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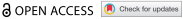
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REVIEW



Spinal AMPA receptors: Amenable players in central sensitization for chronic pain therapy?

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ABSTRACT

The activity-dependent trafficking of AMPA receptors (AMPAR) mediates synaptic strength and plasticity, while the perturbed trafficking of the receptors of different subunit compositions has been linked to memory impairment and to causing neuropathology. In the spinal cord, nociceptive-induced changes in AMPAR trafficking determine the central sensitization of the dorsal horn (DH): changes in AMPAR subunit composition compromise the balance between synaptic excitation and inhibition, rendering interneurons hyperexcitable to afferent inputs, and promoting Ca²⁺ influx into the DH neurons, thereby amplifying neuronal hyperexcitability. The DH circuits become over-excitable and carry out aberrant sensory processing; this causes an increase in pain sensation in central sensory pathways, giving rise to chronic pain syndrome. Current knowledge of the contribution of spinal AMPAR to the cellular mechanisms relating to chronic pain provides opportunities for developing target-based therapies for chronic pain intervention.

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AMPA receptors; AMPAR trafficking; GluA1-4 subunits; the dorsal horn (DH) of the spinal cord; sensory DH neurons; nociceptive circuits; chronic pain

Introduction: Implication of spinal AMPAR for chronic pain

Chronic pain is a major healthcare issue worldwide; it has a serious impact on an individual's life and work activities, as well as on society [1]. It has been estimated that about 20% of the population in Europe suffers from persistent or chronic pain [2,3], with even more in the United States, with a figure ranging from 20% to 40% [4]. This problem is growing each year, in both absolute numbers and in terms of its distribution across the global population [5]. The International Association for the Study of Pain (IASP) has announced the 2020 Global Year for the Prevention of Pain to help focus on disseminating pain prevention strategies among researchers, clinicians, and every involved party for a global view on pain. Through the cooperation between the IASP and the World Health Organization, the definition of chronic pain has been revised and will appear in the upcoming edition of the International Classification of Diseases in the form of a new classification: chronic pain as a disease in its own right [6,7]. In such a context, chronic pain requires specialized, targeted treatment, i.e. with no regard to the trigger(s) that give rise to the development of chronic pain syndrome. To date, however, there is no available therapy for curing chronic pain that is both effective and free of adverse effects.

After decades of studying chronic pain, it has emerged that mechanisms contributing to the development of chronic pain and its maintenance are multifaceted and complex; such mechanisms can involve versatile and intricate signaling pathways engaged in sensory processing, both in the periphery and in the central nervous system (CNS), as has been outlined in a number of exceptional reviews [8-10]. A plethora of alterations have been described in signaling cascades, synaptic and neuronal activity, nerve cell function and network integration - from individual proteins to ion channels and receptors - suggesting that the multiple discovered mechanisms can significantly contribute to perturbed sensory processing, resulting in pathological pain.

Among those, a class of glutamatergic receptors – the α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptors (AMPAR) - is of great interest. This class of ionotropic receptors plays a critical role in synaptic strength and plasticity

[11–13] and is considered to underpin learning and memory formation in the brain. In the spinal cord, these receptors mainly determine the central sensitization of the dorsal horn (DH) a phenomenon which has been thought to represent a specific form of spinal plasticity and which underlies pain states of various origins [14-19] (for a review see [20,21]).

The implications of the role of spinal AMPAR in cellular mechanisms mediating the development and maintenance of chronic pain have emerged from several lines of experimental evidence. Studies from in vivo models of pain have clearly demonstrated that the pharmacological inhibition of spinal AMPAR significantly alleviates nociceptive hypersensitivity in animals experiencing pain of different origin: neuropathic [18], inflammatory [22-24], injury-induced [25,26] or postoperative pain [27-29]. Spinal AMPAR mediates sensory signaling, both at nociceptive and nociceptive circuits, and are heavily involved in the development [23,30,31] and maintenance of chronic pain [14,23,31-33]. Another line of evidence implicating AMPAR in terms of chronic pain originates from genetically engineered mouse lines, in which genetic deletion of a subunit alone (one of the four subunits, GluA1-4, which assemble into a tetramer of various combinations to make up the functional cationpermeable channel) clearly changed pain sensation in knock-out (KO) animals [32,34,35].

