

Hippocampome.org v2.0: a knowledge base enabling data-driven spiking neural network simulations of rodent hippocampal circuits

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Abstract

Hippocampome.org (<https://hippocampome.org/>) is a mature open-access knowledge base of the rodent hippocampal formation focusing on neuron types and their properties. **Hippocampome.org** (<https://hippocampome.org/>) v1.0 established a foundational classification system identifying 122 hippocampal neuron types based on their axonal and dendritic morphologies, main neurotransmitter, membrane biophysics, and molecular expression. Releases v1.1 through v1.12 furthered the aggregation of literature-mined data, including among others neuron counts, spiking patterns, synaptic physiology, in vivo firing phases, and connection probabilities. Those additional properties increased the online information content of this public resource over 100-fold, enabling numerous independent discoveries by the scientific community. **Hippocampome.org** (<https://hippocampome.org/>) v2.0, introduced here, incorporates over 50 new neuron types and extends the functionality to build real-scale, biologically detailed, data-driven computational simulations. In all cases, the freely downloadable model parameters are directly linked to the specific peer-reviewed empirical evidence from which they were derived. Possible research applications include quantitative, multiscale analyses of circuit connectivity and spiking neural network simulations of activity dynamics. These advances can help generate precise, experimentally testable hypotheses and shed light on the neural mechanisms underlying associative memory and spatial navigation.

eLife assessment

The authors have updated a helpful resource for the neuroscience community regarding hippocampal cell types and their interactions, called the "hippocampome." The updates expand content, tools, and potential applications, and therefore the significance is **important**. The strong presentation makes the paper **compelling**.

Introduction

Neuroscience knowledge continues to increase every year (Eke et al., 2022 [↗](#); Yeung et al., 2017 [↗](#)), making it challenging for researchers to keep abreast of mounting data and evolving information even in their own domain of expertise. Large-scale endeavors, such as the Brain Research through Advancing Innovative Neurotechnologies (BRAIN) Initiative (Insel et al., 2013 [↗](#)) and the European Union's Human Brain Project (Amunts et al., 2016 [↗](#)), are contributing to this tremendous growth along with the 'long tail' of independent labs and individual scientists (Ferguson et al., 2014 [↗](#)). A key organizing principle for neuroscience knowledge is the seminal notion of *neuron types* (Petilla Interneuron Nomenclature Group et al., 2008; Zeng and Sanes, 2017 [↗](#)), which constitute the conceptual 'parts list' of functional circuits. The National Institutes of Health launched the BICCN (see abbreviations in Materials and Methods) to help establish a comprehensive reference of the cell type diversity in the human, mouse, and non-human primate brains (BRAIN Initiative Cell Census Network, 2021). This multi-institution collaboration is already producing innovative results (Muñoz-Castañeda et al., 2021 [↗](#)) and actionable community resources (BICCN Data Ecosystem Collaboration, 2022 [↗](#)).

Hippocampome.org (<https://hippocampome.org/>) [↗](#) is an open-access knowledge base of the rodent hippocampal circuit (dentate gyrus, CA3, CA2, CA1, subiculum, and entorhinal cortex) at the mesoscopic level of neuron types (Wheeler et al., 2015 [↗](#)). This resource has proven popular and effective thanks to the adoption of a simple yet powerful classification system for defining neuron types. Specifically, a key property for the identification of neuron types in *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) is the location of axons and dendrites across the subregions and layers of the hippocampal formation. This approach can be broadly extended to classify neurons in other brain regions and neural systems (Ascoli and Wheeler, 2016 [↗](#)). Focusing on axonal and dendritic distributions provides two considerable advantages. First, these features mediate neuronal connectivity, thus immediately revealing the underlying blueprint of network circuitry (Rees et al., 2017 [↗](#)). Second, they are widely used in the neuroscience community as a reliable and concrete anchoring signature to correlate electrophysiological and transcriptomic profiles (DeFelipe et al., 2013 [↗](#)). Therefore, starting from the foundational morphology-based identification of 122 neuron types in the first release (version 1.0 or v1.0), *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) progressively amassed an increasing amount of complementary data, such as firing patterns, molecular expression, cell counts, synaptic communication, in vivo oscillations, and connection probabilities. In all cases, the public repository provided direct links to the specific peer-reviewed empirical evidence supporting the added knowledge.

Having established a web-based integrated storehouse of hippocampal information, *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) also expanded its scope by including data-driven computational models of neuronal excitability and synaptic signaling, as well as ties to community resources such as NeuroMorpho.Org (Akram et al., 2018 [↗](#)), SenseLab ModelDB (McDougal et al., 2017 [↗](#)), CARLsim (Niedermeier et al., 2022 [↗](#)), and the Allen Brain Atlas (Jones et al., 2009 [↗](#)). Altogether, these extensions resulted in the emergence of a complete framework for launching real-scale, biologically detailed computer simulations of the hippocampal formation. The present report thus marks a new phase in the life cycle of this community resource. *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) v2.0, introduced here, also incorporates over 50 newly identified neuron types, updating the classification to the most recent literature findings.

The following "Description of resource" section begins with a concise, referenced overview of the neural properties collated from *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) v1.0 through release v1.12. We then briefly describe the new neuron types and data currently being added in *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) v2.0. Next is an abridged summary of the usage and recognition of this online portal in biomedical research. This is followed by an explanation of

the latest capabilities to search, filter, and download the complete set of computational parameters enabling quantitative connectomic analyses and spiking neural network simulations. The section concludes with an outlook of possible research applications allowed by the expansion of this scientific resource.

Description of resource

Characterizing properties of hippocampal neuron types

Hippocampome.org (<https://hippocampome.org/>) [v1.0](#) (Wheeler et al., 2015 [↗](#)) established the morphological encoding of axonal and dendritic locations and the main neurotransmitter (glutamate or GABA) as the primary determinants of neuron types in the rodent hippocampal formation. For example, a DG Basket cell (with name capitalized to indicate a formally defined neuron type) is a GABAergic cell with axon contained in the granular layer and dendrites spanning all DG layers (**Figures 1A1-4** [↗](#)). In this framework, two neurons releasing the same neurotransmitter belong to different types if the axon or dendrites of only one of them invades any of the 26 layers across 6 subregions of the hippocampal formation (hippocampome.org/morphology). In other words, neurons of the same type share the same potential inputs, outputs, and excitatory vs. inhibitory function. These properties were initially supplemented with additional empirical evidence for molecular expression of major protein biomarkers (**Figure 1A5** [↗](#); hippocampome.org/markers) and membrane biophysics (**Figure 1A6-7** [↗](#); hippocampome.org/electrophysiology).

Many neuronal properties and functionalities were progressively added in 12 subsequent releases (**Table 1** [↗](#)). The numerical sequencing of these *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) versions depended on the order of peer-review and publication of the corresponding scientific reports. Here we will describe them instead in logical groupings. The first two updates enhanced the user functionality of the knowledge base. Specifically, v1.1 integrated a web-based interactive thesaurus mapping of synonyms and definitions (Hamilton et al., 2017a [↗](#); hippocampome.org/find-term) to help disambiguate the many terminological inconsistencies in the neuroscience literature (Shepherd et al., 2019 [↗](#); Yuste et al., 2020 [↗](#)). Release v1.2 introduced the capability to browse, search, and analyze the potential connectivity between neuron types (Rees et al., 2016 [↗](#); *Hippocampome.org* (<https://hippocampome.org/>) [↗](#)/connectivity) as derived from the compiled overlapping locations of all the presynaptic axons and postsynaptic dendrites. Transcriptomic information was greatly expanded in both v1.3 (Hamilton et al., 2017b [↗](#)), which incorporated in situ hybridization data from the Allen Brain Atlas (Lein et al., 2007 [↗](#)), and v1.5 (White et al., 2020 [↗](#)), which leveraged relational inferences interlinking the region-specific expression of two or more genes.

The quantifications of firing pattern phenotypes, such as rapid adapting spiking, transient stuttering, and persistent slow-wave bursting, in v1.6 (Komendantov et al., 2019 [↗](#); hippocampome.org/firing_patterns) were fitted by dynamical systems modeling (Izhikevich, 2003 [↗](#)) in v1.7 (Venkadesh et al., 2019 [↗](#); hippocampome.org/Izhikevich). Although the above properties were largely measured from slice preparations, v1.9 made available measurements from *in vivo* recordings (Sanchez-Aguilera et al., 2021 [↗](#); hippocampome.org/in-vivo). Release v1.10 provided a compendium of cognitive functions linked to specific hippocampal neurons (Sutton and Ascoli, 2021 [↗](#); hippocampome.org/cognome), while the v1.11 neuron type census estimated the population counts for each neuron type (Attili et al., 2022 [↗](#); hippocampome.org/census). Last but not least are a set of properties pertaining not to individual neuron types but to synaptic connections between a pair of pre- and post-synaptic neuron types. In particular, v1.8 calculated the synaptic probabilities and the numbers of contacts per connected pair (Tecuatl et al., 2021b [↗](#); hippocampome.org/syn_probabilities), and v1.4 data mined synaptic physiology (Moradi and Ascoli, 2020 [↗](#); hippocampome.org/synaptome), with conductance, time constant, and short-term

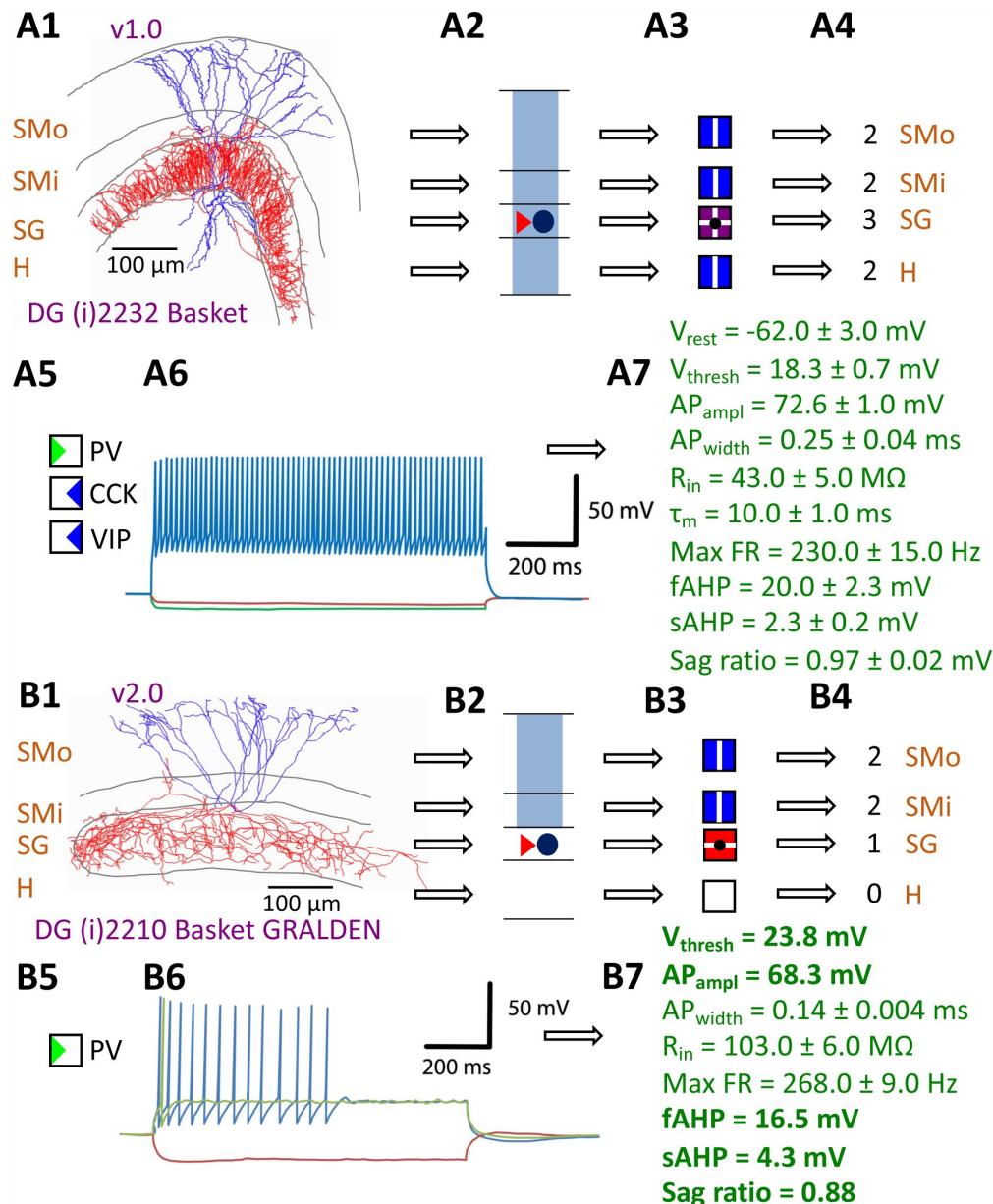


Figure 1.

