‘Silent’ priming of translation-dependent LTP by β-adrenergic receptors involves phosphorylation and recruitment of AMPA receptors

Gustavo Tenorio,1,6 Steven A. Connor,2,6 Diane Guévremont,3 Wickliffe C. Abraham,4 Joanna Williams,3 Thomas J. O’Dell,5 and Peter V. Nguyen1,2,7

1Department of Physiology, University of Alberta School of Medicine, Edmonton, Alberta, T6G 2H7, Canada; 2Centre for Neuroscience, University of Alberta School of Medicine, Edmonton, Alberta, T6G 2H7, Canada; 3Department of Anatomy and Structural Biology, Brain Health and Repair Research Centre, University of Otago, Dunedin 9054, New Zealand; 4Department of Psychology, Brain Health and Repair Research Centre, University of Otago, Dunedin 9054, New Zealand; 5Department of Physiology, David Geffen School of Medicine, Center for the Health Sciences, University of California, Los Angeles, California 90095-1751, USA

The capacity for long-term changes in synaptic efficacy can be altered by prior synaptic activity, a process known as “metaplasticity.” Activation of receptors for modulatory neurotransmitters can trigger downstream signaling cascades that persist beyond initial receptor activation and may thus have metaplastic effects. Because activation of β-adrenergic receptors (β-ARs) strongly enhances the induction of long-term potentiation (LTP) in the hippocampal CA1 region, we examined whether activation of these receptors also had metaplasic effects on LTP induction. Our results show that activation of β-ARs induces a protein synthesis-dependent form of metaplasticity that primes the future induction of late-phase LTP by a subthreshold stimulus. β-AR activation also induced a long-lasting increase in phosphorylation of α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptor (AMPAR) GluA1 subunits at a protein kinase A (PKA) site (S845) and transiently activated extracellular signal-regulated kinase (ERK). Consistent with this, inhibitors of PKA and ERK blocked the metaplastic effects of β-AR activation. β-AR activation also induced a prolonged, translation-dependent increase in cell surface levels of GluA1 subunit-containing AMPA receptors. Our results indicate that β-ARs can modulate hippocampal synaptic plasticity by priming synapses for the future induction of late-phase LTP through up-regulation of transnational processes, one consequence of which is the trafficking of AMPARs to the cell surface.

Long-term potentiation (LTP), an activity-dependent increase in synaptic transmission, is considered the cellular mechanism underpinning learning and memory (Bliss and Lømo 1973; Bliss and Collingridge 1993). Interestingly, the previous history of synaptic activity can prime future changes in synaptic strength through a process known as “metaplasticity” (Abraham and Bear 1996; Abraham 2008; Abraham and Williams 2008). Metaplasticity, by expanding the temporal window for formation of associative memories, may provide the necessary mechanisms for binding temporally separated events. For example, prior activation of ryanodine receptors facilitates the induction and persistence of homosynaptic and heterosynaptic LTP elicited by a subthreshold stimulation protocol applied 30 min later (Mellentin et al. 2007; Sajikumar et al. 2009). Thus, prior cellular activity can alter the threshold for the induction of LTP in a cell-wide manner and extend the time periods typically associated with synaptic integration.

Synaptic activity is subject to modulation through activation of G protein-coupled receptors, which can influence synaptic responses on timescales that extend well beyond the seconds–minutes of initial receptor activation (O’Connor et al. 1994; Cohen et al., 1999). Metaplastic processes can be regulated through neuromodulators by lowering the threshold for induction of LTP or by extending its maintenance. Previous studies have demonstrated that β-adrenergic receptor (β-AR) activation reduces the threshold for the induction of protein synthesis-dependent LTP in the hippocampus, a brain structure critically involved in memory formation (Straube et al. 2003; Gelinas and Nguyen 2005). LTP induction can also be regulated by phosphorylation and insertion of postsynaptic glutamate receptors (GlurRs). Norepinephrine, a stress hormone released in response to emotional and arousing stimuli, induced a β-AR-dependent phosphorylation of GluA1 (also termed GluR1), in mouse hippocampus that facilitated the subsequent synaptic delivery of GluA1. GluA1 phosphorylation correlated with lowered thresholds for both memory formation and LTP induction (Hu et al. 2007). Similarly, PKA-dependent phosphorylation and insertion of GluA1 are increased following chemical potentiation with forskolin (a cAMP agonist), and correlate with the degree of LTP expressed (Oh et al. 2006; Man et al. 2007). Activation of β-ARs, by increasing levels of cAMP (O’Dell et al. 2010), can also engage intracellular signaling cascades involved in translation regulation (Gelinas et al. 2007). In previous studies, we have shown that activation of translational mechanisms by Group 1 metabotropic glutamate receptors leads to priming of local translation-dependent LTP (Raymond et al. 2000). As β-ARs couple to translation regulation mechanisms, we tested whether activation of...
β-ARs engage similar metaplasticity mechanisms for priming LTP through regulation of GluRs and protein synthesis.

