

# Traces of semantization - from episodic to semantic memory in a spiking cortical network model

Nikolaos Chrysanthidis<sup>1</sup> · Florian Fiebig<sup>1</sup> · Anders Lansner<sup>1,2</sup> · Paweł Herman<sup>3</sup>

**1 Abstract** Episodic memory is the recollection of past personal experiences associated with particular times and places. This kind of memory is commonly subject to loss of contextual information or "semantization", which gradually decouples the encoded memory items from their associated contexts while transforming them into semantic or gist-like representations. Novel extensions to the classical Remember/Know behavioral paradigm attribute the loss of episodicity to multiple exposures of an item in different contexts. Despite recent advancements explaining semantization at a behavioral level, the underlying neural mechanisms remain poorly understood. In this study, we suggest and evaluate a novel hypothesis proposing that Bayesian-Hebbian synaptic plasticity mechanisms might cause semantization of episodic memory. We implement a cortical spiking neural network model with a Bayesian-Hebbian learning rule called Bayesian Confidence Propagation Neural Network (BCPNN), which captures the semantization phenomenon and offers a mechanistic explanation for it. Encoding items across multiple contexts leads to item-context decoupling akin to semantization. We compare BCPNN plasticity with the more commonly used spike-timing dependent plasticity (STDP) learning rule in the same episodic memory task. Unlike BCPNN, STDP does not explain the decontextualization process. We also examine how selective plasticity modulation of isolated salient events may enhance preferential retention and re-

sistance to semantization. Our model reproduces important features of episodicity on behavioral timescales under various biological constraints whilst also offering a novel neural and synaptic explanation for semantization, thereby casting new light on the interplay between episodic and semantic memory processes.

**Keywords** Episodic memory · Semantic memory · Semantization · Decontextualization · Bayesian-Hebbian plasticity · BCPNN · STDP · Spiking cortical memory model

## 1 INTRODUCTION

Remembering single episodes is a fundamental attribute of human cognition. A memory, such as with whom you celebrated your last birthday, is more vividly recreated when we can recall contextual information, such as the location of the event (Eichenbaum et al., 2007; Gillund, 2012). The term "episodic memory" was originally introduced by Tulving (1972) to designate memories of personal experiences. Retrieval from episodic memory includes a feeling of mental time travel realized by "I remember". In contrast, semantic memory retrieval encapsulates what is best described by "I know" (Tulving, 1985; Umanath and Coane, 2020). Unlike episodic memories, semantic memories refer to general knowledge about words, items and concepts, lacking spatiotemporal source information, possibly resulting from the accumulation of episodic memories (Schendan, 2012; Gillund, 2012).

Initially, Tulving (1972) proposed that episodic and semantic memory are distinct systems and compete in retrieval. Recent studies suggest, however, that the division between episodic and semantic memory is rather vague (McCloskey and Santee, 1981; Renault et al., 2019), as neural activity reveals interaction between episodic and semantic systems during retrieval (Weidemann et al., 2019). According to Squire and Zola (1998) retrieval of semantic memory depends on the acquisition of the episode in which such information was experienced. Apparently, there is a clear interdependence between the two systems as the content of episodic memory invariably involves semantic representations (Martin-Ordas

Nikolaos Chrysanthidis   
E-mail: nchr@kth.se

Florian Fiebig   
E-mail: fiebig@kth.se

Anders Lansner   
E-mail: ala@kth.se

Paweł Herman   
E-mail: paherman@kth.se

<sup>1</sup> School of Electrical Engineering and Computer Science, KTH Royal Institute of Technology, 10044 Stockholm, Sweden

<sup>2</sup> Department of Mathematics, Stockholm University, 10691 Stockholm, Sweden

<sup>3</sup> School of Electrical Engineering and Computer Science, and Digital Futures, KTH Royal Institute of Technology, 10044 Stockholm, Sweden

70 et al., 2014), and consequently semantic similarity aids  
71 episodic retrieval (Howard and Kahana, 2002).

72 Episodic memory traces are susceptible to transformation  
73 and loss of information (Tulving, 1972), and this loss of episodicity can be attributed to semantization,  
74 which typically takes the form of a decontextualization  
75 process (Duff et al., 2020; Habermas et al., 2013;  
76 Viard et al., 2007). Meeter and Murre (2004) highlight  
77 and review the dynamical nature of memories and neural  
78 interactions through the scope of Transformation theory,  
79 which suggests that all memories start as episodic  
80 representations that gradually transform into semantic  
81 or gist-like representations (Winocur and Moscovitch,  
82 2011; Petrican et al., 2010). Decontextualization can occur  
83 over time as studies suggest that older adults report  
84 fewer episodic elements than younger adults (Petrican  
85 et al., 2010). Yet, could this item-context decoupling rely  
86 on accumulation of episodicity over multiple exposures  
87 of stimuli in various contexts over time? Baddeley (1988)  
88 hypothesized that semantic memory might represent the  
89 accumulated residue of multiple learning episodes, con-  
90 sisting of information which has been semanticized and  
91 detached from the associated episodic contextual detail.  
92 In fact, simple language vocabulary learning implies  
93 that learners encode words in several different contexts,  
94 which leads to semantization and definition-like knowl-  
95 edge of the studied word (Beheydt, 1987; Bolger et al.,  
96 2008).

97 Retrieval from episodic memory has been studied  
98 extensively through the lens of the classical Remember/Know  
99 (R/K) paradigm, in which participants are  
100 required to provide a Know or Remember response  
101 to stimulus-cues, judging whether they are able to re-  
102 call item-only information or additional details about  
103 episodic context, respectively (van den Bos et al., 2020).  
104 Extensions of the classical R/K behavioral experiment  
105 demonstrate that item-context decoupling can occur  
106 rapidly (Opitz, 2010). In these experiments, items are  
107 presented during an encoding phase either in a unique  
108 context, or across several contexts. Low context variabil-  
109 ity leads to greater recollection, whereas context overload  
110 results in decontextualization and a higher fraction of  
111 correctly classified Know responses (Opitz, 2010; Smith  
112 and Manzano, 2010; Smith and Handy, 2014). In the cur-  
113 rent study, we offer and evaluate a Bayesian-hypothesis  
114 about synaptic and network mechanisms underlying the  
115 memory semantization (item-context decoupling).

116 In earlier works, we developed and investigated a  
117 modular spiking neural network model of cortical asso-  
118 ciative memory with respect to memory recall, includ-  
119 ing oscillatory dynamics in multiple frequency bands,  
120 and compared it to experimental data (Lundqvist et al.,  
121 2010, 2011; Herman et al., 2013). Recently we demon-  
122 strated that the same model, enhanced with a Bayesian-  
123 Hebbian learning rule (Bayesian Confidence Propagation  
124 Neural Network, BCPNN) to model synaptic and in-  
125 trinsic plasticity, was able to quantitatively reproduce key  
126 behavioral observations from human word-list learning  
127 experiments (Fiebig and Lansner, 2017), such as serial  
128 order effects during immediate recall. This model per-

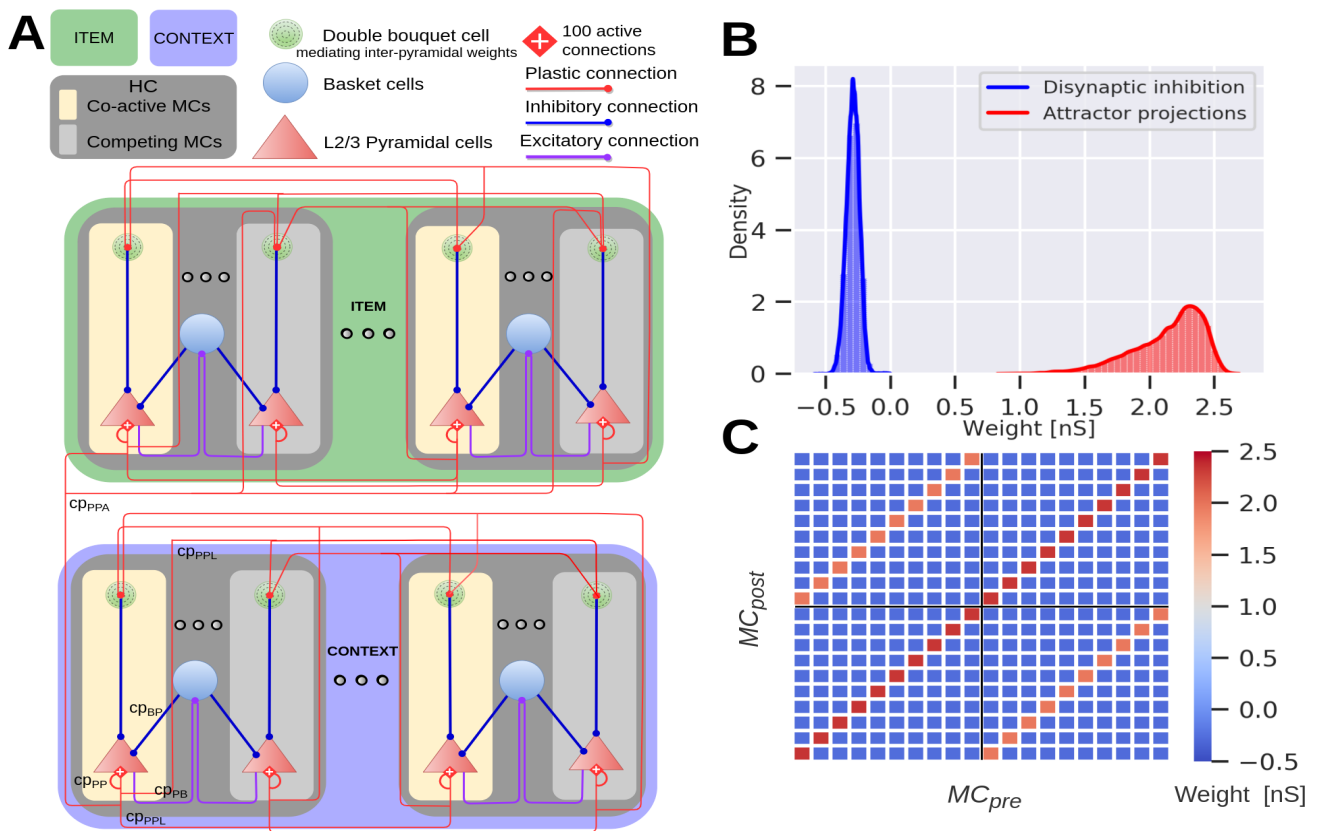
129 formed one-shot memory encoding and was further ex-  
130 panded into a two-area cortical model used to explore  
131 a novel indexing theory of working memory, based on  
132 fast Hebbian synaptic plasticity (Fiebig et al., 2020). In  
133 this context, it was suggested that the underlying naive  
134 Bayes view of association would make the associative  
135 binding between two items weaker if one of them is later  
136 associated with additional items. Thus, if we conceive of  
137 episodicity as an associative binding between item and  
138 context, the BCPNN synaptic plasticity update rule might  
139 provide a mechanism for semantization. In this work, we  
140 test this hypothesis and examine to what extent the re-  
141 sults match available behavioral data on semantization.  
142 We further compare those outcomes of dynamic learn-  
143 ing with a model featuring the more well-known spike-  
144 timing dependent plasticity (STDP) learning rule. We  
145 also demonstrate how selective plasticity modulations of  
146 one-shot learning (tentatively modelling effects of atten-  
147 tion, emotional salience, valence, surprise, etc. on plas-  
148 ticity) may enhance episodicity and counteract semanti-  
149 zation.

