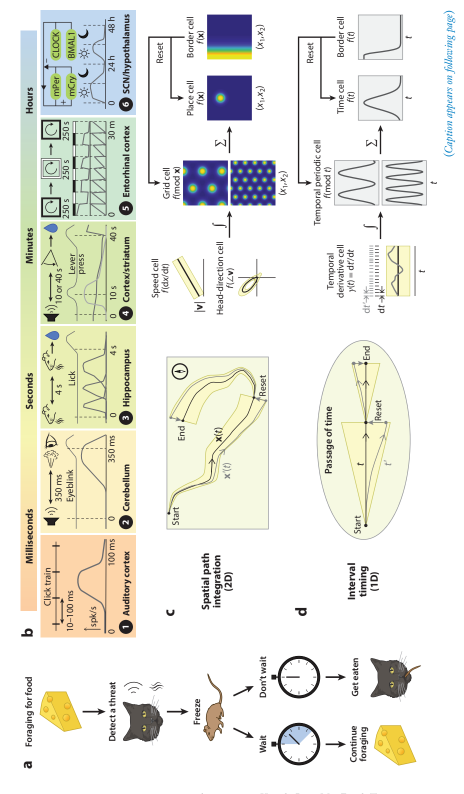
Contents

INTRODUCTION	74
TIMESCALES OF NEURAL TIMING CIRCUITS AND BEHAVIORS	76
DEFINITION OF VARIABLES IN INTERVAL TIMING BY ANALOGY	
TO SPATIAL NAVIGATION	77
Why Compare Seconds–Minutes Timing to Spatial Navigation?	77
Defining the Variables and Concepts	78
Comparison to Cells of the Spatial Navigation System	79
COMPUTATIONAL MODELS OF TIME REPRESENTATION	80
Models for the Sequential Firing of Cells	80
Models for Ramping Cells	84
PROPOSED EXPERIMENTS	85
Behavioral Design to Isolate Timing and Perturbations to Test Causality	85
External Cues	86
Timescales	86
Circuit Structures	86
CONCLUSIONS	87

INTRODUCTION

Perhaps no concept is simultaneously as permeative and as elusive as time. Time pervades our behaviors and is ingrained in our language and interactions, yet it is unclear if time is a real physical quantity (Rovelli 2018). No sensory organ is dedicated to the detection of time, indicating that our percept and concept of time are internally generated, maintained, and understood. As humans, our devotion to clocks, schedules, and ETAs has clouded our ability to describe what is a natural timing behavior in the animal kingdom, and thus we may overestimate the importance of timing.

But consider the alternative, that timing is so innate and natural that we take it for granted. Imagine a foraging animal who detects a nearby predator (Figure 1a). To avoid capture, she freezes and stops foraging until she thinks the danger has passed (Dill & Fraser 1997, Koivula et al. 1995). An optimal strategy would be to have an internal model of how long the predator typically stays in a given area (Cooper & Frederick 2007, Martín & López 1999) and to wait at least that long before venturing onward. Using an internal measure of time, she compares the current duration and the estimated typical duration. Likewise, timing is important for the predator (Hugie 2003), as she can maximize her chances of catching dinner by waiting for the prey to emerge in an open field and then estimating how long it will take to intercept her expected trajectory. But this example illustrates a pitfall in studying time. Rather than measure how long it takes for the prey to reach safety, the predator can optimize her pursuit based upon relative distance (Jennions et al. 2003). This entanglement of time with variables such as distance and velocity often confounds the study of timing. While one approach is to dedicate separate circuits to encoding pure timing and pure spatial information, an alternative implementation is for time to be embedded in an all-inclusive spatial-temporal framework, where a more general process generates behaviorally relevant sequences (Buzsáki & Llinás 2017). The hippocampus has been proposed as the region where these variables are integrated (Eichenbaum 2017; Hasselmo 2012; Hasselmo et al. 2010; Meck et al. 1984, 2013).



www.annualreviews.org • How the Brain May Encode Time 75

Figure 1 (Figure appears on preceding page)

Scales of neural timing and analogy to spatial navigation. (a) Diagram of an animal foraging for food when a predator is detected. The animal can choose to wait for a given amount of time (left) or risk continuing without waiting (right). (b) Diversity of timescales, brain regions, and neural circuits involved in the encoding of time from milliseconds to hours. (1) Neurons in auditory cortex tuned to millisecond interclick intervals (Sadagopan & Wang 2009). (
) In eyeblink conditioning, a conditioned stimulus (tone) precedes an unconditioned stimulus (air puff) by a fixed interval, and animals learn to blink and cerebellar neurons respond just before the air puff (Berthier & Moore 1986, Kotani et al. 2003). () Two odors are presented with a delay, and the animal licks to receive a reward only when the odors are identical. Pyramidal cells of the hippocampus fire sequentially during the delay period (MacDonald et al. 2013). (**④**) Animals press a lever after a delay period to receive a reward. Neurons in the cortex and striatum encode the delay period with a variety of patterns, including tuning to various times or ramping activity (Matell et al. 2003). () Neurons in the lateral entorhinal cortex encode the cumulative duration of traversals in alternating black and white rooms over the course of tens of minutes (Tsao et al. 2018). (•) A transcription-translation feedback loop in neurons of the suprachiasmatic nucleus (SCN) results in 24-h rhythms (Takahashi 2017, Welsh et al. 1995). Data for illustrations in panel b from the studies cited above. (c, left) Spatial path integration in two dimensions (2D). At start, the position is known exactly. As the animal moves toward the target end or goal, the actual position $\mathbf{x}(t)$ is shown in black and the internal estimate of position $\mathbf{x}'(t)$ in gray. Integration of a velocity signal (yellow region, inferred from self-motion cues) updates position but accumulates error until a landmark or border that allows for error correction is reached (reset or end, gray arrow). (c, right) Cells in spatial navigation from left to right: Speed cells and head-direction cells encode speed and orientation of the animal, with uncertainty indicated in yellow; grid cells encode periodic representations of position; place cells encode current position; and border cells indicate proximity to borders or objects. Grid cells integrate incoming speed and orientation signals and are summed to produce single-peaked place cells. Border cells can reset or correct these position signals. Colors indicate firing rate (high rate, yellow; medium, green; low, blue). (d, left) Interval timing through time integration in one dimension (1D). Elapsed time is measured after a start signal. External time t is shown in black, while the internal estimate of time t' in gray indicates the perception that more time (gray line above black line) or less time (gray line below black line) has elapsed relative to external time. The estimate of time is informed by a moment-by-moment temporal derivative that represents the animal's ongoing passage of time signal and runs until a reset (correction cue) or stop is encountered. (d, right) Cells of interval timing include temporal derivative cells, which encode the internal representation of the passage of time, represented here as a pacemaker (*left*) whose interspike interval dt' approximates discrete time steps dt. The continuous-time version of this process is represented by a temporal derivative cell whose deviation from true time (black line) is shown by the gray trace. Output of a temporal derivative cell is integrated by hypothesized periodically firing temporal cells. These are summed to produce well-tuned time cells. Border cells fire at start or stop positions and reset or correct encoding of time in temporal periodic cells and time cells.

While timing is fundamental, it is unclear how explicit or conscious its representation is, whether neural circuits precisely encode time, or how many neural clocks reside in the brain (Howard et al. 2014, Karmarkar & Buonomano 2007, Lusk et al. 2016, Meck 1996). In this review, to frame the discussion in a manner amenable to achieving mechanistic insight, we focus on pure timing in the brain that operates on the range of seconds to minutes. First, we discuss the brain regions and behaviors associated with the encoding of time across a broad range of timescales. Next, we draw an analogy to spatial navigation, as the seconds–minutes scale can be considered as navigating an abstract time dimension as opposed to space, and thus similar mechanisms and circuitry may underlie both. Within this framework, we then explore cellular and circuit mechanisms capable of computing seconds–minutes timing. Finally, we propose experiments that could help uncover the neural circuitry responsible for these computations.

TIMESCALES OF NEURAL TIMING CIRCUITS AND BEHAVIORS

Organisms can accurately track temporal information across many timescales, from milliseconds to days. The circuits must be diverse enough to cover these timescales and also to match a wide range of desired behaviors (**Figure 1***b*). Previous research has sought to establish whether the brain uses a single clock or many independent clocks in order to encode time across scales and modalities and whether different clocks might time using distinct or common neural mechanisms (Ivry & Schlerf 2008, Karmarkar & Buonomano 2007, Lusk et al. 2016, Merchant et al. 2013, Paton & Buonomano 2018). As evidence that the brain appears to use multiple independent clocks

across different time scales, duration judgments at one interval are not disrupted by a distractor at a different timescale (Spencer et al. 2009).

