Trends in Biochemical Sciences

Opinion

Learning of Signaling Networks: Molecular Mechanisms



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Molecular processes of neuronal learning have been well described. However, learning mechanisms of non-neuronal cells are not yet fully understood at the molecular level. Here, we discuss molecular mechanisms of cellular learning, including conformational memory of intrinsically disordered proteins (IDPs) and prions, signaling cascades, protein translocation, RNAs [miRNA and long noncoding RNA (IncRNA)], and chromatin memory. We hypothesize that these processes constitute the learning of signaling networks and correspond to a generalized Hebbian learning process of single, non-neuronal cells, and we discuss how cellular learning may open novel directions in drug design and inspire new artificial intelligence methods.

Learning of Non-neural Cells: Adaptive Molecular Responses Observed When the Same Stimulus is Repeated

Molecular mechanisms of neuronal learning have been well established in the past decades [1]. However, we know relatively little about the molecular details of learning mechanisms of nonneuronal cells. We define cellular learning as an adaptive response to a simple stimulus observed when the same stimulus is repeated in a short time (as compared with the duration of the cell cycle of the given cell). We note that it is crucial to discriminate between molecular changes that are indeed adaptive, and those that are fortuitous byproducts of other, co-occurring adaptive phenomena. To make this distinction, we focus on the molecular mechanisms of adaptation at the unicellular level and do not detail the long-term, multistep processes of cell reprogramming, development, or disease formation. In these latter cases, many individual learning steps may be interwoven and may reflect adaptation to unrelated events. Due to similar reasons, we do not cover the formation of intergenerational, **epigenetic memory** (see Glossary) and do not detail evolutionary processes. Finally, we focus on non-neuronal cells, since we want to concentrate on the adaptation of the **signaling network** of a single, non-neuronal cell and not that of multicellular networks connecting several neuronal cells.

It is well known from many experimental observations that cellular responses change, that is, they can become either faster and/or stronger or weaker upon repeated stimuli. For example, budding yeast cells displayed a faster reactivation of the inositol-3-phosphate synthase (Ino1) and galactokinase (Gal1) enzymes after a previous activating stimulus [2] and developed a **molecular memory** of a previous heat stress lasting several generations [3]. This, and a few other examples, show that the molecular mechanisms we describe here may go beyond the single cell cycle time frame of cellular learning as defined in the current opinion article, and that forming a molecular memory of the cell promotes intergenerational, epigenetic learning. We will compare cellular learning and molecular memory formation in the Concluding Remarks section in detail.

Additional examples have been described. In *Arabidopsis*, exposure to the damage signaling hormone, jasmonic acid, caused a stronger response to consecutive dehydration stress [4];

Highlights

Besides the well-known learning processes of neurons, non-neuronal, single cells are able to learn and show a more robust (and often faster) adaptive response when the same stimulus is repeated.

Known examples of cellular learning are sensitization- or habituation-type responses.

Several molecular mechanisms of neuronal learning, such as conformational memory, protein translocation, signaling cascades, miRNAs, IncRNAs, and chromatin memory, also participate in learning of non-neuronal, single cells.

We propose that these molecular mechanisms form the integrative memory of signaling networks and display a generalized Hebbian learning process by increasing those edge weights through which the signal has been propagated.

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also in *Arabidopsis*, the molecular memory of a previous heat shock was maintained for several days [5]. Similarly, rice developed molecular memory of drought stress [6]. Additionally, mouse fibroblasts showed a faster and stronger response to the second interferon-β stimulus given one day after the first [7]. Murine CD8⁺ memory T cells displayed a stronger response if they were re-exposed to *Listeria monocytogenes* 48 hours after the first infection [8]. Importantly, cellular learning can also result in the repression of a response. In budding yeast cells, the *STL1* sugar transporter gene showed a reduced expression to the second hyperosmotic stress as compared with the first [9]. In *Arabidopsis*, a subset of *MYC*-dependent genes related to multiple abiotic and hormone response networks did not respond to repeated dehydration stress [10]. Finally, mouse macrophages developed an immune tolerance after repeated lipopolysaccharide exposure [11]. The activation and repression described in these examples resemble the classical learning types: **sensitization** and **habituation** [1], respectively. Regrettably, other **hallmarks of learning**, such as **conditioning** [1], better recognition of the signal from its partial representation, or increased tolerance to noise, have not yet been convincingly demonstrated at the single-cell level.

