Synaptic competition in structural plasticity and cognitive function

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Connections between neurons can undergo long-lasting changes in synaptic strength correlating with changes in structure. These events require the synthesis of new proteins, the availability of which can lead to cooperative and competitive interactions between synapses for the expression of plasticity. These processes can occur over limited spatial distances and temporal periods, defining dendritic regions over which activity may be integrated and could lead to the physical rewiring of synapses into functional groups. Such clustering of inputs may increase the computational power of neurons by allowing information to be combined in a greater than additive manner. The availability of new proteins may be a key modulatory step towards activity-dependent, long-term growth or elimination of spines necessary for remodelling of connections. Thus, the aberrant growth or shrinkage of dendritic spines could occur if protein levels are misregulated. Indeed, such perturbations can be seen in several mental retardation disorders, wherein either too much or too little protein translation exists, matching an observed increase or decrease in spine density, respectively. Cellular events which alter protein availability could relieve a constraint on synaptic competition and disturb synaptic clustering mechanisms. These changes may be detrimental to modifications in neural circuitry following activity.

1. Functional and structural plasticity

During development, the nervous system undergoes an important period of synaptogenesis in which the proper connections are established between neurons. This period is hallmarked by robust formation and pruning of synapses, a process that is shaped by activity [1]. Much less is known about the relationship between physical synaptic changes and learning in the mature brain and whether remodelling of connections can occur. The discovery of long-term potentiation (LTP) 40 years ago shed light on a unique and critical ability of neurons to change the efficacy of their connections upon learning [2]. Depending on the nature of the activity, bidirectional modifications of synaptic connections can occur. However, these changes need to be physically stored over long periods of time and much less is known about how this is accomplished; specifically, what is the relationship between the structure and function of synapses. The direct connection between these first became clear following experiments in which new spine growth upon LTP induction was demonstrated [3]. The advent of precise glutamate uncaging, in combination with two-photon imaging at single spines, allowed for the induction of synaptic plasticity at the level of single inputs that showed a direct physical enlargement of the stimulated spines [4]. A linear relationship between spine volume and the amount of current in a spine was also established [5]. Analogously, synaptic depression has been shown to lead to spine shrinkage [6,7], although the specific effects of activity on single spines have not been defined as precisely as for the case of synaptic potentiation. Taken together, these findings support the idea that bidirectional changes in efficacy correlate with bidirectional structural modifications.
2. Cooperation and competition

Notably, there are still many details to be worked out regarding how different forms of activity are stored at the synaptic level, especially with regard to changes that are long lasting. One reason for this is owing to an important mechanistic difference between short-lasting (less than 2 h) and long-lasting (3 h or more) plasticity, in that the former does not require protein synthesis, while the latter does (reviewed in [8]). These changes add an extra dimension of complexity because the availability of new proteins contributes to the facilitation of plasticity between synapses, a phenomenon that was first described by the synaptic tagging and capture hypothesis [9,10]. It describes cooperativity between differentially activated inputs, in which strong stimulation of one pathway contributes to the expression of long-lasting plasticity at the second, weakly stimulated pathway, when new proteins are synthesized and shared. The impact of protein availability on the kinetics of synaptic cooperation is highlighted by experiments probing these interactions at individual spines. When protein independent forms of plasticity are induced, cooperation between inputs, termed ‘cross-talk’, occurs for a limited time and over short distances (10 min/10 μm) [11]. By contrast, if new proteins are produced, cooperation between inputs can occur for up to 70 μm and over the course of 1.5 h [12]. The presence of proteins confers plasticity even to synapses stimulated with subthreshold activity, and can lead to potentiation and spine growth at these sites that last for hours. Indeed, earlier experiments demonstrating that dendritic stimulation-induced growth of multiple spines within a 70 μm distance hinted that such interactions could occur, although the specific location and number of stimulated spines were unknown [13]. Therefore, depending on the nature of the activity, cellular mechanisms may be activated which lead to the growth of neighbours over protracted spatial and temporal parameters. Interestingly, protein cooperation can occur even between synapses expressing different forms of plasticity. A series of experiments by Sajikumar & Frey [14] and Sajikumar et al. [15] showed that induction of LTP can be coopted to enable long-lasting long-term depression (LTD; termed cross-tagging). This implies that a similar set of proteins may be required for different types of plasticity, however, not all the proteins required were the same. The blockade of PKMζ selectively inhibited LTP while still allowing for the conversion of LTD from a short- to a long-lasting form [15]. It will be interesting to see in future studies whether and how specific proteins contribute to activity-dependent structural changes.