150 To our knowledge, there are no previous computa-  
151 tional models of item-context decoupling akin to seman-  
152 tization. Overall, there are rather few computational mod-  
153 els of episodic memory (Norman and O'Reilly, 2003), and those that exist are typically abstract, aimed at pre-  
154 dicting behavioral outcomes without a specific focus on  
155 underlying neural and synaptic mechanisms (Greve et al.,  
156 2010; Wixted, 2007). Our model bridges these perspec-  
157 tives and explains semantization based on synaptic plas-  
158 ticity, while it also reproduces important episodic mem-  
159 ory phenomena on behavioral time scales under con-  
160 strained network connectivity with plausible postsynaptic  
161 potentials, firing rates, and other biological parameters.

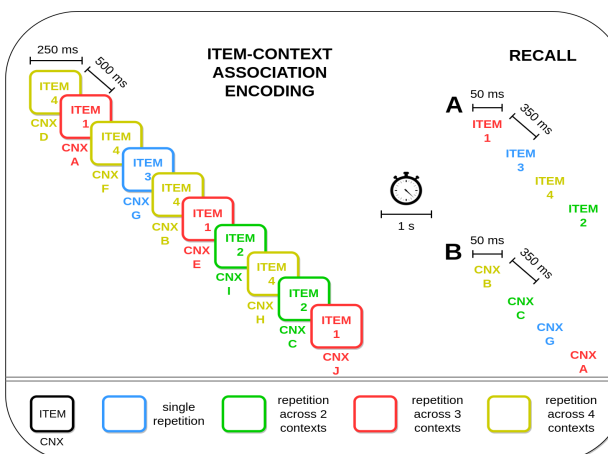
## 2 RESULTS

### 2.1 Semantization of episodic representations in the BCPNN model

164 The network model used here features two reciprocally  
165 connected networks, the so-called Item and Context net-  
166 works. The architecture of each network follows our pre-  
167 vious spiking implementations of attractor memory net-  
168 works (Lansner, 2009; Tully et al., 2014, 2016; Lundqvist  
169 et al., 2011; Fiebig and Lansner, 2017; Chrysanthidis  
170 et al., 2019; Fiebig et al., 2020), and is best understood  
171 as a subsampled cortical layer 2/3 patch with nested hy-  
172 percolumns (HCs) and minicolumns (MCs; Fig. 1A, see  
173 **STAR METHODS** for details). In our model, items are  
174 embedded in the Item network, and context informa-  
175 tion in the Context network as internal long-term mem-  
176 ory representations, derived from prior Hebbian learning  
177 (Fig. 1B,C, **STAR METHODS**). Our episodic memory  
178 task is designed to follow a seminal experimental study  
179 by Opitz (2010). We stimulate some items in a single con-  
180 text and others in a few different contexts establishing  
181 multiple associations (Fig. 2). Stimulus duration during  
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**Fig. 1** Network architecture and connectivity of the Item (green) and Context (blue) networks. **A**) The model represents a subsampled modular cortical layer 2/3 patch consisting of minicolumns (MCs) nested in hypercolumns (HCs). Both networks contain 12 HCs, each comprising 10 MCs. We preload abstract long-term memories of item and context representations into the respective network, in the form of distributed cell assemblies with weights establishing corresponding attractors. Associative plastic connections bind items with contexts. The network features lateral inhibition via basket cells (purple and blue lines) resulting in a soft winner-take-all dynamics. Competition between attractor memories arises from this local feedback inhibition together with disynaptic inhibition between HCAs. **B**) Weight distribution of plastic synapses targeting pyramidal cells. The attractor projection distribution is positive with a mean of 2.1, and the disynaptic inhibition is negative with a mean of -0.3 (we show the fast AMPA weight components here, but the simulation also includes slower NMDA weight components). **C**) Weight matrix between attractors and competing MCs across two sampled HCAs. The matrix displays the mean of the weight distribution between a presynaptic ( $MC_{pre}$ ) and postsynaptic minicolumn ( $MC_{post}$ ), within the same or different HC (black cross separates grid into blocks of HCAs, only two of which are shown here). Recurrent attractor connections within the same HC are stronger (main diagonal, dark red) compared to attractor connections between HCAs (off-diagonals, orange) while inhibition is overall balanced between patterns (blue). Negative inter-pyramidal weights amounts to disynaptic inhibition mediated by double bouquet cells.

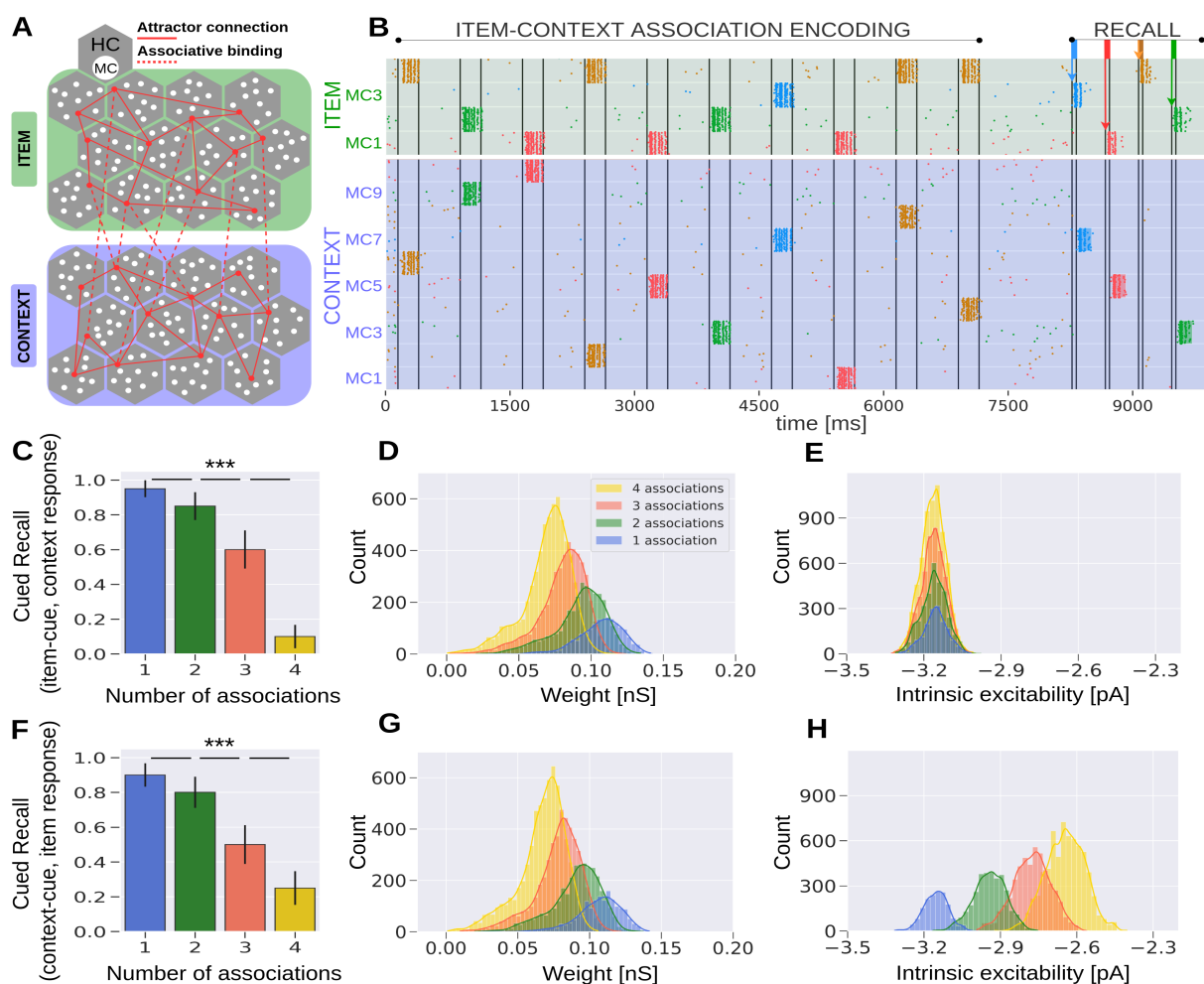


**Fig. 2** Trial structure of the two simulated variants of the episodic memory task. Items are first associated with one or several contexts (CNX) during the encoding phase in 250 ms cue episodes, with an interstimulus interval of 500 ms. The colors of the co-activated contexts are consistent with their corresponding associated item. The recall phase occurs with a delay of 1 s and involves different trials with either brief cues (50 ms) of the **A**) items, or **B**) contexts presented during the item-context association encoding phase.

encoding is  $t_{stim}=250$  ms with a  $T_{stim}=500$  ms interstimulus interval, and a test phase occurs after a 1 s delay period, which contains brief  $t_{cue}=50$  ms cues of previously learned items (Table S2).

Figure 3A illustrates an item-context pair, established by an associative binding through plastic bidirectional BCPNN projections (dashed lines). Item and context attractors (solid red lines) are embedded in each network and remain fixed throughout the simulation, representing well-consolidated long-term memory. We show an exemplary spike raster of pyramidal neurons in HC1 of both the Item and Context networks reflecting a trial simulation (Fig. 3B). Herein, item-3 (blue) establishes a single association, whilst item-4 (yellow) is encoded in four different contexts (Fig. 2A, 3B). We observe evidence of item-context decoupling as the yellow item (but not the blue) is successfully recognized when cued but without any corresponding accompanying activation in the Context network. Successful and complete item recognition without any contextual information retrieval accounts for a Know response, as opposed to Remember judgments, which are accompanied by successful

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**Fig. 3** Semantization of episodic memory traces. **A)** Schematic of the Item (green) and Context (blue) networks. Attractor projections are long-range connections across HCs in the same network and learned associative binding are connections between networks. **B)** Spike raster of pyramidal neurons in HC1 of both the Item and Context networks. Items and their corresponding context representations are simultaneously cued in their respective networks (cf. Fig. 2A). Each item is drawn with a unique color, while contexts inherit their coactivated item's color in the raster (i.e., the yellow pattern in the Item network is repeated over four different contexts, forming four separate associations marked with the same color). The testing phase occurs 1 s after the encoding. Brief 50 ms cues of already studied items trigger their activation. Following item activation, we detect evoked attractor activation in the Context network. **C)** Average cued recall performance in the Context network (20 trials). The bar diagram reveals progressive loss of episodic context information (i.e., semantization) over the number of context associations made by individual cued items (cf. Fig. 2A). **D)** Distribution of plastic connection weights between the Item and Context networks (NMDA component shown here). Weights are noticeably weaker for items which participate in multiple associations. The distributions of synaptic weights exhibit a broader range for the items with multiple context associations, as the sample size is larger. **E)** The distribution of intrinsic excitability currents of pyramidal cells coding for specific context representations. The intrinsic excitability distributions feature similar means because each context is activated exactly once, regardless of whether the associated item forms multiple associations or not. **F)** Average cued recall performance in the Item network (20 trials). Decontextualization over the number of associations is also observed when we briefly cue episodic contexts instead (cf. Fig. 2B, S3). **G)** Distribution of strength of plastic connections from the contexts to their associated items. Analogously to **D**, synapses weaken once an item is encoded in another context. **H)** Intrinsic plasticity distribution of cells in the Item network. Means of the intrinsic excitability distributions are higher for pyramidal cells coding for repeatedly activated items. \*\*\*p<0.001 (Mann-Whitney, N=20 in C, F); Error bars in C, F represent standard deviations of Bernoulli distributions; Means of distributions of one, two, three, and four associations in D, G, H show significant statistical difference (p<0.001, Mann-Whitney, N=2000).