Defining neuron types in *Hippocampome.org* (<https://hippocampome.org/>) [v1.0](#) neuron type. A1) Morphology of a DG (i)2232 Basket cell (NeuroMorpho.Org cell NMO_34300: Hosp et al., 2014 [v1.0](#)) with axons (red) in SG and dendrites (blue) in all four layers. A2) Schematic interpretation of the morphological tracing, where the circle represents the location of the soma in SG, the red triangle the location of the axons in SG, and the blue rectangles the locations of the dendrites in all four layers. A3) *Hippocampome.org* (<https://hippocampome.org/>) representation of the morphology, where a blue square with a vertical line (|) indicates dendritic presence in SMO and SMi and H, a purple square with a cross (+) indicates both axonal and dendritic presence in SG, and a black dot (•) indicates the soma location in SG. A4) *Hippocampome.org* (<https://hippocampome.org/>) numerical coding of the reconstructed neuron, where 2 indicates the presence of dendrites (in SMO, SMi, and H); and 3 indicates the presence of both axons and dendrites (in SG). A5) Biomarker expressions, where a green triangle indicates positive expression, and a blue triangle indicates negative expression. A6) Firing pattern phenotype (NASP; adapted from **Figure 1B** [v1.0](#) in Savanthrapadian et al., 2014 [v1.0](#)). A7) Membrane biophysics values (from **Figure 3C** [v1.0](#) and **Table 1** [v1.0](#) in Lübke et al., 1998 [v1.0](#)) recorded at 35–37 °C. B) Properties for a *Hippocampome.org* (<https://hippocampome.org/>) [v2.0](#) neuron type. B1) Morphology of a DG (i)2210 Basket GRALDEN (NeuroMorpho.Org cell NMO_146159: Vaden et al., 2020 [v2.0](#)) with red axons in SG and blue dendrites in SMO and SMi. B2) Schematic interpretation of the reconstruction (same symbols as in A2). B3–4) *Hippocampome.org* (<https://hippocampome.org/>) representation and numerical coding of the morphology (same symbols as in A3–4). B5) Biomarker expression. B6) Firing pattern phenotype (TSTUT.SLN; adapted from Figure S4 in Markwardt et al., 2011 [v2.0](#)). B7) Membrane biophysics values recorded at room temperature (from **Figure 4D** [v2.0](#) in Vaden et al., 2020 [v2.0](#)), and at 22 °C (from Figure S4 in Markwardt et al., 2011 [v2.0](#)); emboldened values were extracted from the firing pattern trace in B6. Abbreviations are defined in Materials and Methods.

Version	Contribution	Article
v1.1	definitions for terms and phrases relevant to Hippocampome.org	(Hamilton et al., 2017a)
v1.2	- clickable connectivity matrix - interactive connectivity navigator Java applet - searching by connectivity	(Rees et al., 2016)
v1.3	- downloadable list of ABA predictions of marker expressions - utility for viewing the effects of thresholds on ABA marker expression predictions	(Hamilton et al., 2017b)
v1.4	access to the synapse knowledge base	(Moradi and Ascoli, 2020)
v1.5	relational biomarker expression inferences	(White et al., 2020)
v1.6	- firing pattern phenotypes - clickable firing pattern matrix - clickable firing pattern parameters matrix - search by firing pattern - search by firing pattern parameter	(Komendantov et al., 2019)
v1.7	- Izhikevich models - clickable Izhikevich model parameters matrix - downloadable single-neuron parameter files - downloadable single-neuron CARLSim4 simulation files - ability to perform single-neuron simulations of the firing patterns	(Venkadesh et al., 2019)
v1.8	- clickable/downloadable neurite lengths matrix - clickable/downloadable somatic path distances matrix - clickable/downloadable numbers of potential synapses matrix - clickable/downloadable numbers of contacts matrix - clickable/downloadable connection probabilities matrix	(Tecuatl et al., 2021b)
v1.9	clickable matrix for in vivo recordings	(Sanchez-Aguilera et al., 2021)
v1.10	Cognome knowledge base of spiking neural circuit functions and network simulations of the hippocampal formation	(Sutton and Ascoli, 2021)
v1.11	clickable matrix of neuron type census values for rat and mouse	(Attili et al., 2022)
v1.12	clickable/downloadable matrices of synaptic physiology parameter values (g, τ_d, τ_r, τ_f, U) for combinations of species, sex, age, temperature, and recording mode	(Moradi et al., 2022)

Table 1.

Added knowledge and functioning in *Hippocampome.org* (<https://hippocampome.org/>)  releases v1.1-12.

plasticity values normalized by age, temperature, species, sex, and recording method in v1.12 (Moradi et al., 2022 [↗](#); hippocampome.org/synapse), leveraging machine learning and a phenomenological model (Tsodyks et al., 1998 [↗](#)).

Expanding the catalog of neuron types and properties from *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) v1.x to v2.0

The *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) framework to classify neuron types and collate their properties allows agile content updates as new data are continuously reported in the peer-reviewed literature. For example, the description of a parvalbumin-positive DG GABAergic interneuron with axon contained in the granular layer and dendrites invading the molecular layer but not the hilus (Vaden et al., 2020 [↗](#)) supported the definition of a new neuron type (**Figures 1B1-5** [↗](#)), referred to in *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) v2.0 as DG Basket GRALDEN. Moreover, such an identification made it possible to unequivocally ascribe to this neuron type previously reported electrophysiological characteristics (**Figures 1B6-7** [↗](#); Markwardt et al., 2011 [↗](#)). Comprehensive literature mining following the same process expanded the *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) v2.0 catalog with 51 new neuron types¹ [↗](#) across 5 of the 6 subregions of the hippocampal formation (**Figure 2** [↗](#)), including axonal-dendritic morphological patterns (**Figure 2A** [↗](#)), molecular expression (**Figure 2B** [↗](#)), and membrane biophysics (**Figure 2C** [↗](#)).

Besides identifying new neuron types, the *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) classification system also allows the ongoing accumulation of new properties onto existing neuron types as well as the reconciliation of fragmented descriptions from scientific publications (**Figure 3** [↗](#)). For instance, converging evidence indicates that EC Layer III Pyramidal cells have axonal projections in all layers of CA1 (Deller et al., 1996 [↗](#); Takács et al., 2012 [↗](#)), not just in stratum lacunosum-moleculare (SLM) as originally reported (Steward, 1976 [↗](#)). *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) v2.0 captures both the new extracted knowledge and the corresponding experimental evidence (**Figure 3A** [↗](#)). The annotation of neuron type-specific firing phases relative to in vivo oscillations in v1.9 highlighted a clear distinction between Superficial and Deep CA1 Pyramidal cells (Sanchez-Aguilera et al., 2021 [↗](#)). The present release enriches that description with accompanying novel molecular markers (**Figure 3B1** [↗](#)), membrane biophysics values (**Figure 3B2** [↗](#)), and differential connectivity with other subregions and neuron types (**Figure 3B3** [↗](#)). Similarly, numerous additional firing patterns (**Figure 3C** [↗](#)) have been datamined for existing neuron types, such as adapting spiking in CA3 Basket cells and non-adapting spiking in CA3 Bistratified cells (Fidzinski et al., 2015) or transient stuttering in CA1 Radiatum Giant cells (Kirson and Yaari, 2000 [↗](#)). Notably, this includes a novel phenotype, TSTUT.PSTUT in CA1 Interneuron Specific O-targeting QuadD cells (Chamberland et al., 2010 [↗](#)). With this report, we also release new differential connection probabilities to various CA1 neuron type targets from traditional CA3 Pyramidal cells vs. CA3c Pyramidal cells (**Figure 3D** [↗](#)) and from DG Granule cells to mossy fiber CA3 targets (**Table 2** [↗](#)).

Quantifying the content and impact of

Hippocampome.org (<https://hippocampome.org/>) [↗](#)

Over the course of subsequent releases, we have measured *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) content using two metrics. The number of pieces of knowledge (PoK) tallies the distinct units of structured information, such as the statements that DG Granule cell axons invade the hilus or that CA1 Basket cells express parvalbumin. The pieces of evidence (PoE) are specific excerpts of peer reviewed publications (portion of text, figure, or table) or database entries (e.g., from the Allen Brain Atlas) always linked to each PoK. Both PoK and PoE continued to grow with successive releases of *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) (**Figure 4A** [↗](#)). Notably, the largest increases in PoK and PoE were related to synaptic properties (Moradi and

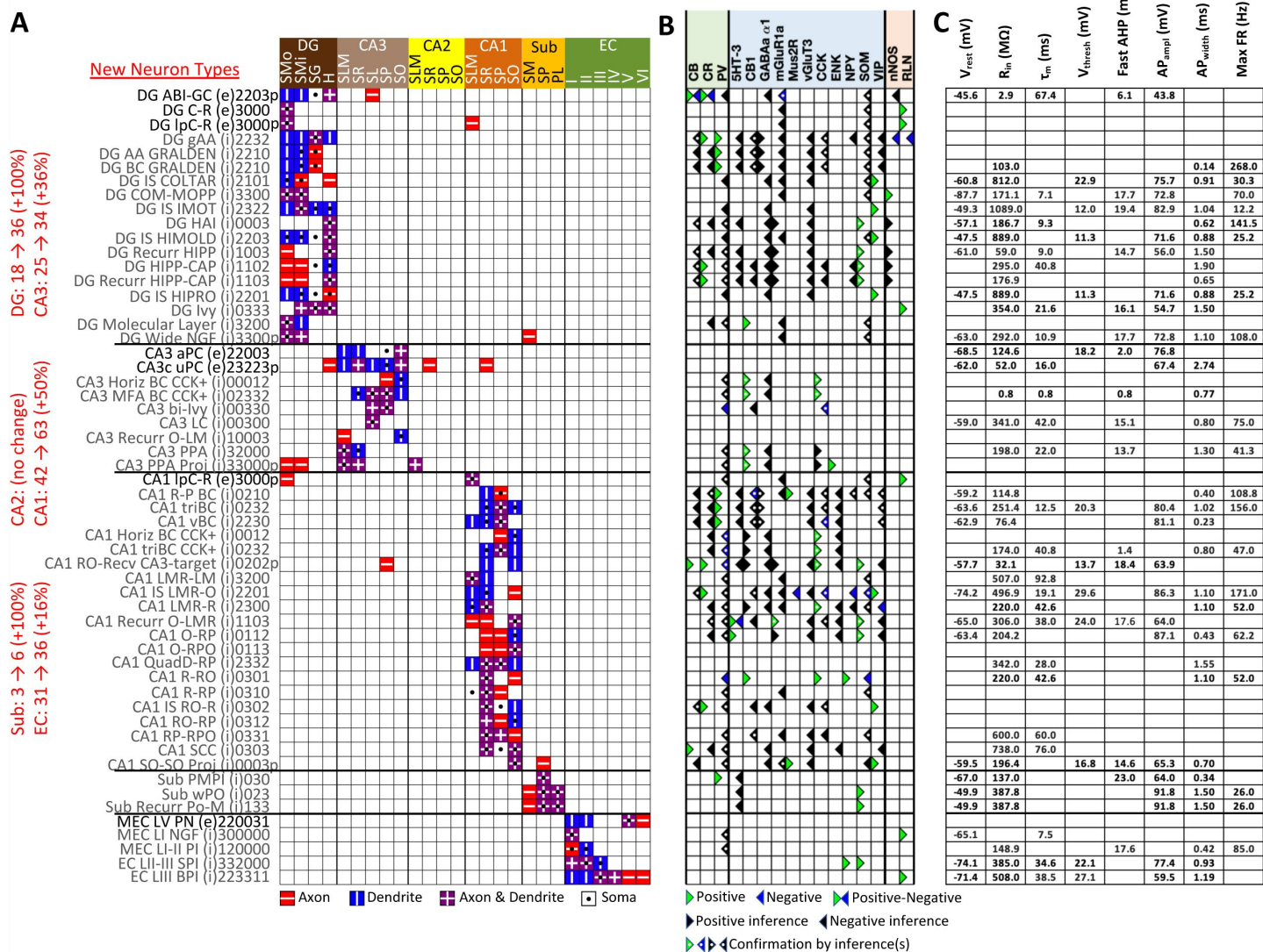


Figure 2.