Specialized circuits can operate with millisecond precision, especially for coordinating motor movements (Sober et al. 2018). The cerebellum has been implicated in a number of timing behaviors on this timescale (Ivry 1997, Mauk & Buonomano 2004), such as for eyeblink conditioning (**Figure 1***b*). The auditory system is also specialized to operate with high temporal precision, with neurons detecting submillisecond interaural time differences (Carr & Konishi 1990) or others tuned to intervals of ~10 ms (Sadagopan & Wang 2009) (**Figure 1***b*). On the other end of the timescale are the circuits involved in 24-h circadian rhythms, particularly neurons of the suprachiasmatic nucleus (Takahashi 2017) (**Figure 1***b*).

Between these two extremes, many behaviors require tracking time at the intermediate scale of seconds to minutes (**Figure 1***b*). While our conscious perception of these timescales is well ingrained in our lexicon (e.g., "just a sec," "in a minute"), the neural mechanisms responsible are not well understood. Nonetheless, the neocortex, basal ganglia, and cerebellum have been implicated in interval timing (Akhlaghpour et al. 2016, Bakhurin et al. 2017, Howard et al. 2014, Jin et al. 2009, Leon & Shadlen 2003, Lusk et al. 2016, Matell et al. 2003, Mello et al. 2015, Mita et al. 2009, Soares et al. 2016, Tiganj et al. 2018).

The hippocampus is also involved in interval timing, as temporal information is crucial to memory and the hippocampus is heavily involved in episodic memory formation (Squire 1992, Tulving 2002). Following upon early efforts (Meck et al. 1984), numerous recent studies have expanded upon the role of the hippocampal formation in the encoding of time. For example, neurons in CA1 fire sequentially during a delay period between the presentation of two odors when animals are immobile (MacDonald et al. 2013) (**Figure 1b**). Neurons in the rodent hippocampus and medial entorhinal cortex (MEC) represent elapsed time during treadmill running (Kraus et al. 2013, 2015; Mau et al. 2018; Pastalkova et al. 2008). In freely foraging rats, neurons in the lateral entorhinal cortex (LEC) ramp their activity over the course of seconds to minutes such that time intervals can be decoded from their population activity (Tsao et al. 2018) (**Figure 1b**). And in one final example, MEC neurons fire sequentially and tile the wait period in mice trained to stop at a door in virtual reality for 6 s (Heys & Dombeck 2018).

Given the vast difference in timing mechanisms at the ends of the spectrum (milliseconds for synaptic mechanisms, 24 h for genetic mechanisms), discovery of the mechanisms responsible for the seconds–minutes scale should prove informative on its own and may also shed light on general mechanisms of neural circuits (Howard et al. 2014, Karmarkar & Buonomano 2007, Tiganj et al. 2015). Different mechanistic frameworks will be useful to guide interpretation of neural data and to design future experiments. In this context, we propose that a way forward for understanding this intermediate seconds–minutes timescale is to draw an analogy to spatial navigation.

DEFINITION OF VARIABLES IN INTERVAL TIMING BY ANALOGY TO SPATIAL NAVIGATION

Why Compare Seconds-Minutes Timing to Spatial Navigation?

The brain regions and neural mechanisms responsible for seconds-minutes time encoding may parallel those of spatial navigation. Many commonalities are shared between the two systems and in the encoded parameters. The timescale for activity patterns in spatially active cells in the rodent hippocampus, for example, spans periods of subsecond to tens of seconds (Bittner et al. 2017, Kjelstrup et al. 2008, Mau et al. 2018, Stensola et al. 2012). The resolution of time and space encoding appears to be similar; for example, the statistics of neural activity in hippocampal neurons during track navigation and duration-specific treadmill running are highly similar as quantified by field width, peak firing rates, and measures of population activity (Kraus et al. 2013, Pastalkova et al. 2008). Lesion studies also show parallels in the types of tasks that are dependent on the hippocampus. In spatial tasks, navigation based on cues is not hippocampus dependent, but navigation reliant on an allocentric map is hippocampus dependent (Maguire et al. 2006, Morris et al. 1982, O'Keefe & Nadel 1978). Similarly, for timing tasks, delay classical conditioning is not hippocampus dependent, but trace conditioning is hippocampus dependent (Bangasser et al. 2006, McEchron et al. 2003). These parallels indicate that the hippocampus is differentially necessary for aspects of spatial and temporal tasks.

Moreover, many brain regions with spatial representations also engage in seconds-minutes timing, and thus similar computational strategies and circuit motifs may be responsible in both cases. For example, alongside place cells in dorsal CA1 in the hippocampus, studies have found cells that encode temporal information or combined spatial and temporal information (Kraus et al. 2013, MacDonald et al. 2011). In the MEC, spatially periodic grid cells coexist with time-encoding cells, while some grid cells can also encode temporal information (Heys & Dombeck 2018, Kraus et al. 2015). Thus, the study of time encoding may be able to follow the successful strategies used in the navigation field, where the tight interplay between theoretical models and experiments has yielded a rich understanding of the circuits involved in spatial navigation. Just as the brain uses idiothetic cues for path integration and sensory landmark cues for error correction through updates of current position during spatial navigation (Bush & Burgess 2014, Jayakumar et al. 2019, Savelli & Knierim 2019, Zhang et al. 2014), we reason that an internal sense of the passage of time may analogously be integrated to compute elapsed time while external sensory cues correct for errors (Figure 1c,d). Our main contention is that the neural computations underlying both systems share the same canonical form with two main components: integration and updates to correct for accumulated errors.

Defining the Variables and Concepts

Each key variable of spatial navigation has a complementary variable in seconds–minutes timing (**Figure 1***c*,*d*): position in space $\mathbf{x}(t)$ [a two-dimensional (2D) vector of spatial coordinates] to position in a time interval *t*; spatial velocity $\mathbf{v} = d\mathbf{x}/dt$ to the passage of time $\gamma = dt'/dt$; and physical boundaries or landmarks in space to sensory cues indicating the start, end, or intermediate position in a time interval. We distinguish actual position $\mathbf{x}(t)$ and the internal representation of position $\mathbf{x}'(t)$. Differences between these quantities indicate an error in the internal representation. Likewise, the internal representation of time *t'* may deviate from *t* (e.g., $t' \neq t$).

In both cases, two key processes are necessitated: a moment-by-moment integration of a velocity signal (path integration or time integration) and the update of the representation based on sensory cues. The first process (time integration) is useful in timing for the same reason path integration is used in spatial navigation: Sensory cues may provide accurate locations in time or space, but they can be infrequent. To attain a continuous sense of time, an integration process is needed to bridge the gap. However, integration suffers from an accumulation of errors, and thus sensory cues can reset or correct the internal representations of time and space positions (Campbell et al. 2018, Jayakumar et al. 2019).

In seconds-minutes timing, the organism begins with a start point. Next, the duration of the timing period must be tracked in the absence of external cues using some internally generated representation of time and using time integration. This internal time t' typically approximates an absolute (external) time t, and its passage is denoted by what we term the temporal derivative, defined as $\gamma = dt'/dt$, where the choice of γ is inspired by the use of the Lorentz factor in physics to relate the change in time between two reference frames (Lorentz 1903) and can be thought

of as the perceived passage of time for an observer relative to absolute time (**Figure 1***d*). At any point, sensory cues carrying information about external (absolute) time can adjust or reset the internal representation. These resets can be thought of as a new start point when the internal time *t'* is synchronized to absolute time *t*. Similarly, in spatial navigation, after a start point, the internal spatial representation of position $\mathbf{x}'(t)$ is continuously updated by an estimate of distance traveled based on idiothetic cues used to estimate heading direction and speed, a process termed path integration (McNaughton et al. 2006) or dead reckoning (**Figure 1***c*). At any point, sensory landmark cues can provide information to reset the internal representation of spatial position $[\mathbf{x}'(t)]$ is synchronized to actual position $\mathbf{x}(t)$].

A pure integrator of an uncorrelated random signal, such as we propose for path integration or time integration in the absence of external cues, accumulates error proportional to the square root of the distance or time interval. This relation represents a lower bound. Observed error rates in interval timing are typically described as linear in accordance with Weber's law and scalar expectancy timing (Gibbon 1977), although error rates in time intervals inferred from population neural activity in at least one study appear to follow a square root relationship (Itskov et al. 2011). In practice, error rates can grow linearly with the time interval through simple changes in the statistics of the temporal derivative signal, such as correlated errors across time or a bias that shifts the signal up or down as exemplified by neuropharmacological manipulations that increase or decrease one's perception of the passage of time (Meck 1996, Soares et al. 2016). External cues play a large role in error correction. In spatial navigation, even in the dark, the animal can use odors and somatosensory cues to guide navigation (Save et al. 2000). For example, interacting with the border of an open field can reset some of the errors accumulated by grid cells (Hardcastle et al. 2015). Thus, it is important to compare spatial path integration and seconds–minutes timing across conditions under which the presence and reliability of external cues or resets are similar.