207 context recall. Cue-based activations are reported using  
208 a detection algorithm (see STAR METHODS). Figure  
209 3C demonstrates the performance of contextual  
210 retrieval when items serve as cues. To elucidate this  
211 observed progressive loss of episodicity, we sample and  
212 analyze the learned weight distributions of item-context  
213 binding recorded after the association encoding period  
214 (Fig. 3D). The item-context weight distribution in the  
215 one-association case has a significantly higher mean than  
216 in the two-, three-, or four-association case ( $p<0.001$ ,  
217 Mann-Whitney,  $N=2000$ ). This progressive weakening of  
218 weights leads to significantly lower mean EPSP amplitudes  
219 for the associative projections ( $p<0.05$  for one vs

220 two associations;  $p<0.001$  for two vs three and three vs  
221 four associations, Mann-Whitney,  $N=300$ , Fig. S1). So,  
222 we attribute the loss of episodicity to a statistically signifi-  
223 cant weakening of means of the associative weight dis-  
224 tributions with the increasing number of associated con-  
225 texts. The associative weight distributions shown here re-  
226 fer to the NMDA component, while the weight distribu-  
227 tions of the faster AMPA receptor connections display a  
228 similar trend (Fig. S2). The gradual trace modification we  
229 observe relies on the nature of Bayesian learning, which  
230 normalizes and updates weights over estimated presynap-  
231 tic (Bayesian-prior) as well as postsynaptic (Bayesian-  
232 posterior) spiking activity (see Sect. 2.3 for details).

233 Our simulation results are in line with related behav-  
234 ioral studies (Opitz, 2010; Smith and Manzano, 2010;  
235 Smith and Handy, 2014), which also reported item-  
236 context decoupling as the items were presented across  
237 multiple contexts. In agreement with our study, Opitz  
238 (2010) concluded that repetition of an item across different  
239 contexts (similar to high context variability) leads to  
240 item-context decoupling. Furthermore, Smith and Man-  
241 zano (2010) demonstrated in an episodic context variabil-  
242 ity task configuration, that episodicity deteriorates with  
243 context overload (number of words per context). Mean  
244 recall drops from  $\sim 0.65$  (one word per context) to 0.50  
245 (three words per context), reaching  $\sim 0.33$  in the most  
246 overloaded scenario (fifteen words per context).

247 In Figure 3E we show the distribution of intrinsic ex-  
248 citability over units representing different contexts. Pyra-  
249 midal neurons in the Context network have a similar in-  
250 trinsic excitability, regardless of their selectivity because  
251 all the various contexts are encoded exactly once.

252 Next, analogously to the previous analysis, we show  
253 that item-context decoupling emerges also when we  
254 briefly cue contexts rather than items during recall testing  
255 (Fig. 2B, Fig. S3). In agreement with experimental data  
256 (Smith and Manzano, 2010; Smith and Handy, 2014) we  
257 obtain evidence of semantization as items learned across  
258 several discrete contexts are hardly retrieved when one of  
259 their associated contexts serves as a cue (Fig. 3F). We fur-  
260 ther sample and present the underlying associative weight  
261 distribution, between the Context and the Item networks  
262 (Fig. 3G). The distributions again reflect the semantiza-  
263 tion effect in a significant weakening of the correspond-  
264 ing weights. In other words, an assembly of pyramidal  
265 neurons representing items encoded across multiple con-  
266 texts receive weaker projections from the Context net-  
267 work. Beyond four or more associations, the item-context  
268 binding becomes so weak that it fails to deliver sufficient  
269 excitatory current to trigger associated representations in  
270 the Item network. At the same time, intrinsic excitability  
271 of item neurons increases with the number of associated  
272 contexts corresponding to how much these neurons were  
273 active during the encoding phase [Fig. 3H; cf. Egorov  
274 et al. (2002), Tully et al. (2014)].

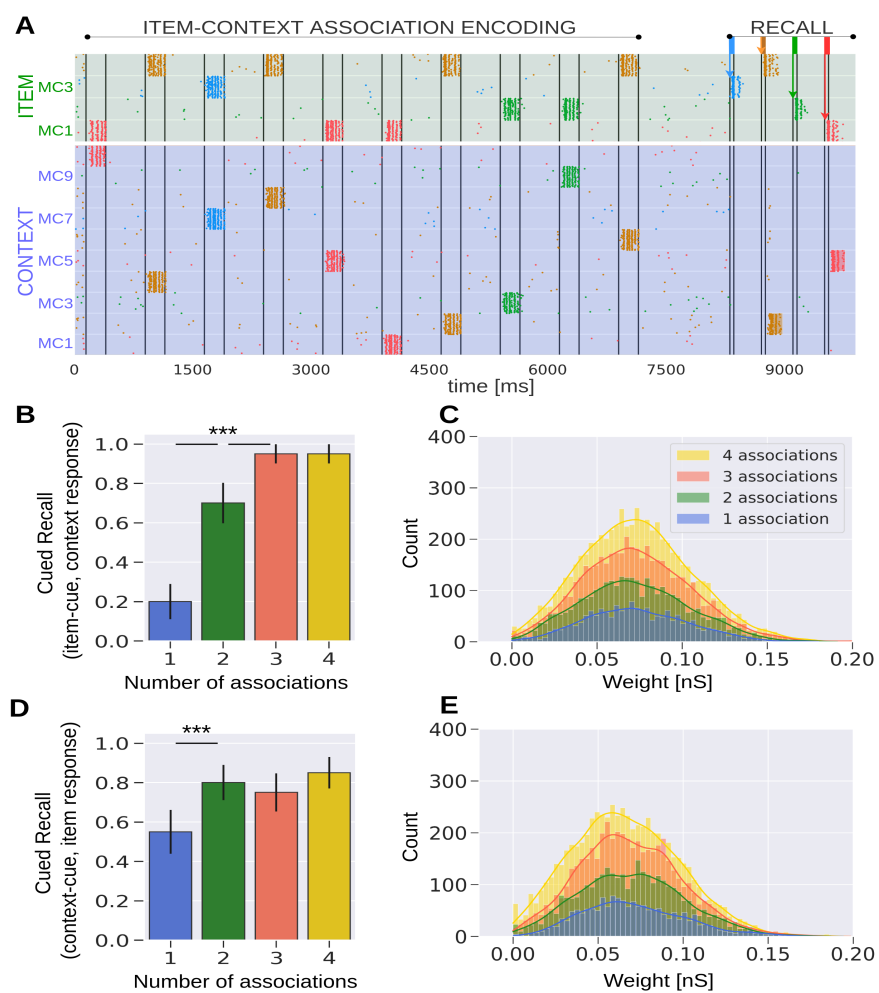
## 275 2.2 Item-context interactions under STDP

276 In this section, we contrast the results obtained  
277 with the BCPNN synaptic learning rule with those de-  
278 riving from the more commonly used STDP learning  
279 rule in the same episodic memory task (Fig. 2, see  
280 STAR+METHODS). The modular network architecture  
281 as well as neural properties and embedded memory pat-  
282 terns remain identical, but associative projections be-  
283 tween networks are now implemented using a standard  
284 STDP synaptic learning rule (Morrison et al., 2008). The  
285 parameters of the STDP model are summarized in Table  
286 S3.

287 Figure 4A shows an exemplary spike raster of pyra-  
288 midal cells in HC1 of both the Item and the Context net-

289 works, based on the first variant of the episodic memory  
290 task described in Figure 2A. As earlier, items are encoded  
291 in a single or in multiple different contexts and they are  
292 briefly cued later during recall. A successful item acti-  
293 vation may lead to a corresponding activation of its as-  
294 sociated information in the Context network. We detect  
295 these activations as before (see STAR+METHODS), and  
296 report the cue-based recall score over the number of as-  
297 sociations (Fig. 4B).

298 Unlike the BCPNN network, we observe no evi-  
299 dence of semantization for high context variability. In-  
300 stead, recollection is noticeably enhanced with an in-  
301 creasing number of associations, which is in fact the op-  
302 posite of what would be needed to explain item-context  
303 decoupling. STDP generates similarly strong associative  
304 binding regardless of context variability (Fig. 4C). The  
305 enhanced recollection in high context variability cases  
306 stems from the multiplicative effect of synaptic augmen-  
307 tation in the Tsodyks-Makram model on the Hebbian  
308 attractor weights. Items stimulated multiple times (e.g.,  
309 four times) have a higher likelihood of being encoded  
310 near the end of the task, leading to more remaining aug-  
311 mentation during testing, thus, effectively boosting cued  
312 recall (see Fig. S4). This effect of the enhanced recall di-  
313 minishes after removing synaptic augmentation from the  
314 model (Fig. S5). As far as the context-cued variant of the  
315 task is concerned, there are also no signs of item-context  
316 decoupling for high context variability (Fig. 4D). The  
317 associative projections between Context and Item net-  
318 works again have distributions with comparable means  
319 over context variability (Fig. 4E). An inclusion of in-  
320 trinsic plasticity dynamics in the model does not explain de-  
321 contextualization either (see Fig. S6). Overall, decon-  
322 textualization is not evident in either variant of the episodic  
323 memory task under the STDP learning rule.



**Fig. 4** Network model where associative projections are implemented using standard STDP synaptic plasticity. **A)** Spike raster of pyramidal neurons in HC1 of both the Item and Context networks (20 trials). **B)** Average item-cued recall performance in the Context network (20 trials). Episodic context retrieval is preserved even for high context variability (as opposed to BCPNN, cf. Fig. 3C). **C)** Distribution of NMDA receptor mediated synaptic weights between the item and context neural assemblies following associative binding. The distributions of item-context weights have comparable means at  $\sim 0.065$  nS regardless of how many context associations a given item forms. Bins merely display a higher count for the four-association case as the total count of associative weights is more extensive compared to items with fewer associations. **D)** Average cued recall performance in the Item network when episodic contexts are cued (20 trials). **E)** Distribution of NMDA component weights between associated context and item assemblies. \*\*\* $p < 0.001$  (Mann-Whitney,  $N=20$  in **B, D**); Error bars in **B, D** represent standard deviations of Bernoulli distributions.