Here, we provide examples of how different levels of cellular architecture, such as **intrinsically disordered proteins (IDPs)**, signaling cascades, translocating proteins, RNAs, and chromatin structure, contribute to cellular learning. **Conformational memory** (including that of IDPs), signal integration by signaling cascades, and **protein translocation** may be considered as a faster phase of cellular learning. RNA-based (such as miRNA or IncRNA) molecular memory and many forms of **chromatin memory** develop more slowly but have a longer duration. We demonstrate, through the example of the **epithelial–mesenchymal transition** network [12], how these elements of cellular learning at single-cell level are all organized in one signaling network that potentially possesses a learning capability at multiple levels. As the major hypothesis of our paper, we propose that, in signaling networks, cellular learning may be interpreted as a generalized **Hebbian learning** process [1] in which weights of **network edges** of signaling networks where the signal has propagated become increased (i.e., molecular connections become stronger) during the adaptive changes. This novel, integrative understanding of cellular learning may lead to new artificial intelligence and drug design technologies.

Conformational Memory

Several proteins display conformational memory, whereby the protein transiently keeps its active conformation after the dissociation from its former binding partner [13]. Examples include the active state of the endocytosed, unliganded integrin receptor [14] as well as the sarcoplasmic/ endoplasmic reticulum Ca²⁺ATPase (SERCA) [15]. Here, we propose that conformational memory may participate in the molecular memory formation of single cells (Figure 1). Importantly, the process of increased binding affinity of 'protein B', having a conformational memory, to its signaling neighbor 'protein A', is the same as the signaling network representation of the Hebbian learning process [1], where the network edge weight of two signal-transducing neighbors (characterizing the strength of their association) will increase because of the signaling networks and may have key roles in cellular learning processes.

IDPs are proteins that lack an organized 3D structure, whereas intrinsically disordered regions (IDRs) are disordered segments (i.e., loops, linkers) of at least 20 amino acids in length located in otherwise ordered proteins. Intrinsic disorder can be found in 85% of human signaling proteins [13,16]. Importantly, IDRs regulate organized protein cores in several protein kinases [17], and often act as **molecular switches** that can change the direction of signal propagation [18]. IDRs are enriched in sites of post-translational modifications and are often alternatively spliced

Glossary

Anti-Hebbian learning: a learning process where edge weights are not increasing (as in Hebbian learning) but decreasing.

Chromatin memory: altered 3D structure of the chromatin which provides different accessibility of genes for transcription after a repeated signal; histone and DNA modifications may play a role in its development.

Conditioning: a learning procedure in which stimulus A is paired with stimulus B, and the response to stimulus B becomes activated after stimulus A alone.

Conformational memory: an 'active' (e.g., high-affinity, binding-competent) conformation of a protein which is adopted upon binding to its partner (which may be a substrate, another protein, or a membrane) and does not relax back to the original, 'inactive' conformation before the next activation of the same protein.

Epigenetic memory: a heritable, intergenerational change in gene expression or behavior that is induced by a previous signal.

Epithelial-mesenchymal transition: a process by which epithelial cells lose their cell polarity and cell-cell adhesion, and gain migratory and invasive properties to become mesenchymal cells; occurring (among others) in wound healing, organ fibrosis, and initiation of metastasis in cancer progression.

Habituation: a form of nonassociative learning where a response to a stimulus decreases after repeated or prolonged presentations of that stimulus.

Hallmarks of learning: sensitization, habituation, or conditioning are the major hallmarks of learning processes; in addition, discrimination of different signals, reconstruction of the signal from its partial representation, and noise tolerance are also considered to be hallmarks.