Comparison to Cells of the Spatial Navigation System

Given these parallels, it is reasonable to hypothesize that similar cell types and encoding strategies may be encountered across the two systems and be found in similar brain regions (Figure 1c,d). For example, many hippocampal place cells have single-peaked, well-tuned, and sparse firing fields (O'Keefe & Dostrovsky 1971) that are highly context dependent and flexible (Anderson & Jeffery 2003). Time cells in the hippocampus with similarly single-peaked, well-tuned, and sparse fields have already been described (see below). Grid cells in the MEC are coherent over different contexts, and their multi-peaked firing fields appear unstructured over small environments (Fyhn et al. 2004), but the hexagonal structure is more apparent over larger environments (Hafting et al. 2005). These properties are consistent with specialized circuitry that yields a metric for space (Moser & Moser 2008). We expect that time-encoding cells in the MEC similarly form a metric for timing, with periodically active multi-peaked time cells emerging for longer time intervals (Kraus et al. 2015). Place cells can also show multi-peaked firing in larger environments (Fenton et al. 2008), so multiple fields may be expected for hippocampal time cells over longer times. We also expect start and stop cells that are similar to border cells in the spatial navigation system (Solstad et al. 2008). Lastly, analogous to head-direction cells of the postsubiculum (Taube et al. 1990) and speed cells within the MEC (Kropff et al. 2015), the input to the timing system should be a temporal derivative cell whose moment-by-moment firing rate represents the temporal derivative γ and can account for changes in the perception of the speed of time, a phenomenon dependent on emotional state (Schirmer 2011) or disease state (Densen 1977) and modifiable with certain drugs (Meck 1996) or optogenetic manipulations (Soares et al. 2016). There are, however, some notable nuances to this analogy. Since time is one-dimensional (1D) and unidirectional, a direction-of-time cell is not needed, so there is no analogy to head-direction cells. Also, certain network architectures such as attractor networks could move neural activity at a constant speed using asymmetric connectivity and thus avoid the need for a temporal derivative input. Lastly, the spatial encoding system stops updating when the animal stops, as speed inputs drop to zero, but in time encoding the temporal derivative input is presumably constant and will persist. An explicit stop or reset signal is required to halt time encoding.

Experimental evidence for many of these cell types already exists. Timing cells have been observed in the hippocampus over the 10–20-s timescale (Kraus et al. 2013, MacDonald et al. 2011) and in the MEC over the 6–8-s timescale (Heys & Dombeck 2018, Kraus et al. 2015). Temporal periodic cells with multiple activity peaks might be expected in the MEC over longer timescales, which have been observed in grid cells during treadmill running (Kraus et al. 2015) and rest active cells during periods of immobility (J.G. Heys, J.B. Issa & D.A. Dombeck, unpublished manuscript). Cells with ramping activity over timescales ranging from seconds to hours have been observed in the LEC (Tahvildari et al. 2007, Tsao et al. 2018), the hippocampus (Hampson et al. 1993) and other cortical structures (Quintana & Fuster 1999). An interoceptive signal that provides the temporal derivative signal would imply the presence of parallel mechanisms between time and space; for example, time cells in the MEC might be expected to perform integration of a temporal derivative signal using a similar circuit to grid cells integrating speed and head-direction signals. This signal could arise in the LEC or other regions such as the insula (Craig 2009, Wittmann 2009).

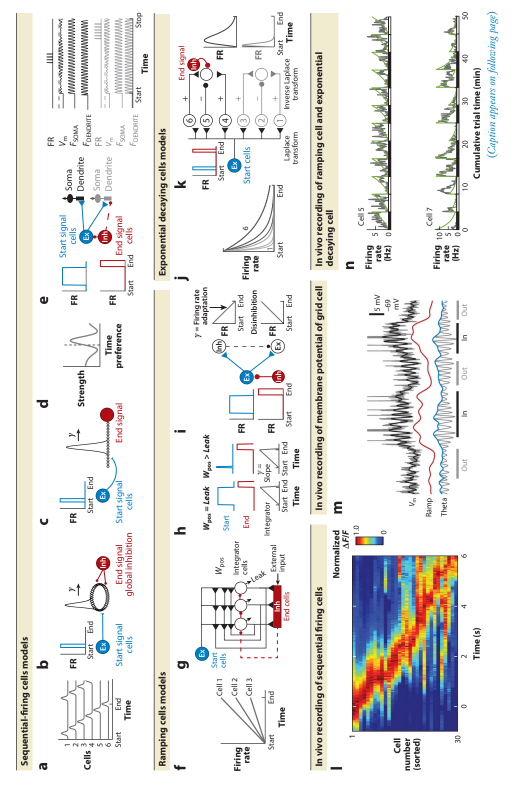
COMPUTATIONAL MODELS OF TIME REPRESENTATION

Various patterns of activity have been described by which neurons can encode time. These patterns of activity include cells that fire sequentially during a time interval, especially in the hippocampus and the MEC (Heys & Dombeck 2018; Kraus et al. 2013, 2015; MacDonald et al. 2011; Pastalkova et al. 2008; Salz et al. 2016), and cells that monotonically increase or decrease their firing rate during a time interval (Hampson et al. 1993, Tahvildari et al. 2007, Tsao et al. 2018). Why does the brain encode time in such different ways? Do the different activity patterns implement the same algorithm? Are the algorithms biologically plausible? Computational models that explain mechanisms underlying various patterns of time encoding could help provide answers.

Here, we focus on models of sequentially active cells and of ramping cells (**Figure 2**). A ramping code allows for any given individual cell to carry information about elapsed time throughout the entire interval, and therefore fewer cells are needed. On the other hand, each moment in the ramping code is highly correlated to other moments, making it difficult to associate one moment with a memory of an item or an object. In contrast, with sequential activity, each cell is active at one specific moment, allowing association of this specific moment with an item and/or an object by associative plasticity (Rolls & Mills 2019, Hasselmo 2012). Furthermore, the code is energetically efficient, as only one cell needs to be active at any given time point (Barlow 1972), but such a sparse code demands a larger population of cells for long intervals. Thus, each code may be better suited for certain demands of the neural computation. Conversely, the biology of a given network or neuron may determine which coding strategy it can support.

Models for the Sequential Firing of Cells

In the sequential firing of timing cells, each cell fires at a specific moment in time (time field), and together the population tiles the entire interval (**Figure 2***a*,*l*). What network or cellular mechanisms could generate such a code? Several theoretical models of spatial path integration have suggested means through which grid and place fields could be formed. In many of these



www.annualreviews.org • How the Brain May Encode Time 81

Figure 2 (Figure appears on preceding page)

Models for time cells. (a) Sequential firing of time cells. Each row represents the firing rate (FR) of one cell. Each cell is active at a specific moment within the time interval, and together the sequence of firing cells tiles the entire interval. (b) Ring attractor, continuous attractor network (CAN) model. Start population of cells fires a burst at the beginning of the time interval (left, blue) and input to a ring attractor (middle, black). Global inhibition provides the end signal (right, red). (c) Line attractor CAN model. Start population fires a burst at the beginning of the time interval (left, blue) and inputs to a line attractor (middle, black). The end signal is simply cells at the end of the sequence (right, red). (d) Mean connectivity pattern of cells in the CAN model. The y-axis represents the strength of outgoing weights, and the x-axis represents the time preference of cells that receive the inputs (aligned to the time preference of the output cell at 0, dashed line). (e) Sequence of two time cells created by the oscillatory interference model. The start cell fires during the entire interval (top left, blue). The end signal cell inhibits start cell or integrator cells at the end of the interval (bottom left). On the right is a scheme of two cells (one in *black* and one in gray). Each cell is excited by a start cell. The dendritic frequency oscillation of each cell ($F_{dendrite}$) is different than its somatic frequency oscillation (F_{soma}, which differs in the two cells) and results in an interference pattern (Vm, membrane potential). The membrane potential crosses the firing threshold every beat oscillation. (f) Scheme of three ramping cells. Each cell has a different slope. (g) A network model that uses a positive feedback mechanism to integrate inputs over time and generate ramping cells. All integrator cells receive feedforward input from start cells (blue). Output integrator cells (black) are connected by recurrent excitation and also send excitation to end neurons. End cells (red) inhibit integrator cells once the rate of inputs they receive crosses a threshold or by external input (Seung et al. 2000). (b) Activity of each cell type (start cells, blue; end cells, red; integrator cells, *black*) when recurrent excitation is equal to the leak current (*left*, $W_{\text{pos}} = Leak$) or when recurrent excitation is slightly higher than the leak current (right, W_{pos} > Leak). (i) A model network generating ramping activity by an adaptation process. A population of excitatory persistent firing cells (which fire at a constant rate during the time interval) excite a population of inhibitory (Inh) and excitatory (Ex) neurons. Due to adaptation of the firing rate in the Inh neurons, their firing rate decreases. This disinhibits the Ex population, which in turn increases its firing rate and creates a ramping firing pattern. (i) Exponentially decaying firing rates of six cells with different time constants in response to a start burst, analogous to the Laplace transform of a delta function (a start burst). (k) Start cells (blue) give a pulse input to a cell population with exponentially decaying firing rates, each with different time constants (the cells are arranged according to their time constants in increasing order from bottom to top). The exponentially decaying cell population projects to the output layer cells (right) through a center-off, surround-on connectivity pattern. Thus, each cell in the output layer receives a weighted sum of several exponential decaying cells, implementing an inverse Laplace transform, which in turn produces time cells in the output layer (Howard et al. 2014, Liu et al. 2019). (1) In vivo imaging of sequential activation of neuronal population in the medial entorhinal cortex (MEC) during a door-stop timing task; a similar sequence is schematized in panel a. Each row represents an individual neuron's mean $\Delta F/F$ value versus wait time during the 6-s door-stop wait interval. Mean $\Delta F/F$ is normalized to peak. Panel *l* adapted with permission from Heys & Dombeck (2018). (m) In vivo intracellular recording of membrane potential oscillations from a grid cell in the MEC during navigation. The membrane potential (gray) is decomposed into ramp (red) and theta components (oscillations, gray; envelope, blue). Time cells also reside in the MEC and could have similar oscillations in their membrane potential, similar to those depicted in panel e. Panel m adapted with permission from Domnisoru et al. (2013). (n, top) An example of a ramping cell recorded from the lateral entorhinal cortex (LEC) in vivo during behavior, similar to ramps depicted in panel f. In gray is the actual firing rate and in green is the predicted firing rate by a general linear model. (Bottom) Example of an exponential decaying cell recorded from the LEC in vivo during behavior, similar to exponential decay depicted in panel *j*. Panel *n* adapted with permission from Tsao et al. (2018).