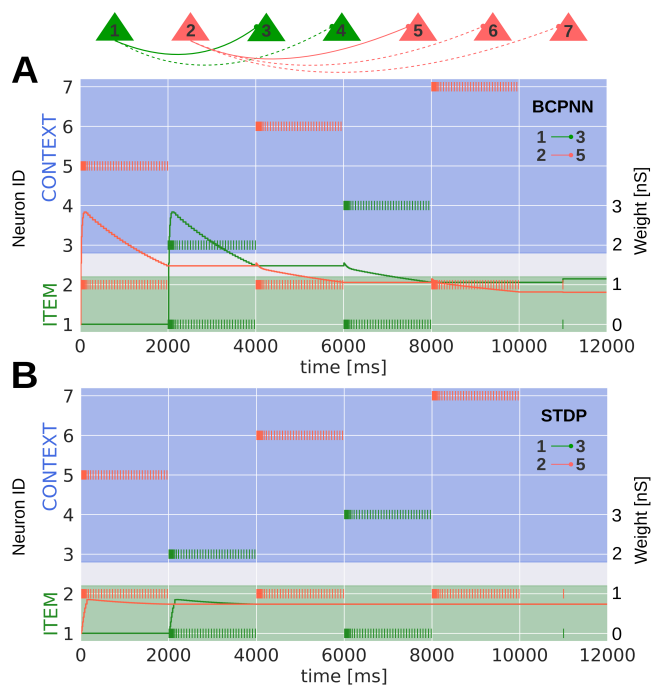
### 324 2.3 BCPNN and STDP learning rule in a microcircuit 325 model

326 To better elucidate the emergent synaptic changes  
327 of the BCPNN and STDP model, we also apply these  
328 learning rules in a highly reduced microcircuit of spiking  
329 neurons. To this end, we now track the synaptic weight  
330 changes continuously.

331 First, we apply the BCPNN learning rule to the mi-  
332 crocircuit model. We consider two separate item neu-  
333 rons (ID=1 and 2), which form two or three associa-  
334 tions with context neurons (ID=3,4, or 5,6,7), respec-  
335 tively (Fig. 5A). We display the synaptic strength de-  
336 velopment of the synapse between item neuron-1 and  
337 context neuron-3 (two associations, green), as well as  
338 the synapse between item neuron-2 and context neuron-5  
339 (three associations, red) over the course of training these  
340 associations via targeted stimulation. BCPNN synapses  
341 get strengthened when the item-context pairs are sim-  
342 ultaneously active and weaken when the item in question  
343 is activated with another context. Therefore, synapses of  
344 the item neuron that is encoded in three different con-

345 texts converge on weaker weights (Fig. 5A, 12 s), than  
346 those of the item neuron with two associated contexts.  
347 Weight modifications in the microcircuit model reflect  
348 the synaptic alterations observed in the large-scale net-  
349 work. BCPNN weights are shaped by traces of activation  
350 and co-activation (Eq. 7.8, STARMETHODS), which  
351 also get updated during the activation of an item within  
352 another context. For example, the item neuron-1 and con-  
353 text neuron-3 are not stimulated together between 6 s and  
354 8 s, but neuron-1 and context neuron-4 are. Thus, the  
355 traces of the item activation ( $P_i$ ) increase, while the ones  
356 linked to context-3 ( $P_j$ ) decay with a time constant of 15  
357 s (Table S1). Since the item and context neuron (ID=1, 3)  
358 are not stimulated together, their coactivation traces ( $P_{ij}$ )  
359 decay between 6 s and 8 s. Overall, this leads to a weak-  
360 ening of the weight and hence, to a gradual decoupling  
361 (Eq. 8, STARMETHODS).

362 In the same manner, we keep track of weight change  
363 in a microcircuit with the STDP learning rule (Fig. 5B).  
364 Unlike the microcircuit with BCPNN presented in Figure  
365 5A, the STDP weights corresponding to the associations  
366 made by both item neurons converge to similar values,



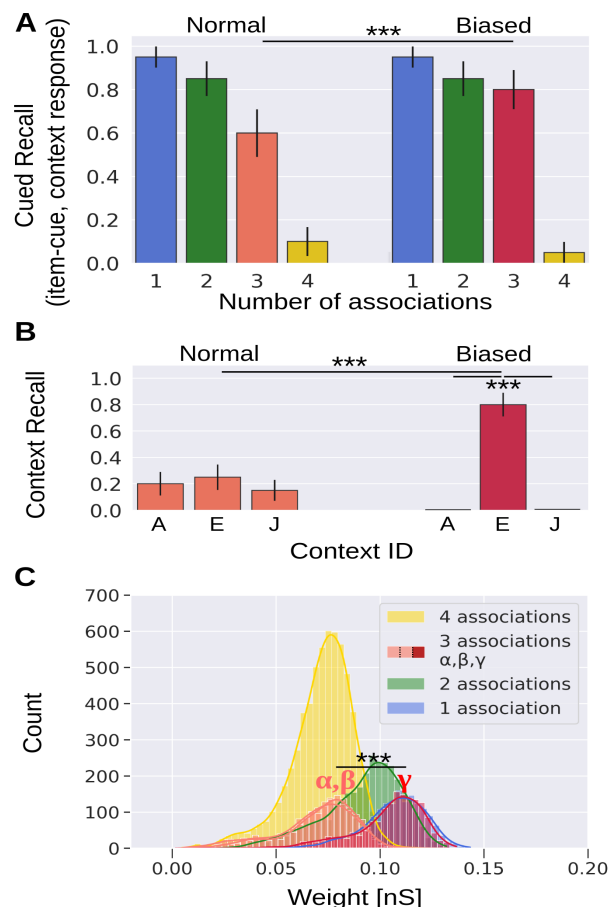
**Fig. 5** Continuous weight recordings in a microcircuit model with plastic synapses under the **A)** BCPNN or **B)** STDP learning rule. Neural and synaptic parameters correspond to those in the scaled model. In both cases, two item neurons (ID=1,2) are trained to form two or three associations, respectively (dashed connections are simulated but their weight development is not shown here). During training, neurons are stimulated to fire at 20 Hz for 2 s. We display the developing synaptic weight between specific item-context pairs, (ID=1 and 3 in the two-association scenario) and (ID=2 and 5 in the three-association scenario), and compare the converged weight values between the two- and three-association case under both learning rules, following a final readout spike at 11 s.

367 even though they are associated with different number of  
 368 contexts. As before, the synapse between an item neuron  
 369 and an associated context neuron strengthens when this  
 370 pair is simultaneously active, but remains stable when  
 371 the item neuron is encoded in another context. For in-  
 372 stance, the synapse between item neuron-2 and context  
 373 neuron-5 strengthens when this pair is encoded (0-2 s),  
 374 yet remains unaffected when item neuron-2 is activated  
 375 in another context (i.e., context neuron-6, 4-6 s). This  
 376 synaptic behavior explains the observed differences be-  
 377 tween the BCPNN and STDP large-scale model.

#### 378 2.4 Preferential retention

379 Several studies propose that one-shot salient events  
 380 promote learning, and that these memories can be re-  
 381 tained on multiple time scales ranging from seconds to  
 382 years (Petrican et al., 2010; Gruber et al., 2016; Frank-  
 383 land et al., 2004; Panoz-Brown et al., 2016; Eichenbaum,  
 384 2017; Sun et al., 2018). Hypothetical mechanisms behind  
 385 these effects are dopamine release and activation of DR1  
 386 like receptors, resulting in synapse-specific enhancement  
 387 (Otmakhova and Lisman, 1996; Kuo et al., 2008), and  
 388 systems consolidation (McClelland et al., 1995; Fiebig  
 389 and Lansner, 2014). On the whole, salient or reward  
 390 driven events may be encoded more strongly as the result  
 391 of a transient plasticity modulation. Recall from long-  
 392 term memory is often viewed as a competitive process

393 in which a memory retrieval does not depend only on its  
 394 own synaptic strength but also on the strength of other  
 395 components (Shiffrin, 1970). In view of this, we study  
 396 the effects of plasticity modulation on encoding specific  
 397 items within particular contexts, with the aim of investi-  
 398 gating the role of enhanced learning for semantization in  
 399 our model.



**Fig. 6** Plasticity modulation of a specific item-context pair enhances recollection and counteracts semantization. **A)** Context recall performance. One of the pairs (context-E, item-1) presented in the episodic memory task (cf. Fig. 2A) is subjected to enhanced plasticity during encoding, resulting in the boosted recall rate (3 associations, Normal vs Biased). **B)** Individual context retrieval contribution in the overall recall (3 associations). Retrieval is similar among the three contexts since plasticity modulation is balanced (left: Normal,  $\kappa=\kappa_{normal}$ , cf. Table S1). However, when context-E is encoded with enhanced learning (with item-1), its recall increases significantly (right: Biased,  $\kappa=\kappa_{boost}$ , cf. Table S1). **C)** Weight distributions of the NMDA weight component. Encoding item-1 with context-E under modulated plasticity yields stronger synaptic weights [3 association,  $\alpha, \beta$  (light red, highly overlapping distributions) vs  $\gamma$  (dark red)]. \*\*\*p<0.001 (Mann-Whitney, N=20 in A, B, N=2000 in C); Error bars in A, B represent standard deviations of Bernoulli distributions; Means of the weight distributions of one, two, three- $\alpha, \beta$ , and four associations in C show significant statistical difference (p<0.001, Mann-Whitney, N=2000).

