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This need for multiple modifiable parameters arises even with a single statistic expressed as the synaptic weight. In addition, synapses may usefully store statistics accumulated over different timescales (e.g. transient and consolidated memory expressed through binary and graded mechanisms in series (1)), while variation of the parameters of synaptic dynamics (4) offers scope to express several statistics, requiring additional modifiable mechanisms.

Three distinct concepts:

1. How many (and which) local parameters are involved in causing long-lasting synaptic modifications?
   Many - e.g. pre- and post-synaptic electrical and chemical conditions, precise timings of activity, and neuromodulators.

2. How many (and which) independently modifiable mechanisms operate within a modifiable synapse?
   Several - often with different timecourses and conditions for modification, and with complex interaction.

3. How many parameters are needed to characterise changes of expressed function in a modified synapse?
   Possibly (and certainly in many models) as few as one - a synaptic ‘efficacy’ or ‘weight’ - though varying synaptic dynamics (e.g. facilitation and fatigue, etc.) may usefully express more than one.

Local synaptic computations

Modifiable synaptic parameters can only depend on the history of local conditions - not, for example, on patterns of activity across many cells. This presents an interesting and profound constraint on neural network computations, but from a theoretical standpoint there is no constraint on how many local parameters may be involved and how complex the functions may be (1, above).

A puzzle and a challenge:

Where modification is expressed by variation of a single synaptic weight (3, above) then it is tempting to think that a single modifiable storage mechanism (2, above) would suffice. Is this correct?

*This poster aims to show that the answer is NO! Multiple independent storage mechanisms are sometimes necessary within a synapse to compute potentially important functions, even when these are expressed through only a single parameter.*

This argument is not the only reason why synapses might require multiple modifiable mechanisms - models have suggested useful roles for independent mechanisms that have different timecourses of memory retention and for ways in which the dynamics, as well as the strength, of a synapse may be varied. But the issue addressed here is particularly interesting because it may seem counter-intuitive.

What does a Hebb synapse compute?

The Hebb synapse (and its many variants) strengthen:

“when the presynaptic terminal contributes to firing the postsynaptic cell”

Such potentiation is often said to depend on pre- and post- association. But strictly, it depends not on a statistical association of pre- and post-synaptic firing, but on temporal coincidence (within some time-frame) of such firing, which may be due to chance. The distinction can be crucial when learning is to be used for inference.
Modulation of pre- & post-synaptic firing coincidences without statistical association

Blue lines show the probabilities of independent pre- and post-synaptic firing and the conjoint firing ($P_{A&B} = P_A \cdot P_B$). Black lines show running synaptic frequency estimates based on single weight parameters that undergo fixed increments when the events occur, and exponential relaxation at other times (time constant 20 units). The graph based on coincidences ($A&B$) is analogous to a Hebb synapse, with substantial coincidence-dependent potentiation despite the absence of any pre & post-synaptic association.

What is an appropriate synaptic measure of statistical association?

Neurons make a ‘decision’ about when to fire on the basis of ‘evidence’ in the activity of their afferent axons. In many learning situations a Bayesian approach to this decision seems appropriate, where the evidence is used to establish the conditional probability that, with such evidence in the past, the postsynaptic cell has actually fired. When there is no association (i.e. pre- and post-synaptic firing have been statistically independent) then the pre-synaptic firing provides no evidence about whether firing should currently be elicited.

Since simple dendrites tend often to sum synaptic currents approximately linearly, the appropriate synaptic strength should on this basis be an evidence function ($\varepsilon$) that sums linearly for different (sufficiently independent) pieces of evidence, to compute a conditional probability. This is the log likelihood ratio: -

$$\varepsilon = \text{Evidence for firing of B, given firing of A} = \log \left( \frac{P(A | B)}{P(A | \text{not-B})} \right)$$ [1]

where $P(A|B)$ means the conditional probability of A, given B. The summed synaptic influence, given such a measure of association, is the increment (above an a priori level without any evidence) for $\log(P/(1-P))$ for the firing of cell B, known as a belief function $\beta$ or log-odds: -

$$\beta = \log \left( \frac{P(B)}{1-P(B)} \right) = \sum (\varepsilon \text{ from afferent fibres}) + \beta_0$$ [2]