Hebbian learning: in the current opinion article, the classical, neuronal Hebbian learning is generalized as an intracellular process of single, nonneuronal cells, which increases those edge weights (i.e., strengths of molecular connections) of signaling networks where the signal has propagated.

Intrinsically disordered proteins (IDPs): proteins that do not have an ordered 3D structure; in many proteins, structural disorder only extends to a





Figure 1. Conformational Memory of Signaling Proteins as a Potential Form of Cellular Learning. Conformational memory, whereby proteins transiently keep their binding competent state after dissociation, is a well-established phenomenon [13–15]. For example, the integrin receptor (β1 subunit) [14], sarcoplasmic/endoplasmic reticulum Ca²⁺ATPase (SERCA) [15], and prion-like proteins [3,20,21] all possess conformational memory and participate in cellular learning. Here, we illustrate the steps of a conformational memory-mediated learning process on the example of an intrinsically disordered protein (IDP). Importantly, 85% of human signaling proteins contain intrinsically disordered regions (IDRs), which opens the possibility of the transient stabilization of their signal-induced folding [13,16]. (1) The first signal induces the association of neighboring proteins A and B in a signaling cascade, which induces a binding-competent conformation as a conformational memory. (3) If the first signal is repeated soon, the second signal finds protein B (e.g., via folding of an IDR of protein B) [58]. (2) After the IDR of protein B), protein B keeps its binding-competent conformation as a conformational memory. (3) If the first signal is repeated soon, the second signal finds protein B still in a binding-competent state, which causes a faster and more robust signal transmission. The signal-induced conformational memory of protein B increases the binding affinity between protein A and protein B. Note, this is exactly the same as the signal inducer representation of the Hebbian learning rule [1], where the edge weight of two signal transducing neighbors increases because of the signaling process.

[13], they may have conserved molecular features, such as subcellular localization, membrane transport, motor activity, ribosomal function, etc. [19]. Thus, IDPs/IDRs are good candidates for the conformational memory providing fast cellular learning (Figure 1).

Prion proteins are enriched for structural disorder and represent another form of conformational memory. A conformational switch may convert prions to a β -sheet-enriched form, making extensive aggregates. Chaperones are required for prion formation but may also erase prion memory in cases of severe stress [13]. For example, in budding yeast cells, the prion form of Pin1 maintained the molecular memory of a previous heat stress for subsequent generations [3]. Additionally, the neuronal cytoplasmic polyadenylation element binding (CPEB) protein of the mollusk *Aplysia* can undergo a prion-like conformational transformation and behave as a molecular switch perpetuating molecular memory for years [20]. These observations confirmed the earlier hypothesis that prions may participate in memory formation [21].

segment (intrinsically disordered region, IDR) of the protein.

Molecular memory: molecular mechanisms inducing cellular learning either within a single cell cycle or by developing intergenerational, epigenetic memory.

Molecular switch: a molecule that can be reversibly shifted between two or more stable states.

Moonlighting proteins: proteins that perform multiple functions, often in different cellular locations, and/or participate in different protein complexes.

Network edge: connection between two network nodes (i.e., basic elements of the network); network edges may be weighted, directed, and may have sign (i.e., they may encode activation or inhibition).

Oja's rule: introduces a 'forgetting' term to the Hebbian learning rule making sure that the sum of total edge weights should not increase.

Prion proteins: misfolded proteins capable of transmitting their misfolded conformation to normal variants of the same protein.

Protein translocation: signal-induced relocalization of proteins between subcellular compartments.

Sensitization: a form of nonassociative learning where a repeated stimulus results in the amplification of its response.

Signaling network: a directed network of proteins and RNAs participating in cellular signaling processes.

System-level memory: a form of molecular memory which is not provided by individual signaling molecules but by the concerted activation of signaling pathways.

Transcriptional memory: a set of modifications of DNA and DNA-binding proteins (primarily histones) regulating the accessibility of genes for transcription.