400 Using the same network and episodic memory task  
 401 as before (Fig. 2A), we modulate plasticity during the  
 402 encoding of item-1 (red) in context-E via  $\kappa=\kappa_{boost}$  (Eq.  
 403 7, STAR+METHODS, Table S1). This results in an in-  
 404 creased cued recall probability for the item associated  
 405 with three episodic contexts relative to the unmodulated  
 406 control (Fig. 6A, Normal vs Biased scenario, 3 associ-  
 407 ations). Episodic retrieval improves from 0.6 (Normal,  
 408

408 Fig. 6A, left) to 0.8 (Biased; modulated plasticity, Fig. 463  
409 6A, right) when item-1 is cued, which now performs 464  
410 more similarly to item with just two associated contexts. 465  
411 We further analyze and compare the recall of each 466  
412 context when its associated item-1 is cued (Fig. 6B, 3 467  
413 associations). The control scenario (Normal, Fig. 6B, left) 468  
414 without transient plasticity modulation shows that the 469  
415 three contexts (ID=A, E and J) are all recalled with 470  
416 similar probabilities (20 trials). In contrast, encoding a 471  
417 specific pair with enhanced learning (upregulated  $\kappa=\kappa_{boost}$ ) 472  
418 yields higher recall for the corresponding context. In 473  
419 particular, the plasticity enhancement during associative 474  
420 encoding of the context-E (with item-1) results in an 475  
421 increased recall score to 0.8 (0.25 control), while the other 476  
422 associated contexts, ID=A and J, are suppressed (Fig. 477  
423 6B), primarily due to soft winner-take-all competition 478  
424 between contexts (Fig. 1A). 479  
425

426 We attribute these changes to the stronger weights 480  
427 due to enhanced learning (Fig. 6c, dark red distribution, 481  
428  $\gamma$ ). Weights between unmodulated item-context pairs 482  
429 (item-1 and context-A,-J) show mostly unaltered weight 483  
430 distributions ( $\alpha,\beta$ , light red), while the biased associative 484  
431 weight distribution between item-1 and context-E is 485  
432 now comparable to the weight distribution of the one- 486  
433 association case. Performance does not exactly match 487  
434 that case though due to some remaining competition 488  
435 among the three contexts. Overall, these results demon- 489  
436 strate how a single salient episode may strengthen mem- 490  
437 ory traces and thus impart resistance to semantization 491  
(Rodríguez et al., 2016). 492

### 438 3 DISCUSSION

439 The primary objective of this work was to explore the 495  
440 interaction between synaptic plasticity and context vari- 496  
441 ability in the semantization process. To cast new light 497  
442 on the episodic-semantic interplay, we built a memory 498  
443 model of two spiking neural networks coupled with plas- 499  
444 tic connections, which collectively represent distributed 500  
445 cortical episodic memory. Our results suggest that some 501  
446 forms of plasticity offer a synaptic explanation for the 502  
447 cognitive phenomenon of semantization, thus bridging 503  
448 scales and linking network connectivity and dynamics 504  
449 with behavior. In particular, we demonstrated that with 505  
450 Bayesian-Hebbian (BCPNN) synaptic plasticity, but not 506  
451 with standard Hebbian STDP, the model can reproduce 507  
452 traces of semantization in the learning outcomes. Not- 508  
453 ably, this was achieved with biologically constrained 509  
454 network connectivity, postsynaptic potential amplitudes, 510  
455 firing rates and oscillatory dynamics compatible with 511  
456 mesoscale recordings from cortex and earlier models. 512  
457 Nevertheless, our hypothesis of the episodic-semantic 513  
458 interplay at a neural level requires further experimental 514  
459 study of synaptic strength dynamics in particular. We also 515  
460 demonstrated how a transient plasticity modulation (re- 516  
461 reflecting known isolation effects) may preserve episodic- 517  
462 ity, staving off decontextualization. 518

463 Our study conforms to related behavioral exper- 464  
464 iments reporting that high context variability or con- 465  
465 text overload leads to item-context decoupling (Opitz, 466  
466 2010; Smith and Manzano, 2010; Smith and Handy, 467  
467 2014). These studies suggest that context-specific mem- 468  
468 ory traces transform into semantic representations while 469  
469 contextual information is progressively lost. Memory 470  
470 traces remain intact but fail to retrieve their associated 471  
471 context. Semantization is typically described as a de- 472  
472 contextualization process that occurs over time. How- 473  
473 ever, several experiments, including this study, proposed 474  
474 that exposures of stimuli in different additional contexts 475  
475 (rather than time itself) is the key mechanism advancing 476  
476 semantization (Opitz, 2010; Smith and Manzano, 2010; 477  
477 Smith and Handy, 2014). Admittedly, our hypothesis 478  
478 cannot exclude other seemingly coexisting phenomena 479  
479 that may benefit semantization over time (e.g., reconsol- 480  
480 idation or systems consolidation due to sleep or aging). 481

482 To our knowledge, there is no other spiking bio- 483  
483 physical computational model of comparable detail that 484  
484 captures the semantization of episodic memory explored 485  
485 here, whilst simultaneously offering a neurobiological 486  
486 explanation of this phenomenon. Unlike other dual- 487  
487 process episodic memory models, which require repeated 488  
488 stimulus exposures to support recognition (Norman and 489  
489 O'Reilly, 2003), our model is able to successfully recall 490  
490 events learned in "one shot" (a distinctive hallmark of 491  
491 episodicity). We note that the attractor-based theory pro- 492  
492 posed in this study does not exclude the possibility of a 493  
493 dual-process explanation for recollection and familiarity 494  
(Yonelinas, 2002; Yonelinas et al., 2010). 495

### 494 3.1 Related models of familiarity and recollection

495 Perceptual or abstract single-trace dual-process com- 496  
496 putational models based on signal detection theory ex- 497  
497 plain episodic retrieval but the potential loss of contextual 498  
498 information is only implied as it does not have its own 499  
499 independent representation (Greve et al., 2010; Wixted, 500  
500 2007). These computational models often aim to explain 501  
501 traditional R/K behavioral studies. As discussed earlier, 502  
502 participants in such studies are instructed to give a Know 503  
503 response if the stimulus presented in the test phase is 504  
504 known or familiar without any contextual detail about its 505  
505 previous occurrence. Conversely, Remember judgments 506  
506 are to be provided if the stimulus is recognized along with 507  
507 some recollection of specific contextual information per- 508  
508 taining to the study episode. This results in a strict cri- 509  
509 terion for recollection, as it is possible for a subject to 510  
510 successfully recall an item but fail to retrieve the source 511  
511 information (Ryals et al., 2013). Numerous studies sug- 512  
512 gest that recollection contaminates Know reports because 513  
513 recalling source information sensibly assumes prior item 514  
514 recognition (Wais et al., 2008; Johnson et al., 2009). 515  
515 Mandler (1979, 1980), and Atkinson and Juola (1973) 516  
516 treat familiarity as an activation of preexisting memory 517  
517 representations. Our results are compatible with this no- 518  
518 tion because our model proposes to treat item-only activa- 519  
519 tions as Know judgments, while those accompanied by 520  
520 the activation of context representations best correspond

521 to a Remember judgment. Item activation is a faster process and precedes context retrieval (Yonelinas and Ja-  
522 coby, 1994), and our model reflects this finding by ne-  
523 cessity, as item activations are causal to context retrieval.  
524

525 To us, familiarity recognition is simply characterized  
526 by a lack of contextual information, yet the distinction  
527 we make between Context and Item networks is arbitrary.  
528 Any item can be a context and vice versa, so the networks  
529 are interchangeable. While sparse interconnection is suf-  
530 ficient for our model's function, both networks may just  
531 as well be part of the same modality and cortical brain  
532 areas. A more specific scenario might assume that items  
533 and contexts share part of the same local network. In prin-  
534 ciple, our model should be capable of replicating similar  
535 results in a single modality scenario.

### 536 3.2 Semantization on longer time scales

537 Source recall is likely supported by multiple inde-  
538 pendent, parallel, interacting neural structures and pro-  
539 cesses since various parts of the medial temporal lobes,  
540 prefrontal cortex and parts of the parietal cortex all con-  
541 tribute to episodic memory retrieval including informa-  
542 tion about both where and when an event occurred (Diana  
543 et al., 2007; Gilboa, 2004; Watrous et al., 2013). A related  
544 classic idea on semantization is the view that it is in fact  
545 an emergent outcome of systems consolidation. Sleep-  
546 dependent consolidation in particular has been linked to  
547 advancing semantization of memories and the extraction  
548 of gist information. (Friedrich et al., 2015; Payne et al.,  
549 2009).

550 Models of long-term consolidation suggest that richly  
551 contextualized memories, become more generic over  
552 time. Without excluding this possibility, we note that this  
553 is not always the case, as highly salient memories often  
554 retain contextual information (which our model speaks  
555 to). Instead, our model argues for a much more imme-  
556 diate neural and synaptic contribution to semantization  
557 that does not require slow multi-area systems level pro-  
558 cesses that have yet to be specified in sufficient detail  
559 to be tested in neural simulations. We have previously  
560 shown, however, that an abstract simulation network of  
561 networks with broader distributions of learning time con-  
562 stants can consolidate memories across several orders  
563 of magnitude in time, using the same Bayesian-Hebbian  
564 learning rule as used here (Fiebig and Lansner, 2014).  
565 That model included representations for prefrontal cor-  
566 tex, hippocampus, and wider neocortex, implementing an  
567 extended complementary learning systems theory (Mc-  
568 Clelland et al., 1995), which is itself an advancement of  
569 systems consolidation (Squire and Alvarez, 1995). We  
570 consequently expect that the principled mechanism of se-  
571 mantization explored here can be scaled along the tempo-  
572 ral axis to account for lifelong memory, provided that the  
573 plasticity involved is itself Bayesian-Hebbian. Our model  
574 does not advance any specific anatomical argument as  
575 to the location of the respective networks (Diana et al.,  
576 2007; Yonelinas, 2002).

577 The model purposefully relies on a generic cortical  
578 architecture focused on a class of synaptic plastic-

579 ity mechanisms which may well serve as a substrate of  
580 a wider system across brain areas and time.

### 581 3.3 Biological plausibility and parameter sensitivity

582 We investigate and explain behavior and macroscale  
583 system dynamics with respect to neural processes, bio-  
584 logical parameters of network connectivity, and electro-  
585 physiological evidence. Our model consequently builds  
586 on a broad range of biological constraints such as in-  
587trinsic neuronal parameters, cortical laminar cell densi-  
588 ties, plausible delay distributions, and network connectiv-  
589 ity. The model reproduces plausible postsynaptic poten-  
590 tials (EPSPs, IPSPs) and abides by estimates of connec-  
591 tion densities (i.e., in the associative pathways and pro-  
592 jections within each patch), axonal conductance speeds,  
593 typically accepted synaptic time constants for the vari-  
594 ous receptor types (AMPA, NMDA, and GABA), with  
595 commonly used neural and synaptic plasticity time con-  
596 stants (i.e., adaptation, depression). We reproduce oscil-  
597 latory dynamics in multiple frequency ranges, that were  
598 previously studied in the same modular spiking network  
599 implementations (Lundqvist et al., 2010, 2011; Herman  
600 et al., 2013).

601 The model synthesizes a number of functionally  
602 relevant processes, embedding different components to  
603 model composite dynamics, hence, it is beyond this study  
604 to perform a detailed sensitivity analysis for every pa-  
605 rameter. Instead, we provide insightful observations for  
606 previously unexplored parameters that may critically af-  
607 fect semantization. Importantly, a highly related modular  
608 cortical model already investigated sensitivity to impor-  
609 tant short-term plasticity parameters (Fiebig and Lansner,  
610 2017). After extensive simulation testing, we conclude  
611 that the model is generally robust to a broad range of  
612 parameter changes and degrades gracefully. Small net-  
613 works like this are typically more sensitive to parameter  
614 changes, so conversely, we expect even lower sensitivity  
615 to parameter variations in a full scale system.