New neuron types added to *Hippocampome.org* (<https://hippocampome.org/>) v2.0. A) Morphology encodings of the 56 new neuron types. (Left) Increase in number of neuron types for each subregion. B) Biomarker expressions of the neuron types. C) Membrane biophysics values for the neuron types. Abbreviations are defined in Materials and Methods.

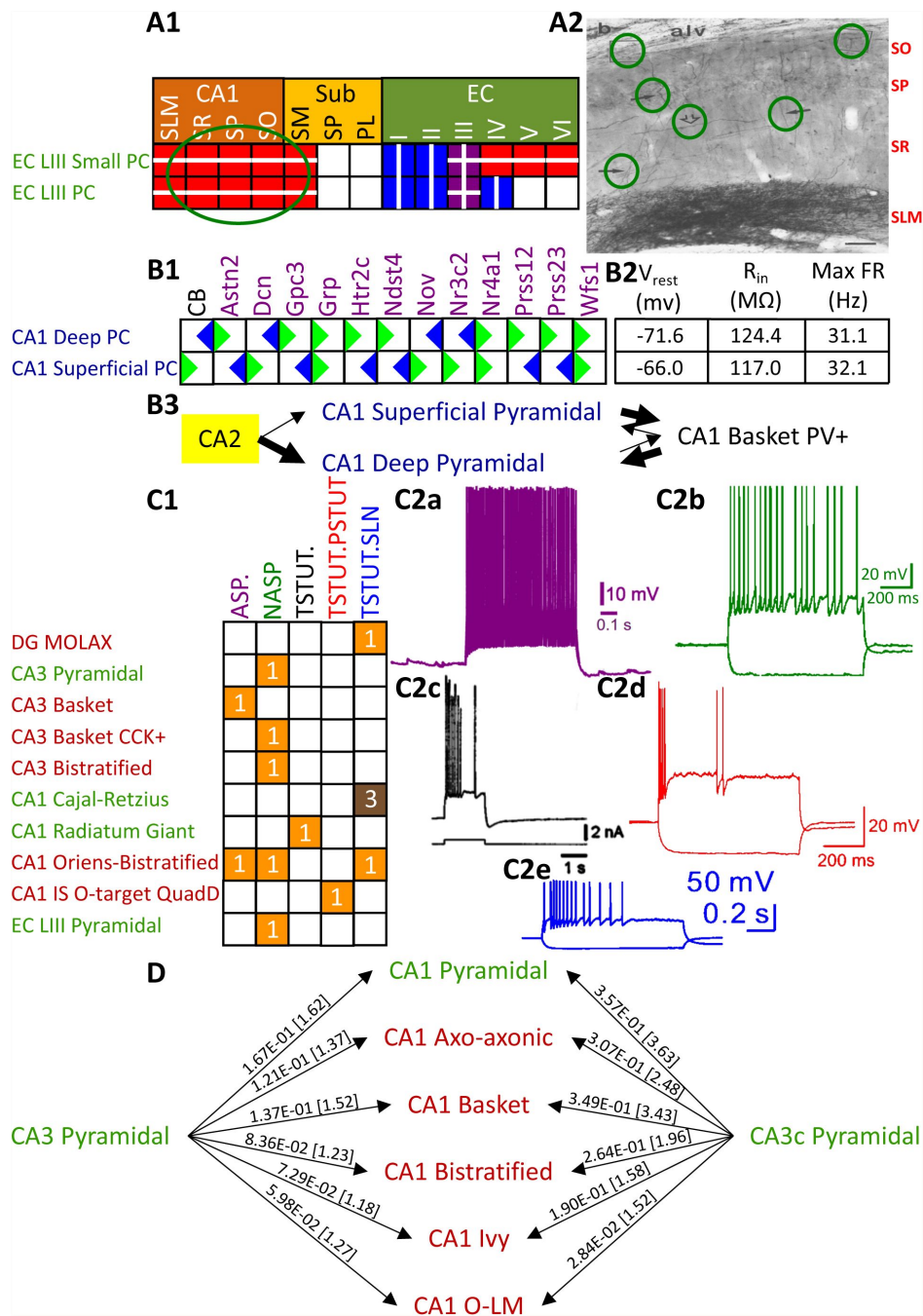


Figure 3.

Extensions to the neuronal properties of *Hippocampome.org* (<https://hippocampome.org/>) v1.x. A1) Additions to the axonal projections (circled in green) for two v1.0 neuron types. A2) Evidence for the axonal extensions in strata oriens, pyramidal, and radiatum from A1 are circled in green (adapted from **Figure 2b** in Deller et al., 1996). Scale bar: 75 μ m. B1) Biomarker expressions for the two CA1 Pyramidal sub-types added in *Hippocampome.org* (<https://hippocampome.org/>) v1.9 (Sanchez-Aguilera et al., 2021). B2) Membrane biophysics values for the two sub-types. B3) CA2 projects preferentially to the deep sublayer of CA1 (Kohara et al., 2014). More perisomatic PV+ GABAergic boutons are found at CA1 Deep Pyramidal cells (Valero et al., 2015). CA1 Superficial Pyramidal cells form more frequent connections to PV+ CA1 Basket cells, and PV+ CA1 Basket cells form significantly more perisomatic axon terminals on CA1 Deep Pyramidal cells (Lee et al., 2014). C1) Additions to the firing pattern phenotypes of v1.0 neuron types. C2a) Example of ASP. in a CA1 Oriens-Bistratified cell (adapted from **Figure 4B** in Craig and McBain, 2015). C2b) Example of NASP in a CA3 Basket CCK+ cell (adapted from **Figure 3A** in Szabadics and Soltesz, 2009). C2c) Example of TSTUT. in a CA1 Radiatum Giant cell (adapted from **Figure 2Bb** in Kirson and Yaari, 2000). C2d) Example of TSTUT.PSTUT in a CA1 Interneuron Specific O-targeting QuadD (adapted from **Figure 2D** in Chamberland et al., 2010). C2e) Example of TSTUT.SLN in a DG MOLAX cell (adapted from Figure S2c in Lee et al., 2016). neuron types. Abbreviations are defined in Materials and Methods.

Postsynaptic neuron type	Probability	# contacts
CA3 Pyramidal	1.11E-04	1.08
CA3c Pyramidal	3.91E-04	1.31
CA3 Spiny Lucidum Dentate-Projecting	5.89E-04	1.69
CA3 Mossy Fiber-Associated ORDEN	4.44E-04	1.27
CA3 Basket	6.55E-04	1.50
CA3 Basket CCK+	2.14E-04	1.16
CA3 Ivy	3.35E-04	1.29
CA3 Mossy Fiber-Associated	3.78E-05	1.04
CA3 LMR-Targeting	1.31E-04	1.21
CA3 Lucidum ORAX	2.62E-04	1.19
CA3 Lucidum-Radiatum	3.25E-04	1.13
CA3 Axo-Axonic	7.56E-04	1.50
CA3 Bistratified	8.25E-04	1.45
CA3 QuadD-LM	2.91E-04	1.25

Table 2.

Probabilities of connection and number of contacts per connected pair from DG Granule cell to mossy fibers targets in CA3.

Ascoli, 2020 [↗](#); Tecuatl et al., 2021b [↗](#); Moradi et al., 2022 [↗](#)). Specifically, the data underlying synaptic physiology and connection probabilities were supported by over 23,000 PoE and yielded a remarkable 500,000 PoK thanks to the normalized collection of signaling and short-term plasticity modeling parameters for multiple combinations of experimental conditions.

To assess community usage of *Hippocampome.org* (<https://hippocampome.org/>) [↗](#), we tracked the number of citations of the original publication (Wheeler et al., 2015 [↗](#)) and of the subsequent versions (**Figure 4B** [↗](#)), separating simple references from actual employment of information extracted from *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) for secondary analyses (**Table 3** [↗](#)). At the time of this writing, year 2021 proved to be the most prolific citation-wise; however, more than a third of the releases (v1.8-12) appeared after 2021 and most PoK were added in 2022, so usage could potentially accelerate further in coming years. An early application of *Hippocampome.org* (<https://hippocampome.org/>) [↗](#)-sourced data used subthreshold biophysical measures, such as input resistance and membrane time constant, for multicompartmental models of signal integration and extracellular field generation (Gulyas et al., 2016). That study concluded that somatic and proximal dendritic intracellular recordings in pyramidal cells and calretinin-positive interneurons, in particular, do not capture a sizable portion of the synaptic inputs. As a recent usage example, another lab employed *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) as the primary information resource for neuron types in DG, CA3, and CA1 (Schumm et al., 2022 [↗](#)). They discovered that mild traumatic brain injury, in the form of alterations in spike-timing-dependent plasticity, may affect the broadband power in CA3 and CA1 and the phase coherence between CA3 and CA1.

From experimental data to biologically realistic computational models

Several key neural properties collated into *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) have gradually transformed the site from an organized repository of hippocampal knowledge to a computational framework for launching real-scale neural network simulations. Specifically, building a data-driven circuit model of a neural system (such as the hippocampal formation or portion thereof) requires four essential quantities besides the full list of neuron types (Bahmer et al., 2023 [↗](#); DePasquale et al., 2023 [↗](#)): (i) the number of neurons in each type; (ii) the input-output response function for each neuron type; (iii) the connection probability for each pair of interacting neuron types; and (iv) the unitary synaptic signals for each pair of connected neuron types (**Figure 5** [↗](#)). Of those quantities, (i) and (ii) are neuron type properties, while (iii) and (iv) are properties of directional connections, defined as a distinct pair of a presynaptic and a postsynaptic neuron type. Moreover, (i) and (iii) are structural features, while (ii) and (iv) are electrophysiological ones.

Hippocampome.org (<https://hippocampome.org/>) [↗](#) v2.0 provides estimates of the number of neurons in each neuron type (i) for both rats and mice (**Figure 5A** [↗](#)). These values were derived in a two-step process (Attili et al., 2020 [↗](#)): first, literature mining extracted suitable quantitative relations such as the cellular density in a given layer (Attili et al., 2019 [↗](#)), the total count of neurons expressing a certain gene, or the fraction of sampled cells with a particular morphology; second, numerical optimization of the corresponding equations yielded a complete census for all neuron types. *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) represents the neuronal input-output response function (ii) in the form of single- and multi-compartment Izhikevich models (**Figure 5B** [↗](#)) fitted by evolutionary algorithms to accurately reproduce the observed firing behavior of each neuron type (Venkadesh et al., 2018 [↗](#)). The connection probability (iii) from one neuron type to another (**Figure 5C** [↗](#)) was computed from measurements of the appropriate axonal and dendritic lengths in each invaded subregion and layer (hippocampome.org/A-D_lengths). Additionally, users can also access the presynaptic and postsynaptic path distances from the respective somata (hippocampome.org/soma_distances) and the number of contacts per connected neuron pairs (hippocampome.org/num_contacts). As for the synaptic communication