Spinal AMPAR: Are they any different from those across the brain?

The key role that AMPAR plays in synaptic transmission and plasticity makes it difficult to use ablation of this class of receptors in order to manage chronic pain because it can cause severe dysfunctions in the CNS. In a wider context, studies using generalized genetic manipulations offer a basic picture of the specific roles of all four AMPAR subunits (GluA1-4). Using gene targeting to harbor the expression of the AMPAR subunits, it has been found that mice deficient in GluA2 exhibit impaired motor coordination and behavioral abnormalities [36] and died of seizures during the first weeks postnatal [37]. Interestingly, mice with the genetic deletion of GluA2 were able to express long-term potentiation (LTP) [36,38] - an experimental model of synaptic plasticity for learning and memory formation. Likewise, double KO animals lacking GluA2/A3 expressed LTP, but displayed a deficit in basal transmission [39]. Conversely, the genetic deletion of GluA1 resulted in impaired LTP [40] and robust short-term memory deficit [41-43], pointing to a central role of the GluA1 subunit in synaptic plasticity and LTP.

In the spinal cord pain pathways, LTP at the C-fiber synapses was documented following a high-frequency stimulation of primary afferents [44,45], a mode of presynaptic stimulation similar to that used for the hippocampal LTP induction [46,47], or in response to noxious peripheral stimuli [48,49]. Such LTP caused hyperalgesia that could last for several days [50]. Although it highlights the causality between LTP at the C-fiber synapses and chronic pain, whether and to what degree the GluA1-containing AMPAR are engaged in nociceptive-induced potentiation remains open to question.

A common stoichiometry of AMPAR – identified in the forebrain neurons - is the receptor composition of the two GluA1 and two GluA2 subunits [51]. The hetero-tetrameric composition has been confirmed using a single-cell genetic approach coupled with electrophysiology, and it was found that most receptors in the hippocampal pyramidal neurons are GluA1/A2 heteromers (~80%), with minor GluA2/A3 ones [52,53]. The GluA4 subunit is amply expressed by interneurons and is not involved in the fast transmission in the "mature" excitatory neurons of adults [54,55]. GluA2 homomers are not thought to occur naturally, and only a few GluA1 homomers (approximately 8%) were identified [39]. Conversely, most of the knowledge, if not all, with regard to the AMPAR subunit composition in the spinal cord has emerged from immunostaining with quantification analysis of the immunoreactivity of GluA1-4.

According to the immunofluorescence regional analysis, all four AMPAR subunits are expressed in the spinal cord [56,57]. Their distribution, however, is not uniform: a higher immunoreactivity of both GluA1 and GluA2 was found in the DH, showing the highest GluA1 density in the superficial DH (laminae I and II) - an area where most C-and A δ -nociceptive afferents terminate to form synapses with the laminae I-II DH neurons for nociceptive information integration and processing (for a review see [8,58]) - while the GluA2 immunoreactivity is equally distributed throughout the DH [59-61]. In contrast, GluA3 was detected in deeper laminae (III to V), an area of low-threshold afferent the projections. Consequently, it can be interpreted that AMPAR in the DH predominantly consist of GluA1/A2 heteromers. Intriguingly, the GluA4 immunostaining was noticed in the lamina I NK1 receptorpositive projection neurons [60], a small group of large neurons which convey chronic nociceptive transmission from the periphery [62]. Studies identifying the GluA1-4 stoichiometry in different neuronal types, especially in nociceptive vs. nonnociceptive circuits, will be of great importance.