models, neurons path-integrate speed and direction of movement to form fields (Burak & Fiete 2009, Burgess et al. 2007, Fuhs & Touretzky 2006, McNaughton et al. 2006, O'Keefe & Recce 1993, Samsonovich & McNaughton 1997, Zhang 1996). These models can be modified for time integration, resulting in the formation of time fields instead of place fields and, at the population level, in the formation of sequences of firing fields.

Continuous attractor network models. In continuous attractor network (CAN) models (Burak & Fiete 2009, Couey et al. 2013, Fuhs & Touretzky 2006, Guanella et al. 2007, McNaughton et al. 2006, Samsonovich & McNaughton 1997), a population of neurons is arranged on a 2D sheet and connected using so-called Mexican hat connectivity (**Figure 2***d*). Excitation of this network generates a regular pattern of activity bumps. Inputs reflecting the animal's velocity cause this pattern to move on the sheet, creating a place or grid-firing pattern in single neurons. For time encoding, speed and direction inputs are not needed, and only a 1D network and excitatory input supporting 1D bump generation are needed. Cells at opposite ends of the 1D network can be connected to form a periodic boundary condition, resulting in a ring attractor (**Figure 2***b*), as in models of head direction (Blair et al. 2014, Sharp et al. 2001, Skaggs et al. 1995, Zhang 1996).

Alternatively, the cells need not be connected at the ends, resulting in a line attractor (**Figure 2***c*). One mechanism to generate movement of the bump at a constant speed is to use asymmetric connectivity (e.g., Mexican hat offset in one direction) (**Figure 2***d*) or threshold adaptation that creates effective connection asymmetry (Itskov et al. 2011). The temporal derivative (γ) in such a model is defined by the speed of the sequential firing of cells (**Figure 2***b*,*c*) and is controlled by the level of asymmetry (**Figure 2***d*). Ideally, these connections, and therefore the bump speed, would remain constant to represent external time consistently; however, this speed could be modified through plasticity or via external inputs. The speed of the sequential firing could also be used to encode different time intervals: Longer intervals could be encoded by a slower-moving bump and shorter intervals by a faster-moving bump. Alternately, the bump could move at a constant speed for all time intervals, and the number of cells participating in the sequence could change.

How and where could an activity bump be initiated? A start signal could be generated by a population of neurons that are driven by an external cue, which in turn excite the neurons at a specific location on the 1D network (**Figure** 2b,c), from which point time integration begins. Different contexts could be encoded by starting the bump at different locations on the 1D network (**Figure** 2b,c). For time encoding, as opposed to space encoding, since the temporal derivative signal is internally generated, a distinct stop signal would likely need to act directly on the integrator. For a ring attractor, global (rather than focal) inhibition would be needed to suppress the bump(s) and prevent more from emerging (**Figure** 2b). Ambiguity in the amount of time encoded will arise if the duration of the interval is longer than the duration of one period around a ring attractor. This ambiguity could be solved using multiple modules, each with different periods. Interestingly, a modular grid code was found to be more efficient than a place cell code for encoding space (Burak et al. 2006, Mathis et al. 2012, Mosheiff et al. 2017, Sreenivasan & Fiete 2011). It remains to be seen whether time encoding is governed by similar rules.

Other similar models can also generate sequential activity for time encoding. For example, chaotic networks can generate the sequential firing of cells similar to those found in the posterior parietal cortex (Rajan et al. 2016). Time-dependent changes in the state of a complex neural network can also provide a reliable readout of elapsed time (Karmarkar & Buonomano 2007, Laje & Buonomano 2013, Sussillo & Abbott 2009).

Oscillatory interference models. Oscillatory interference models (Burgess et al. 2007, O'Keefe & Recce 1993) could also generate sequential firing of time-selective cells. Each cell receives two oscillatory input signals, either through oscillating inputs or by different cell compartments possessing different resonant frequencies that oscillate when depolarized (Giocomo et al. 2007) (**Figure 2***e*,*m*). These signals interfere with each other, creating beat oscillations in the membrane potential. By modulating the frequency of one oscillator proportional to running speed, these models can generate 1D place fields (O'Keefe & Recce 1993) and, with proper modifications, also 2D grid-firing fields (Burgess et al. 2007). A simple version of this model could generate time cells (Hasselmo 2008). For a constant temporal derivative, the two oscillatory signal frequencies should be fixed, resulting in temporal beat oscillations of the membrane potential and firing at every beat, creating time fields (**Figure 2***e*). This model also resembles the use of multiple oscillators for encoding the order of words in serial recall (Brown et al. 2000) and temporal intervals in the timing of behavioral responses (Matell & Meck 2004, Miall 1989).

To generate a sequence of firing cells, various mechanisms could be used. Multiple cells could fire at the same beat frequency but with different phases. This is similar to grid cell networks, where grid cells belonging to the same module share the same spatial frequencies but differ in their phases (Barry et al. 2007, Stensola et al. 2012). This could be implemented by presynaptic cells triggering oscillations in time cells with different delays. The problem, however, is that these time delays must

be on the order of the sequences they are meant to generate. Thus, the task of generating a time sequence is passed to the input layer, making such a scheme implausible. Alternatively, sequence generation could be implemented by different time cells oscillating at a range of different baseline frequencies (Hasselmo 2014) so that a single input frequency would result in different phases of firing across the population. In this model, if time cells fire periodically in time, the population firing sequence would not repeat and instead the cell firing order would change at each repetition. This contrasts with CAN models, which predict the preservation of the order of cells' firing in each repetition. To circumvent these issues, combining CAN and oscillatory interference models has been used to model robust grid cell firing (Bush & Burgess 2014) and could also be useful for time encoding.

The start signal could be provided by cells that fire following certain internal or external cues and persist during the time interval (**Figure 2***e*). Such persistent firing has been observed in the entorhinal cortex (Tahvildari et al. 2007, Yoshida et al. 2008). Like CAN models, the temporal derivative (γ) in oscillatory interference models is defined by the speed of sequential firing. Encoding different time intervals could be implemented either by changing the speed of sequential firing (changing the frequency of one of the oscillators) or by using a constant speed of sequential firing but recruiting a different number of cells according to the interval. Time integration could be terminated by inhibiting the start population or the integrator (time cells) directly (**Figure 2***e*).

Models for Ramping Cells

Recordings in several brain areas have revealed cells that monotonically increase or decrease in firing rate during task time intervals (Hampson et al. 1993, Quintana & Fuster 1999, Tsao et al. 2018) (Figure 2f,n). This change in firing rate can appear linear or exponential, and therefore elapsed time is encoded by an instantaneous firing rate (Figure 2f). Though there are no clear analogies to ramping cells in spatial navigation, we consider ramping models since these cells have been widely described in the timing literature.