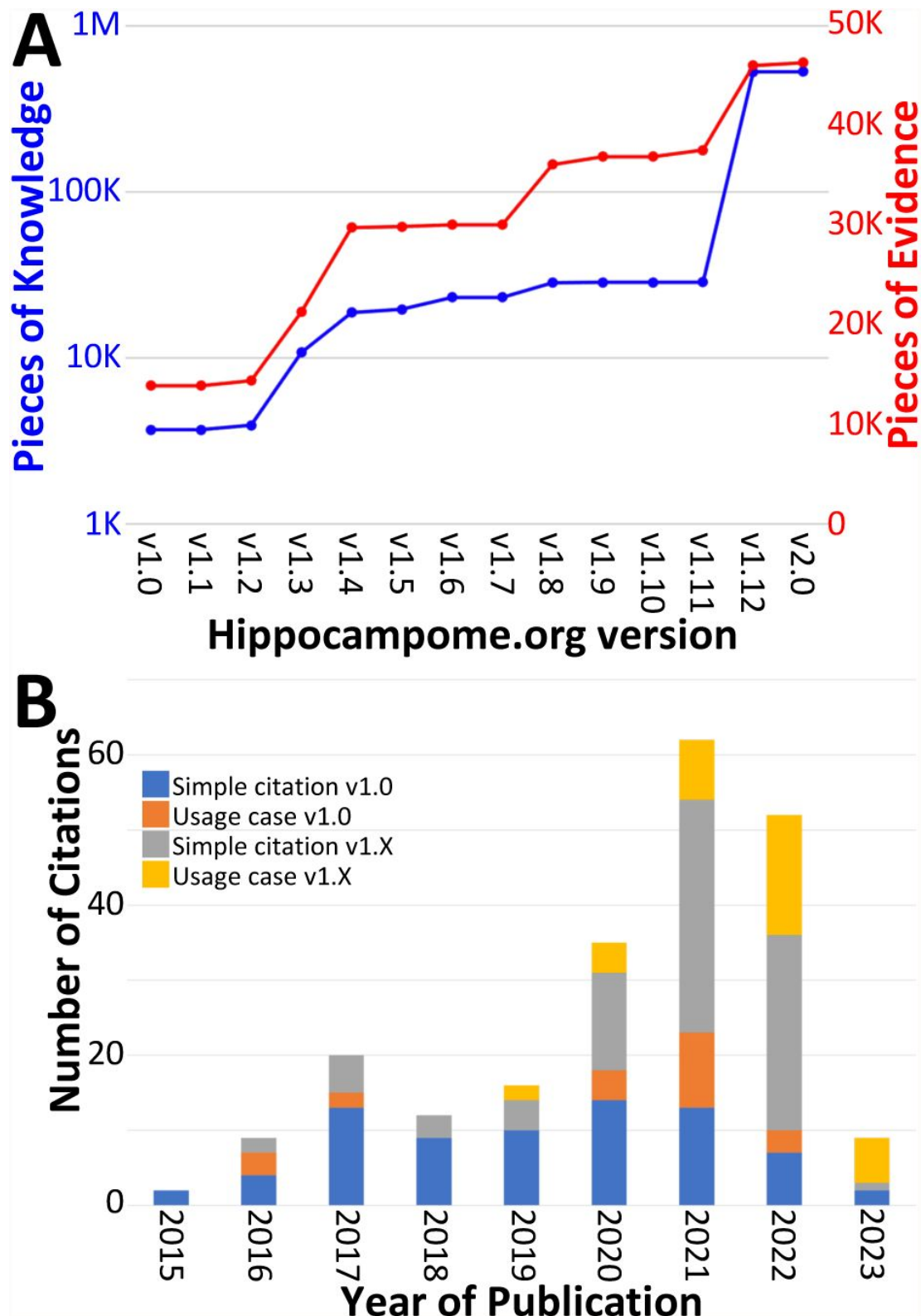


Figure 4.

Trends in *Hippocampome.org* (<https://hippocampome.org/>) data, knowledge, citations, and usage since v1.0. A) Increase in pieces of knowledge (blue) and evidence (red) with *Hippocampome.org* (<https://hippocampome.org/>) version number. B) Number of citations, in which the publication is simply referenced (blue and gray portions), and usage cases, in which the citing work makes use of the information contained within the *Hippocampome.org* (<https://hippocampome.org/>)-related work (orange and yellow portions), by year.

Article	Usage
(Gulyás et al., 2016)	Lists of subthreshold physiological properties for multicompartmental modeling
(Skene and Grant, 2016)	Catalog of CA1 Interneuron types
(Faghihi and Moustafa, 2017)	Diversity of hippocampal neuron types and morphological neuronal features
(Puighermanal et al., 2017)	Biomarker expression in CA1 interneurons
(Depannemaecker et al., 2020)	Parameter values for a model of synaptic neurotransmission
(Ecker et al., 2020)	Evidence that CA1 interneurons express multiple overlapping chemical markers
(Hunsberger and Mynlieff, 2020)	Cell identification based on firing properties
(Schumm et al., 2020)	Directionality of connections in the hippocampus
(Aery Jones et al., 2021)	Local connectivity of CA1 PV+ interneurons
(Ciarpella et al., 2021)	Lists of hippocampal genes
(Luo et al., 2021)	Confirmation of multiple hippocampal neuron types
(Mehta et al., 2021)	Connectome model inspired by entorhinal-CA1 circuit
(Obafemi et al., 2021)	Principal channels of information processing are DG Granule cells and CA1-3 Pyramidal cells
(Sáray et al., 2021)	Membrane biophysics values for CA1 Pyramidal cells
(Smith et al., 2021)	Omni-directionality of axons of CA1 Pyramidal cells
(Venkadesh and Van Horn, 2021)	Example of a brain region's mesoscopic structural connectivity
(Walker et al., 2021)	Reference to morphological and molecular characteristics of hippocampal principal cells and interneurons
(Wynne et al., 2021)	Example brain region with a variety of cell types
(Kopsick et al., 2022)	Utilize accumulated knowledge as the basis for simulations
(Schumm et al., 2022)	Hippocampal morphology, biomarker expression, connectivity, and typing of neurons
(Zagrean et al., 2022)	Diversity of hippocampal neuronal types and their properties

Table 3.

Examples of independent studies utilizing unique neuronal properties from *Hippocampome.org* (<https://hippocampome.org/>)  v1.0.

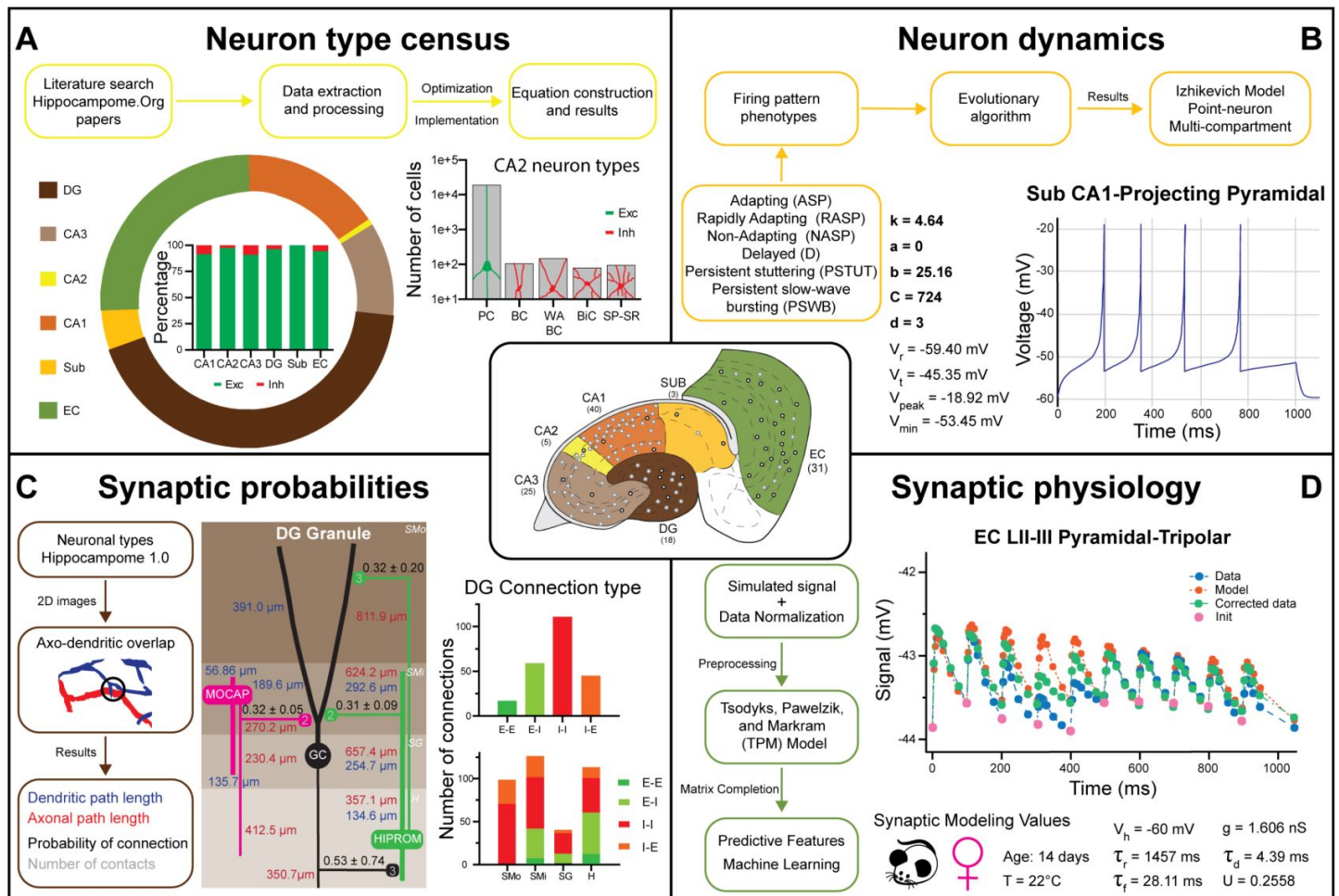


Figure 5.

Transitional knowledge enabling *Hippocampome.org* (<https://hippocampome.org/>) to support spiking neural network simulations. Center: General diagram of the hippocampal formation and the number of cell types in Hippocampome v1.0. A) Neuron type census. Top: General pipeline for obtaining cell counts for specific collections of neurons from the peer reviewed literature. Left: Neuron count proportions for the different subregions of the hippocampal formation. Insert: Normalized neuron counts for the inhibitory vs. excitatory balance by subregion. Right: Neuron counts for five identified CA2 neuron types (green schematic: excitatory, red schematic: inhibitory). B) Neuron dynamics. Top: General pipeline for obtaining Izhikevich models to reproduce the firing pattern phenotypes from peer reviewed data. Right: Simulated firing pattern from a Sub CA1-Projecting Pyramidal cell in response to a 250-pA current injection pulse lasting 1 s. Izhikevich model parameters are shown in bold and the membrane biophysics properties are shown in regular font. C) Synaptic probabilities. Left: General pipeline for obtaining the connection probabilities, number of contacts, and dendritic and axonal path lengths from 2D reconstructions. Middle: Example of a connectivity diagram of a DG Granule cell and two interneurons across the different parcels of DG. Probabilities of connection (mean ± SD) are shown in black, numbers of contacts in gray, dendritic path lengths in blue, and axonal lengths in red. Right top: Total number of connections within DG by connection type. Right bottom: Breakdown of the total number of connections by parcel and connection type. D) Synaptic physiology. Left: General pipeline for obtaining normalized synaptic parameters from paired recordings with a TPM model. Right top: Digitized synaptic data between two EC LII-III Pyramidal-Tripolar cells. Experimental data are shown in blue, initiation synaptic points in pink, model data in orange, and corrected data in green. Right bottom: Simulated modeling conditions, electrophysiological parameters, and TPM parameters. Abbreviations are defined in Materials and Methods.

between neurons (iv), *Hippocampome.org* (<https://hippocampome.org/>)  adopts the Tsodyks-Pawelzik-Markram formulation, representing unitary signals and short-term plasticity with five constants for each directional pair of interacting neuron types: the synaptic conductance, decay time, recovery time, facilitation time, and the utilization ratio (Tsodyks et al., 1998 ; Moradi et al., 2022 ). Once again, these parameters were fitted from the experimental data (Moradi and Ascoli, 2018 ) employing deep learning to account for (and predict the effects of) numerous experimental variables (**Figure 5D** ) , including species (rat vs. mouse), sex (male vs. female), age (young vs. adult), recording temperature (room vs. body), and clamping configuration (voltage vs. current).