Despite the fact that the AMPAR subunit composition in the DH remains largely elusive, electron microscopy studies, combined with immunofluorescence, have provided insights into the subcellular distribution of GluA1/A2. Both GluA1 and GluA2 have been identified in the postsynaptic and extrasynaptic membranes of the superficial DH neurons [34,57,63]. At synapses, the receptors are concentrated in varied densities; their number depends on spine geometry, and ranges from tens to hundreds, and correlates with the synaptic strength [64,65]. For instance, an individual "mature" (mushroomshaped) spine of a CA1 pyramidal neuron can host up to 150 AMPAR [66]. Likewise, the number of AMPAR in the extrasynaptic plasma membranes varies greatly. The density of extrasynaptic AMPAR has been, however, many times lower than that of the synaptic ones, based on immunogold labeling assessments [64] or studies of functional AMPAR using the two-photon uncaging of glutamate combined with non-stationary fluctuation analysis [65,67]. Estimates have shown an AMPAR density in immature Purkinje cells of approximately 910/µm² at the synapses compared with 19/µm² in the extrasynaptic membranes [65]. Similarly, the AMPAR count was several times less in the extrasynaptic membranes than at the synapses in the superficial DH [57,63]. Studies

have also identified presynaptic AMPAR, both in the form of functional receptors [68] and as GluA1-4 immunoreactivity in primary afferent terminals [69]. The AMPAR subunit distribution revealed in the DH terminals closely resemble the distribution patterns of spinal GluA1-4, showing the predominant expression of presynaptic GluA4-containing CP-AMPAR in the laminae I-III (~70%), preferentially localized in terminals of unmyelinated (nociceptive) fibers, and the GluA2/A3 expression in the laminae III-IV, on myelinated fibers.

All four AMPAR subunits are highly homologous (~70% amino acid identity), having conand served transmembrane extracellular domains, with only diverse C-terminal intracellular tails [70,71]. Most GluA2 subunits undergo RNA editing that replaces glutamine with arginine (Q/R editing) in the pore-forming region of the channel [72]; this ultimately prevents Ca²⁺influx through the assembled channel [73]. Thus, almost all AMPAR in the adult brain (around 99%) are GluA2-containing, hence forming Ca² +-impermeable channels [74,75], whereas small amounts of GluA2-lacking AMPAR are Ca2+permeable (CP). Consistently, AMPAR in the are predominantly GluA2superficial DH(Ca²⁺-impermeable) containing channels. Electrophysiological studies have confirmed this by recording a weak blocking effect of intracellular polyamines on the AMPAR-mediated currents (i.e., an almost linear I-V curve) induced by pharmacological activation (total AMPAR pool) in the lamina II DH neurons [63,76]. Meanwhile, the postsynaptic AMPAR at nociceptive synapses - between primary nociceptive afferents and the DH neurons - consist of a mixed population of GluA2-containing (Ca²⁺impermeable) receptors, with a relatively large proportion of GluA2-lacking, CP-AMPAR. This appears due to the compelling blocking effects of i) intracellular polyamines (i.e., the inwardly rectified I-V curve) and ii) selective antagonists of CP-AMPAR on the excitatory postsynaptic AMPAR-mediated currents induced by primary afferent stimulation nociceptive (EPSC) [22,34,77]. Aside from functional studies,

quantitative estimates of the GluA1-labeled particles at the synapses of the superficial DH were found to exceed those of GluA2 [63].

Nociceptive-induced AMPAR trafficking: Broken recycling of GluA1/A2 and chronic pain

AMPAR are highly dynamic and mobile in plasma membrane: the receptors shuttle into and out of synapses and diffuse laterally (Figure 1a). AMPAR diffuse almost freely due to Brownian forces, either individually or within a cluster, before being trapped at the synapses. This is how a large fraction of AMPAR is recruited from their extrasynaptic pool to synapses for canonical LTP and hippocampal learning [78,79]. Another route for recruiting AMPAR is exocytosis of the receptors from the intracellular stores via recycling endosomes transporting AMPAR to the plasma membrane [80-82].

