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Hebbian, correlational learning provides a memory-less mechanism for Statistical Learning irrespective of implementational choices: Reply to Tovar and Westermann (2022)

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## Author Note

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#### Abstract

Statistical learning relies on detecting the frequency of co-occurrences of items and has been proposed to be crucial for a variety of learning problems, notably to learn and memorize words from fluent speech. Endress and Johnson (2021) (hereafter EJ) recently showed that such results can be explained based on simple memory-less correlational learning mechanisms such as Hebbian Learning. Tovar and Westermann (2022) (hereafter TW) reproduced these results with a different Hebbian model. We show that the main differences between the models are whether temporal decay acts on both the connection weights and the activations (in TW) or only on the activations (in EJ), and whether interference affects weights (in TW) or activations (in EJ). Given that weights and activations are linked through the Hebbian learning rule, the networks behave similarly. However, in contrast to TW, we do not believe that neurophysiological data are relevant to adjudicate between abstract psychological models with little biological detail. Taken together, both models show that different memory-less correlational learning mechanisms provide a parsimonious account of Statistical Learning results. They are consistent with evidence that Statistical Learning might not allow learners to learn and retain words, and Statistical Learning might support predictive processing instead.


Keywords: Statistical Learning; Implicit Learning; Transitional Probabilities; Neural Networks; Chunking

Hebbian, correlational learning provides a memory-less mechanism for Statistical Learning irrespective of implementational choices: Reply to Tovar and Westermann

Statistical learning relies on detecting the frequency of co-occurrences of items, and has been proposed to be crucial for a variety of learning problems (e.g., Aslin, Saffran, \& Newport, 1998; Kirkham, Slemmer, \& Johnson, 2002; Morgan, Fogel, Nair, \& Patel, 2019; Saffran, Aslin, \& Newport, 1996; Saffran, Newport, \& Aslin, 1996; Saffran \& Griepentrog, 2001; Stalinski \& Schellenberg, 2010; Turk-Browne \& Scholl, 2009; Verosky \& Morgan, 2021), notably learning words from fluent speech (Aslin et al., 1998; Aslin \& Newport, 2012; Saffran, Aslin, \& Newport, 1996; Saffran, Newport, \& Aslin, 1996). We recently showed that such results can be explained based on simple correlational learning mechanisms such as Hebbian Learning (Endress \& Johnson, 2021) (hereafter EJ). Tovar and Westermann (2022) (hereafter TW) reproduced these results with a slightly different model (with temporal decay acting on both the connection weights and the activations, rather than on only the activations, and interference affecting weights rather than activations), and offering different interpretations of some network parameters (e.g., conceiving of forgetting as decay).

Here, we first stress the common theoretical implications of both models: While Statistical Learning is often assumed to help learners learn (and thus memorize) words from fluent speech (e.g., Erickson, Thiessen, \& Estes, 2014; Graf-Estes, Evans, Alibali, \& Saffran, 2007; Isbilen, McCauley, Kidd, \& Christiansen, 2020; Karaman \& Hay, 2018; Shoaib, Wang, Hay, \& Lany, 2018), results from the tasks used to explore Statistical Learning can be explained by a memory-less correlational learning model. As a result, Statistical Learning might be more useful for predictive processing than for learning words per se (e.g., Endress \& de Seyssel, under review; Morgan et al., 2019;

Sherman \& Turk-Browne, 2020; Turk-Browne, Scholl, Johnson, \& Chun, 2010; Verosky \& Morgan, 2021). Following this, we briefly discus the differences between EJ's and TW's models. As already argued by EJ, we agree that different implementations of correlational learning are likely to result in fairly similar results. ${ }^{1}$ However, we also show that, contrary to TW's characterization of their model, activation decay is critical to their model's performance, and argue that models of psychological phenomena should be evaluated by their psychological predictions rather than by reference to their putative "biological plausibility" when neither model includes biophysical attributes.