The above description underscores the crucial interconnectedness of individually measured neuronal properties forming a cohesive whole in *Hippocampome.org* (<https://hippocampome.org/>)  (**Figure 6** ). In particular, normalized simulation parameters (e.g., the sensitivity of recovery variable in Izhikevich models) are derived from quantitative experimental measurements, such as the spiking adaptation rate (**Figure 6a** ). Those in turn are linked to an identified neuron type based on qualitative features, like calbindin expression or laminar distribution of axons and dendrites. In addition to enabling computational applications as described below, such integration also allows the meta-analysis of correlations between morphological features, molecular profiles, electrophysiological properties, and dynamic circuit functions. At the same time, several components of *Hippocampome.org* (<https://hippocampome.org/>)  are also synergistically linked to external community resources (**Figure 6B** ). For example, each neuron page links out to all three-dimensional morphological reconstructions of the same cell type available in NeuroMorpho.Org (Ascoli et al., 2007 ) , and selected data from NeuroMorpho.Org were used to compute axonal and dendritic length and connection probabilities. Each neuron page also links out to all computational models (including Hodgkin-Huxley, stochastic diffusion, mean firing rate, etc.) involving the same cell type on ModelDB (McDougal et al., 2017 ) , while conversely ModelDB includes the Izhikevich models for all *Hippocampome.org* (<https://hippocampome.org/>)  neuron types. Moreover, simulation parameters from *Hippocampome.org* (<https://hippocampome.org/>)  are exportable to the CARLsim simulation environment (Nageswaran et al., 2009 ) , enabling fast execution of spiking neural network models optimized for GPUs. Furthermore, *Hippocampome.org* (<https://hippocampome.org/>)  harnessed data from the Allen Brain Atlas (Lein et al., 2007 ) to infer gene expression for principal neurons and cell densities for use in the neuron type census.

To facilitate construction of spiking neural network simulations, *Hippocampome.org* (<https://hippocampome.org/>)  v2.0 also includes a new graphical user interface (GUI) . With this GUI, users can download sets of simulation parameter values for arbitrarily selected neuron types, a subregion of interest, or the whole hippocampal formation (**Figure 7** ). The sets consist of files for the instantiation of CARLsim simulations and a CSV spreadsheet of parameters for use in a different simulation environment of the user's choice. For the convenience of users interested in simplified circuit models, *Hippocampome.org* (<https://hippocampome.org/>)  informally ranks the importance of each neuron type from 1 (essential) to 5 (dispensable). For instance, a user may choose to simulate only the canonical, or rank 1, neuron types of the tri-synaptic circuit and entorhinal cortex, consisting of DG granule, CA3 pyramidal, CA1 Pyramidal, and MEC LII Stellate cells. When *Hippocampome.org* (<https://hippocampome.org/>)  is missing a parameter value due to insufficient experimental evidence, the GUI exports a default value clearly indicating so in the downloadable files. For missing Izhikevich and synaptic signaling parameters, the default values are those provided by the CARLsim simulation environment. For missing synaptic probabilities, *Hippocampome.org* (<https://hippocampome.org/>)  precomputes values averaged by connection type, namely excitatory-excitatory (0.0117), excitatory-inhibitory (0.0237), inhibitory-excitatory (0.00684), and inhibitory-inhibitory (0.00423).

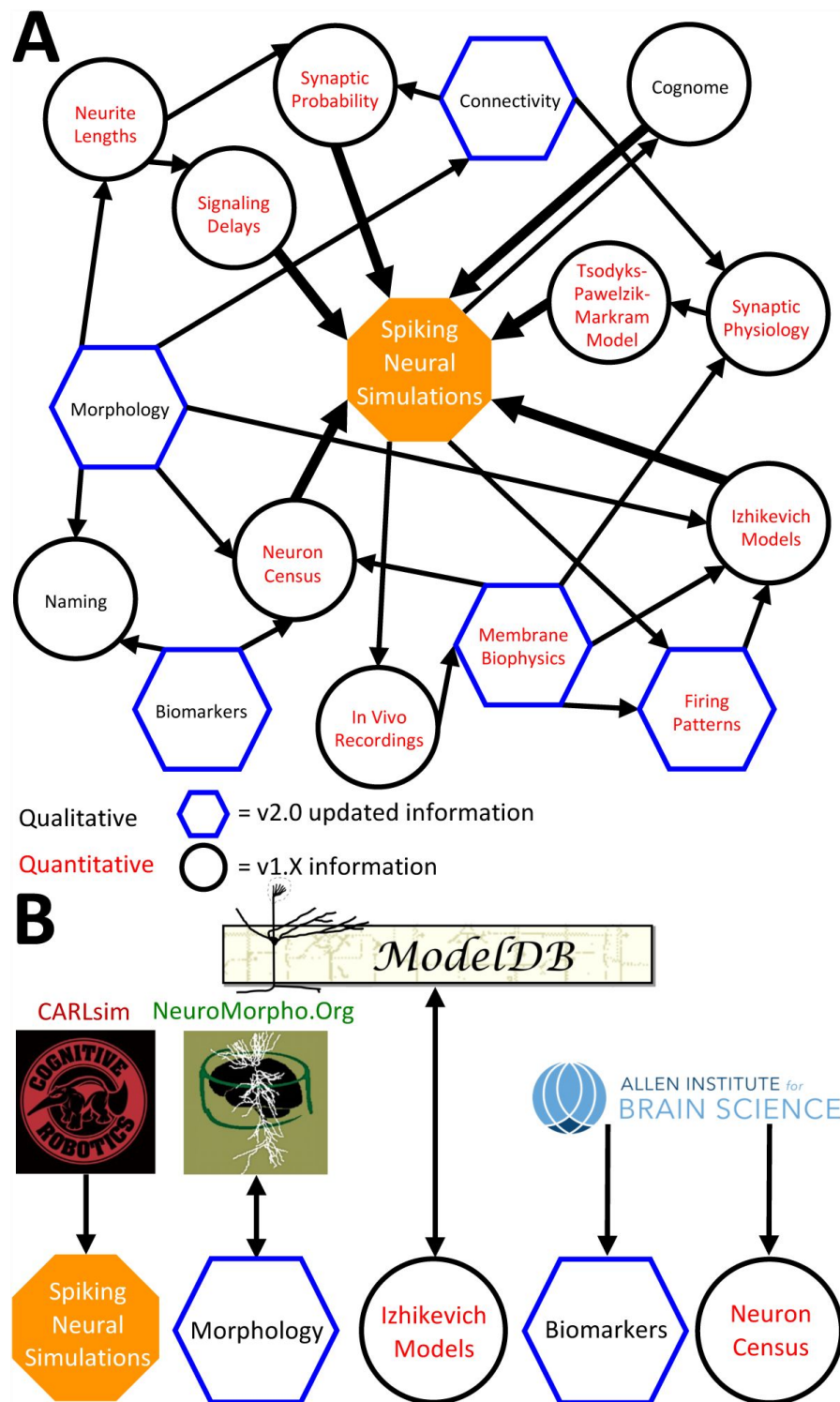


Figure 6.

Hippocampome.org (<https://hippocampome.org/>) data provenance. A) The internal web of constituent neuron-type properties (thin arrows) that ultimately contribute to the instantiation of spiking neural simulations (thick arrows). Properties described qualitatively, such as morphological presence of axons in a layer or molecular biomarker expressions, are in black font. Properties described by quantitative values, such as membrane biophysics and neurite lengths, are in red font. Properties with v2.0 updated information, such as connectivity and firing pattern phenotypes, are depicted by blue hexagons, and v1.x information, such as Izhikevich modeling parameter values and neuron-type census values, is visualized by black circles. B) External resources that contribute data to and receive data from *Hippocampome.org* (<https://hippocampome.org/>) (the ModelDB logo has been modified from the original).

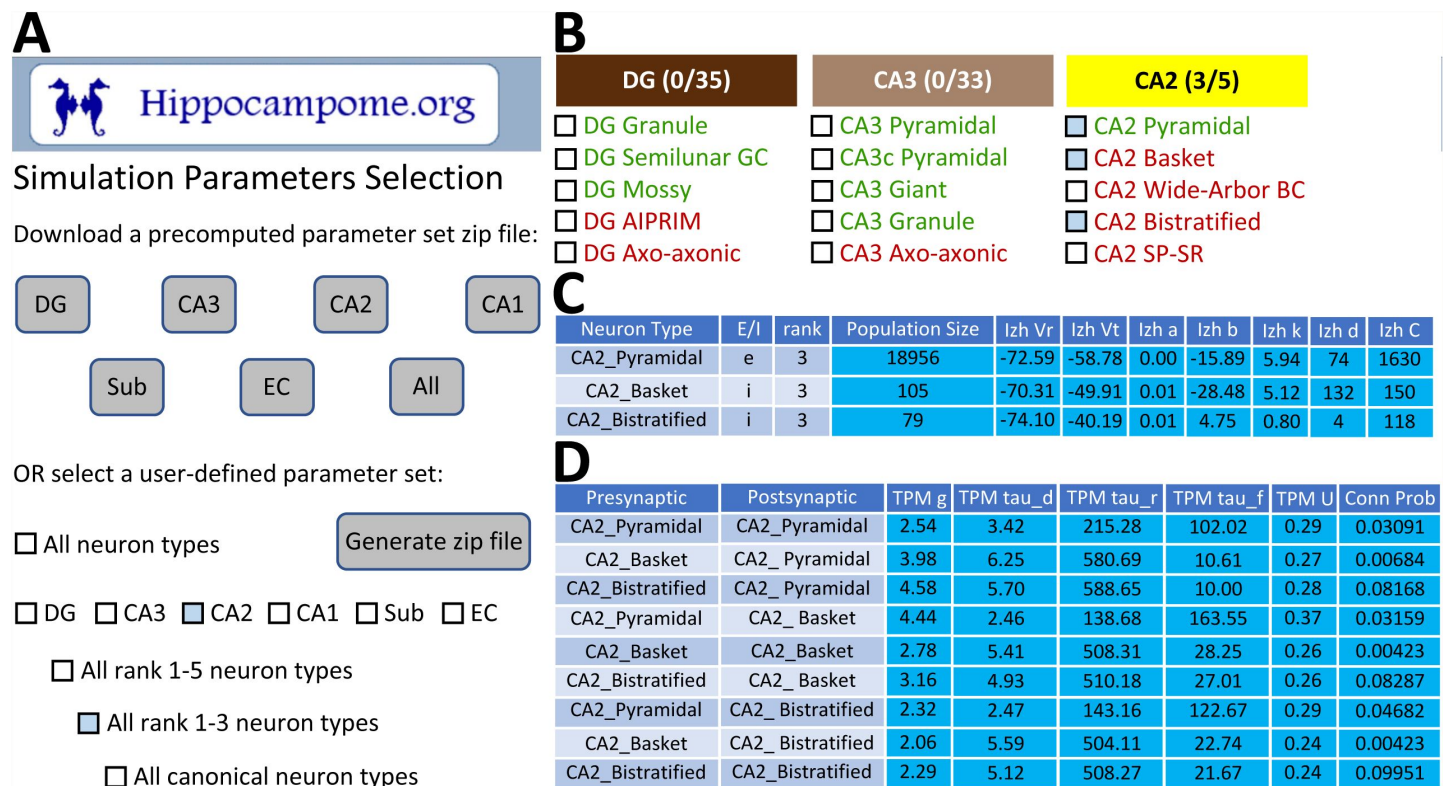


Figure 7.

CARLsim simulation parameters selection and file generation interface. A) The user chooses which subset of the available neuron types to include in the generated downloadable parameter file. Neuron types can be selected (check boxes and gray highlights) either individually or by groupings, such as by subregion and/or by importance rank. B) Representative user selection. C) Downloadable neuron-level parameters. D) Downloadable connection-level parameters.

Potential applications to connectomic analyses and spiking neural networks simulations

Hippocampome.org (<https://hippocampome.org/>)  v2.0 enables the multiscale analysis of circuit connectivity (**Figure 8** ). At the highest echelon are the connections between hippocampal subregions, which are comprised of the mesoscopic level potential connectivity between individual neuron types (**Figure 8A** ). Expanding, for example, upon the 147 connections between DG and CA3 neuron types reveals all the connections every individual neuron type forms with the other neuron types within and across the subregions (**Figure 8B** ). Zooming in onto a single neuron type each from DG and CA3, it is possible to quantify the efferent and afferent connections with other neuron types from throughout the hippocampal formation in terms of synaptic probabilities and number of neuronal partners (**Figure 8C** ). Diving even deeper into the isolated connection between two neuron types, such as the mossy fiber contacts from DG Granule cells to CA3 Basket cells, expands the connectivity analysis to several physiological factors affecting neuronal communication: the subcellular location of the synaptic contact (e.g., soma in stratum pyramidale and proximal dendrites in stratum lucidum), the transfer function (product of synaptic conductance and decay time constant), the in vivo firing rate of the presynaptic neuron type, and the relationship between input current and resulting output spiking frequency (F-I curve) of the post-synaptic neuron (**Figure 8D** ).