616 The P trace decay time constant,  $\tau_p$ , of the BCPNN  
617 model is critical for the learning dynamics modelled in  
618 this study because it controls the speed of learning in as-  
619 sociative connections. High values of  $\tau_p$  lead to slower  
620 and more long-lasting learning. Varying  $\tau_p$  by  $\pm 30\%$   
621 does not change the main outcome, that is, episodicity  
622 still deteriorates with a higher context variability. Slower  
623 weight development may result in weaker associative  
624 binding and overall lower recall though (and vice versa  
625 for faster learning). To compensate for this loss of episod-  
626 icity, an additional increase in the unspecific input is usu-  
627 ally sufficient to trigger comparable recall rates. Alter-  
628 natively, the recurrent excitatory gain can be amplified  
629 to complete noisy inputs towards discrete embedded at-  
630 tractors. Unspecific background input during recall plays  
631 a critical role as well. We use a low such noise input to  
632 model cue-association responses, however, when boosted  
633 by  $+40\%$ , the model operates in a free replay regime in-  
634 stead, where cues become unnecessary as the network re-  
635 trievies content without them by means of intrinsic back-  
636 ground noise.

637 This study also demonstrated how a selective transient  
638 increase of plasticity can counteract semantization.  
639 The plasticity of the model can be modulated via the  
640 parameter  $\kappa$  (Eq. 7, STAR METHODS). Typically,  $\kappa$   
641 is set to 1 ( $\kappa=\kappa_{normal}$ ), whereas we double plasticity  
642 ( $\kappa=\kappa_{boost}$ ), when modelling salient episodic encoding.  
643 We noticed that by selectively tripling or quadrupling  
644 plasticity (relative to baseline) during encoding of a specific  
645 pair whose item component forms many other associations,  
646 the source recall improves progressively (data shown only for  $\kappa=\kappa_{boost}$  in Sect. 2.4).

647 Finally, in Section 2.3 we compared STDP and  
648 BCPNN plasticity in a highly reduced model. We bind  
649 items with contexts to form different number of associations  
650 and keep track of the weight development per time  
651 step. STDP plasticity generated same magnitude item-  
652 context binding regardless of how many associations an  
653 item forms. A detailed parameter analysis for every critical  
654 synaptic parameter ( $\pm 30\%$ ) did not yield any behaviorally  
655 significant changes to the converged weights.

### 657 3.4 Conclusions

658 We have presented a computational mesoscopic network  
659 model to examine the interplay between episodic and  
660 semantic memory with the grand objective to explain  
661 mechanistically the semantization of episodic traces.  
662 Compared to other models of episodic memory, which  
663 are typically abstract, our model, built on various bi-  
664 ological constraints (i.e., plausible postsynaptic potentials,  
665 firing rates, etc.) accounting for neural processes  
666 and synaptic mechanisms, emphasizes the role of synaptic  
667 plasticity in episodic forgetting. Hence it bridges micro-  
668 and mesoscale mechanisms with macroscale behavior  
669 and dynamics. In contrast to standard Hebbian learning,  
670 our Bayesian version of Hebbian learning readily repro-  
671 duced prominent traces of semantization.

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679 KTH Royal Institute of Technology.

### 680 STAR METHODS

681 Detailed methods are provided in the online version of  
682 this paper and include the following:

- 683 • KEY RESOURCES TABLE

- 684 • METHODS DETAILS

- 685 • Spike-based BCPNN plasticity
- 686 • Spike-based STDP learning rule
- 687 • Two-network architecture and connectivity
- 688 • Axonal conduction delays
- 689 • Stimulation Protocol
- 690 • Attractor activation detector
- 691 • Simulation Tools

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## 996 4 STAR METHODS

### 997 4.1 KEY RESOURCES TABLE

Table 1

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Software and Algorithms		
NEST simulator	(Gewaltig and Diesmann, 2007)	nest-simulator.org
Programming Language Python 2.7		python.org; SCR_008394
BCPNN learning rule module	(Tully et al., 2014)	Zenodo
PDC High Performance Computing		www.pdc.kth.se

## 998 4.2 METHODS DETAILS

### 999 4.2.1 Neuron and synapse model

1000 We use adaptive exponential integrate-and-fire point  
 1001 model neurons, which feature spike frequency adaptation,  
 1002 enriching neural dynamics and spike patterns, especially  
 1003 for the pyramidal cells (Brette and Gerstner, 2005).  
 1004 The neuron model offers a good phenomenological  
 1005 description of typical neural firing behavior, but it is limited  
 1006 in predicting the precise time course of the sub-threshold  
 1007 membrane voltage during and after a spike or the underlying  
 1008 biophysical causes of electrical activity (Gerstner  
 1009 and Naud, 2009). We slightly modified it for compatibility  
 1010 with the BCPNN synapse model (Tully et al., 2014) by  
 1011 integrating an intrinsic excitability current.

1012 Development of the membrane potential  $V_m$  and the  
 1013 adaptation current  $I_w$  is described by the following equations:

$$C_m \frac{dV_m}{dt} = -g_L(V_m - E_L) + g_L \Delta \tau e^{\frac{V_m - V_t}{\Delta \tau}} - I_w + I_{ext} + I_{syn} \quad (1)$$

$$\frac{dI_w}{dt} = \frac{-I_w}{\tau_{I_w}} + b \delta(t - t_{sp}) \quad (2)$$

1015 Equation 1 describes the dynamics of the membrane  
 1016 potential  $V_m$  including an exponential voltage dependent  
 1017 activation term. A leak current is driven by the leak re-  
 1018 versal potential  $E_L$  through the conductance  $g_L$  over the  
 1019 neural surface with a capacity  $C_m$ . Additionally,  $V_t$  is the  
 1020 spiking threshold, and  $\Delta \tau$  shapes the spike slope factor.  
 1021 After spike generation, membrane potential is reset to  $V_r$ .  
 1022 Spike emission upregulates the adaptation current by  $b$ ,  
 1023 which recovers with time constant  $\tau_{I_w}$  (Table S1). We  
 1024 neglect subthreshold adaptation, which is part of some  
 1025 AdEx models.

1026 Besides a specific external input current  $I_{ext}$ , model  
 1027 neurons receive synaptic currents  $I_{syn_j}$  from conductance  
 1028 based glutamatergic and GABA-ergic synapses. Gluta-  
 1029 matergic synapses feature both AMPA/NMDA receptor  
 1030 gated channels with fast and slow conductance decay dy-  
 1031 namic, respectively. Current contributions for synapses  
 1032 are described as follows:

$$I_{syn_j} = \sum_{syn} \sum_i g_{ij}^{syn}(t) (V_m - E_{ij}^{syn}) = I_j^{AMPA}(t) + I_j^{NMDA}(t) + I_j^{GABA}(t) \quad (3)$$

1033 The glutamatergic synapses are also subject to synap-  
 1034 tic depression and augmentation with a decay factor  $\tau_D$   
 1035 and  $\tau_A$ , respectively (Table S1), following the Tsodyks-  
 1036 Markram formalism (Tsodyks and Markram, 1997). The  
 1037 utilization factor  $U$ , encodes variations in the release  
 1038 probability of available resources:

$$\frac{du_{ij}}{dt} = -\frac{u_{ij}}{\tau_A} + U(1 - u_{ij}) \sum_{sp} \delta(t - t_{sp}^i - t_{ij}) \quad (4)$$

$$\frac{dx_{ij}}{dt} = \frac{1 - x_{ij}}{\tau_D} - U x_{ij} \sum_{sp} \delta(t - t_{sp}^i - t_{ij}) \quad (5)$$

### 4.2.2 Spike-based BCPNN plasticity

1039 We implement synaptic plasticity of AMPA and  
 1040 NMDA connection components using the BCPNN learn-  
 1041 ing rule (Lansner and Ekeberg, 1989; Wahlgren and  
 1042 Lansner, 2001; Tully et al., 2014). BCPNN is derived  
 1043 from Bayes rule, assuming a postsynaptic neuron em-  
 1044 ploys some form of probabilistic inference to decide  
 1045 whether to emit a spike or not. In general, it is considered  
 1046 more complex than the standard STDP learning rule (Ca-  
 1047 porale and Dan, 2008), and it reproduces the main fea-  
 1048 tures of STDP plasticity. As other spiking synaptic learn-  
 1049 ing rules, it is so far insufficiently validated against quan-  
 1050 titative experimental data on biological synaptic plastic-  
 1051 ity.

1052 The BCPNN synapse continuously updates three  
 1053 synaptic biophysically plausible local memory traces,  $P_i$ ,  
 1054  $P_j$  and  $P_{ij}$ , implemented as exponentially moving aver-  
 1055 ages (EMAs) of pre-, post- and co-activation, from which  
 1056 the Bayesian bias and weights are calculated. EMAs pri-  
 1057 oritize recent patterns, so that newly learned patterns  
 1058 gradually replace old memories. Specifically, learning  
 1059 implements a three-level procedure of exponential filters  
 1060 which defines Z, E and P traces. E traces, which enable  
 1061 delayed reward learning, are not used here because such  
 1062 conditions are not applicable to the modelled task.

1063 To begin with, BCPNN receives a binary sequence of  
 1064 pre- and postsynaptic spiking events ( $S_i, S_j$ ) to calculate  
 1065 the traces  $Z_i$  and  $Z_j$ :

$$\begin{cases} \tau_{z_i} \frac{dZ_i}{dt} = \frac{S_i}{f_{max} t_{spike}} - Z_i + \epsilon \\ \tau_{z_j} \frac{dZ_j}{dt} = \frac{S_j}{f_{max} t_{spike}} - Z_j + \epsilon \end{cases} \quad (6)$$

1067  $f_{max}$  denotes the maximal neuronal spike rate,  $\varepsilon$  is the  
 1068 lowest attainable probability estimate,  $t_{spike}$  denotes the  
 1069 spike duration while  $\tau_{z_i} = \tau_{z_j}$  are the pre and postsynaptic  
 1070 time constants, respectively (5 ms for AMPA, and 100 ms  
 1071 for NMDA components, Table S1).

1072 P traces are then estimated from the Z traces as fol-  
 1073 lows:

$$\begin{cases} \tau_p \frac{dP_i}{dt} = \kappa(Z_i - P_i) \\ \tau_p \frac{dP_j}{dt} = \kappa(Z_j - P_j) \\ \tau_p \frac{dP_{ij}}{dt} = \kappa(Z_i Z_j - P_{ij}) \end{cases} \quad (7)$$

1074 The parameter  $\kappa$  adjusts the learning rate, reflect-  
 1075 ing the action of endogenous modulators of learning ef-  
 1076 ficacy (i.e., activation of a D1R-like receptor). Setting  
 1077  $\kappa=0$  freezes the network's weights and biases, though  
 1078 in our simulations the learning rate remains constant  
 1079 ( $\kappa=1$ ) during encoding (Sect. 2.1, 2.2). However, we trig-  
 1080 ger a transient increase of plasticity in specific scenar-  
 1081 os to model preferential retention, assuming encoding of  
 1082 salient events (Sect. 2.4 and Table S1).