Nociceptive stimuli driven by long-lasting afferent activation break down the constituent recycling of spinal AMPAR. Experimental findings have made clear that the nociceptive-induced changes in dynamic AMPAR trafficking in the superficial DH neurons relate to the maintenance of persistent pain of different origins. Molecular biology studies have documented a rapid increase in the amount of GluA1, and a decrease in GluR2 in the plasma membrane (surface fraction) induced by peripheral injection of an inflammatory agent (complete Freund's adjuvant, CFA); both changes were persistent over the time course of long-lasting pain in the CFAinduced model of persistent peripheral inflammation [23,34,63]. A similar re-distribution of GluA1containing CP-AMPAR, namely, an increased number of GluA1 in the plasma membrane while a decrease in the cytosol, has been observed in different animal models of pain, such as neurogenic peripheral pain induced by capsaicin [83,84], and inflammatory pain induced by carrageenan [85] or

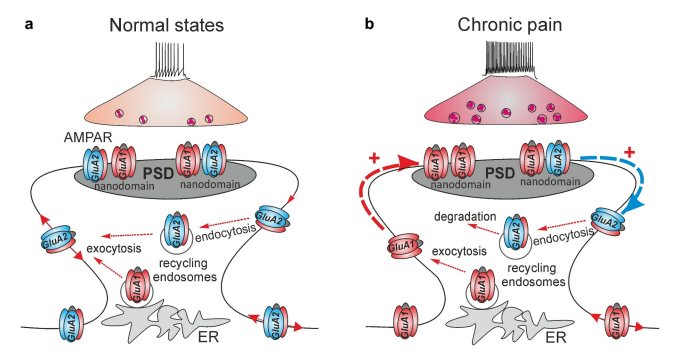


Figure 1. Spinal AMPAR trafficking for nociceptive signaling in normal (healthy) and chronic pain states. (a) After exocytosis from the intracellular stores such as the endoplasmic reticulum (ER), AMPAR laterally diffuse across the extrasynaptic plasma membrane to and from the postsynaptic density (PSD), undergoing constituent endocytosis/exocytosis cycles via recycling endosomes until are immobilized within the nanodomains of the receptors at PSD. (b) In chronic pain, the long-lasting activation of nociceptive afferents triggers the mobilization of the GluA2-containing AMPAR at PSD and the receptors undergo internalization from the synapses (blue arrow) followed by degradation (endocytosis via recycling endosomes). The GluA1-containing AMPAR replenish a pool of surface receptors and are mobilized to synapses (red arrow) to replace the GluA2-containing AMPAR within the nanodomains at PSD. Such a re-arrangement of the spinal AMPAR of different subunit compositions represents nociceptive plasticity in central pain pathways for the chronic pain maintenance.

formalin [86]. The nociceptive-induced upregulation of CP-AMPAR (GluA2-lacking receptors) in the superficial DH neurons was further confirmed through recordings of the augmented AMPARmediated currents, either alone or in combination with Ca²⁺ imaging technique, in persistent inflammatory pain [30,63,76], or following acute noxious stimulation [84] or after being induced by proinflammatory mediators [87,88].

Nociceptive stimuli facilitate the membrane insertion of GluA1-containing AMPAR at the extrasynaptic sites (Figure 1b). In the case of persistent inflammatory pain, the relative ratio of GluA1 at the postsynaptic and extrasynaptic membranes was estimated as 0.62 and 2.54, respectively, and 0.79 in the cytoplasm [63]. Neuroinflammatory mediators also enhanced the recycling of GluA1 homomers from the cytosol back to the extrasynaptic membrane, thereby promoting lateral diffusion of GluA1/GluA2 heteromers to the synapses [89,90].