Release of v2.0 makes the original objective of *Hippocampome.org* (<https://hippocampome.org/>) , to enable data-driven spiking neural network simulations of rodent hippocampal circuits (Ascoli, 2010), finally achievable. An ongoing line of research in this regard focuses on a real-scale mouse model of CA3, with the eventual goal of investigating the cellular mechanisms of pattern completion. Initial work included excitatory Pyramidal cells and seven main inhibitory interneuron types: Axo-axonic, Basket, Basket CCK+, Bistratified, Ivy, Mossy Fiber-Associated ORDEN, and QuadD-LM (Kopsick et al., 2022 ). Use of *Hippocampome.org* (<https://hippocampome.org/>)  parameters for cell census, Izhikevich models, synaptic signals, and connection probabilities resulted in robust, realistic, rhythmic resting state activity for all neuron types (**Figure 9A** ). Another pursuit seeks to replicate the spatial representation in grid cells (Sargolini et al., 2006 ), modeled utilizing the *Hippocampome.org* (<https://hippocampome.org/>)  MEC LII Stellate cells and supported by various GABAergic interneuron types (Dhillon and Jones, 2000 ). This study aims to reproduce the in vivo firing of these neuron types as a virtual rodent explores an open field (**Figure 9B** ). While preliminary, these illustrative examples highlight the potential of *Hippocampome.org* (<https://hippocampome.org/>) -enabled data-driven spiking neural network simulations to investigate computational theories of cognitive functions in hippocampal circuits at the level of biologically detailed mechanisms (Sutton and Ascoli, 2021 ).

Discussion

Hippocampome.org (<https://hippocampome.org/>) , through its continuous updates and conspicuous usage, has established itself prominently amongst other readily accessible, evidence-based, expert-curated bioscience public resources of note, such as FlyBase for *Drosophila* molecular biology (The FlyBase Consortium, 1994; dos Santos et al., 2015), WormBase for nematode genomics (Stein et al., 2001 ), the Blue Brain Project for somatosensory cortex (Markram, 2006 ), SynGO for synaptic functions (Koopmans et al., 2019 ), and RegenBase for spinal cord injury biology (Callahan et al., 2016 ).

The growth of *Hippocampome.org* (<https://hippocampome.org/>)  since the initial release of v1.0 (Wheeler et al., 2015 ) has been prodigious. To date, the site has been visited over 136,000 times with over 33,000 unique visits, and the original publication has been cited more than hundred times. Each successive release of *Hippocampome.org* (<https://hippocampome.org/>)  has added new dimensions of knowledge and/or functionality and has been building toward assembling all the

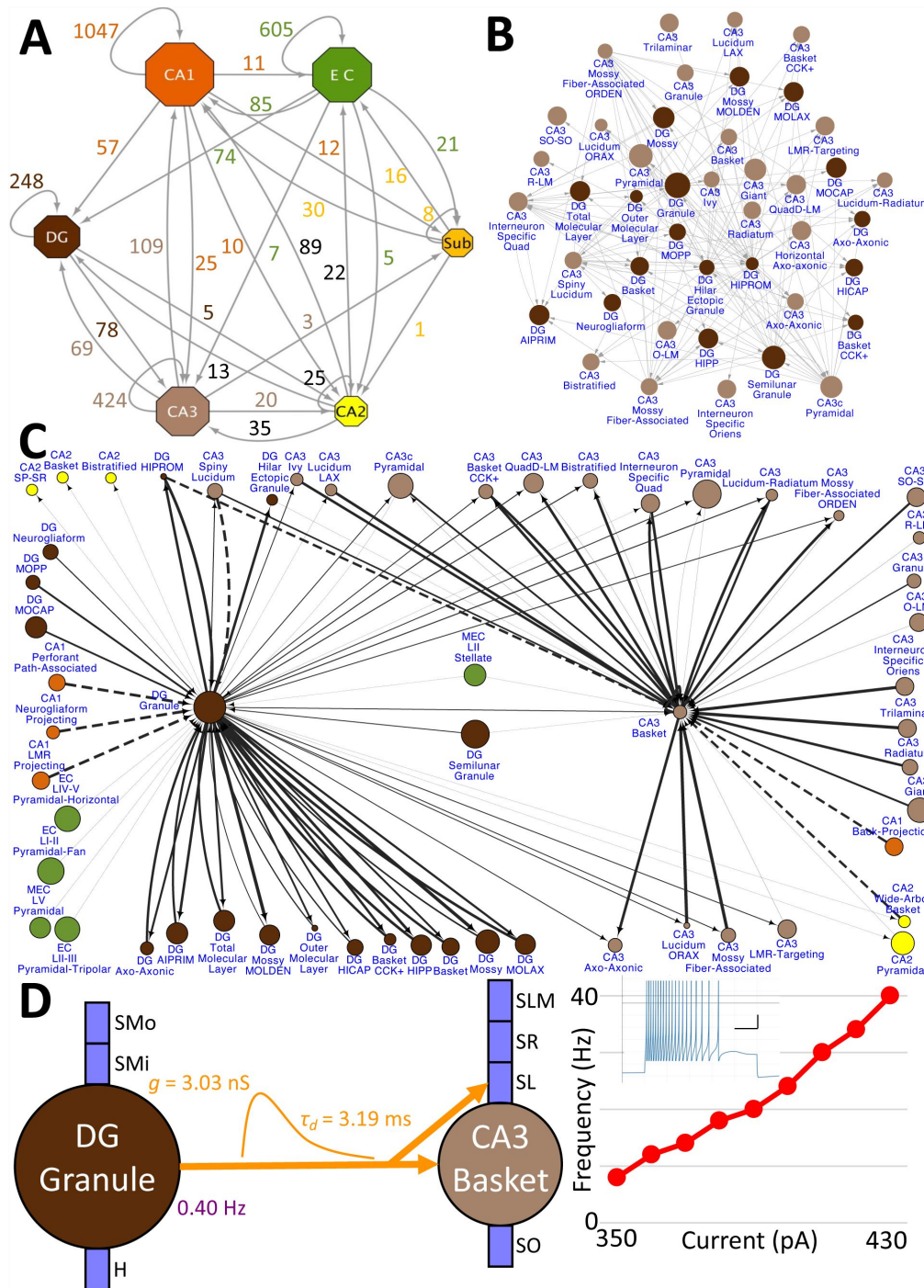


Figure 8.

Hierarchy of neuronal connectivity in *Hippocampome.org* (<https://hippocampome.org/>). **A**) Subregional connectivity, where the number of connections between subregions is shown, and the node size is proportional to the number of neuron types in each subregion. **B**) The reciprocal connectivity between DG and CA3 neuron types consists of 147 connections. The node size is proportional to the census size for each neuron type. **C**) The full connectivity involving DG Granule and CA3 Basket neuron types consists of 98 connections. The node size is proportional to the census size for each neuron type, and the thicknesses of the connecting arrows are proportional to the synaptic probability. The dashed lines are connections for which the synaptic probability has been approximated based on the means of known values. **D**) The electrophysiological connection between a DG Granule cell and a CA3 Basket cell. The in vivo firing rate is shown for the presynaptic neuron. The transfer function between the two neuron types is proportional to the synaptic conductance times the single-exponential decay time constant ($g \cdot \tau_d$; rat, male, P56, 32°C, current clamp). The frequency-current (F-I) curve of the single-compartment Izhikevich model of a CA3 Basket cell was obtained with 10 pA current steps. Inset: Izhikevich model firing pattern of a CA3 Basket cell simulated with 430 pA of current applied for 500 ms (vertical and horizontal scale bars, respectively).

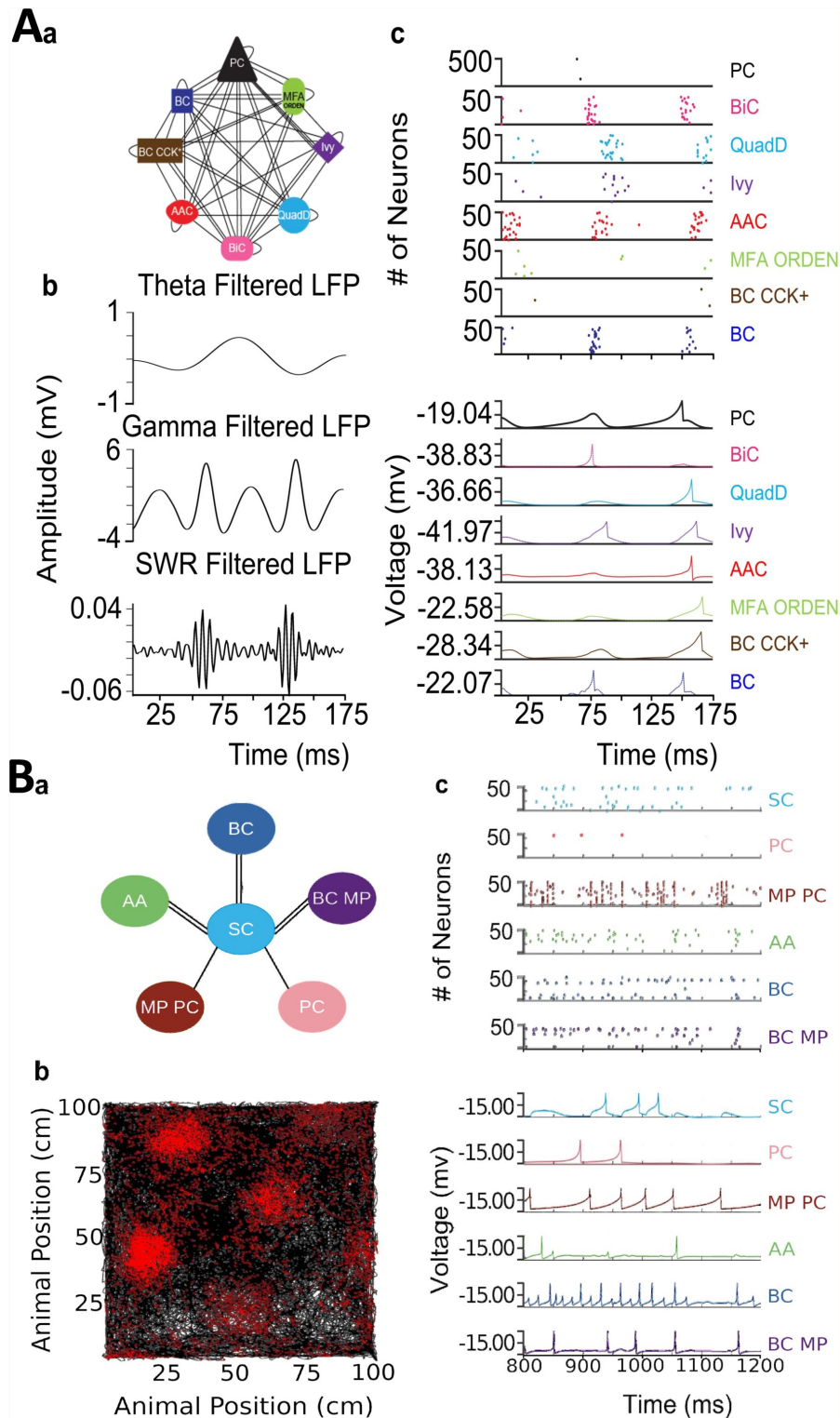


Figure 9.