We report that activation of β-ARs with the β-AR agonist isoproterenol (ISO) facilitates the subsequent induction of LTP by a subthreshold stimulation protocol. Blocking β-ARs with propranolol during ISO application prevented the subsequent induction of LTP. Additionally, ISO significantly increased GluA1 phosphorylation at serine 845 (S845), as well as increasing GluA1 surface expression. Inhibition of cAMP-dependent protein kinase (PKA), and extracellular signal-regulated kinase (ERK) similarly blocked metaplasticity, as did protein synthesis inhibition. Our findings thus provide evidence that β-ARs prime metaplastic mechanisms that support the induction of protein synthesis-dependent LTP.

**Results**

**Activation of β-ARs primes the future induction of LTP**

Activation of β-ARs in the CA1 region of mouse hippocampal slices enhances the induction of LTP by stimulation protocols that are normally subthreshold for generating persistent enhancement of extracellular field EPSPs (fEPSPs) (Thomas et al. 1996; Katsuki et al. 1997). We sought to determine whether β-AR activation could engage metaplastic mechanisms that enhance the future induction of LTP. Application of a β-AR agonist, ISO (1 μM), to area CA1 of hippocampal slices for 15 min caused a transient facilitation of baseline synaptic responses that rapidly decayed to control levels (Fig. 1B; fEPSPs were 103% ± 4% of baseline slopes 60 min after ISO application). Brief high-frequency stimulation (HFS; 1 × 100 Hz, 1 sec) applied after 80 min of baseline stimulation induced decremental LTP which returned to baseline in <2 h (104% ± 6% 120 min after HFS). However, the 15-min ISO application facilitated the induction and persistence of LTP when HFS was delivered 1 h after washout (Fig. 1A; 144% ± 6% 120 min after HFS; *P < 0.01 compared with 100 Hz alone). These results suggest that prior β-AR receptor activation can facilitate L-LTP induction up to 1 h after β-AR stimulation.

To determine the duration of the temporal window for β-AR-mediated metaplastic effects, we extended the length of time between ISO and HFS to 2 h. Under these conditions, LTP was not induced (Fig. 1B; fEPSPs were 102% ± 5% 120 min after HFS). Thus, β-AR activation enhances the ability of future stimuli to induce LTP.

![Figure 1](link)  
**Figure 1.** β-ARs activation induces metaplasticity by reducing the threshold for future LTP induction. (A) ISO application facilitates the induction of LTP (filled diamonds) by subthreshold (HFS: 1 × 100 Hz, 1 sec) stimulation 1 h after washout, whereas HFS stimulation alone results in LTP that decays to baseline in <2 h (open triangles). (B) The metaplastic effect of ISO is temporally limited, as extending the interval between ISO and HFS to 2 h prevented the metaplastic facilitation of LTP. (C) Summary histogram comparing fEPSP slopes obtained 120 min after HFS. Sample traces were taken 10 min after commencement of baseline recordings and 120 min after HFS. Results in C represent means ± SEM; **P < 0.01.
to induce LTP in a temporally restricted (&lt;2 h) fashion, without significantly altering baseline responses. These data suggest that β-ARs can regulate the modifiability of synaptic plasticity for at least 1 h after their activation, and these effects result in enhanced synaptic efficacy in response to stimuli that are normally subthreshold for inducing LTP.

**Metaplasitcity induced by ISO application requires β-adrenergic and NMDA receptors**

Noradrenaline (NA) has been shown to have a crucial role in some forms of synaptic plasticity (Gelinas and Nguyen 2005) and in the formation and retrieval of memories (Berridge and Waterhouse 2003). NA alters synaptic plasticity through activation of β-ARs, an effect mediated primarily through the β1-AR subtype (Winder et al. 1999). Induction of LTP by HFS applied 1 h after ISO application was inhibited by coapplication of a β-adrenergic antagonist, propranolol (50 μM), overlapping with ISO (Fig. 2A; fEPSPs were 98% ± 8% of baseline levels 120 min after HFS; \( P < 0.01 \) compared with controls). Shifting application of propranolol to 40 min after ISO application, however, had no effect on the induction of LTP (Fig. 2B; fEPSPs were potentiated to 143% ± 3% 120 min after HFS; \( P > 0.05 \)). This indicates that the long-lasting enhancement of LTP induction produced by β-AR activation is not due to slow washout of ISO from the slice. Instead, our results suggest that only a transient activation of β-ARs is needed to engage the metaplastic processes that enable future LTP induction.