Facilitative interactions are not the only ones influenced by the presence of newly made proteins. The dendritic availability of new proteins, or limited availability, sets up a competitive environment for the expression of plasticity if these proteins need to be shared. Indeed, field recording experiments demonstrated competition for the functional expression of LTP when multiple pathways are stimulated under protein synthesis blockade [16]. On a different scale, competition for structural plasticity is observed between neighbouring spines (approx. 20 μm apart) when protein availability is limited. After coactivation of multiple spines under these conditions, their physical growth was negatively correlated with one another [12]. Thus, the availability of proteins during long-lasting forms of plasticity may contribute to the physical organization of synapses by providing a constraint on how many coactive inputs ultimately cooperate with one another at a particular time and within a particular space. Determining the specific parameters over which competition functions will be key to understanding the learning rules that underlie structural plasticity and neuronal function.

3. Spine shrinkage and elimination with long-term depression

While protein-driven cooperative and competitive interactions during potentiation have begun to be characterized, much less is known about whether such interactions, and their resulting structural consequences, occur during protein synthesis-dependent synaptic depression. There is some evidence to support the idea that bidirectional structural changes can occur. Upon the induction of NMDA-mediated synaptic depression, which does not require protein synthesis, shrinkage or retraction of spines can occur [6,7,17]. As the presence of proteins has been shown to facilitate spine growth, it was unclear how they would impact structural changes following synaptic depression. A recent study by Ramiro-Cortés and Israelly examines this question by studying the structural correlates of a well-known protein synthesis-dependent form of synaptic depression mediated by metabotropic glutamate receptors (mGluRs) [18,19]. Inducing depression with the group I mGluR agonist DHPG led to the robust shrinkage and elimination of spines in a protein synthesis-dependent manner, which importantly, could be observed for up to 24 h [19]. An interesting question still to be addressed is whether structural correlates of synaptic depression exist at the single spine level, and if so, whether proteins can facilitate functional and structural plasticity between multiple synapses. Thus far, the induction of structural LTD through glutamate uncaging at single spines has been biased towards occurring through NMDA-dependent mechanisms [17] and does not necessarily lead to a reduction in spine volumes [17]. It remains to be determined whether protein synthesis-dependent depression can be induced structurally at the level of single spines or whether it requires the activation of several inputs.

It is worth mentioning that the bidirectional protein synthesis-dependent structural plasticity discussed thus far, appears to be more universal in its ability to induce structural changes at spines of all sizes, compared with its shorter lived counterpart. Specifically, it has been reported that protein synthesis-independent synaptic potentiation is easier to induce at smaller spines [4], and likewise these smaller spines are more likely to shrink following NMDA-mediated synaptic depression [17] (figure 1). Conversely, strong potentiation leads to growth of spines of different sizes when new proteins are made [12]. Similarly to the case of potentiation, protein synthesis-dependent LTD leads to spine shrinkage and elimination that is independent of spine size [19] (figure 1). Although spine shrinkage or elimination induced by mGluR-LTD requires protein synthesis [19] and the pool of proteins required seems to be similar to those needed for late-LTP [14], it is unknown which or how new proteins can lead to these structural changes. One candidate is Arc/Arg3.1, which is translated upon mGluR-LTD induction, leading to the endocytosis of AMPA receptors on postsynapses [20,21]. Endocytosis may be one way by which to reduce spine surface area, and this could provide a mechanism by which to effect concurrent spine shrinkage and depression of synaptic conductances following activity [22]. It remains to
be determined how specific molecular pathways are involved in implementing the functional consequences of LTD. These findings highlight protein synthesis-dependent synaptic plasticity as a potentially robust mechanism by which to remodel spines of various sizes, a mechanism that may be useful in order to rewire neuronal connections, as discussed below.

Overall, the data summarized above indicate that the initiation and maintenance of long-lasting, bidirectional structural changes requires protein synthesis, and that within a dendritic domain, spines can compete or cooperate in order to express the different forms of plasticity. These interactions may comprise a mechanism by which to remodel synaptic connectivity depending on the nature of the activity. In order for such remodelling to occur, certain inputs would be potentiated while others would be reduced or lost, through growth or elimination of spines, respectively. Thus, plasticity mechanisms which extend beyond the level of a single synapse may have significant implications for how a neuron’s structure is shaped in the long term.