Temporal integrators. Cellular (Durstewitz 2003) and network (Shankar & Howard 2012) models have been proposed to explain ramping cell activity. In some models (Balcı & Simen 2016, Lim & Goldman 2013, Simen et al. 2011), the cells act as temporal integrators of their inputs. Leak currents counteract the input-driven depolarization. To counterbalance the leak current and effectively integrate and store inputs over time, cells require either a precise positive feedback mechanism (Seung et al. 2000, Simen et al. 2011) or a negative derivative feedback mechanism (Seung et al. 2000, Simen et al. 2011) or a negative derivative feedback mechanism (Lim & Goldman 2013). In both cases, input depolarization is preserved over time through stable firing (**Figure 2g,b**). To generate ramping activity, an excitatory step input, which defines the temporal derivative (γ), could serve as the start signal (**Figure 2b**). Timing cells integrate this input by using positive feedback to generate a linear increase in firing rate (**Figure 2b**). The temporal derivative (γ) is represented by the slope of the ramp (**Figure 2b**). To terminate integration, inhibitory cells that are activated once the ramping cells reach a certain threshold could send negative feedback (**Figure 2g,b**), or the stop or reset signal could be provided by internal or external cues to drive inhibition of the timing cells (Durstewitz 2003, Grossberg & Merrill 1992, Grossberg & Schmajuk 1991) (**Figure 2g**).

Alternatively, the positive feedback could be larger than the leak currents (Durstewitz 2003), making the cells no longer perfect integrators. A start signal to such cells simply requires an excitatory burst to begin the positive feedback loop and time integration (**Figure 2***b*), resulting in a gradual increase in firing rate during the interval (**Figure 2***b*). The temporal derivative (γ) is defined by the increasing slope of the ramp, which is determined by the excess of positive feedback

over the leak currents. The end mechanism could be the same as in the preceding section. Once the cells stop firing, inhibition at rest would be required to prevent spurious integration.

Adaptation models. Another model (Reutimann et al. 2004) makes use of adaptation to generate ramping. This entails three populations of cells. The first is composed of cells that fire at a constant rate from the beginning to end of the interval. During the interval, these cells excite a second population of inhibitory neurons and a third population of excitatory neurons (**Figure** 2i). The excitatory cells are also inhibited by the inhibitory cells. Adaptation in the inhibitory cells leads to a gradual decrease in their firing rate during the interval. The decrease in firing rate leads to disinhibition of the excitatory cells, which then gradually increase their firing rate, creating a ramping pattern. Thus, the time interval is encoded by both increasing (ramping) and decreasing (ramping) firing rates. The first population of persistent activity cells defines the start signal, and the temporal derivative (γ) is defined by the firing rate adaptation (firing rate slope).

Using exponential firing rate or ramping cells to generate sequential firing. Another class of models uses the exponential decay of firing rate, rather than linear changes in rate, with a range of decay rates across cells (Howard et al. 2014, Liu et al. 2019, Tiganj et al. 2015). The first population of cells performs a Laplace transform on a start input burst to generate different timescales of exponentially decaying firing rates across the population. A second layer of cells then performs an inverse Laplace transform to generate timing fields in sequentially activated cells. This model allows for coding time over a wide range of timescales (**Figure** *2j,k,n*) and accounts for the decrease in numerical density and increase in variance often observed in time cell responses over longer intervals (Mau et al. 2018, Tiganj et al. 2018). Similarly, it is also possible to generate sequential firing by adjusting ramping cell input through a Hebbian learning rule together with mutual inhibition between cells to ensure sparse activity (Rolls & Mills 2019).

PROPOSED EXPERIMENTS

Behavioral Design to Isolate Timing and Perturbations to Test Causality

As time is entangled in most behaviors in one way or another, studying timing circuitry during immobility helps isolate time from other variables. Interestingly, in the hippocampal-entorhinal region, different circuits appear to be recruited depending on whether animals are mobile or immobile (Arriaga & Han 2017, Heys & Dombeck 2018, Kay et al. 2016). Recent studies in the rodent LEC found that neural activity patterns recorded during open field foraging could be used to decode time during the experience (Tsao et al. 2018). However, it is unclear if the neural representations indeed relate to the animal's perception or memory of time or whether the neural representation serves to encode other sensory stimuli or motor patterns that were systematically changing. Similarly, hippocampal sequences recorded during wheel running (Pastalkova et al. 2008) can carry temporal information; however, with animals running at a fairly constant pace, it is difficult to distinguish time from distance. Treadmill running with variations in speed was later used to distinguish these possibilities and demonstrate time encoding during locomotion in the MEC (Kraus et al. 2015), demonstrating the importance of experimental design when investigating time encoding.

Engaging animals in timing tasks in which temporally informative sensory and motor cues are kept to a minimum, and thus the passage of time is the dominant variable, could alleviate some of these issues (Bright et al. 2019, Heys & Dombeck 2018, Leon & Shadlen 2003, MacDonald et al. 2013). However, even for immobile animals in a static environment, internally generated

cues, such as heartbeat and breathing, could still provide temporal information. Furthermore, such tasks limit the study of many natural timing behaviors. Even using tasks in which time is the dominant variable, the question remains as to whether changing neural activity patterns over time relate to the animal's representation, perception, or memory of time. Here, neural perturbation experiments provide an excellent means to test for causality. For example, by inactivating a brain region or circuit thought to contain a time representation using optogenetic or chemogenetic methods (Armbruster et al. 2007, Boyden et al. 2005) and then testing for altered timing behavior or learning, it should be possible to determine the necessity of a region or circuit for the perception or learning of different time intervals (Soares et al. 2016). Optogenetics and chemogenetics may have advantages over lesions. For example, recent seemingly contradictory reports about whether sequential time cell activity in the MEC is necessary for downstream time cell activity in the hippocampus (Robinson et al. 2017, Sabariego et al. 2019) may be explained by compensatory mechanisms allowing for hippocampal activity pattern recovery after MEC lesions, while such recovery does not take place during transient optogenetic MEC inactivation.

External Cues

Similar to the representation of space with path integration, the brain's representation of time is predicted to drift with respect to external time (nonunitary temporal derivative) (**Figure 1***d*), but this drift can be corrected by using external cues. This process could be investigated by adding temporally informative cues to the delay period of a long timing task. For example, animals could first be trained on an instrumental task to wait after a cue (light flash) and then report when an amount of time had elapsed (say 20 s) by licking for reward. On probe trials, a metronome-like sound cue (i.e., a temporal landmark every few seconds) could be added. Behaviorally, the error in lick times would be expected to decrease if the external sensory cues are used to correct for integration errors. Imaging (Harvey et al. 2012, Heys & Dombeck 2018, Mau et al. 2018) or electrophysiology (Jun et al. 2017, Tiganj et al. 2018) methods could be used to identify differences in representations from large populations of time-encoding neurons. For example, in the case of sequential firing representations, time tuning curves would be expected to narrow near the sound cues (Mau et al. 2018, Tiganj et al. 2018), and in the case of ramping representations, slope (trial-trial) variability would be expected to be reduced near the cues (Howard et al. 2014, Liu et al. 2019).

Timescales

Models for sequential firing networks make several testable predictions about repeating firing fields over long intervals. For example, ring attractors are expected to produce repeating sequences when the encoding time exceeds the ring period, while line attractors would not generate repeating sequences and instead would recruit more neurons to the end of the sequence to encode longer intervals. For oscillatory interference models, different beat frequencies across the population are needed to generate sequences, and therefore, while individual cells would display repetitive firing fields over long intervals, population sequences would become incoherent over repeated cycles. Differentiating between these predictions could then be accomplished by engaging an animal in a timing task in which the timing period exceeded the duration of sequences that have been previously described (>20 s) (Cameron et al. 2019; Heys & Dombeck 2018; Kraus et al. 2013, 2015; MacDonald et al. 2013; Pastalkova et al. 2008; Salz et al. 2016).

Circuit Structures

Finally, next-generation circuit-cracking techniques should prove useful in distinguishing between the models and determining the exact mechanisms underlying time integration. Influence mapping, for example, uses single-cell optogenetic activation along with population Ca²⁺ imaging to determine the (direct or indirect) circuit connectivity between neurons in vivo (Chettih & Harvey 2019). This approach could be used to unveil the predicted Mexican hat connectivity between sequential firing cells in CAN models, the predicted connectivity between cells with exponentially decaying firing rates and sequential firing cells in the Laplace transform model (Liu et al. 2019), or the predicted inhibitory connectivity between negative and positive ramping cells in ramping cell models. Dendritic synaptic input mapping might also be useful to establish mechanisms of time encoding and integration (Marvin et al. 2018, Wilson et al. 2018). These methods involve using dendritic imaging of functional fluorescence indicators, such as GCaMP (Dana et al. 2019) or iGluSNFr (Marvin et al. 2013, 2018), in vivo to record synaptic input patterns. This method could be used to discover the asymmetric excitatory tuning curve inputs (with respect to somatic tuning) predicted for CAN models, the exponentially decaying inputs predicted by the Laplace transform model, the pulse of excitatory input predicted for the positive feedback ramping integrator, or the persistent excitatory input predicted for the perfect integrator ramping model. Lastly, retrograde circuit-tracing techniques (Tervo et al. 2016) could be used to hunt down the source and nature of the temporal derivative (γ) signal.