## 1 A memory-less interpretation of Statistical Learning

One of the primary motivations of Statistical Learning is that it might allow learners to extract (and memorize) words from fluent speech (e.g., Aslin et al., 1998; Aslin \& Newport, 2012; Saffran, Aslin, \& Newport, 1996; Saffran, Newport, \& Aslin, 1996). Speech is often thought to be a continuous signal (but see Brentari, González, Seidl, \& Wilbur, 2011; Christophe, Mehler, \& Sebastian-Galles, 2001; Endress \& Hauser, 2010; Johnson \& Jusczyk, 2001; Johnson \& Seidl, 2009; Pilon, 1981; Shukla, Nespor, \& Mehler, 2007; Shukla, White, \& Aslin, 2011). As a result, to acquire any word, learners first need to learn where words start and where they end. To this end, they might use Transitional Probabilities (TPs) among syllables, that is, the conditional probability of a syllable $\sigma_{i+1}$ given a preceding syllable $\sigma_{i}, P\left(\sigma_{i} \sigma_{i+1}\right) / P\left(\sigma_{i}\right)$. Unpredictable transitions might indicate a word boundary, while relatively predictable transitions are likely located inside words. Humans are sensitive to TPs (Aslin et al.,

[^0]1998; Kirkham et al., 2002; Morgan et al., 2019; Saffran, Aslin, \& Newport, 1996; Saffran, Newport, \& Aslin, 1996; Saffran \& Griepentrog, 2001; Stalinski \& Schellenberg, 2010; Turk-Browne \& Scholl, 2009), and might use this sensitivity to memorize words (e.g., Erickson et al., 2014; Graf-Estes et al., 2007; Isbilen et al., 2020; Karaman \& Hay, 2018; Shoaib et al., 2018).

However, the evidence that Statistical Learning leads to memory for words is mixed at best (see Endress, Slone, \& Johnson, 2020 for a critical review). For example, when exposed to statistically structured sequences, participants are sometimes more familiar with high-TP items than with low-TP items, even when they have never encountered either of them and thus could not have memorized them (because the items are played backwards with respect to the familiarization sequence; Endress \& Wood, 2011; Jones \& Pashler, 2007; Turk-Browne \& Scholl, 2009). In other cases, participants are more familiar with high-TP items they have never heard or seen than with low-TP items they have encountered (Endress \& Langus, 2017; Endress \& Mehler, 2009). Further, when instructed to repeat back the items they remember from a statistically structured familiarization sequences, participants are unable to do so even when they learned the statistical structure of the stream (Endress \& de Seyssel, under review).

Such results thus suggest that Statistical Learning abilities do not necessarily support the formation of declarative memories for words. This interpretation mirrors earlier demonstrations of dissociations between Statistical Learning and declarative memory (e.g., Cohen \& Squire, 1980; Finn et al., 2016; Graf \& Mandler, 1984; Poldrack et al., 2001; Squire, 1992), and suggests that Statistical Learning might be more useful for predictive processing rather than declarative memory formation (e.g., Endress \& de Seyssel, under review; Morgan et al., 2019; Sherman \& Turk-Browne,

2020; Turk-Browne et al., 2010; Verosky \& Morgan, 2021). ${ }^{2}$ To the extent that Statistical Learning has a computational function (in Marr and Nishihara's (1992) sense, and is not a spandrel, Gould, Lewontin, Maynard Smith, \& Holliday, 1979), we thus surmise that its function is the prediction of future events.

Both EJ's and TW's models are consistent with this view. EJ simulated the results of a number of Statistical Learning results with a fully connected network where the strength of excitatory connections among neurons was tuned by Hebbian learning. That is, if two neurons are active simultaneously, their connection becomes strengthened ("what fires together wires together"). The network also comprised inhibitory connections among neurons. Further, the network had a "forgetting" mechanism, where activity decayed as time passed. After familiarization with a speech stream, the network was tested by recording the total activation when presented with different types of test items.