4. Synapse clustering

One such outcome could be the clustering of synapses, which has been the subject of growing interest over recent years owing to its implications for the storage capacity of a neuron and a network. Synapses do not necessarily act as independent, linearly summed units on the dendritic arbour; instead, if multiple nearby synapses are activated in concert, supralinear processing can occur (for review, see [23]). This supralinear processing is caused by a local depolarization giving rise to a dendritic spike, facilitated by voltage-gated channels in the cell membrane. This spike will travel efficiently to the soma and has a high probability of making the cell fire. Thus, synapses which are activated concurrently and within a small spatial range will have a disproportionately large effect on the cell’s output, because only a small number need to be activated to cause firing [24–26]. If a neuron’s inputs could be rewired to achieve a clustered organization, the cell would be able to distinguish a larger number of patterns and have a higher processing power [24,25,27]. Furthermore, the potential to disconnect and reconnect the cells can change the overall wiring diagram of the network, allowing it to store more information [28].

Recently, there have been various findings which have shown functional clustering of synapses—i.e. the concurrent activation of inputs closely together on a dendritic branch—both in vitro and in vivo [29–31]. These studies used either calcium imaging to directly observe synapse activation [29,31] or the presence of tagged GluA1 receptors in the synapse as a marker of recent LTP [30], to conclude that in pyramidal neurons of the hippocampus or neocortex, synapses which are close together have an increased likelihood of being activated within a short time frame of one another. In these studies, such clustering was observed at different stages of development, from just after birth to young adult, and over distances of 8–20 μm. Interestingly, these spatial parameters fit well within the bounds of cooperative and competitive plasticity mechanisms that occur during synaptic potentiation. It will be interesting to determine whether the spatial organization of synapses influences how specific proteins act, depending on the type of plasticity and the type of cooperativity that is induced [32].

If the functional clustering discussed above could drive the physical creation of anatomically distinct groups of spines, then it should be possible to find evidence of these structural clusters in vivo. Indeed, Yadav et al. [33] found that anatomical spine clusters (groups of three or more) occur significantly more frequently than chance within apical dendrites of layer III cells of the monkey prefrontal cortex. This study reveals the capability of neurons to spatially organize spines, but it does not prove a causal relationship between functional and structural changes. This latter point requires that the formation of clusters be driven by activity or learning. To begin to address this issue, two studies have looked for cluster formation in neural regions that are involved in encoding a specific task; they find that learning increases the prevalence of clustered synapses in those particular areas. In one case, rearing owls wearing prism goggles lead to an increase in clustered spines specifically in fields of the barn owl auditory system that correspond to the newly shifted topographic map [34]. Subsequently, learning a forelimb reaching task in mice was shown to correlate with the appearance of clustered spines in the motor cortex [35]. These studies suggest that activity in a given region could lead to structural clustering. It will be important to determine which activity patterns lead to branch specific clustering, and if concurrent activation of spines during learning precedes their structural organization. This

![Figure 1. Structural plasticity mediated by LTD.](http://rstb.royalsocietypublishing.org/)

NMDA-mediated LTD

![LTD](http://rstb.royalsocietypublishing.org/)

mGluR-mediated LTD

![LTD](http://rstb.royalsocietypublishing.org/)

(protein synthesis)
question will require the monitoring of both the activity and the structure of dendritic regions throughout the learning process.

5. Synaptic interactions leading to spine clustering

The findings detailed above make a plausible argument for the clustering of dendritic spines in response to activity. Such remodelling could benefit the system by increasing the computational capacity of a neuron and circuit. For example, clustered synapses could more efficiently summate weak inputs and lead to neuronal firing. Remaining to be determined is what type of activity could lead to cluster formation; over what time scale would the formation of clusters be advantageous as a learning mechanism and what are the exact events by which activity could promote clusters. As outlined in figure 2, we propose that the following steps, which combine synaptic tagging and capture with spine remodelling and turnover, may be an avenue by which synaptic clustering can occur on the dendrite. Some of these have experimental support, whereas others will require further investigation in order to ascertain their biological relevance.