CONCLUSIONS

Time is as abstract as it is pervasive in animal behavior and neural circuits. For the same reasons that time is elusive and difficult to study, it may serve as a powerful window into general mechanisms of cognition and neural computations. The study of spatial navigation was once viewed similarly, but the unraveling of the place code has opened the door to a goldmine of insights into how the brain works. Understanding the time code for interval timing likewise carries great potential; by leveraging what we already know about spatial navigation, perhaps we are not too far from realizing this potential.

DISCLOSURE STATEMENT

The authors are not aware of any affiliations, memberships, funding, or financial holdings that might be perceived as affecting the objectivity of this review.

ACKNOWLEDGMENTS

The authors thank Brad Radvansky and Marc Howard for comments on this manuscript and members of the Dombeck Lab for helpful discussions. This work was supported by the Office of Naval Research (N00014-16-1-2832 and N00014-19-1-2571 to M.E.H.), the McKnight Foundation (D.A.D.), the Simons Collaboration on the Global Brain Postdoctoral Fellowship (J.G.H.), the Chicago Biomedical Consortium with support from the Searle Funds at The Chicago Community Trust (D.A.D.), and the National Institutes of Health (2R01MH101297 to D.A.D. 1. and R01 MH120073 to M.E.H.).

LITERATURE CITED

Akhlaghpour H, Wiskerke J, Choi JY, Taliaferro JP, Au J, Witten IB. 2016. Dissociated sequential activity and stimulus encoding in the dorsomedial striatum during spatial working memory. *eLife* 5:e19507

- Anderson MI, Jeffery KJ. 2003. Heterogeneous modulation of place cell firing by changes in context. J. Neurosci. 23(26):8827–35
- Armbruster BN, Li X, Pausch MH, Herlitze S, Roth BL. 2007. Evolving the lock to fit the key to create a family of G protein–coupled receptors potently activated by an inert ligand. PNAS 104(12):5163–68

- Arriaga M, Han EB. 2017. Dedicated hippocampal inhibitory networks for locomotion and immobility. 7. Neurosci. 37(38):9222-38
- Bakhurin KI, Goudar V, Shobe JL, Claar LD, Buonomano DV, Masmanidis SC. 2017. Differential encoding of time by prefrontal and striatal network dynamics. J. Neurosci. 37(4):854–70

Balci F, Simen P. 2016. A decision model of timing. Curr. Opin. Behav. Sci. 8:94-101

- Bangasser DA, Waxler DE, Santollo J, Shors TJ. 2006. Trace conditioning and the hippocampus: the importance of contiguity. J. Neurosci. 26(34):8702–6
- Barlow HB. 1972. Single units and sensation: a neuron doctrine for perceptual psychology? *Perception* 1(4):371–94
- Barry C, Hayman R, Burgess N, Jeffery KJ. 2007. Experience-dependent rescaling of entorhinal grids. *Nat. Neurosci.* 10(6):682–84
- Berthier NE, Moore JW. 1986. Cerebellar Purkinje cell activity related to the classically conditioned nictitating membrane response. *Exp. Brain Res.* 63(2):341–50
- Bittner KC, Milstein AD, Grienberger C, Romani S, Magee JC. 2017. Behavioral time scale synaptic plasticity underlies CA1 place fields. *Science* 357(6355):1033–36
- Blair HT, Wu A, Cong J. 2014. Oscillatory neurocomputing with ring attractors: a network architecture for mapping locations in space onto patterns of neural synchrony. *Philos. Trans. R. Soc. Lond. B Biol. Sci.* 369(1635):20120526
- Boyden ES, Zhang F, Bamberg E, Nagel G, Deisseroth K. 2005. Millisecond-timescale, genetically targeted optical control of neural activity. *Nat. Neurosci.* 8(9):1263–68
- Bright IM, Meister MLR, Cruzado NA, Tiganj Z, Howard MW, Buffalo EA. 2019. A temporal record of the past with a spectrum of time constants in the monkey entorhinal cortex. bioRxiv 688341. https://doi. org/10.1101/688341

Brown GD, Preece T, Hulme C. 2000. Oscillator-based memory for serial order. Psychol. Rev. 107(1):127-81

- Burak Y, Brookings T, Fiete I. 2006. Triangular lattice neurons may implement an advanced numeral system to precisely encode rat position over large ranges. arXiv:q-bio/0606005 [q-bio.NC]
- Burak Y, Fiete IR. 2009. Accurate path integration in continuous attractor network models of grid cells. *PLOS Comput. Biol.* 5(2):e1000291
- Burgess N, Barry C, O'Keefe J. 2007. An oscillatory interference model of grid cell firing. *Hippocampus* 17(9):801–12
- Bush D, Burgess N. 2014. A hybrid oscillatory interference/continuous attractor network model of grid cell firing. J. Neurosci. 34(14):5065–79
- Buzsáki G, Llinás R. 2017. Space and time in the brain. Science 358(6362):482-85
- Cameron CM, Murugan M, Choi JY, Engel EA, Witten IB. 2019. Increased cocaine motivation is associated with degraded spatial and temporal representations in IL-NAc neurons. *Neuron* 103(1):80–91.e7
- Campbell MG, Ocko SA, Mallory CS, Low IIC, Ganguli S, Giocomo LM. 2018. Principles governing the integration of landmark and self-motion cues in entorhinal cortical codes for navigation. *Nat. Neurosci.* 21(8):1096–106
- Carr CE, Konishi M. 1990. A circuit for detection of interaural time differences in the brain stem of the barn owl. J. Neurosci. 10(10):3227–46
- Chettih SN, Harvey CD. 2019. Single-neuron perturbations reveal feature-specific competition in V1. *Nature* 567(7748):334–40
- Cooper WE, Frederick WG. 2007. Optimal time to emerge from refuge. Biol. J. Linn. Soc. 91(3):375-82
- Couey JJ, Witoelar A, Zhang S-J, Zheng K, Ye J, et al. 2013. Recurrent inhibitory circuitry as a mechanism for grid formation. *Nat. Neurosci.* 16(3):318–24
- Craig ADB. 2009. Emotional moments across time: a possible neural basis for time perception in the anterior insula. *Philos. Trans. R. Soc. Lond. B Biol. Sci.* 364(1525):1933–42
- Dana H, Sun Y, Mohar B, Hulse BK, Kerlin AM, et al. 2019. High-performance calcium sensors for imaging activity in neuronal populations and microcompartments. *Nat. Methods* 16(7):649–57
- Densen ME. 1977. Time perception and schizophrenia. Percept. Mot. Skills 44(2):436-38
- Dill LM, Fraser AHG. 1997. The worm re-turns: hiding behavior of a tube-dwelling marine polychaete, Serpula vermicularis. Behav. Ecol. 8(2):186–93