System-Level Memory of Signaling Cascades and Protein–Protein Interaction Networks

Francis Crick proposed in 1984 that a signaling complex as simple as a protein kinase and a phosphoprotein phosphatase pair may display molecular memory, preserving its active or inactive state despite the turnover of its constituent proteins [22]. Later studies defined a prominent role of molecular switches in molecular memory formation (including the role of IDPs) [18]. A recent model uncovered how larger segments of the signaling network develop cooperation-based, **system-level memory**. The mitogen-activated protein kinase (MAPK) cascade displays a rich repertoire of transient adaptive responses characterized by both frequency and amplitude modulations. Different relaxation rates of cascade components lead to 'postactivation bursts', keeping the cascade in an 'activation-competent' state. This can form a system-level memory of the first activation, making later responses faster and more robust [23]. Such a short-term molecular memory was demonstrated in the yeast osmotic stress response too. If osmotic stress was repeated within several minutes, members of the Hog1 signaling pathway were still phosphorylated and thus 'awaited' the next signal in a preactivated state [24]. These examples show how the concerted activation of signaling cascades may contribute to cellular learning.

Protein–protein interactions constitute an essential element of signaling networks. Yet, many of them are not directly involved in building up signaling networks but rather function in modifying their behavior. Weak protein–protein interactions give rise to 'noise' that diminishes the efficiency of information transmission. Increased interaction strength helps information transmission but slows down response dynamics, showing a tradeoff between efficiency and responsiveness [25]. Molecular chaperones increase the frequency of out-of-equilibrium states and help the 'disorganization' of protein segments [26]. Thus, chaperones may act both as facilitators of molecular memory formation and as a 'forgetting mechanism.' These examples show how protein–protein interactions may fine-tune cellular learning of signaling networks.

Subcellular Protein Translocation

Signal-induced protein translocation (triggered by, e.g., phosphorylation) between two cellular compartments is a widespread phenomenon potentially affecting thousands of human proteins [27]. Protein translocation is actively involved in the reconfiguration of signaling networks in cellular learning processes. For example, inhibition of NF- κ B p65 nuclear translocation disrupted the formation of both CD8⁺ memory T cells and memory B cells [28,29]. Further, protein kinase C β II-induced upregulation and mitochondrial translocation of the adaptor protein p66SHC, was associated with the formation of hyperglycemic molecular memory of human aortic endothelial cells (Figure 2) [30]. Protein translocation may also occur between subcompartments of a cellular organelle, such as in the nucleus or in the form of the formation of biomolecular condensates by liquid–liquid phase separation [31]. Protein translocation establishes a whole set of novel protein–protein interactions, increasing their edge weights in signaling networks.

Moonlighting proteins perform multiple functions, often in different locations, resulting from protein translocation [27]. For example, multiple interactions between the moonlighting immuno-modulatory activities of acute phase proteins and monocyte-derived dendritic cells play a key role in forming immunological memory [32].

RNA-Based Molecular Memories

Various types of RNAs were also shown to participate in cellular learning processes. Since RNAs have a short lifetime, their *de novo* transcription is needed to initiate their effects. MiRNAs decrease the protein expression noise of hundreds of lowly expressed proteins [33], increasing the noise tolerance and thus robustness of cellular learning by reducing gene expression





Figure 2. Subcellular Translocation as a Form of Cellular Learning. Signal-induced translocation of proteins between subcellular compartments is a widespread phenomenon [27]. Many of these processes may participate in cellular learning. As an example, here we show the protein kinase C ßII (PKCßII)-induced upregulation and mitochondrial translocation of the adaptor and reactive oxygen species (ROS) sensor protein, p66Shc, which is associated with forming hyperglycemic molecular memory of human aortic endothelial cells [30]. (1) Hyperglycemia leads to PKCßII-induced phosphorylation and mitochondrial translocation of p66Shc, which induces ROS production. (2) Persistent hyperglycemia upregulates ROS and, consequently, PKCßII and translocated p66Shc, which leads to a vicious circle. (3) After restored normoglycemia, p66Shc remains in the mitochondria, causing a hyperglycemic molecular memory characterized by increased production of ROS.