However, increasing the number of GluA1containing CP-AMPAR in the plasma membrane is insufficient to potentiate synapses [91]. The newly added receptors shuttling into the synapses have to be trapped and stabilized at postsynaptic density (PSD); this occurs via the interaction of an individual AMPAR with the scaffold proteins. approaches combined with superresolution imaging for single-molecule localization and tracking have enabled the visualization of AMPAR in hippocampal neurons. Their stabilization inside synapses, along with the receptors' compactness within nanodomains, was reversibly regulated by the expression level of the scaffold protein PSD-95 [92], or the auxiliary subunit stargazin which binds to PDZ domain scaffolds [91,93]. Electrophysiological recordings of the postsynaptic AMPAR-mediated EPSC in the lamina II DH neurons have documented the nociceptive-induced upregulation of CP-AMPAR at nociceptive synapses in chronic inflammatory pain conditions [19,34,77]. Such upregulation can last up to 21 days after the induction of inflammatory pain [94]. A rapid increase in GluA1, but not GluA2/A3, in the synaptosomal fraction [83] and at the nociceptive synapses established by C-fibers [84], was also confirmed in a model of capsaicin pain. Changes in the trafficking of spinal AMPAR persistent pain conditions include

complementary boosted internalization of GluA2containing, Ca²⁺-impermeable AMPAR from the synapses between the nociceptive afferents and the DH neurons (Figure 1b). Experimental evidence consists of electrophysiological recordings showing the inwardly rectified (GluA2-lacking) AMPARmediated EPSC in the lamina II DH neurons [19,34,77] and the quantification of the relative ratio of GluA2 across cellular compartments, which dropped to 0.61 at synapses, while increased to 1.2 in the cytoplasm, as rapidly as within 24 h of the onset of inflammatory pain [63]. Studies utilizing the latest advances involving super-resolution imaging in the superficial DH are in high demand to provide new insights into the dynamic redistribution of AMPAR following nociceptive stimuli for single-particle tracking with nanoscale resolution. Using such an approach has allowed, to date, the identification of the GluA1 nanodomains at nociceptive terminals on peptidergic afferent fibers in the conditions of inflammatory pain [95]. However, whether it mirrors the organization and the distribution of the nanodomains of the receptors at the postsynaptic membrane at nociceptive synapses remains enigmatic. Likewise, the trafficking of spinal AMPAR of various subunit composition regarding different neuronal subtypes is still one of the major outstanding questions in normal states, let alone in cases of chronic pain.

AMPAR phosphorylation as a molecular mechanism for broken trafficking

The phosphorylation of AMPAR subunit(s) is a key regulatory factor that determines not only biophysical properties but also the trafficking of the receptors, leading to the modulation of synaptic plasticity. GluA1 and/ or GluA2 can undergo phosphorylation at different substrates (amino acid residues) at the C-tail, which contains multiple phosphorylation sites and proteinbinding motifs. Phosphorylated receptors thus exhibit changes in single-channel properties [96] and interaction with scaffold proteins [96,97].

The mechanistic basis of the internalization of GluA2containing AMPAR from nociceptive synapses, as deciphered by Park et al. [34], includes the phosphorylation of GluA2 at Serine (S) 880 by the protein kinase C (PKC) - the most heavily studied protein kinase - in

the postsynaptic compartments of DH neurons; it disrupts the binding of the receptor to synaptic proteins, such as the AMPAR-binding protein (ABP)/glutamate receptor-interacting protein (GRIP), stargazing, and protein interacting with C kinase 1 (PICK1) [98]. The phosphorylated receptor consequently becomes capable of diffusing and shuttling out the synapse. The upstream trigger in this mechanism has been the activation of PKC subtype α, which is Ca²⁺-dependent process, induced by the activation of postsynaptic NMDA receptors (NMDAR), thereby NMDAR-mediated Ca²⁺ influx, following the neuronal depolarization by nociceptive afferent stimuli [34]. In summary, the PKCα-dependent phosphorylation of GluA2 sets off a cascade of intracellular events, resulting in GluA2 internalization from nociceptive synapses, and eventually pain hypersensitivity [14]. Using the targeted mutation of the GluA2 phosphorylation site (by editing S882 to alanine, K882A) prevents the phosphorylation of GluA2 at S880 by PKC, and alleviates pain hypersensitivity in animals with persistent peripheral inflammation [34].