Computation of an evidence function

Evidence [1, above] is fairly easily computed, but depends on the full 3 degrees of freedom of the contingency table for the combined probabilities of two random variables (pre- and post-synaptic firing). It requires either 3 or (with loss of information about the rate at which data has been collected - ok if associations are assumed to be unvarying) at least 2 modifiable synaptic mechanisms for storage of independent variables. Simply storing the current evidence function $\varepsilon$ itself is not sufficient, because the way it changes in response to a particular contingency, like the joint firing of A and B, depends not just on the current value of $\varepsilon$, but on the separate values of other parameters, such as the conditional probabilities $P(A/B)$ and $P(A/\text{not-B})$. 
Simulation results
(mean ± s.d. from 10 simulations)

1. Estimates of pre-, post- and paired firing probability per time unit, with a relaxation time constant of 100 units. True probabilities shown in blue. Each estimate would require one modifiable mechanism and one stored parameter.

2. Evidence for firing of B conveyed by firing of the pre-synaptic axon A, calculated from the above 3 parameters.

\[ \varepsilon = \ln \left( \frac{P(A\&B)(1-P(B))}{(P(A)-P(A\&B))P(B)} \right) \]

Note reduced s.d. during periods with more information.

Evidence estimated with just 2 modifiable parameters

In a steady state, evidence can be computed from just 2 independently variable parameters. One way to do this uses one parameter \( \omega \) that is pre- and post- dependent while the other \( \gamma \) is purely post- dependent.

\[ \Delta \omega = (1-\omega)\delta \] if A&B active
\[ \Delta \omega = -\omega \delta \] if A active alone
\[ \Delta \omega = 0 \] if B active alone

(\( \delta = .025 \))

The simulation uses the odds ratio for firing of B given A [\( \omega = P(A\&B)/(P(A)-P(A\&B)) \)] and the odds ratio for firing of B itself [\( \gamma = P(B)/(1-P(B)) \)]. Computation equations are above. Note that 1/\( \gamma \) rather than \( \gamma \) is graphed, to be analogous to a component of synaptic efficacy, though \( \gamma \) itself could be modelled by spine conductance.

Evidence to be summed across active synapses is computed as \( \ln(\omega/\gamma) \), or approximated by simpler functions. With only 2 stored parameters, changes of probabilities, even with no statistical association, can lead to marked transient errors of evidence estimation, as at *.
Summary
- Every expressed synaptic parameter that is modifiable during learning may (depending on an aspect of the complexity of its computation*) require two or more separately variable storage mechanisms within the synapse for its computation and correct updating.
- A Bayesian approach to synaptic computation, in which the manipulated parameters are probabilities, can give insight into the possible nature and complexity of elementary synaptic learning processes.
- Appropriate manipulation of probability estimates depends on the statistical model of underlying causes (especially in a changing environment) and may require modulation of elementary synaptic computation for its optimisation.
- Constraints of realistic physiology (for example the fact that synapses probably do not switch between excitation and inhibition - analogous to evidence for and against activation) provide interesting challenges for efficient design.
- There is seldom talk of ways that modifiable synapses might adaptively change the effect they have on dendrites when they are not active. This might be:-
  (i) a trick that evolution missed (failing to convey useful evidence based on when an axon is silent),
  or (ii) quantitatively unimportant (because axons are silent most of the time),
  or (iii) something simply experimentally less tractable than changes of the response to stimulation.
- Can anyone put this in more precise mathematical terminology?

HOW MANY MODIFIABLE MECHANISMS DO MODIFIABLE SYNAPSES NEED?
Gardner-Medwin, AR, Dept of Physiology, University College London, London WC1E 6BT, UK

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takes two (at least) to tango