variability. Molecular memory formation was shown by miRNA-156, which post-transcriptionally downregulated SPL transcription factor genes in the plant *Arabidopsis thaliana*, causing the development of thermotolerance and thus conferring a molecular memory of a previous heat shock. This molecular memory was maintained for several days (Figure 3A) [5]. Additionally, miRNA-221 and miRNA-222-induced inhibition of macrophage activity during the development of lipopolysaccharide tolerance [11]. The miRNA cluster 17–92 was transiently induced after T cell activation. Both the induction and later silencing of the miRNA-17–19 cluster were mandatory to the development of CD8⁺ memory T cells [34]. Further, miRNA-21 was involved in the development of fibrotic mechanical memory of mesenchymal stem cells [35]. These examples show the widespread involvement of miRNAs in both sensitization- and habituation-type cellular learning processes. MiRNA induction can be perceived in signaling networks as an increased edge weight of participating miRNAs.

Other types of RNA have been shown to contribute to cellular learning. In particular, IncRNAs participated in forming molecular memory of rice drought stress [6] and the development of CD8⁺ memory T cells after viral infection [36]. In contrast, a IncRNA originating at –2700 upstream of the budding yeast HO endonuclease erased previous molecular memory of nutrient deprivation- or pheromone-induced cell cycle arrest [37]. These examples elucidate the





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Figure 3. Various Forms of RNA-Based and Chromatin Memory. (A) RNA-based molecular memory. Many types of RNA molecules, such as miRNA and long noncoding RNA (IncRNA), participate in cellular learning. As an example, (1) the priming heat shock upregulated miRNA-156 (with the help of the argonaute RNA-induced silencing complex, AGO1), which (2) post-transcriptionally downregulated the SPL2/11 transcriptional repressor, allowing the synthesis of heat

(Figure legend continued at the bottom of the next page.)



RNA-dependent regulation of molecular memory formation, showing the richness of the contribution of various RNAs to cellular learning processes in signaling networks.

Chromatin Memory

Histone modifications (including histone-methylation, -phosphorylation, -acetylation, -ubiquitylation, and -sumoylation) as well as DNA methylation (occurring at adenine and cytosine nucleotides and often forming CpG dinucleotides, especially in mammals) also play key roles in cellular learning. These processes are called **transcriptional memory**. Changes in histone acetylation can occur on a much faster timescale than those in DNA methylation [38]. Lysine methylation of histone H3 participates in both sensitization and habituation of *Arabidopsis* [4,10], in sensitization of mouse fibroblasts and human HeLa cells by interferon- β and - γ , respectively [7,39], as well as in CD8⁺ memory T cell formation [8]. In the study of Komori *et al.* [40], 466 CpG dinucleotides of 132 genes displayed differential DNA methylation between naive and memory CD4⁺ T lymphocytes. Erasure of DNA methylation ('forgetting') can be performed via ten–eleven translocation (TET) DNA demethylases [41].

The 3D chromatin structure also plays an important role in cellular learning. Sensitization to hyperosmotic stress was abrogated if the reporter gene was placed to a pericentromic chromatin domain in yeast cells [9]. Nup2-mediated association of the *INO1* and *GAL1* genes with the nuclear pore complex and histone modifications led to the rapid reactivation of *INO1* and *GAL1* genes after a repeated signal. Both the Set1/COMPASS methyl-transferase complex and the Mediator complex were remodeled in these processes (Figure 3B) [2]. The human MHC class II gene DRA was persistently relocated to promyelocytic leukemia nuclear bodies after interferon- γ treatment, causing a sensitization to a subsequent interferon- γ stimulus [39]. Increased transcription by changes in chromatin organization can be perceived as increasing edge weights of transcription factors in signaling networks. It is a question for future studies, whether shape fluctuations [42] or rotation of cell nucleus [43] also play a role in cellular learning processes.