The phosphorylation of GluA1, at S831 and S845, has been found in neurogenic pain induced by capsaicin [59,99]. This was effectively blocked by the pharmacological inhibition of PKC [100]. Using genetic approaches (gene-silencing) combined with the spinal delivery of genetic materials, the PKCa-dependent upregulation of GluA1containing CP-AMPAR has been validated in persistent inflammatory pain; moreover, the transient knockdown of spinal PKCa recovered both upregulated AMPAR-mediated currents and Ca²⁺ influx in the lamina II DH neurons, returning them back to normal levels [76] and alleviating persistent inflammatory pain [22].

Among other PKC isoforms, PKC gamma was found to be capable of phosphorylating GluA1 at S831, but not at S845, in cases of neuropathic pain [101], and to enhance the membrane insertion of GluA1 in post-surgical pain [102]. In addition to S831 and S845, the other substrate on the GluA1 subunit - the S818 phosphorylation site (the PKC substrate only) - has been found to be critical for the synaptic incorporation of GluA1-containing CP-AMPAR, and the hippocampal LTP [103]. However, the implications of this highly conserved GluA1 residue for nociceptive-induced trafficking of the receptors remains unknown.

Apart from PKC, other protein kinases also phosphorylate the GluA1 subunit. The role of protein kinase A (PKA) in promoting the membrane insertion of GluA1 is fairly notable. PKA targeted the S831 and S845 sites on the GluA1 in inflammatory pain [85] and capsaicin-induced pain models [104]. Meanwhile, the engagement of Ca² +-calmodulin-dependent kinase II (CaMKII) in the phosphorylation of spinal GluA1 remains, to some degree, debatable, with conflicting evidence across the literature. Some of the studies have reported no link between the phosphorylation of spinal GluA1, at S831 and/nor S845 sites, and the activation of CaMKII in chronic injury [101] or thermal pain models [105], while others have shown an increased expression of CaMKII that triggered the membrane insertion of GluA1 in a capsaicin pain model [83,106], as well as the CaMKII alphadependent GluA1 phosphorylation at S831 in postoperative pain [107]. In comparison, the insertion of CP-AMPAR into the synapses in the brain was found to depend upon CaMK activation [108], while the CaMKII-dependent phosphorylation of GluA1 at S831 clearly potentiated synaptic transmission [109]. Consequently, knock-in mutations in GluA1 phosphorylation sites targeted by CaMKII and PKA produced deficits in hippocampal LTP and unambiguous memory defects [110], indicating that the PKA-driven phosphorylation of GluA1 is not sufficient for the receptors' incorporation into synapses, but requires also CaMKII activation [111].

Perturbed trafficking of spinal AMPAR in chronic pain: To what degree does it underlie nociceptive plasticity in the DH?

Nociceptive-induced changes in functional AMPAR lead to the altered excitability of the DH neurons; they become over-excited, causing overall hyperexcitability of the DH, a state that is associated with chronic pain in persistent peripheral inflammation [112], or after spinal cord injury [89,113]. The re-distribution of spinal AMPAR across neuronal compartments implies nociceptive neural plasticity evoked by a long-lasting presynaptic drive from nociceptive afferents. This AMPAR-mediated plasticity largely contributes to

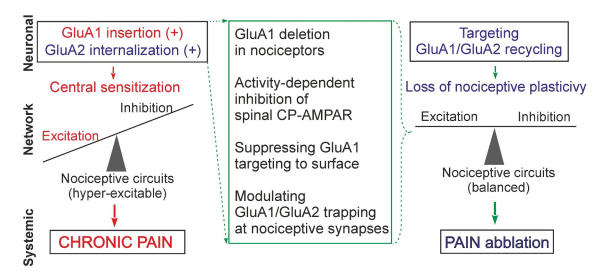


Figure 2. Nociceptive-induced changes in the trafficking of spinal AMPAR (neuronal level) cause a broken balance between synaptic excitation and inhibition that leads to overall hyper-excitability and central sensitization of the DH (network level); aberrant signaling in central pain pathways turns pain chronic (systemic level). Various scenarios for targeting the broken GluA1 and/or GluA2 recycling can be implemented to suppress the AMPAR-mediated nociceptive plasticity, and recover the relative balance between neuronal excitation and inhibition in the DH circuits, thereby alleviating chronic pain.

central sensitization and pain hypersensitivity (Figure 2).