The basic result was that this fairly generic network accounted for a number of Statistical Learning results. Critically, given that all learning resided in the connection strengths, it could do so without any memory representations at all. In fact, just as in human participants (Endress \& Langus, 2017; Endress \& Mehler, 2009), the network activation was determined by the associative strength of the syllables in an item, irrespective of whether the network had encountered the item or not. As a result, the network had no memory representation of either item (or one would need to conclude that the network remembered items it has never encountered).

[^1]EJ also found that, to account for these Statistical Learning results, the forgetting rate needed to be reasonable. Rather unsurprisingly, if forgetting was so fast that neurons were never active together, no learning ensued. Conversely, if forgetting was so slow that all neurons were active simultaneously, all neurons formed connections, making these indiscriminate connections useless as an indicator of learning.

## 2 Differences between EJ's and TW's model

TW reproduced these results in a similar network, confirming that basic Hebbian learning mechanisms can explain Statistical Learning results, to some extent independently of how they are implemented. As far as we can see, there are four main differences between TW's and EJ's models. First, TW take issue with our characterization of decay as forgetting. Second, TW stress the importance of spreading activation. Third, TW evaluate learning by inspecting connections rather than activations. Fourth, instead of including separate inhibitory and decay/forgetting components that affect activations (and thus indirectly connection weights through the Hebbian learning rule), their model uses a modified Hebbian learning rule (with an additional parameter) where decay/forgetting affects weights (and thus indirectly activations); this learning rule also comprises a thresholding mechanism that presumably mimics the effects of mutual inhibition.

### 2.1 Forgetting vs. decay

Regarding the interpretation of EJ's "forgetting" parameter, TW "argue that [interpreting decay as forgetting] may be a misleading interpretation. Activation values from external stimuli in both artificial and biological networks are non-persistent but are constantly updated in response to changes in the environment (Huber \& O'Reilly, 2003)." We agree that, in our specific implementation, decay would also be a reasonable description of the phenomenon we tried to capture. That said, given the controversy over
whether decay plays a role in forgetting (see below), we believe that forgetting is a theoretically more neutral term, especially because the same phenomenon can likely be captured by manipulating the interference parameter rather than the decay parameter (see below). However, we would question to what extent results from single neuron recordings are relevant for psychological models that are not particularly plausible biologically to begin with; for example both EJ's and TW's "neurons" code for speaker-independent, phonological representations of syllables, which would presumably be encoded by some fairly abstract population code in actual brains (Pouget, Dayan, \& Zemel, 2000). Further, while decay has certainly been widely documented, so has persistent neural activity, which exists in various brain areas and taxa (e.g., Major \& Tank, 2004). As a result, neurophysiological findings may not be informative about psychological theories.

In fact, the question of whether time-based decay exists in memory is a controversial one in cognitive psychology. Under some circumstances, humans can remember thousands of items for hours or weeks (Brady, Konkle, Alvarez, \& Oliva, 2008; Standing, 1973); under other circumstances, very similar pictures disappear from memory after a few seconds but can be reviewed through repeated exposure (Endress \& Potter, 2014; Pertzov, Avidan, \& Zohary, 2009; Thunell \& Thorpe, 2019). Further, it is controversial whether there is any decay in Short-Term Memory at all, or whether all decreases in memory are due to interference (e.g., Baddeley \& Scott, 1971; Berman, Jonides, \& Lewis, 2009; Lewandowsky, Oberauer, \& Brown, 2009; Nairne, Whiteman, \& Kelley, 1999). We are thus open to different psychological interpretations of the forgetting parameter, and EJ already acknowledged the possibility that the effects of their forgetting parameter could likely be mimicked by tuning inhibition (see below).