Learning of Signaling Networks

All the processes we described so far constitute changes of signaling networks playing a role in integrative cellular learning and molecular memory formation. We illustrate the potential cellular learning mechanisms of signaling networks by the epithelial–mesenchymal transition network of hepatocellular carcinoma cells as described by Reka Albert and her group (Figure 4, Key Figure) [12]. As shown in Figure 4, many nodes of this signaling network may participate in one or more mechanisms of cellular learning (i.e., possessing intrinsic disorder, participating in translocation, being an RNA, or being a protein regulated by chromatin changes). Note that these adaptive changes all recalibrate the edge weights of signaling networks. Increasing the edge weights of those connections of the signaling network, which have been activated by the incoming signal, corresponds to a Hebbian learning process of the signaling network of single, non-neuronal

shock proteins (HSPs) and ascorbate peroxidase (APX2) in the plant *Arabidopsis thaliana*. These molecular mechanisms caused the development of thermotolerance, thus (3) induced an increased survival upon a second, larger, lethal heat shock due to the previous, priming heat shock. This molecular memory was maintained for several days [5]. (B) Chromatin memory. Chromatin segments remain open and accessible long after the repeated signal due to persistent histone acetylation and DNA demethylation [8]. Gene demarcation and gene association with the nuclear pore complex are forms of global chromatin rearrangements leading to the development of molecular memory. As an example, (1) the yeast genes of inositol-3-phosphate synthase (*INO1*) and galactokinase (*GAL1*) associate with the nuclear periphery via the nuclear pore complex component, Nup2. (2) This association, together with histone methylation, acetylation, the incorporation of the specific histone variant, H2A.Z, as well as a modified preinitiation complex (mPIC) lacking the Kin28 CTD kinase, (3) led to a rapid reactivation of *INO1* and *GAL1* after a repeated signal [2]. (B) was modified with permission using Figure 8 from [2].



Key Figure

Learning of Signaling Networks



Figure 4. We illustrate various potential cellular learning mechanisms in the signaling network regulating epithelial–mesenchymal transition of hepatocellular carcinoma cells [12]. The large pale blue rectangle represents the hepatocellular carcinoma cell. Grey arrows and black dot-head arrows mark activations and inhibitions, respectively. Intrinsically disordered proteins (IDPs), identified using the DisProt database [59], which may possess conformational memory, are marked with starbursts. Proteins potentially participating in subcellular translocation (identified as high confidence translocating proteins in the Translocatome database [27]) are marked with light green rectangles. The participating miRNA is marked with a dark green rectangle. Targets of chromatin regulators in hepatocellular carcinoma (collected from the CR2Cancer database [60]) are marked with dark green edges. Note that epithelial–mesenchymal transition (EMT) has many more participating RNAs [61] than miRNA-200 of the original network [12]. The addition of more RNAs and chromatin regulators (like histone modifiers or DNA methylases) will be a logical step of future work. Though the examples depicted are not complete, it is obvious that many nodes of this signaling network may participate in one or more mechanisms of cellular learning. Purple arrows highlight those nodes, which have already been identified as part of the mechanisms inducing epithelial–mesenchymal transition memory [44–46]. All of these nodes possess one or more features identified as potential mechanisms of cellular learning in our opinion article.



cells. We note that, for example, in case of a habituation response, certain edge weights may also decrease, which is a molecular example of **anti-Hebbian learning**. The demonstration of learning of signaling networks and the extension of the Hebbian learning process to the molecular level of single cells are the major novel concepts of our opinion paper.

Several observations showed that epithelial–mesenchymal transition has a molecular memory [44–46]. Purple arrows in Figure 4 point to those nodes, which have already been identified as participants in these processes. Importantly, all of these nodes possess one or more features that pertain to the potential mechanisms of cellular learning discussed herein.

Signaling networks may be extended by cytoskeletal [47] and interorganelle [48] networks as well as by intercellular signaling [49], filamental [50], and membrane [51] networks. These networks may all have a potential role, heretofore not exactly described, in promoting cellular learning of non-neuronal cells.

Concluding Remarks

In this opinion article, we have shown how conformational memory of proteins, signaling cascades, subcellular protein translocation, various RNA molecules, and chromatin memory can result in integrative learning of signaling networks in single, non-neuronal cells. We hypothesize that signaling networks of non-neuronal cells display features of Hebbian learning [1] by increasing the strength of molecular connections between signaling molecules involved. We believe the examples outlined herein demonstrate that various molecular mechanisms develop two major types of cellular learning: sensitization and habituation. However, the direct demonstration of more complex forms of cellular learning remained notoriously difficult in non-neuronal cells.