Induction of standard LTP in CA1 requires activation of N-methyl-D-aspartate receptors (NMDARs) (Collingridge et al. 1983; Morris et al. 1990). To determine whether NMDARs are required for induction of the primed LTP, or whether β-AR priming promotes subsequent induction of an NMDAR-independent form of LTP, we applied the NMDA receptor antagonist 2-amino-5-phosphonovaleric acid, APV (50 μM), overlapping with HFS after prior application of ISO. The induction of LTP primed by prior β-AR stimulation was blocked when APV was paired with HFS (Fig. 3A; fEPSPs were 105% ± 3% of baseline levels 120 min after HFS; \( P < 0.01 \) compared with controls). Thus, metaplasticity engaged by prior β-AR activation facilitates the NMDAR-dependent form of LTP.

**β-AR-primed LTP is immune to depotentiation (DPT)**

DPT is the activity-induced reversal of LTP, which may play a role in maintaining synaptic homeostasis (Staubli and Lynch 1990; Fujii et al. 1991; Huang et al. 1999). Low-frequency stimulation of translation-dependent LTP

![Figure 2. β-ARs are required for priming of LTP. (A) Application of the β-AR antagonist propranolol during β-AR activation inhibited metaplasticity (filled triangles). (B) Shifting propranolol application to overlap with HFS did not impair LTP (black diamonds). (C) Summary histogram comparing fEPSP slopes obtained 120 min after HFS. Sample traces were taken 10 min after commencement of baseline recordings and 120 min after HFS. Results in C represent means ± SEM; \( ** P < 0.01 \).](https://www.learnmem.org)
Metaplasitc of translation-dependent LTP

Protein synthesis is required for β-AR-dependent metaplasticity

LTP and long-lasting forms of memory share a similar requirement for protein synthesis. Previously, we have shown that β-AR activation facilitates the generation of translation-dependent LTP, which is immune to reversal by depotentiating stimuli (Gelinas and Nguyen 2005). As the maintenance of LTP is critically dependent upon de novo protein synthesis (Frey et al. 1988; Nguyen and Kandel 1996; Kandel 2001), we hypothesized that translation is required for the metaplastic enhancement of LTP triggered by β-ARs.

To address this question, we used the translational inhibitor anisomycin (ANI) at a concentration that inhibits >80% of protein synthesis (40 μM; Frey et al. 1988). When ANI was coapplied with ISO (1 h prior to HFS), LTP was no longer induced (Fig. 5A; fEPSPs were 101% ± 3% of baseline levels 120 min after HFS; P < 0.05 compared with controls). Shifting application of ANI to overlap with HFS had no effect on the induction of LTP (Fig. 5B; fEPSPs were potentiated to 151% ± 8% 120 min after HFS). These results suggest that β-AR activation alone is sufficient for inducing the translation that is necessary for subsequent induction of protein synthesis-dependent LTP. The weak HFS used in this study does not by itself initiate the critical protein synthesis, even after priming, as ANI failed to block the metaplastic effects of β-AR activation when applied during HFS.

The validity of ANI as a translational inhibitor has recently been questioned due to the possibility of negative side effects that are independent of translational inhibition (Routtenberg and Rekart 2005; but see Alberini 2008). To reconcile this issue, we used a second inhibitor of translation, emetine (EME, 20 μM), which targets translational mechanisms separate from those affected by ANI. Similar to our results with ANI, we found that when EME was applied during β-AR stimulation, subsequent LTP was blocked (Fig. 5D; fEPSPs were 100% ± 6% of baseline levels 120 min after HFS; P < 0.01 compared with controls). An analysis of variance (ANOVA) comparing fEPSPs 120 min after ISO followed by HFS with ISO coapplied with either ANI or EME, followed by HFS demonstrated significant differences between groups (F[2,15] = 23.404; P < 0.01). Subsequent Tukey-Kramer post-hoc tests revealed that both ANI and EME significantly inhibited LTP (P < 0.05). Furthermore, the ANI and EME groups did not significantly differ from each other in their impairment of LTP (P > 0.05). Taken together, these data indicate that priming of LTP by β-AR activation is critically dependent upon protein synthesis, and that the disruption of metaplasticity is not due to nonspecific side effects of translation inhibitors.