- Domnisoru C, Kinkhabwala AA, Tank DW. 2013. Membrane potential dynamics of grid cells. *Nature* 495:199–204
- Durstewitz D. 2003. Self-organizing neural integrator predicts interval times through climbing activity. *J. Neurosci.* 23(12):5342–53
- Eichenbaum H. 2017. On the integration of space, time, and memory. Neuron 95(5):1007-18
- Fenton AA, Kao H-Y, Neymotin SA, Olypher A, Vayntrub Y, et al. 2008. Unmasking the CA1 ensemble place code by exposures to small and large environments: more place cells and multiple, irregularly arranged, and expanded place fields in the larger space. *J. Neurosci.* 28(44):11250–62
- Fuhs MC, Touretzky DS. 2006. A spin glass model of path integration in rat medial entorhinal cortex. *7. Neurosci.* 26(16):4266–76
- Fyhn M, Molden S, Witter MP, Moser EI, Moser M-B. 2004. Spatial representation in the entorhinal cortex. Science 305(5688):1258–64
- Gibbon J. 1977. Scalar expectancy theory and Weber's law in animal timing. Psychol. Rev. 84(3):279-325
- Giocomo LM, Zilli EA, Fransén E, Hasselmo ME. 2007. Temporal frequency of subthreshold oscillations scales with entorhinal grid cell field spacing. *Science* 315(5819):1719–22
- Grossberg S, Merrill JW. 1992. A neural network model of adaptively timed reinforcement learning and hippocampal dynamics. *Brain Res. Cogn. Brain Res.* 1(1):3–38
- Grossberg S, Schmajuk NA. 1991. Neural dynamics of adaptive timing and temporal discrimination during associative learning. In *Pattern Recognition by Self-Organizing Neural Networks*, ed. GA Carpenter, S Grossberg, pp. 637–74. Cambridge, MA: MIT Press
- Guanella A, Kiper D, Verschure P. 2007. A model of grid cells based on a twisted torus topology. *Int. J. Neural* Syst. 17(4):231–40
- Hafting T, Fyhn M, Molden S, Moser M-B, Moser EI. 2005. Microstructure of a spatial map in the entorhinal cortex. Nature 436(7052):801–6
- Hampson RE, Heyser CJ, Deadwyler SA. 1993. Hippocampal cell firing correlates of delayed-match-tosample performance in the rat. *Behav. Neurosci.* 107(5):715–39
- Hardcastle K, Ganguli S, Giocomo LM. 2015. Environmental boundaries as an error correction mechanism for grid cells. *Neuron* 86(3):827–39
- Harvey CD, Coen P, Tank DW. 2012. Choice-specific sequences in parietal cortex during a virtual-navigation decision task. *Nature* 484:62–68
- Hasselmo ME. 2008. Grid cell mechanisms and function: contributions of entorhinal persistent spiking and phase resetting. *Hippocampus* 18(12):1213–29
- Hasselmo ME. 2012. How We Remember: Brain Mechanisms of Episodic Memory. Cambridge, MA: MIT Press
- Hasselmo ME. 2014. Neuronal rebound spiking, resonance frequency and theta cycle skipping may contribute to grid cell firing in medial entorhinal cortex. *Philos. Trans. R. Soc. Lond. B Biol. Sci.* 369(1635):20120523
- Hasselmo ME, Giocomo LM, Brandon MP, Yoshida M. 2010. Cellular dynamical mechanisms for encoding the time and place of events along spatiotemporal trajectories in episodic memory. *Behav. Brain Res.* 215(2):261–74
- Heys JG, Dombeck DA. 2018. Evidence for a subcircuit in medial entorhinal cortex representing elapsed time during immobility. Nat. Neurosci. 21(11):1574–82
- Howard MW, MacDonald CJ, Tiganj Z, Shankar KH, Du Q, et al. 2014. A unified mathematical framework for coding time, space, and sequences in the hippocampal region. *7. Neurosci.* 34(13):4692–707
- Hugie DM. 2003. The waiting game: a "battle of waits" between predator and prey. *Behav. Ecol.* 14(6):807–17
- Itskov V, Curto C, Pastalkova E, Buzsáki G. 2011. Cell assembly sequences arising from spike threshold adaptation keep track of time in the hippocampus. J. Neurosci. 31(8):2828–34
- Ivry R. 1997. Cerebellar timing systems. Int. Rev. Neurobiol. 41:555-73
- Ivry RB, Schlerf JE. 2008. Dedicated and intrinsic models of time perception. Trends Cogn. Sci. 12(7):273– 80
- Jayakumar RP, Madhav MS, Savelli F, Blair HT, Cowan NJ, Knierim JJ. 2019. Recalibration of path integration in hippocampal place cells. *Nature* 566(7745):533–37

- Jennions MD, Backwell PRY, Murai M, Christy JH. 2003. Hiding behaviour in fiddler crabs: How long should prey hide in response to a potential predator? *Animal Behav*. 66(2):251–57
- Jin DZ, Fujii N, Graybiel AM. 2009. Neural representation of time in cortico-basal ganglia circuits. *PNAS* 106(45):19156–61
- Jun JJ, Steinmetz NA, Siegle JH, Denman DJ, Bauza M, et al. 2017. Fully integrated silicon probes for highdensity recording of neural activity. *Nature* 551:232–36
- Karmarkar UR, Buonomano DV. 2007. Timing in the absence of clocks: encoding time in neural network states. Neuron 53(3):427–38
- Kay K, Sosa M, Chung JE, Karlsson MP, Larkin MC, Frank LM. 2016. A hippocampal network for spatial coding during immobility and sleep. *Nature* 531(7593):185–90
- Kjelstrup KB, Solstad T, Brun VH, Hafting T, Leutgeb S, et al. 2008. Finite scale of spatial representation in the hippocampus. *Science* 321(5885):140–43
- Koivula K, Rytkonen S, Orell M. 1995. Hunger-dependency of hiding behaviour after a predator attack in dominant and subordinate willow tits. *Ardea* 83(2):397–404
- Kotani S, Kawahara S, Kirino Y. 2003. Purkinje cell activity during learning a new timing in classical eyeblink conditioning. *Brain Res.* 994(2):193–202
- Kraus BJ, Brandon MP, Robinson RJ, Connerney MA, Hasselmo ME, Eichenbaum H. 2015. During running in place, grid cells integrate elapsed time and distance run. *Neuron* 88(3):578–89
- Kraus BJ, Robinson RJ, White JA, Eichenbaum H, Hasselmo ME. 2013. Hippocampal "time cells": time versus path integration. *Neuron* 78(6):1090–101
- Kropff E, Carmichael JE, Moser M-B, Moser EI. 2015. Speed cells in the medial entorhinal cortex. Nature 523(7561):419–24
- Laje R, Buonomano DV. 2013. Robust timing and motor patterns by taming chaos in recurrent neural networks. Nat. Neurosci. 16(7):925–33
- Leon MI, Shadlen MN. 2003. Representation of time by neurons in the posterior parietal cortex of the macaque. Neuron 38(2):317–27
- Lim S, Goldman MS. 2013. Balanced cortical microcircuitry for maintaining information in working memory. Nat. Neurosci. 16(9):1306–14
- Liu Y, Tiganj Z, Hasselmo ME, Howard MW. 2019. A neural microcircuit model for a scalable scale-invariant representation of time. *Hippocampus* 29(3):260–74
- Lorentz HA. 1903. Electromagnetic phenomena in a system moving with any velocity smaller than that of light. In Proceedings of the Royal Netherlands Academy of Arts and Sciences, Vol. 6, pp. 809–31. Amsterdam: KNAW
- Lusk NA, Petter EA, MacDonald CJ, Meck WH. 2016. Cerebellar, hippocampal, and striatal time cells. *Curr*: *Opin. Behav. Sci.* 8:186–92
- MacDonald CJ, Carrow S, Place R, Eichenbaum H. 2013. Distinct hippocampal time cell sequences represent odor memories in immobilized rats. *7. Neurosci.* 33(36):14607–16
- MacDonald CJ, Lepage KQ, Eden UT, Eichenbaum H. 2011. Hippocampal "time cells" bridge the gap in memory for discontiguous events. *Neuron* 71(4):737–49
- Maguire EA, Nannery R, Spiers HJ. 2006. Navigation around London by a taxi driver with bilateral hippocampal lesions. *Brain* 129(Pt. 11):2894–907
- Martín J, López P. 1999. When to come out from a refuge: risk-sensitive and state-dependent decisions in an alpine lizard. *Behav. Ecol.* 10(5):487–92
- Marvin JS, Borghuis BG, Tian L, Cichon J, Harnett MT, et al. 2013. An optimized fluorescent probe for visualizing glutamate neurotransmission. Nat. Methods 10:162–70
- Marvin JS, Scholl B, Wilson DE, Podgorski K, Kazemipour A, et al. 2018. Stability, affinity, and chromatic variants of the glutamate sensor iGluSnFR. Nat. Methods 15(11):936–39
- Matell MS, Meck WH. 2004. Cortico-striatal circuits and interval timing: coincidence detection of oscillatory processes. Brain Res. Cogn. Brain Res. 21(2):139–70
- Matell MS, Meck WH, Nicolelis MAL. 2003. Interval timing and the encoding of signal duration by ensembles of cortical and striatal neurons. *Behav. Neurosci.* 117(4):760–73