Nociceptive-induced changes in spinal AMPAR appear to be cell-type specific. A prominent upregulation of CP-AMPAR has been documented in lamina II DH neurons characterized by intrinsic tonic firing properties in persistent inflammatory pain [30,63,76]. Neurons exhibiting tonic firing properties are a mixed population of heterogeneous cell types, overlapping inhibitory and, to a lesser extent, excitatory interneurons [114,115]. Different patterns of GluA1-4 expression were detected in the superficial DH, even in normal conditions: a majority of GABA and NK1 receptor-positive neurons expressed CP-AMPAR [116], with a major GluA1-immunoreactivity (78%) identified for inhibitory lamina I-II neurons, while almost all GluR2/3-immunoreactivity (97%) was identified for excitatory neurons [117].

Electrophysiological studies have revealed a shift in the relative balance between synaptic excitation and inhibition within the superficial DH, by documenting an increased synaptic excitation accompanied by a decreased synaptic inhibition that same neurons receive in persistent inflammatory pain [112] and after spinal cord injury [113]. The disrupted balance in terms of boosted neuronal excitation leads to nociceptive circuits becoming hyper-excitable and involved in aberrant sensory processing (Figure 2). Although the identity of neurons with perturbed AMPAR trafficking remains largely unclear, the excitatory drive was found to be impaired in GABAergic neurons in case of neuropathic pain [118]. The loss of GABAergic inhibition in the superficial DH has been established in pain states of a diverse nature: neuropathic [119], injuryinduced [120,121] or inflammatory pain [122]. A recent study has revealed that such a loss may be due to a loss of GABAergic interneurons [123], suggesting a morphological basis for the disrupted balance between synaptic excitation and inhibition in chronic pain.

The mechanisms of nociceptive plasticity in the DH circuits also include presynaptic changes. It has been shown that ectopic firing by primary nociceptive afferents is associated with the enhanced release of glutamate [124,125]. Thus, boosted release, often pertaining to "spillover", prompts the activation of perisynaptic/extrasynaptic receptors. In such a scenario, extrasynaptic glutamate escapes from the nociceptive terminals lead to the activation of an amount of extrasynaptic AMPAR, which are highly expressed in the lamina I-II DH neurons. The activation of extrasynaptic AMPAR consolidates changes in the postsynaptic AMPAR-mediated transmission, amplifies the hyper-activation of the DH neurons,

causing over-excitation within nociceptive circuits. Studies of the brain circuits over the last few decades have postulated an increased release probability in LTP [47,126,127]. The implementation of recent advances in genetically engineered optical sensors for the detection of glutamate release at the single synapse level would enable the detection of changes in the quantal content of glutamate released by nociceptive afferents in cases of chronic pain.

The extrasynaptic escape of glutamate plays an important role in inter-synaptic cross-talk [128], potentially boosting robust cross-talk between nociceptive synapses and their neighbors within highly interconnected DH circuits. Furthermore, an elegant hypothesis of "silent synapses" depicts the AMPAR role in synaptic plasticity in the brain [129,130], serving as a form of morphological plasticity, in which a proportion of synapses with absent or nonfunctional AMPAR in the superficial DH neurons can be switched on upon a longlasting nociceptive afferent drive. In addition to the nociceptive-driven subcellular re-arrangement of AMPAR, such a morphological re-arrangement between non-active and active synapses would extend nociceptive signaling, and lead to an pain hypersensitivity increase in a complex of changes in the central pain pathways.