Spiking neural network simulations. A) Full-scale CA3 model. (Aa) Neuron type connectivity schematic. (Ab) Theta (4-12 Hz; top), Gamma (25-100 Hz; middle), and Sharp-Wave Ripple (150-200 Hz; bottom) filtered local field potentials from 175 ms of the simulation. (Ac) Raster plot of 500 Pyramidal cells and 50 interneurons of each type (top), and representative voltage traces for each neuron type (bottom) during the same 175 ms of the simulation in (Ab). B) Spatial representation through grid cell firing. (Ba) Neuron type connectivity schematic. (Bb) Simulated animal trajectory (black) with red dots indicating the firing of a neuron in those locations. (Bc) Raster plot of 50 randomly selected neurons from each type (top), and representative voltage traces for each neuron type. Abbreviations are defined in Materials and Methods.

components necessary to produce real-scale computational models of the rodent hippocampal formation. The culmination of all this work is the release of v2.0, which introduces a framework for launching computer simulations directly from the accumulated knowledge. However, achieving simulations does not mark the end point for this project, because *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) will continue to aggregate new knowledge as it is published in the peer-reviewed literature. Gradually, the focus of this resource will shift from development to exploitation through the in silico emulation of complex dynamics observed in vivo and in vitro, with the goal of shedding light on the underlying synaptic-level computational mechanisms.

The creation of real-scale spiking neural network models of the hippocampal formation and its subregions can foster biologically realistic, data-driven, mesoscopic simulations of cognitive function and dysfunction (Sutton and Ascoli, 2021 [↗](#)). For instance, simulations with *Hippocampome.org* (<https://hippocampome.org/>) [↗](#)'s real-scale model of the dentate gyrus can build on previous network models of epileptogenesis (Dyhrfeld-Johnsen et al., 2007 [↗](#)) by providing further clarity to the roles of all documented neuron types and their corresponding potential connections in seizure initiation and propagation. A real-scale model of CA1 can aim to further the insights into the spatiotemporal dynamics of the circuit during theta oscillations (Bezaire et al., 2016 [↗](#); Navas-Olive et al., 2020 [↗](#); Romani et al., 2023 [↗](#)). Furthermore, network models involving multiple subregions can open new vistas on unexplored territories, such as the use of real-scale models of the entorhinal cortex and CA2 to simulate the neuron- and connection-type specific mechanisms of social memory (Lopez-Rojas et al., 2022 [↗](#)). Moreover, open source sharing of the real-scale models replicating those functions (Gleeson et al., 2017 [↗](#)) will facilitate cross-talk within the systems neuroscience community to better understand the role of distinct neuron types in hippocampal function.

A notable aspect of *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) is that all freely downloadable model parameters are directly linked to the specific peer-reviewed empirical evidence from which they were derived. Thus, if users disagree with a specific interpretation, or are not fully convinced by an individual experimental measurement, they maintain control in selecting the information sources. Conversely, researchers can choose to reuse the collated experimental data to constrain different computational models they may prefer, such as adopting the Hodgkin-Huxley formalism instead of Izhikevich dynamics. At the same time, *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) is not only a collection of model parameters and corresponding empirical evidence, but it also provides an opportunity to unearth knowledge gaps, as facilitated by an intuitive search functionality (hippocampome.org/find-neuron). Missing data can serve to guide the design of targeted “low hanging fruit” experiments or to generate new hypotheses.

Another important element of *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) is the careful annotation of the experimental metadata for each piece of evidence, including the species (rat or mouse), sex (male or female), age (young or adult) as well as any and all reported details that could affect the recorded neuronal property. Examples of these confounding factors abound especially for in vitro electrophysiological data, such as the exact chemical composition of the solution in the electrode and in the bath, slice thickness and orientation, clamping configuration, recording temperature, and animal weight. Because these covariates, when reported by the original investigators, are also stored in the database, it is possible to account for them in subsequent analyses and simulations. *Hippocampome.org* (<https://hippocampome.org/>) [↗](#) therefore constitutes a considerably rich one-stop resource to compare and “translate” key parameters, such as the amplitude and duration of a synaptic signal between two specifically identified neuron types, for instance, from 14 day old male rat at 22 °C in voltage clamp to a 56 day old female mouse at 32 °C in current clamp. When fed into spiking neural network simulations, these differential parameter values can foster intuition while attempting to reconcile neuroscience theories and observations.

Hippocampome.org (<https://hippocampome.org/>)  is yet poised for the onset of an information deluge from current and future big science projects, which will need to be integrated into a complete cohesive picture (de La Prida and Ascoli, 2021 ). Although morphological identification will continue to play a fundamental role in defining neuron types and circuit connectivity, the manner in which knowledge is cross-referenced in this resource will allow its effective linkage to rapidly accumulating molecular and imaging data. The ongoing spatial transcriptomics revolution is already transforming the frontiers of cellular neuroscience, often using the hippocampus as its favorite sandbox (Lein et al., 2017 ; Yao et al., 2021 ; Zeisel et al., 2015 ). Single-cell transcriptomics via scRNAseq can bolster the current morphological information by offering distinct transcription factor codes for existing neuron types and assist in defining new ones (Cembrowski and Spruston, 2019 ; Winnubst et al., 2020 ; Yuste et al., 2020 ). From the functional side, optical imaging via genetically encoded voltage indicators (Knöpfel and Song, 2019 ) will provide in vivo voltage traces for defined neuron types that can greatly enhance the repertoire of firing pattern phenotypes to utilize in simulations (Adam et al., 2019 ). Data-driven computational models can provide a useful conceptual bridge between molecular sequencing and activity imaging by investigating the effects of specific subcellular distributions of voltage- and ligand-gated conductances on neuronal excitability (Migliore et al., 2018 ). With the converging maturation of these young techniques and the advent of others yet on the horizon, *Hippocampome.org* (<https://hippocampome.org/>)  will be able to integrate multidimensional knowledge on the solid foundation of neuronal classification.

Materials and Methods

***Hippocampome.org* (<https://hippocampome.org/>) v2.0 vs. the legacy status of v1.12**

With the release of v2.0 of *Hippocampome.org* (<https://hippocampome.org/>)  upon publication of this article, v1.12 of the website will no longer be updated and will transition to legacy status (hippocampome.org/legacy_v1). In this way, users may avail themselves of the full benefits of the new content and functionality of v2.0, while maintaining access to reference content as published through v1.12. In the near term, neuron types new to v2.0 are tagged with an asterisk on the web site to differentiate them from v1.1x types.

Linking neuron types to NeuroMorpho.Org morphological reconstructions

Hippocampome.org (<https://hippocampome.org/>)  neuron types are regularly linked to appropriately identified digital reconstructions of neuronal morphology from NeuroMorpho.Org (Ascoli et al., 2007 ). Identification of suitable reconstructions with individual neuron types depends on the correspondence of dendritic and axonal locations across hippocampal subregions and layers, as they appear in the reference publication. Alternatively, direct cell typing by the authors in the reference publication text is accepted as evidence for canonical (principal cell) types, such as CA1 pyramidal cells or DG granule cells. Reconstructions are not linked to a neuron type if the experimental conditions are inconsistent with the inclusion criteria of *Hippocampome.org* (<https://hippocampome.org/>) , as in the case of cell cultures or embryonic development. Lack of either axonal or dendritic tracing also disqualifies reconstructions of non-canonical neurons from being linked.

Connections from DG Granule cells to CA3

To compute estimates of connection probabilities and numbers of contacts per connected pair for the rat mossy fiber-CA3 circuit, we used previously calculated average convex hull volume (Tecuatl et al., 2021b ) and several measurements from a seminal anatomical study (Acsády et al.,

1998): DG Granule cell axonal length within CA3 (3,236 μm), inter-bouton distances for mossy boutons on Pyramidal cell targets in CA3c (162 μm) and in the rest of CA3 (284 μm), and inter-bouton distances for en-passant and filipodia boutons onto CA3 interneurons (67.4 μm , considering that 48 interneurons can be contacted by a single GC). Given that the mossy fibers innervate mainly CA3 SL, and due to the lack of information regarding the exact proportion of axons innervating CA3 SP, these calculations assume that GCs only innervate SL. The probabilities of connection and numbers of contacts per connected pair (Table 2) are estimated as previously described (Tecuatl et al., 2021a) utilizing the CA3 dendritic lengths reported in *Hippocampome.org* (<https://hippocampome.org/>).

Connections from CA3 and CA3c Pyramidal cells to CA1

To compute estimates of connection probabilities and numbers of contacts per connected pair for the rat Schaffer collaterals-CA1 circuit, we utilized previously reported values for the distinct axonal innervation patterns (Ropiredy et al., 2011; Sik et al., 1993; Wittner et al., 2007) in CA1 SR and SO from CA3 Pyramidal cells (27.5% of total axonal length: 64% to SR, 15% to SP, 21% to SO) and CA3c Pyramidal cells (64.1% of total axonal length: 94% to SR, 3% to SP, 3% to SO). In addition, we used the average inter-bouton distance reported for the Schaffer collaterals (Li et al., 1994) in SR (4.47 μm) and SO (5.8 μm). Total axonal length was measured with L-Measure (Scorcioni et al., 2008) from three NeuroMorpho.Org reconstructions for CA3c (NMO_00187, NMO_00191) and CA3b (NMO_00931). We extracted parcel-specific convex hull volumes from Janelia MouseLight (Winnubst et al., 2019) Pyramidal cell reconstructions (AA0304, AA0307, AA0420, AA0960, AA0997, AA0999, AA1548) mapped to the 2022 version of the Allen Institute Common Coordinate Framework (CCF). The probabilities of connection and number of contacts per connected pair were estimated as previously described (Tecuatl et al., 2021a) using CA1 dendritic lengths from *Hippocampome.org* (<https://hippocampome.org/>). We used separate values for inter-bouton distances in CA1 SR for CA3c Pyramidal cells (5.5 μm : Wittner et al., 2007) and CA3 Pyramidal cells (3.7 μm : Shepherd et al., 2002; 4.4 μm : Li et al., 1994; 4.29 μm : Sik et al., 1993; averaged as 4.1 μm).

Constructing *Hippocampome.org* (<https://hippocampome.org/>) spiking neural simulations

Hippocampome.org (<https://hippocampome.org/>) utilizes CARLsim (Nageswaran et al., 2009) as its default simulation environment (socsci.uci.edu/~jkrichtma/CARLsim/). CARLsim is a graphics processing unit (GPU)-accelerated library of functions for simulating spiking neural networks based on Izhikevich neuron models (Izhikevich, 2003). The current version is CARLsim 6 (Niedermeier et al., 2022), and the most up-to-date *Hippocampome.org* (<https://hippocampome.org/>)-optimized code base, including features not yet released in the main CARLsim version, can be found at hippocampome.org/CARLsim (Kopsick et al., 2022).

Web portal, database, and source code

Hippocampome.org (<https://hippocampome.org/>) runs on current versions of Chrome, Safari, and Edge web browsers, and it is deployed on a CentOS server running Apache. The website runs off of PHP from a MySQL database. The code for *Hippocampome.org* (<https://hippocampome.org/>) is available open source at github.com/Hippocampome-Org. This includes all code for displaying the pages of the website, all scripts for importing spreadsheets into the database, code for using evolutionary algorithms to optimize Izhikevich model parameters, code for the graph theory analysis of the potential connectome, code for the implementation of the firing pattern classification algorithm, and code for analyzing network simulations in CARLsim.