6. Activity-dependent structural correlates in vivo

A crucial mechanistic step to achieving activity-dependent structural clustering would be the ability to physically re-model connections, potentially through spine loss and gain. Chronic two-photon imaging in mice has allowed for the examination of spine dynamics in vivo over the course of many months (for reviews, see [35,36]). These data indicate that in the adult brain, different pools of spines exist—some of which turn over and some of which appear stable over the lifetime of the animal. These two elements provide essential components of a system which retains the ability to learn into adulthood, while being able to store long-lasting memories. Further studies have investigated the link between learning and spine remodelling in vivo. Some of the observations include: (i) spine formation—following a successful reaching task [37]; (ii) spine loss—associated with Pavlovian fear conditioning [38] and (iii) increased spine turnover-related to songbird learning [39]. All of these demonstrate structural flexibility of neurons in the respective brain areas involved in the encoding of behaviours. They also show that not all learning necessarily results in the same types of structural modifications. These results add weight to the idea that spine dynamics, and hence structural plasticity, may be the substrates for the storage of information.

7. Behavioural implications

The availability of proteins for cooperative and competitive interactions may have interesting outcomes for how information is processed by neurons during the encoding of behaviours. Indeed, the necessity for new protein synthesis has previously been established to be important for long-lasting memory formation [40]. Therefore, the coincidence of a salient event that leads to the production of new proteins may facilitate the subsequent encoding of otherwise weakly relevant information, potentially within the same engram. Indeed, experiments in which such ‘behavioural tagging’ was tested recently demonstrated the facilitation of memory storage [41]. When animals were trained in an inhibitory avoidance paradigm, which normally produces short-term learning, they were able to perform the task for up to an hour. However, when they were exposed to a novel open field prior to the training, the inhibitory avoidance learning lasted for over 24 h. This effect, which required dopamine receptors, was protein synthesis dependent because the infusion of the inhibitor anisomycin into the hippocampus prevented this facilitation. Thus, the exposure to novelty, which probably releases dopamine-induced protein synthesis allowing the facilitation of subsequent learning [41]. An additional study similarly found that novelty-induced
behavioural tagging could occur during the learning of a
non-fear inducing spatial learning task in which animals
learned to locate a food reward within an arena [42]. Although
it is not known which neural circuits encode this information, it
is intriguing to consider that the various aspects of such events
may be encoded within the same synaptic clusters. In this way,
the availability of proteins during information processing
would facilitate cognitive function by enhancing the ability to
bind divergent, yet relevant, information together. Importantly,
our brain is continuously presented with new information, yet
not all of this is incorporated into any given memory. Perhaps a
limited window of protein availability could serve to include
timely experiences into a common learning trace, while avoiding
the random incorporation of irrelevant information into any active circuit.

8. What is the relevance of synaptic competition
for brain function?

We have discussed the fact that synaptic and structural plasticity
can occur over a defined region when multiple inputs are
coactive, by competing for limiting proteins that are necessary
for the expression of plasticity. The physical boundaries
over which these processes function could delineate a region
for cooperative and competitive interactions, whereby spines
could either grow or shrink. In addition to these potentiation
mechanisms, competition for long-lasting synaptic depression
(requiring protein availability) may also be involved in establishing
which spines are lost or maintained, although this has
yet to be explored experimentally. In this way, protein avail-
ability could select a subset of synapses to be incorporated
into a physical cluster through either synaptic potentiation or
depression. Therefore, normal levels of protein synthesis
could provide an optimal functional range for: (i) the integra-
tion of information over long time scales and (ii) the refi-
nement of neuronal connections through competitive inter-
actions. An imbalance of protein translation could therefore
affect the optimal range of protein concentration and result in
altered spine densities (figure 3).

Figure 3. Hypothetical structural outcomes depending on protein availability. The distribution of spines on a dendritic arbour may be shaped by the balance between cooperation and competition among synapses for the expression of plasticity. The overall level of activity which a neuron experiences may contribute to maintaining an optimal balance of protein synthesis. In cases where mutations drive an increase or decrease in the general amount of proteins made, a reciprocal increase or decrease in spine density may result in a scenario which could change synaptic connectivity and neural circuits. (Online version in colour.)