- Mathis A, Herz AVM, Stemmler MB. 2012. Resolution of nested neuronal representations can be exponential in the number of neurons. *Phys. Rev. Lett.* 109(1):018103
- Mau W, Sullivan DW, Kinsky NR, Hasselmo ME, Howard MW, Eichenbaum H. 2018. The same hippocampal CA1 population simultaneously codes temporal information over multiple timescales. *Curr. Biol.* 28(10):1499–508.e4
- Mauk MD, Buonomano DV. 2004. The neural basis of temporal processing. Annu. Rev. Neurosci. 27:307-40
- McEchron MD, Tseng W, Disterhoft JF. 2003. Single neurons in CA1 hippocampus encode trace interval duration during trace heart rate (fear) conditioning in rabbit. *J. Neurosci.* 23(4):1535–47
- McNaughton BL, Battaglia FP, Jensen O, Moser EI, Moser M-B. 2006. Path integration and the neural basis of the "cognitive map." *Nat. Rev. Neurosci.* 7(8):663–78
- Meck WH. 1996. Neuropharmacology of timing and time perception. Brain Res. Cogn. Brain Res. 3(3-4):227-42
- Meck WH, Church RM, Matell MS. 2013. Hippocampus, time, and memory—a retrospective analysis. *Behav. Neurosci.* 127(5):642–54
- Meck WH, Church RM, Olton DS. 1984. Hippocampus, time, and memory. Behav. Neurosci. 98(1):3-22
- Mello GBM, Soares S, Paton JJ. 2015. A scalable population code for time in the striatum. *Curr. Biol.* 25(9):1113-22
- Merchant H, Harrington DL, Meck WH. 2013. Neural basis of the perception and estimation of time. Annu. Rev. Neurosci. 36:313–36
- Miall C. 1989. The storage of time intervals using oscillating neurons. Neural Comput. 1(3):359-71
- Mita A, Mushiake H, Shima K, Matsuzaka Y, Tanji J. 2009. Interval time coding by neurons in the presupplementary and supplementary motor areas. *Nat. Neurosci.* 12(4):502–7
- Morris RG, Garrud P, Rawlins JN, O'Keefe J. 1982. Place navigation impaired in rats with hippocampal lesions. Nature 297(5868):681–83
- Moser EI, Moser M-B. 2008. A metric for space. Hippocampus 18(12):1142-56
- Mosheiff N, Agmon H, Moriel A, Burak Y. 2017. An efficient coding theory for a dynamic trajectory predicts non-uniform allocation of entorhinal grid cells to modules. PLOS Comput. Biol. 13(6):e1005597
- O'Keefe J, Dostrovsky J. 1971. The hippocampus as a spatial map. Preliminary evidence from unit activity in the freely-moving rat. *Brain Res.* 34(1):171–75
- O'Keefe J, Nadel L. 1978. The Hippocampus as a Cognitive Map. Oxford, UK: Clarendon Press
- O'Keefe J, Recce ML. 1993. Phase relationship between hippocampal place units and the EEG theta rhythm. *Hippocampus* 3(3):317–30
- Pastalkova E, Itskov V, Amarasingham A, Buzsáki G. 2008. Internally generated cell assembly sequences in the rat hippocampus. *Science* 321(5894):1322–27
- Paton JJ, Buonomano DV. 2018. The neural basis of timing: distributed mechanisms for diverse functions. *Neuron* 98(4):687–705
- Quintana J, Fuster JM. 1999. From perception to action: temporal integrative functions of prefrontal and parietal neurons. *Cereb. Cortex* 9(3):213–21
- Rajan K, Harvey CD, Tank DW. 2016. Recurrent network models of sequence generation and memory. *Neuron* 90(1):128–42
- Reutimann J, Yakovlev V, Fusi S, Senn W. 2004. Climbing neuronal activity as an event-based cortical representation of time. 7. Neurosci. 24(13):3295–303
- Robinson NTM, Priestley JB, Rueckemann JW, Garcia AD, Smeglin VA, et al. 2017. Medial entorhinal cortex selectively supports temporal coding by hippocampal neurons. *Neuron* 94(3):677–88.e6

Rolls ET, Mills P. 2019. The generation of time in the hippocampal memory system. *Cell Rep.* 28(7):1649–58.e6 Rovelli C. 2018. *The Order of Time*. London: Allen Lane

- Sabariego M, Schönwald A, Boublil BL, Zimmerman DT, Ahmadi S, et al. 2019. Time cells in the hippocampus are neither dependent on medial entorhinal cortex inputs nor necessary for spatial working memory. *Neuron* 102(6):1235–48.e5
- Sadagopan S, Wang X. 2009. Nonlinear spectrotemporal interactions underlying selectivity for complex sounds in auditory cortex. J. Neurosci. 29(36):11192–202

- Salz DM, Tiganj Z, Khasnabish S, Kohley A, Sheehan D, et al. 2016. Time cells in hippocampal area CA3. *7. Neurosci.* 36(28):7476–84
- Samsonovich A, McNaughton BL. 1997. Path integration and cognitive mapping in a continuous attractor neural network model. J. Neurosci. 17(15):5900–20
- Save E, Nerad L, Poucet B. 2000. Contribution of multiple sensory information to place field stability in hippocampal place cells. *Hippocampus* 10(1):64–76
- Savelli F, Knierim JJ. 2019. Origin and role of path integration in the cognitive representations of the hippocampus: computational insights into open questions. J. Exp. Biol. 222:jeb188912
- Schirmer A. 2011. How emotions change time. Front. Integr. Neurosci. 5:58
- Seung HS, Lee DD, Reis BY, Tank DW. 2000. Stability of the memory of eye position in a recurrent network of conductance-based model neurons. *Neuron* 26(1):259–71
- Shankar KH, Howard MW. 2012. A scale-invariant internal representation of time. *Neural Comput.* 24(1):134– 93
- Sharp PE, Blair HT, Cho J. 2001. The anatomical and computational basis of the rat head-direction cell signal. *Trends Neurosci.* 24(5):289–94
- Simen P, Balci F, de Souza L, Cohen JD, Holmes P. 2011. A model of interval timing by neural integration. J. Neurosci. 31(25):9238–53
- Skaggs WE, Knierim JJ, Kudrimoti HS, McNaughton BL. 1995. A model of the neural basis of the rat's sense of direction. Adv. Neural Inf. Process. Syst. 7:173–80
- Soares S, Atallah BV, Paton JJ. 2016. Midbrain dopamine neurons control judgment of time. *Science* 354(6317):1273–77
- Sober SJ, Sponberg S, Nemenman I, Ting LH. 2018. Millisecond spike timing codes for motor control. *Trends* Neurosci. 41(10):644–48
- Solstad T, Boccara CN, Kropff E, Moser M-B, Moser EI. 2008. Representation of geometric borders in the entorhinal cortex. Science 322(5909):1865–68
- Spencer RMC, Karmarkar U, Ivry RB. 2009. Evaluating dedicated and intrinsic models of temporal encoding by varying context. *Philos. Trans. R. Soc. Lond. B Biol. Sci.* 364(1525):1853–63
- Squire LR. 1992. Memory and the hippocampus: a synthesis from findings with rats, monkeys, and humans. *Psychol. Rev.* 99(2):195–231
- Sreenivasan S, Fiete I. 2011. Grid cells generate an analog error-correcting code for singularly precise neural computation. Nat. Neurosci. 14(10):1330–37
- Stensola H, Stensola T, Solstad T, Frøland K, Moser M-B, Moser EI. 2012. The entorhinal grid map is discretized. Nature 492(7427):72–78
- Sussillo D, Abbott LF. 2009. Generating coherent patterns of activity from chaotic neural networks. *Neuron* 63(4):544–57
- Tahvildari B, Fransén E, Alonso AA, Hasselmo ME. 2007. Switching between "On" and "Off" states of persistent activity in lateral entorhinal layer III neurons. *Hippocampus* 17(4):257–63
- Takahashi JS. 2017. Transcriptional architecture of the mammalian circadian clock. *Nat. Rev. Genet.* 18(3):164–79
- Taube JS, Muller RU, Ranck JB. 1990. Head-direction cells recorded from the postsubiculum in freely moving rats. I. Description and quantitative analysis. *J. Neurosci.* 10(2):420–35
- Tervo DGR, Hwang B-Y, Viswanathan S, Gaj T, Lavzin M, et al. 2016. A designer AAV variant permits efficient retrograde access to projection neurons. *Neuron* 92(2):372–82
- Tiganj Z, Cromer JA, Roy JE, Miller EK, Howard MW. 2018. Compressed timeline of recent experience in monkey lateral prefrontal cortex. *J. Cogn. Neurosci.* 30(7):935–50
- Tiganj Z, Hasselmo ME, Howard MW. 2015. A simple biophysically plausible model for long time constants in single neurons. *Hippocampus* 25(1):27–37
- Tsao A, Sugar J, Lu L, Wang C, Knierim JJ, et al. 2018. Integrating time from experience in the lateral entorhinal cortex. *Nature* 561(7721):57–62
- Tulving E. 2002. Episodic memory: from mind to brain. Annu. Rev. Psychol. 53:1-25
- Welsh DK, Logothetis DE, Meister M, Reppert SM. 1995. Individual neurons dissociated from rat suprachiasmatic nucleus express independently phased circadian firing rhythms. *Neuron* 14(4):697–706

- Wilson DE, Scholl B, Fitzpatrick D. 2018. Differential tuning of excitation and inhibition shapes direction selectivity in ferret visual cortex. *Nature* 560(7716):97–101
- Wittmann M. 2009. The inner experience of time. Philos. Trans. R. Soc. B Biol. Sci. 364(1525):1955-67
- Yoshida M, Fransén E, Hasselmo ME. 2008. mGluR-dependent persistent firing in entorhinal cortex layer III neurons. Eur. J. Neurosci. 28(6):1116–26
- Zhang K. 1996. Representation of spatial orientation by the intrinsic dynamics of the head-direction cell ensemble: a theory. *J. Neurosci.* 16(6):2112–26
- Zhang S, Schönfeld F, Wiskott L, Manahan-Vaughan D. 2014. Spatial representations of place cells in darkness are supported by path integration and border information. *Front. Behav. Neurosci.* 8:222