In a network description, a neuron corresponds to a single node at the neuronal network level, while the same neuron contains, as one of its segments, its own signaling network (thus, in the pre- and postsynaptic neurons, two of these signaling networks become joined). We would like to emphasize that learning at both levels, the single-cell signaling network and the multicellular neuronal network, uses the same underlying molecular mechanisms elucidated here (such as conformational memory, signaling cascades, protein translocation, miRNA, and chromatin memory). However, neuronal learning displays several molecular forms even in a single neuron (such as synaptic densities, changes of membrane potentials, etc.), which are not characteristic of the cellular learning process of a single, non-neuronal cell. Obviously, neuronal learning also mobilizes the enormous potential of multicellular, neuronal networks, which, by definition, cannot be reached at the single-cell level. Thus, evidently, multicellular, neuronal networks allow the development of incomparably more sophisticated learning processes than those of single, non-neuronal cells described in this paper.

The formation of transgenerational (epigenetic) memory also uses many of the molecular mechanisms of the intragenerational, cellular learning listed here (e.g., DNA and histone methylation and related chromatin rearrangements [2,40,41,52] as well as protein compartmentalization [53], miRNAs [54], and prions [3]). These mechanisms build up the molecular memory of the individual cell, lasting for single or multiple cell cycles. However, short-term changes, such as conformational memory of IDPs and changes of signaling cascades, may not be extended for multiple cell cycles and thus may only participate in the *sensu stricto* cellular learning we defined in this opinion paper and not in epigenetic memory formation. We note that later experiments will certainly provide a solid basis to extend the molecular mechanisms of cellular learning far beyond a single cell cycle.

Outstanding Questions

Are there different types of molecular mechanisms for different types of cellular learning, like sensitization or habituation?

How are the elements of the Hebbian learning process of cellular learning, highlighted in this paper, coordinated at the signaling network level?

Are the mechanisms of cellular forgetting (i.e., erasure of molecular memory) coordinated at the signaling network level?

What is conformational memory's contribution to cellular learning?

How many intrinsically disordered regions (IDRs) do actually fold after a signal-dependent binding event to the signaling partner protein or to the membrane?

How long does the folded IDR stay folded after the signal's termination, thus (presumably), the dissociation of the signal-induced protein complex?

What are the roles of cytoskeletal, interorganelle, and intercellular networks as well as liquid–liquid phase separation in the formation of molecular memory?

How do nuclear dynamics contribute to cellular learning?



It is an open question (see Outstanding Questions) whether different types of cellular learning (e.g., sensitization and habituation) proceed via different or similar mechanisms. Current data have not yet been examined in detail to elucidate the molecular mechanism(s) of these phenomena in the same system. Another important open question is how forgetting of cellular learning proceeds. Forgetting introduces the **Oja's rule** to Hebbian learning, preventing the 'overexcitation' of the network due to the continuous growth of its edge weights.

A better understanding of cellular learning processes will inspire progress in several areas of science. A recent paper on non-Markovian chemical reaction networks on gene expression showed that molecular memory of protein synthesis and degradation may induce feedback, bimodality, switch behavior, and may fine-tune gene expression noise [55]. These findings open the possibility that our concept of generalized Hebbian learning may be extended to metabolic and other types of molecular networks in the future. Chemically induced proximity between two adjacent signaling molecules by a drug became a recent drug design paradigm [56]. Enhanced proximity in these therapeutic approaches may also mimic the effect of cellular learning. Chromatin modifier drugs are already used in anticancer therapy [57]. We expect a much wider use of drugs targeting the cellular learning mechanisms described in this paper in the future. In the analogy of genetic algorithms and neural networks, cellular learning may also inspire novel artificial intelligence methods. Cellular learning is a research area that will show dramatic progress in the coming years, and we are happy to invite our colleagues to join these efforts.

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