In contrast, although TW argue that their "simulation results . . . challenge E\&J's
notion of activation decay as the key ingredient for Hebbian statistical learning"(p. ???), forgetting/decay is critical to their model. They use decay in two places. First, the activation of each input is maintained only for two time steps (at $90 \%$ for the second time step); given that the current input is likely the strongest activation at each time step, the effects are similar to a global forgetting parameter. Second, TW consider only activation greater than a certain threshold. While the effect of the latter seems to be a reduced overall magnitude of the weights, the former is critical for the results. To illustrate this fact, we exposed the network to the familiarization stream from Saffran, Aslin, and Newport's (1996) Experiment 2, and then recorded the weights in high-TP items ("words") and low-TP items (part-words, of BC:D and C:DE type, a difference that is irrelevant for the current purposes). We ran 1000 simulations with three version of TW's model: With the original decay function from TW ("Standard" in Figure 1), no forgetting at all (i.e., the input to each neurons was the cumulative sum of prior inputs; "Never" in Figure 1) and immediate forgetting (i.e., the activation decays immediately after presentation; "Immediate" in Figure 1). As shown in Figure 1, the network discriminated between words and part-words only using TW's decay function; as in EJ's simulations, all weights reach the maximum of 1.0 in the absence of decay, and reached zero with immediate forgetting. A suitably chosen decay parameter is thus crucial to TW's model. Be that as it might, we believe that the merits of psychological models should be evaluated by their empirical adequacy, and links between psychological parameters and neurobiological findings should be investigated empirically.

### 2.2 The role of spreading activation

TW stress the importance for spreading activation for network performance. We certainly agree, and, in their Section 2, EJ explained the role of spreading activation in detail. In fact, we suggested that spreading activation might be a more parsimonious


Figure 1. Average connection weights of the test items in a simulation of Saffran, Aslin, and Newport's (1996) Experiment 2, using TW's model. (Left) Simulations using decay parameters from TW's model. (Middle) Simulations with no activation decay. (Right) Simulations with immediate decay. High-TP items (words) are discriminated from low-TP items (part-words of different types) only with a suitable decay function. With no decay, all weights are maximal; with immediate forgetting, no connections are formed.
account of previous claims that Statistical Learning might lead to word-like memory representations (see Erickson et al., 2014; Graf-Estes et al., 2007; Isbilen et al., 2020; Karaman \& Hay, 2018; Shoaib et al., 2018 vs. Endress \& Langus, 2017; Endress et al., 2020).

Given the importance of spreading activation, it is surprising that TW evaluate their model by inspecting connections weights rather than by measuring activations. In fact, even in a network with uniform connections and no learning, it is hard to describe the network dynamics mathematically without resorting to simulation (Endress \& Szabó, 2020). Given that, in TW's model, interference and decay act on weights rather than activations, this problem might be somewhat reduced in their model. Still, just relying on the pattern of weights, it is hard to evaluate the dynamic interplay of first and higher order associations or the dynamic aspects of statical learning such as those revealed by electrophysiological recordings (Endress \& Fló, in preparation).

### 2.3 The effects of different learning rules

2.3.1 One vs. two component learning rules. The most critical difference between EJ and TWs models is the learning rule. TW's learning rule has two components. First, all weights undergo decay. This decay is proportional to the current weight and the product of the activations connected by that weight, that is

$$
\Delta_{\text {Decay }} W_{A B} \propto-W_{A B} \times \text { activation }_{A} \times \operatorname{activation}_{B},
$$

where $A$ and $B$ are two neurons. However, given that, even in the simple Hebbian learning rule

$$
\Delta W_{A B} \propto \operatorname{activation}_{A} \times \operatorname{activation}_{B}
$$

the weight change is proportional to product of the activations, the effects of decay on learning will be very similar irrespective of whether decay originates from weights, activations or, as in TW's model, both. However, in the absence of targeted experiments investigating the empirical adequacy of weight-based vs activation-based decay, the key result is that both formalisms account for Statistical Learning results in the absence of a memory mechanism.

The second component of TW's learning rule is the strengthening of associations according to the simple Hebbian learning rule above. Critically, however, TW's model strengthens connections only when the product of the activation exceeds an arbitrary threshold ( activation $_{A} \times$ activation $_{B}>\theta$ ). However, the effect of this thresholding is similar to inhibitory connections. To see why this is the case, consider two pairs of
neurons. The activations in each pair are roughly similar to each other, but the activation in the first pair is somewhat larger than in the second pair (i.e, activation $_{A} \approx$ activation $_{B}>$ activation $_{C} \approx$ activation $_{D}$ ). If there is inhibition, the first pair will reduce the activation of the second pair as long as the inhibitory input exceeds their excitatory input (though the difference does not necessarily disappear; Endress \& Szabó, 2020). Given that weight changes are proportional to the product of the corresponding activations, connections between neurons with greater coactivation will be strengthened to a greater extent, irrespective of whether this is implemented through inhibitory connections or through an co-activation-based threshold for learning.