1083 Finally,  $P_i$ ,  $P_j$  and  $P_{ij}$  are used to calculate intrinsic  
 1084 excitability  $\beta_j$  and synaptic weights  $w_{ij}$  with a scaling  
 1085 factor  $\beta_{gain}$  and  $w_{gain}^{syn}$  respectively (Table S1):

$$\begin{cases} w_{ij} = w_{gain}^{syn} \log \frac{P_{ij}}{P_i P_j} \\ \beta_j = \beta_{gain} \log(P_j) \end{cases} \quad (8)$$

#### 1086 4.2.3 Spike-based STDP learning rule

1087 In our study, we examine the impact on semantiza-  
 1088 tion when the STDP learning rule replaces BCPNN as-  
 1089 sociative connectivity in the same episodic memory task.  
 1090 Synapses under STDP are developed and modified by a  
 1091 repeated pairing of pre- and postsynaptic spiking activ-  
 1092 ity, while their relative time window shapes the degree of  
 1093 modification (Ren et al., 2010). The amount of trace mod-  
 1094 ification depends on the temporal difference ( $\Delta_t$ ) between  
 1095 the time point of the presynaptic action potential ( $t_i$ ) and  
 1096 the occurrence of the postsynaptic spike ( $t_j$ ) incorporat-  
 1097 ing a corresponding transmission delay from neuron i to  
 1098 j ( $\tau_d$ ):

$$\Delta t = t_j - (t_i + \tau_d) \quad (9)$$

1099 After processing  $\Delta t$ , STDP updates weights accord-  
 1100 ingly:

$$\Delta w_{ij}(\Delta t) = \begin{cases} \lambda(1-w)^{\mu_+} e^{(-|\Delta t|/\tau_+)} & \text{if } \Delta t \geq \tau_d \\ -\lambda \alpha w^{\mu_-} e^{(-|\Delta t|/\tau_-)} & \text{if } \Delta t < \tau_d \end{cases} \quad (10)$$

1101 Here,  $\lambda$  corresponds to the learning rate,  $\alpha$  reflects  
 1102 a possible asymmetry between the scale of potentiation  
 1103 and depression,  $\tau_{\pm}$  control the width of the time win-  
 1104 dows, while  $\mu_{\pm} \in \{0,1\}$  allows to choose between dif-  
 1105 ferent versions of STDP (i.e., additive, multiplicative),  
 1106 (Morrison et al., 2008). Synapses are potentiated if the  
 1107 synaptic event precedes the postsynaptic spike and get

1108 depressed if the synaptic event follows the postsynaptic  
 1109 spike (Van Rossum et al., 2000).

1110 Associative weights  $w_{ij}$  are initialized to  $w_0$ , and their  
 1111 maximum allowed values are constrained according to  
 1112  $w_{max}$  to ensure that synaptic weights are always posi-  
 1113 tive and between  $[w_0, w_{max}]$  (Table S3). The resulting as-  
 1114 sociative weight distributions are generally comparable  
 1115 in strength to the BCPNN model weights, but to make  
 1116 them match, we adjust  $w_{max}$  in conjunction with a reason-  
 1117 ably small learning rate  $\lambda$ . To obtain a stable competitive  
 1118 synaptic modification, the integral of  $\Delta w_{ij}$  must be nega-  
 1119 tive (Song et al., 2000). To ensure this, we choose  $\alpha=1.2$ ,  
 1120 which introduces an asymmetry between the scale of po-  
 1121 tentiation and depression along with a symmetric time  
 1122 window resulting in a ratio of  $\alpha \tau_- / \tau_+ > 1.0$  (Ren et al.,  
 1123 2010). We set  $\mu_{\pm}=1$  resulting in multiplicative STDP (in-  
 1124 between values lead to rules which have an intermediate  
 1125 dependence on the synaptic strength). Pyramidal cells re-  
 1126 ceive an unspecific background noise at 420 Hz during  
 1127 recall.

#### 1128 4.2.4 Two-network architecture and connectivity

1129 The network model includes two reciprocally con-  
 1130 nected networks, the Item and Context networks. For  
 1131 simplicity, we assumed that item and context infor-  
 1132 mation engage different modalities and cortical areas and  
 1133 thus the corresponding networks are located at a substan-  
 1134 tial distance (Table S2). Both networks span a regular-  
 1135 spaced grid of 12 HCs (Table S2), each with a diameter  
 1136 of 500  $\mu\text{m}$  (Mountcastle, 1997). Our model employs dis-  
 1137 tributed orthogonal representations with one active MC  
 1138 per HC, approximating the exceedingly sparse neocorti-  
 1139 cal activity patterns with marginal overlap. Each minicol-  
 1140 umn is composed of 30 pyramidal cells with shared selec-  
 1141 tivity, forming a functional (not strictly anatomical) col-  
 1142 umn. In total, the 24 HCs of the model contain 7200 ex-  
 1143 citatory and 480 inhibitory cells, significantly downsam-  
 1144 pling the number of MC per HC ( $\sim 100$  MC per HC in  
 1145 biological cortex). The high degree of recurrent connec-  
 1146 tivity within MCs (Thomson et al., 2002; Yoshimura and  
 1147 Callaway, 2005) and between them link coactive MCs  
 1148 into larger cell assemblies (Eyal et al., 2018; Binzegger  
 1149 et al., 2009; Muir et al., 2011; Stettler et al., 2002). Long-  
 1150 range bidirectional inter-area connections (item-context  
 1151 bindings or associative connections) are plastic (shown  
 1152 in Fig. 1A only for MC1 in HC1 of the Context net-  
 1153 work), binding items and contextual information (Ran-  
 1154 ganath, 2010). Recurrent connectivity establishes 100 ac-  
 1155 tive plastic synapses on average onto each pyramidal cell  
 1156 from other pyramids with the same selectivity, due to  
 1157 a sparse inter-area connectivity ( $cp_{PPA}$ ) and denser local  
 1158 connectivity ( $cp_{PP}$ ,  $cp_{PPL}$ ; connection probabilities are  
 1159 indicated in Fig. 1A only for MC1 in HC1 of the Con-  
 1160 text network). The model yields biologically plausible  
 1161 excitatory postsynaptic potentials (EPSPs) for connec-  
 1162 tions within HCs ( $0.45 \pm 0.13$  mV), measured at resting  
 1163 potential  $E_L$  (Thomson et al., 2002). Densely recurrent  
 1164 non-specific monosynaptic feedback inhibition mediated  
 1165 by fast spiking inhibitory cells (Kirkcaldie, 2012) imple-  
 1166

1166 ments a local winner-take-all structure (Binzegger et al.,  
1167 2009) amongst the functional columns. Inhibitory post-  
1168 synaptic potentials (IPSPs) have an amplitude of -1.160  
1169 mV ( $\pm 0.003$ ) measured at -60 mV (Thomson et al.,  
1170 2002). These bidirectional connections between basket  
1171 and pyramidal cells within the local HCs are drawn with  
1172 a 70% connection probability. Notably, double bouquet  
1173 cells shown in Figure 1A, are not explicitly simulated,  
1174 but their effect is nonetheless expressed by the BCPNN  
1175 rule. A recent study based on the same basic model archi-  
1176 tecture demonstrated that learned mono-synaptic inhibi-  
1177 tion between competing attractors is functionally equiva-  
1178 lent to the disynaptic inhibition mediated by double bou-  
1179 quet and basket cells (Chrysanthidis et al., 2019). Param-  
1180 eters characterising other neural and synaptic properties  
1181 including BCPNN can be found in Table S1.

1182 Figure 1B shows the weight distributions of em-  
1183 bedded distributed cell assemblies, representing different  
1184 memories stored in the Item and Context networks. At-  
1185 tractor projections can be further categorized into strong  
1186 local recurrent connectivity within HCs, and slightly  
1187 weaker long-range excitatory projections across HCs  
1188 (Fig. 1C).

#### 1189 4.2.5 Axonal conduction delays

1190 Conduction delays ( $t_{ij}$ ) between a presynaptic neuron  
1191  $i$  and a postsynaptic neuron  $j$  are calculated based on their  
1192 Euclidean distance,  $d$ , and a conduction velocity  $V$  (Eq.  
1193 11). Delays are randomly drawn from a normal distribu-  
1194 tion with a mean according to distance and conduction  
1195 velocity, with a relative standard deviation of 30% of the  
1196 mean. In addition, a minimal delay of 1.5 ms ( $t_{min}^{syn}$ , Ta-  
1197 ble S2) is added to reflect synaptic delays due to effects  
1198 that are not explicitly modelled, e.g. diffusion of neuro-  
1199 transmitters over the synaptic cleft, dendritic branching,  
1200 thickness of the cortical sheet and the spatial extent of  
1201 columns. Associative inter-area projections have a ten-  
1202 fold faster conduction speed than those within each net-  
1203 work, reflecting axonal myelination.

$$\bar{t}_{ij} = \frac{d}{V} + t_{min}^{syn}, \quad t_{ij} \sim N(\bar{t}_{ij}, .30 \bar{t}_{ij}) \quad (11)$$

#### 1204 4.2.6 Stimulation Protocol

1205 Noise input to pyramidal cells and fast spiking in-  
1206 hibitory basket cells is generated by two independent  
1207 Poisson generators with conductances of opposing signs.  
1208 Pyramidal cells coding for specific items and contexts are  
1209 stimulated with an additional specific excitation during  
1210 encoding and cued recall (all parameters in Table S2).  
1211 Item-context association encoding is preceded by a brief  
1212 period of background noise excitation to avoid initializa-  
1213 tion transients.

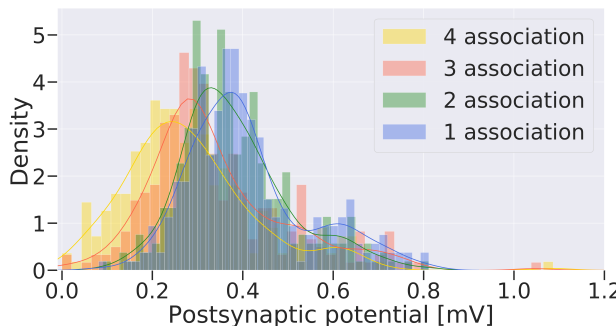
#### 1214 4.2.7 Attractor activation detector

1215 We detect and report cue-based activation of items or  
1216 contexts by utilizing an attractor activation detection al-  
1217 gorithm based on EMAs of spiking activity. Pattern-wise  
1218 EMAs are calculated using Equation 12, where the delta  
1219 function  $\delta$  denotes the spike events of a pattern-selective  
1220 neural population of  $n_{pop}=30$  pyramidal cells. The filter  
1221 time constant  $\tau=40$  ms is much larger than the sampling  
1222 time interval  $\Delta T=1$  ms.