Glossary of abbreviations

- 5HT-3
serotonin receptor 3
- AA
EC LII Axo-axonic cell
- AAC
Axo-axonic cell
- ABA
Allen Brain Atlas
- AP_{ampl}
action potential amplitude
- AP_{width}
action potential width
- ASP.
adapting spiking
- Astn2
astrotactin 2
- BC
Basket cell
- BC MP
EC LII Basket-Multipolar cell
- BiC
Bistratified cell
- BICCN
BRAIN Initiative Cell Census Network
- C
Capacitance

- CB
calbindin
- CB1
cannabinoid receptor type 1
- CCK+
cholecystokinin-positive
- CR
calretinin
- CSV
comma-separated value
- Dcn
decorin
- DG
dentate gyrus
- E-E
excitatory to excitatory
- E-I
excitatory to inhibitory
- EC
entorhinal cortex
- ENK
enkephalin
- fAHP
fast after-hyperpolarizing potential
- g
conductance constant
- GABA α 1

GABA-a alpha 1 subunit

- GC

granule cell

- Gpc3

glypican 3

- GPUs

graphical processing units

- GRALDEN

GRAnule-Like DENdrites Garci grid cell

- Grp

gastrin releasing peptide

- GUI

graphical user interface

- H

hilus

- HIPROM

Hilar Interneuron with PROjections to the Outer Molecular layer

- Htr2c

5-hydroxytryptamine receptor 2c

- I-E

inhibitory to excitatory

- I-I

inhibitory to inhibitory

- IS

interneuron specific

- LI-II

layers 1-2

layer 2

- LIII

layer 3

- Max FR

maximum firing rate

- MEC

medial entorhinal cortex

- MFA

Mossy Fiber-Associated

- mGluR1a

metabotropic glutamate receptor 1 alpha

- MOCAP

cell with MOlecular Commissural-Associational Pathway-related axons and dendrites

- MOLAX

MOlecular Layer Axons

- MP PC

EC LI-II Multipolar Pyramidal cell

- Mus2R

muscarinic type 2 receptor

- NASP

non-adapting spiking

- Ndst4

N-deacetylase and N-sulfotransferase 4

- nNOS

neuronal nitric oxide synthase

- Nov

nephroblastoma overexpressed

neuropeptide Y

- Nr3c2

nuclear receptor subfamily 3 group C member 2

- Nr4a1

nuclear receptor subfamily 4 group A member 1

- ORAX

ORiens AXons

- ORDEN

ORiens DENdrites

- PC

pyramidal cell / CA1 Pyramidal cell (**Fig. 9B** [↗](#))

- PL

polymorphic layer

- PoE

pieces of evidence

- PoK

pieces of knowledge

- Prss12

serine protease 12

- Prss23

serine protease 23

- PV+

parvalbumin-positive

- QuadD-LM

quadrilaminar dendrites – lacunosum-moleculare

- R_{in}

input resistance

reelin

- sAHP

slow after-hyperpolarizing potential

- SC

MEC LII Stellate cell

- SD

standard deviation

- SG

stratum granulosum

- SL

stratum lucidum

- SLM

stratum lacunosum-moleculare

- SMi

inner stratum moleculare

- SMO

outer stratum moleculare

- SO

stratum oriens

- SOM

somatostatin

- SP

stratum pyramidale

- SR

stratum radiatum

- Sub

subiculum

sharp-wave ripple

- SynGO

synaptic gene ontologies

- T

Temperature

- τ_d

synaptic decay constant

- τ_f

facilitation time constant

- τ_m

membrane time constant

- τ_r

recovery time constant

- TPM

Tsodyks-Pawelzik-Markram

- TSTUT.

transient stuttering

- TSTUT.PSTUT

transient stuttering followed by persistent stuttering

- TSTUT.SLN

silence preceded by transient stuttering

- U

utilization ratio

- vGluT3

vesicular glutamate transporter 3

- V_h

holding potential

vasoactive intestinal polypeptide

- $V_{\min}$

post-spike reset potential

- V_{peak}

spike cutoff potential

- V_r / V_{rest}

resting membrane potential

- V_t / V_{thresh}

firing threshold potential

- WA BC

Wide-arbor Basket cell

- Wfs1

wolframin ER transmembrane glycoprotein

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Competing Interests

The authors have no competing interests.

Contributions

DWW contributed to the conceptualization, methodology, software, investigation, data curation, writing – original draft, visualization, supervision, and project administration. JDK contributed to the methodology, software, formal analysis, investigation, data curation, writing – review and editing, and visualization. NS contributed to the methodology, software, investigation, data curation, writing – review and editing, and visualization. CT contributed to the methodology, validation, formal analysis, investigation, data curation, writing – review and editing, visualization, and supervision. AOK contributed to the formal analysis, writing – review and

editing, and visualization. KN contributed to the software and writing – review and editing. GAA contributed to the conceptualization, methodology, resources, investigation, data curation, writing – review and editing, visualization, supervision, project administration, and funding acquisition.

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Reviewer #1 (Public Review):

Summary:

The authors made significant updates to Hippacampome.org including 50 new cell types.

Strengths:

The authors have been thorough in basing their views on peer-reviewed literature. They have made the data highly accessible and the user has the ability to control what is included.

Weaknesses:

There are many inconsistencies in the literature regarding cell types and how these are incorporated into hippocampome.org is not clear.

Properties are often a result of modeling and not biological data, and caveats to this approach, and other assumptions are unclear.

Several interneuron subtypes in the dentate gyrus do not appear to be listed, such as neurogliaform cells.

The nomenclature HIPROM should be distinguished or made synonymous with HIPP. Same for MOCAP and MOPP/HICAP.

Dorsal ventral and sex differences are not mentioned.

Reviewer #2 (Public Review):

Summary and strengths:

The authors have developed a helpful resource for the community regarding hippocampal cell types and their interactions from many perspectives. There have been many updates to hippocampome v1.0 to v1.12, that are nicely summarized and explained (e.g., Table 1). The content and impact are also presented (Fig. 4).

Weaknesses:

My main comment is that it is not completely clear and/or it is a bit buried as to what makes this v2.0 (rather than v1.13). The title would seem to encompass it ('... enabling data-driven spiking neural network simulations...'), but in the introduction, the authors seem to emphasize "50 newly identified neuron types...". Is it the case that launching network simulations (using CARLsim) was not possible up to v1.12? I don't think so? I think that this research advance is to announce and summarize the various updates and to demonstrate how network simulations can be easily done? If so, this should and could be made more clear so that the reader does not necessarily have to go through all the previous versions to understand what is 'special' or different about v2.0. This could perhaps be achieved by situating their tool and its goals relative to other efforts (e.g., blue brain project) that are mentioned in the Discussion?

Reviewer #3 (Public Review):

Summary:

The authors aim to provide a multidisciplinary resource on the structural and physiological organization of the hippocampal system and make the available experimental data available for further theoretical work, providing tools to do so in a very flexible and user-friendly way. Since this is a new version of an already existing data-resource, the authors certainly reach their aim and fulfil expectations that the reader might have. The content of the database is as good as the original data, collected from the published knowledge-database, sometimes with the help of the original authors, and the overall quality depends further on how the data are curated by the team of authors and many others who helped them. That process is briefly described and more details are available in descriptions of previous versions and on the website. The data extraction, examples of how data can be used, and the part on attempts to model the hippocampus are exciting and open doors to new and exciting research opportunities.

Strengths:

Excellent description with many outlined opportunities. Nicely illustrated and inviting to explore the online database.

Weaknesses:

The figures are complex, containing a heavy information load with many abbreviations. You need some general knowledge of the system in order to grasp the enormous potential of what is provided.

Author Response:

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Strengths: The authors have been thorough in basing their views on peer-reviewed literature. They have made the data highly accessible and the user has the ability to control what is included.

Weaknesses: There are many inconsistencies in the literature regarding cell types and how these are incorporated into hippocampome.org is not clear.

We agree with the Reviewer that there can be inconsistencies in the literature, especially when it comes to nomenclature. This is why for Hippocampome.org v1.0 we decided to focus on the morphologies, the distributions of axons and dendrites across the layers and parcels of the hippocampal formation, rather than the names authors have applied to the neurons they are studying. We have also clarified our stance on nomenclature in our Brain Informatics manuscript that accompanied v1.1. We will revise the manuscript to make these points explicit.

Properties are often a result of modeling and not biological data, and caveats to this approach, and other assumptions are unclear.

The foundation for Hippocampome.org has always been the data that are published in the literature. Those include, among others, the axonal and dendritic spans in each layer and subregion, the molecular expression patterns, the total neuron count by layer and subregion, the membrane properties, firing patterns, and experimental synaptic signals and corresponding covariates. For all of those, we do not depend on how the data are modeled, although there is always some level of interpretation of the data to make them machine readable and ready for incorporation into our database. However, some of the simulation-ready parameters now also included in Hippocampome.org are indeed the result of modeling, such as the neuronal input/output functions (Izhikevich model) and the unitary synaptic values (Tsodyks-Markram model). Other simulation-ready parameters are the result of specific analysis approaches, including the connection probabilities (axonal-dendritic spatial overlaps) and the neuron type census (numerical optimization of all constraints). We plan to explicitly distinguish among these various cases in the revised manuscript.

Several interneuron subtypes in the dentate gyrus do not appear to be listed, such as neurogliaform cells.

The neuron types listed in Figure 2 of the current manuscript are only the new additions to the catalog of neuron types at Hippocampome.org v2.0. DG Neurogliaform cells were included in our original eLife manuscript, which described the deployment of v1.0 of the website. We will clarify this in the revisions.

The nomenclature HIPROM should be distinguished or made synonymous with HIP. Same for MOCAP and MOPP/HICAP.

The Reviewer has referred to 5 separate neuron types in Hippocampome.org. Each neuron type has a unique distribution of axonal and dendritic invasions of the 26 layers and parcels of the hippocampal formation. For example, HIPROM cells have dendrites in the inner one-third of stratum moleculare, stratum granulosum, and hilus and axons in all four layers of the dentate gyrus in addition to axonal projections into CA3 stratum radiatum, stratum lucidum, stratum pyramidale, and stratum oriens. HIP cells in contrast have dendrites only in the hilus and axons only in the outer two-thirds of stratum moleculare with no cross-

subregional projections. Similar considerations distinguish MOPP, MOCAP, and HICAP cells in Hippocampome.org. In expanding the nomenclature to include the neuron types we first described at Hippocampome.org, we attempted to mimic the styling of the already established neuron types of the DG: HIPROM (Hilar Interneuron with PROjections to the Outer Molecular layer), HIP (Hilar Perforant Path-associated), MOCAP (MOlecular Commissural-Associational Pathway-related axons and dendrites), MOPP (MOlecular layer Perforant Path-associated), and HICAP (Hilar Commissural-Associational Pathway-related). We intend to insert a paragraph in the revised version to clarify these issues.

Dorsal ventral and sex differences are not mentioned.

We thank the Reviewer for pointing this out. As a result of the dearth of literature describing differences between dorsal and ventral hippocampus when we first assembled Hippocampome.org v1.0, we made the decision to focus solely on the distributions of the axons and dendrites along the depth, or layers, of the hippocampal formation. As the amount of literature concerning relating to the other axes of the hippocampus continues to grow, we will gradually incorporate information along the added dimensions into our knowledge base. In the revised manuscript we intend to note this, and also stress the fact that Hippocampome.org contains knowledge from a mixture of sexes, and that whenever the original papers report the animal sex, so does our knowledge base. The revised manuscript will also mention that, whenever possible (e.g. synaptic physiology parameters), values are reported separately for males and females.

Reviewer #2 (Public Review):

Summary and strengths: The authors have developed a helpful resource for the community regarding hippocampal cell types and their interactions from many perspectives. There have been many updates to hippocampome v1.0 to v1.12, that are nicely summarized and explained (e.g., Table 1). The content and impact are also presented (Fig. 4).

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We thank the Reviewer for their helpful suggestions. Hippocampome.org v1.12 included the final piece needed, the synaptic physiology parameter values, to start fully simulating the hippocampal formation. In the revised manuscript, we will endeavor to emphasize more the specialness of v2.0 over the various v1.X in the Abstract, Introduction, and Discussion, in part by more fully describing the differences between our work and that of other efforts, such as the Blue Brain Project.

Reviewer #3 (Public Review):

Summary: The authors aim to provide a multidisciplinary resource on the structural and physiological organization of the hippocampal system and make the available

experimental data available for further theoretical work, providing tools to do so in a very flexible and user-friendly way. Since this is a new version of an already existing data-resource, the authors certainly reach their aim and fulfil expectations that the reader might have. The content of the database is as good as the original data, collected from the published knowledge-database, sometimes with the help of the original authors, and the overall quality depends further on how the data are curated by the team of authors and many others who helped them. That process is briefly described and more details are available in descriptions of previous versions and on the website. The data extraction, examples of how data can be used, and the part on attempts to model the hippocampus are exciting and open doors to new and exciting research opportunities.

Strengths: Excellent description with many outlined opportunities. Nicely illustrated and inviting to explore the online database.

Weaknesses: The figures are complex, containing a heavy information load with many abbreviations. You need some general knowledge of the system in order to grasp the enormous potential of what is provided.

We agree with the Reviewer that we generously used abbreviations throughout our figures as a means of conserving limited space. We have attempted to balance that by providing a complete glossary of all the abbreviations used throughout the manuscript. However, we will make an effort to supply definitions of the abbreviations in the figure captions and at their first use in the manuscript, or even replacing the abbreviations altogether in key places in the figures.