Again, we believe that targeted psychological experiments are necessary to gauge the empirical adequacy of activation-based decay and inhibition (as in EJ) vs. weight-based decay and inhibition (as in TW). In fact, there is evidence for both kinds of processes. One the one hand, the type of lateral, activation-based inhibition assumed in EJ has been proposed as a psychological mechanism for phenomena from perception to attention to response inhibition (e.g., Desimone \& Duncan, 1995; Hampshire \& Sharp, 2015). On the other hand, to the extent that knowledge of TPs resides in connection weights, the finding that knowledge of TPs is forgotten after a few minutes (e.g., Karaman \& Hay, 2018; Vlach \& DeBrock, 2019) suggests that "weights" can be forgotten over time, which is consistent with TW's proposal of weight-based decay (though this forgetting might still occur through interference or decay).

For the current purposes, we just assume that some normalization mechanism keeps activations at a reasonable level, and believe that the question of whether normalization occurs through weight decay, weight interference or lateral inhibition is best answered through experimental rather than computational investigations. To the extent that biological plausibility is relevant for psychological models, the ubiquity of
lateral inhibition across domains and taxa certainly suggest that activation-based inhibition is no less plausible than weight-based inhibition.
2.3.2 Connection weights do not grow excessively in the absence of weight
forgetting. TW justified their two component learning rule in part by arguing that "it is not clear their [EJ's] model prevents excessive growth of connections" (p. ???). However, it is easy to see from EJ's Hebbian learning rule that the final weight of the connection between two neurons after $t$ time steps is proportional to the average coactivation of the neurons, $W_{A B}(t) \propto t \times\left\langle\right.$ activation $_{A} \times$ activation $\left._{B}\right\rangle$ (for $W_{A B}(0)=0$ ). As a result, if the activations remain in a reasonable range, so will the weights. This is confirmed when examining the connection weights after familiarization with a stream modeled after Saffran, Aslin, and Newport's (1996) Experiment 2. As shown in Figure 2, connection weights diverge for slow decay rates of up to .2 , but generally stay below or around 1 for faster decay rates. In other words, weights stay in a reasonable range for decay rates that led to learning in EJ's simulations; for decay rates that were too slow for learning to occur, weights diverge as well. This confirms our point above that qualitatively similar results can be achieved by controlling weights (and thus indirectly activations, as in TW's simulations) or by controlling activations (and thus indirectly weights, as in EJ's simulations).

### 2.4 Decay vs. interference

TW questioned EJ's rationale for not varying their interference parameter (p. ???). However, and as mentioned above, EJ argued that their "interference parameter might well mimic the role of forgetting," and thus simply sought to limit the number of moving parts in their model. To see why this is the case, consider a network of $N$ neurons that receive external stimulation in a regular sequence. In the absence noise, the activation change between times $t$ and $t+1$ is given by (exponential) decay (first term), spreading

## Average and maximum weights



Figure 2. Final weights after simulation of stream from Saffran, Aslin, and Newport's (1996), using EJ's model. Mean (left) and maximal (right) weights for slow (top) and fast (bottom) forgetting rates.
activation (second term), inhibition (third term) and external stimulation (fourth term).

$$
\begin{equation*}
x_{i}(t+1)-x_{i}(t)=-\lambda_{a} x_{i}(t)+\alpha \sum_{j \neq i} w_{i j} F\left(x_{j}\right)-\beta \sum_{j \neq i} F\left(x_{j}\right)+I(t) \tag{1}
\end{equation*}
$$