Would targeting spinal AMPAR provide relief in chronic pain without side effects? Challenges and perspectives

Up-to-date knowledge of broken AMPAR trafficking as a cellular mechanism that turns pain chronic in central sensory pathways has identified new therapeutic targets and favors developing target-based strategies, which can be predicted as an effective treatment against chronic pain. The current understanding of deciphered signaling cascades highlights a critical step - potentially best suited for reliable alleviation of chronic pain - that focuses on restoring the broken GluA1/GluA2 recycling in the superficial DH neurons to a normal state. A crucial role that GluA1containing CP-AMPAR plays in LTP and synaptic plasticity precludes the deletion of GluA1 (in order to prevent systemic side effects), but it can be applicable when utilized exclusively in nociceptive pathways (Figure 2). Such an approach – the deletion of GluA1 in primary nociceptors - results in the ablation of GluA1-containing AMPAR in the DH nociceptive terminals, and reduces mechanical hypersensitivity in mice with chronic pain of an inflammatory or arthritic origin [17]; no changes in pain sensation were found after a similar deletion of GluA2. A major challenge, however, remains with regard to suppressing the activityinduced upregulation of GluA1-containing CP-AMPAR in the superficial DH neurons.

Several strategies focusing on spinal AMPAR have been probed in parallel. Experimental attempts comprise, as of now, a number of different strategies. These can be subdivided into two main directions: pharmacological or genetic modulation of the GluA1, and/or GluA2 recycling. pharmacological Amongst the promising approaches is the combination of the advantages of using activity-dependent antagonists of CP-AMPAR, such as dicationic compounds, with localized spinal administration (e.g., intrathecal delivery to a targeted spinal cord region) [31]. The exact mechanism of activity-dependent inhibition has been described in terms of the exceptional ability of dicationic compounds to modulate glutamatergic transmission in the brain [131,132] and to restore the nociceptive-induced upregulation of AMPAR-mediated currents in lamina II DH neurons in persistent inflammatory pain [63,76]. In rodents, dicationic compounds reduced hyperalgesia and shortened the period of inflammatory pain maintenance [31]. Using various tests by addressing potential changes in the animals' sensitivity to thermal or mechanical stimuli, locomotion activity, or exploratory behavior (anxiety), no adverse effects were observed following treatment.

Novel therapeutic strategies have focused on targeting the broken GluA1/GluA2 recycling via interfering with intracellular signaling cascades. One of those can act to suppress the upstream trigger, which is the activation of PKCa. The pharmacological inhibition of spinal PKCa (using a selective PKCa inhibitor peptide, C2-4) or genetic inhibition of PKCα (with antisense oligonucleotides) has both effectively reduced nociceptive hypersensitivity in animals, and also provided relief in pain-induced locomotive defianxiety in persistent

inflammation [22,76]. At the cellular level, therapeutic effects were manifested in the form of restored AMPAR-mediated currents in lamina II DH neurons and at nociceptive synapses [22,76]. Targeting scaffold proteins regulating the postsynaptic trapping of AMPAR at PSD can be another alternative (Figure 2). This has been recently validated using an engineered peptide inhibitor of PICK1 to alleviate mechanical hyperalgesia in injury-induced neuropathic pain [133]. Likewise, the peptide GluA2-3y, which inhibits endocytosis of GluA2-containing AMPAR, revealed an antinociceptive effect on neuropathic pain [134]. Targeting transmembrane AMPAR regulatory proteins (TARP), whose specific expression patterns were identified in the superficial DH [135], seemed a promising direction for modulating CP-AMPAR-mediated plasticity at nociceptive synapses, prone to fewer side effects. A recent study has suggested selective TARP y-8 blockage for the treatment of chronic pain [136].

In conclusion, the ongoing development of reliable and well-predictable strategies against chronic pain, further to the current approaches and directions noted here, remains active. Emerging advances have been made for targeted treatment options based on the recently discovered signaling cascades and newly identified proteins regulating AMPAR trafficking; this will help focus on refining approaches for finetuning the AMPAR-mediated nociceptive plasticity in central pain pathways in such a way to identify the most powerful therapies against chronic pain, with great care being taken to reduce (or prevent) any side effects.

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Disclosure statement

The authors declare that there is no conflict of interest.

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