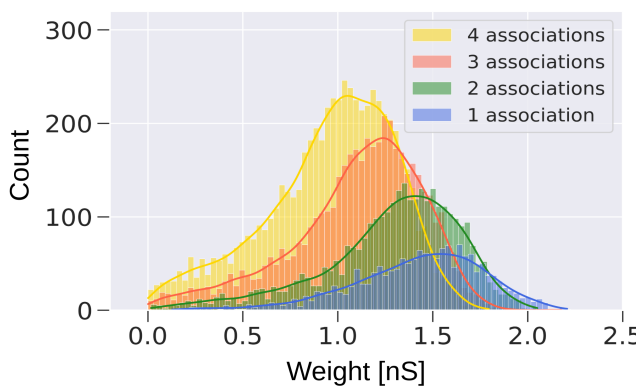
$$e_0 = 0, \quad e_t = \frac{\Delta T}{\tau} e_{t-\Delta T} + \delta_t \frac{1}{\tau n_{pop}} \quad (12)$$

1223 Pattern activations are detected by a simple threshold  
1224 ( $r_{th}$ ) at about tenfold the baseline activity with a small  
1225 caveat: To avoid premature offset detection due to syn-  
1226 chrony in fast spiking activity, we only count activations  
1227 as terminated if they do not cross the threshold again in  
1228 the next 40 ms. Despite the complications of nested os-  
1229 cillations, this method is highly robust due to the explo-  
1230 sive dynamics of recurrent spiking activity for activated  
1231 attractors in the network. Any attractor activation that  
1232 crosses this threshold for at least 40 ms is considered a  
1233 successful recall.

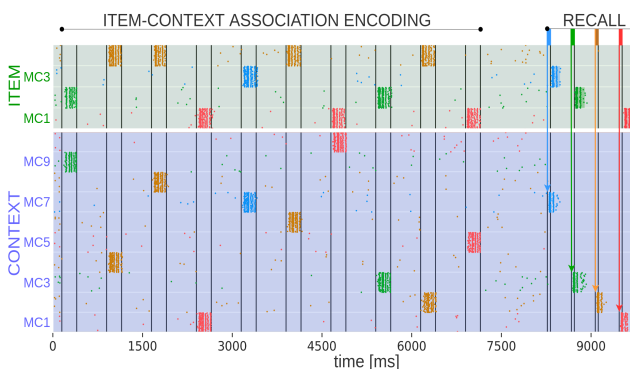
1234 **5 SUPPLEMENTARY MATERIAL**



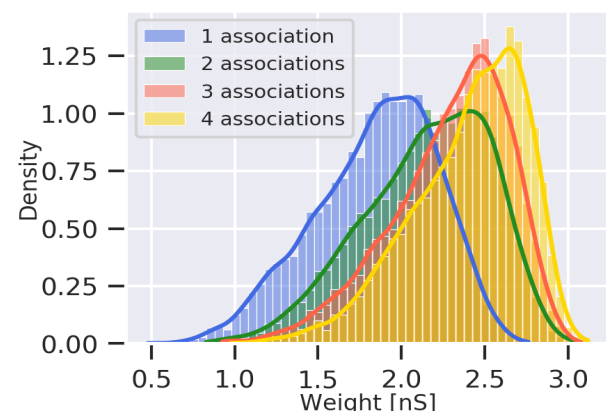
**Fig. S1** Excitatory postsynaptic potentials (EPSPs) for the binding between Item and Context networks. EPSPs were recorded (at resting potential  $E_L$ ) after item-context association encoding phase. We stimulate individually all the neurons in HC1 of an item which forms one, two, three or four associations and record the postsynaptic potential onto their associated context neurons. Means of the EPSP distributions show significant statistical difference ( $p<0.05$  for one vs two associations;  $p<0.001$  for two vs three and three vs four associations, Mann-Whitney,  $N=300$ ).



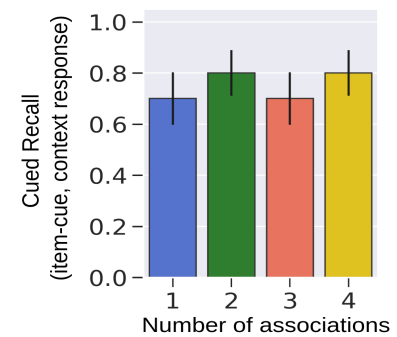
**Fig. S2** Distributions of the AMPA component weights between Item and Context networks. Slower NMDA receptor weights follow a similar pattern of weakening for items which participate in multiple associations. Means of the weight distributions of one, two, three, and four associations show significant statistical difference ( $p<0.001$ , Mann-Whitney,  $N=2000$ ).



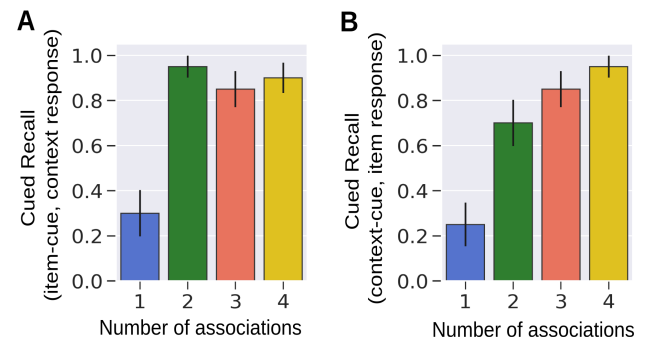
**Fig. S3** Spike raster of pyramidal cells in HC1 of both the Item and Context networks in the BCPNN model. Items and their corresponding context representations are simultaneously cued in their respective networks. The testing phase occurs 1 s after the encoding and triggers activations via partial cues of contexts (50 ms cues). Repetition of items across various contexts leads to progressive item-context decoupling. Item-4 is repeated across four different contexts, and while its associated context gets activated when cued (context-B), item-4 is not retrieved.



**Fig. S4** Weight distribution of AMPA component weights of the Item network including synaptic augmentation. The multiplicative effect of synaptic augmentation on the consolidated Items features stronger combined synaptic strength for items with higher context variability. Slower NMDA receptor weights follow a similar pattern. Means of the weight distributions of one, two, three, and four associations show significant statistical difference ( $p<0.001$ , Mann-Whitney,  $N=2000$ ).



**Fig. S5** Cued recall under STDP after removing synaptic augmentation. Average item-cued recall performance in the Context network (20 trials). To compensate for the removal of augmentation, we increased the stimulation rates and the synaptic gain leading to comparable elicited spiking activity. Error bars represent standard deviations of Bernoulli distributions.



**Fig. S6** Cued recall under STDP including intrinsic plasticity. **A)** Average item-cued recall performance in the Context network (20 trials). **B)** Average item-cued recall performance in the Context network. Episodic context retrieval is enhanced for high context variability predominantly because of intrinsic excitability dynamics and synaptic augmentation. We observe an opposite trend to the decontextualization effect seen in Figure 3C. Error bars represent standard deviations of Bernoulli distributions.

**Table S1** Model and BCPNN parameters

Parameter	Symbol	Value	Parameter	Symbol	Value	Parameter	Symbol	Value
Adaptation current	$b$	86 pA	Utilization factor	$U$	0.2	BCPNN AMPA gain	$w_{gain}^{AMPA}$	0.76 nS
Adaptation decay time constant	$\tau_{I_w}$	280 ms	Augmentation decay time constant	$\tau_A$	5 s	BCPNN NMDA gain	$w_{gain}^{NMDA}$	0.07 nS
Membrane capacitance	$C_m$	280 pF	Depression decay time constant	$\tau_D$	280 ms	BCPNN bias current gain	$\beta_{gain}$	40 pA
Leak reversal potential	$E_L$	-70.6 mV	AMPA synaptic time constant	$\tau^{AMPA}$	5 ms	BCPNN lowest spiking rate	$f_{min}$	0.2 Hz
Leak conductance	$g_L$	14 nS	NMDA synaptic time constant	$\tau^{NMDA}$	100 ms	BCPNN highest spiking rate	$f_{max}$	25 Hz
Upstroke slope factor	$\Delta_T$	3 mV	GABA synaptic time constant	$\tau^{GABA}$	5 ms	BCPNN lowest probability	$\epsilon$	0.01
Spike threshold	$V_t$	-55 mV	AMPA reversal potential	$E^{AMPA}$	0 mV	BCPNN Spike event duration	$t_{spike}$	1 ms
Spike reset potential	$V_r$	-60 mV	NMDA reversal potential	$E^{NMDA}$	0 mV	P trace time constant	$\tau_p$	15 s
Refractory period	$\tau_{ref}$	5 ms	GABA reversal potential	$E^{GABA}$	-75 mV	Modulated plasticity	$\kappa_{boost}$	2
			Regular plasticity	$\kappa_{normal}$	1			

**Table S2** Network layout, connectivity and stimulation protocol

Layout	Symbol	Value	Connectivity	Symbol	Value	Stimulation	Symbol	Value
Cortical patch size	$C_{ps}$	2.0 x 1.5 mm	Axonal Conduction Speed	$V$	0.2 m/s	Background noise PYR (encoding)	$r_{bg\_encoding}^{PYR}$	650 Hz
Simulated HCs	$n_{HC}$	12	Myelinated axonal speed	$V_{myel}$	2 m/s	Background noise PYR (recall)	$r_{bg\_recall}^{PYR}$	450 Hz
Simulated MCs	$n_{MC}$	120	Minimal synaptic delay	$t_{min}^{syn}$	1.5 ms	Background noise BA	$r_{bg}^{BA}$	75 Hz
Simulated MCs per HC	$n_{MC}^{HC}$	10	Hypercolumn diameter	$d_{HC}$	0.5 mm	Background conductance	$g_{bg}^{PYR,BA}$	$\pm 1.5$ nS
No. of items	$n_{ITEM}$	4 (from 10)	Distance between networks	$d_{CONTEXT}^{ITEM}$	10 mm	Stimulation duration	$t_{stim}$	250 ms
No. of contexts	$n_{CONTEXT}$	10 (from 10)	PYR-PYR recurrent cp	$cp_{pp}$	0.2	Stimulation rate	$r_{stim}$	500 Hz
Layer 2/3 pyramidal per MC	$n_{MC}^{PYR-L23}$	30	PYR-PYR long-range cp	$cp_{ppl}$	0.25	Cue stimulation length	$t_{cue}$	50 ms
Basket cells per MC	$n_{MC}^{Basket}$	2	PYR-PYR associative cp	$cp_{ppa}$	0.02	Cue stimulation rate	$r_{cue}$	400 Hz
MC grid size (Item + Context)	$G_{MC}^{TOTAL}$	24 x 10	PYR-BA cp, BA-PYR cp	$cp_{pb}, cp_{ba}$	0.7	Stimulation and cue conductance	$g_{stim}$	+1.5 nS
			PYR-BA cc	$g_{pb}$	3 nS	Interstimulus interval	$T_{stim}$	500 ms
			BA-PYR cc	$g_{bp}$	-7 nS	Attractor detection threshold	$r_{th}$	10 Hz

PYR, Pyramidal cell; BA, Basket cell.

cp, connection probability; cc, connection conductance.

**Table S3** STDP synaptic model parameters

Parameter	Symbol	Value
Weight initialization	$w_0$	0 nS
AMPA maximum allowed weight	$w_{max}^{AMPA}$	13.5 nS
NMDA maximum allowed weight	$w_{max}^{NMDA}$	3.5 nS
Learning rate	$\lambda$	0.01
Asymmetry parameter	$\alpha$	1.2
Weight dependence exponent, potentiation	$\mu_+$	1
Weight dependence exponent, depression	$\mu_-$	1
Symmetric time window	$\tau_{\pm}$	20 ms