To see the relationship between decay and inhibition, we assume that excitatory connectivity is relatively sparse, and partition the neurons into a set of $K$ neurons with excitatory connections with target neuron $i$, and $N-K$ neurons with negligeable excitatory input to neuron $i$.

$$
\begin{align*}
x_{i}(t+1)-x_{i}(t)= & -\lambda_{a} x_{i}(t)+\alpha \sum_{j=1, j \neq i}^{K} w_{i j} F\left(x_{j}\right) \\
& -\beta \sum_{j=1, j \neq i}^{K} F\left(x_{j}\right)-\beta \sum_{j=K+1, j \neq i}^{N} F\left(x_{j}\right)  \tag{2}\\
& +I(t)
\end{align*}
$$

In the absence of external stimulation, the $N-K$ neurons not providing excitatory input to $i$ will provide periodic inhibitory input (Endress \& Fló, in preparation) that is generally unrelated to the activation of $i$. Averaged across time, this input thus mimics the effect of linear (rather than exponential) decay. For the $K$ neurons that provide excitatory input to $i$, their excitatory input is proportional to their inhibitory input. As the activation of $i$ is also an increasing function of this excitatory input, the inhibition might thus mimic exponential decay (though the specific functional form is more complex). Further, given that $i$ is presumably most active when closely associated neurons are active as well (and assuming sparse activations), this exponential-like inhibition is likely the dominant inhibitory input when $i$ has noteworthy activation.

In the presence of external stimulation, the excitatory input to $i$ from other neurons is no longer related to $i$ 's activation. However, given the symmetry of the Hebbian learning rule, $i$ will also excite the neurons it has excitatory connections with. As a result, these neurons will again provide inhibitory input that is an increasing function of $i$ 's activation, albeit with a time-lag. Consequently, the effects of inhibition and time-based decay can likely mimic one another. Critically, however, given that EJ's objective was to make the conceptual point that Statistical Learning results can be reproduced by a simple, memory-less correlational learning mechanism, they did not explore alternative implementations of this idea. Be that as it may, TW's model confirms that EJ's results can be reproduced with different implementations.

## 3 Conclusions

In sum, both EJ and TW show that a memory-less correlational learning mechanism can account for results from Statistical Learning studies, despite differences in implementation, irrespective of whether decay and inhibition affect activations or weights. ${ }^{3}$ As a result, to the extent that Statistical Learning supports declarative memory formation for words, relevant evidence is still required.

[^2]
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[^0]:    ${ }^{1}$ Historically, many authors have stressed the importance of correlational learning mechanisms (if not exactly Hebb's rule), from Hume's (1739/2003) theory of causation to collocation detection in natural language processing (Manning \& Schütze, 1999), though other authors questioned whether what appears to be correlational learning (e.g., conditioning) really reflects correlational learning mechanisms (Gallistel \& Gibbon, 2000). Our main assumption is that learning mechanisms that show properties of correlational learning (e.g., the effects of Hebbian learning) are psychologically plausible even though there are many other methods of detecting co-occurrences.

[^1]:    ${ }^{2}$ Under the premise that Statistical Learning is used for word learning, the finding that Statistical Learning might not require declarative memory might lead to the conclusion that word learning does not require declarative memory either. However, given the prima facie plausibility of the view that words need to be retrieved from memory (at least for production), it seems more plausible that declarative memory is involved in word learning, and that Statistical Learning abilities might thus not support the formation of memories for words. We thank an anonymous reviewer for suggesting this possibility.

[^2]:    ${ }^{3}$ A stronger argument for the implementation independence of our conclusions would rely on analytic results for classes of activation functions and learning rules. However, given that mathematical treatments of neural network properties (e.g., using statistical mechanics) usually assume some learning rule (e.g., Hopfield, 1982; Amit, 1989; Storkey, 1997; Huang, 2021), and that an analytic derivation of the asymptotic network behavior is challenging even for a simplified version of the current model with less complex stimuli (Endress \& Szabó, 2020), we believe that testing different implementations is important to strengthen the generality of our argument.