Is there evidence to suggest that competitive mechanisms
are involved in information storage? One clue may come
from systems in which the balance of protein availability
is biased towards one direction or another, shifting the
ideal level necessary for neuronal function. Indeed, several
mental retardation disorders have in common mutations
that lead to an upregulation of protein synthesis [43,44].
A well-known example of this can be seen in the case of
Fragile X, the most prevalent genetic disorder leading to cog-
nitive deficits, when loss of the FMR1 gene product leads to
an increase in protein synthesis through overactive mGluR
signalling [45]. A key route to initiating translation is through the
protein mammalian target of Rapamycin (mTOR) [46].
Mutations in several proteins, which regulate this pathway
through PTEN/P13K/AKT signalling, have also been linked
to autism spectrum disorders, and show cognitive impair-
ments. Specifically, these include mutations of TSC1 and
TSC2 in tuberous sclerosis, NF1 in neurofibromatosis type
1, and PTEN hamartoma tumour syndrome [47]. For each
of these cases, the proteins involved negatively regulate the
mTOR pathway and their loss of function results in an
increase in protein synthesis. Importantly, these mutations
also result in structural alterations of increased spine density
[48–51]. Thus, the resulting increase in protein levels in these
disorders may be responsible for both the functional and
structural abnormalities observed, perhaps by relieving a
constraint on competitive processes that occur in neurons
during the encoding of information.

The above described disorders contain mutations which
lead to increased protein synthesis, and hence protein
availability. How would a reciprocal decrease in protein trans-
lation affect cognitive function and dendritic structure? Some
clues can be gleaned from the following two examples of
Down syndrome and Rett syndrome, in which indeed cogni-
tive dysfunction and reduced dendritic spine density are observed.
In both of these disorders, perhaps owing to the
many genes affected in each, the underlying mechanisms are
not fully understood; however, there is evidence to suggest
that the downregulation of protein translation could be
partially responsible. In Down syndrome, chromosomal tripli-
cation of 21q results in the increased gene dosage of over 300
genes. However, a critical region was identified in rare cases
with truncated duplications, from which the DSCR1 gene
was identified (DSCR1) [52]. This protein was recently shown
to interact with Fragile X mental retardation 1 protein (FMRP)
and enhance its function in translational repression [53]. Thus,
a mutation which leads to the opposite cellular effect of Fragile
X, also demonstrates the opposite structural changes. In the case
of Rett syndrome, mutations in the DNA methyltransferase
MeCP2, which regulates the transcriptional silencing of genes,
were found to be responsible for the disorder [54]. Recent evi-
dence indicates that MeCP2 regulation correlates with the
activity-dependent induction of BDNF transcription, and results
in reduced dendritic growth and spine maturation [55]. As
BDNF is known to trigger long-lasting synaptic plasticity and
protein synthesis [56], chronic reductions in its transcription
could lead to a state of decreased protein availability in the
neuron, particularly in regions undergoing synaptic plasticity.
Reductions in protein translation may affect the strengthening
of inputs and predispose neurons to undergo dendritic spine
pruning. Thus, assessing competitive plasticity mechanisms in
these backgrounds will provide important insights regarding
the role of protein availability during plasticity.
9. Concluding remarks

Since the original discovery of LTP, we have learned much about the complexity involved in modifying synaptic weights. We have discussed above how varied plasticity mechanisms requiring the synthesis of new proteins can effect long-lasting changes in synaptic and structural plasticity. These include events that lead to the strengthening of some synapses or that induce the shrinkage and elimination of others. In both cases, the presence of newly made proteins is critical for long-lasting structural changes. These processes may specifically delineate dendritic regions over which inputs can become integrated, and therefore may drive the physical creation of spine clusters. Such physical rewiring into clusters has implications for short-term processing of information, enhancing the subsequent efficacy of activity by optimizing dendritic integration. Importantly, such interactions may require an optimal balance of protein availability to allow for synaptic competition to occur between the inputs. In fact, interfering with this balance, either through blockade of protein synthesis or blockade of protein degradation, is detrimental to synaptic function [57,58]. In the absence of such plasticity constraints, the physical organization of spines may become compromised. Indeed, when too many proteins are made, for example in the case of several mental retardation disorders, spine density is aberrantly increased, while the counterpart is also true (figure 3). The production of too few proteins leads to reduced dendritic spine density in certain examples of cognitive dysfunction. Thus, in the long term, the connectivity of neural circuits may be impacted if spines cannot properly cooperate or compete. Further defining the learning rules for structural plasticity will be important for understanding how activity can shape connectivity in the brain.

Funding statement. This work was supported by the Champalimaud Foundation, Gulbenkian Foundation, Bial Foundation, Fundação para Ciência e Tecnologia and CONACyT.

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