Figure 3. NMDAR activation is required for the induction of β-adrenergic receptor-primed LTP. (A) When the NMDAR antagonist APV was applied during HFS and after prior application of ISO, the metaplastic facilitation of LTP was blocked (open diamonds). Control slices not exposed to APV (black triangles) displayed significantly enhanced expression of LTP compared with APV-treated slices. (B) Summary histogram comparing fEPSP slopes 120 min after HFS. Sample traces were taken 10 min after commencement of baseline recordings and 120 min after HFS. Results in B represent means ± SEM; **P < 0.01.

Figure 4. β-AR-dependent metaplasticity generates LTP that is immune to DPT. (A) Application of DPT stimuli (5 Hz, 3 min) 10 min after LTP induction affected potentiation levels when β-AR are previously activated (black diamonds). Note gradual recovery of fEPSPs to potentiated levels. (B) Summary histogram comparing fEPSP slopes 120 min after HFS. Sample traces were taken 10 min after commencement of baseline recordings and 120 min after HFS stimulation. Results in B represent means ± SEM; **P < 0.01.
Figure 5. Translation regulation required for metaplasticity is engaged specifically during β-AR stimulation. (A) Coapplication of ANI with ISO prevented the metaplastic facilitation of LTP (open triangles) when compared with controls not exposed to ANI (black diamonds). (B) Shifting the application of ANI to overlap with HFS (black diamonds) did not impair the metaplastic enhancement of LTP by prior β-AR activation. These results suggest that translation regulation critical for metaplasticity takes place during β-AR stimulation but not during subsequent HFS. (C) Comparisons of fEPSP slopes of these experiments are shown in this summary histogram. (D) A second translation inhibitor, EME similarly prevented the expression of metaplasticity when applied during application of ISO (black diamonds). (E) Summary histogram comparing fEPSP slopes 120 min after HFS stimulation. Sample traces were taken 10 min after commencement of baseline recordings and 120 min after HFS stimulation. Results in C and E represent means ± SEM (n = 6); *P < 0.05; **P < 0.01.
β-AR-dependent metaplasticity requires PKA and ERK but not mammalian target of rapamycin (mTOR)

To further define the cellular mechanisms mediating β-AR-dependent metaplasticity, we asked whether cAMP-dependent PKA plays a role in priming LTP. PKA has been implicated in LTP and long-term memory formation (Abel et al. 1997; Nayak et al. 1998; Oh et al. 2006) and can be recruited by β-AR activation (Thomas et al. 1996). Is PKA required for the enhancement of LTP by prior β-AR activation? To address this question, the PKA antagonist KT5720 was coapplied during ISO bath application. We found that activation of PKA was necessary for the metaplastic increase in synaptic response, as KT5720 blocked the subsequent induction of LTP (Fig. 6A; fEPSPs were 103% ± 8% of baseline levels 120 min after HFS; P < 0.01 compared with ISO plus HFS).

β-ARs can also recruit ERK, as well as another kinase, mTOR (Gelinas and Nguyen 2005; Gelinas et al. 2007), both of which have been implicated in local protein synthesis (Gelinas et al. 2007) and the long-term stability of LTP (Tang et al. 2002; Kelleher et al. 2004; Sweatt 2004). ERK regulates translation through Mnk1, which phosphorylates the eukaryotic translation initiation factor eIF4E (Pyronnet et al. 1999; Waskiewicz et al. 1999; Banko et al. 2004). ERK-mediated phosphorylation of eIF4E has been implicated in translation-dependent LTP (Gelinas et al. 2007) and memory formation (Kelleher et al. 2004). To determine if ERK is involved in ISO-triggered metaplasticity, we applied U0126, an inhibitor of mitogen-activated protein kinase kinase (MEK), which is the only known kinase upstream of ERK. Slices treated with U0126 exhibited significantly reduced levels of potentiation following stimulation (Fig. 6C; fEPSPs were 100% ± 6% of baseline levels 120 min after HFS; P < 0.01 compared with controls). These results suggest that ERK is required for metaplasticity induced by activation of β-ARs.

Another important component of the translational signaling pathway implicated in β-AR-dependent LTP is mTOR, which regulates protein synthesis by phosphorylating and inactivating a repressor of mRNA translation, eukaryotic initiation factor 4E-binding protein, 4E-BP (Beretta et al. 1996). Beretta et al. (1996) have shown that mTOR is blocked by rapamycin. Phosphorylation of rapamycin-sensitive sites on 4E-BP disrupts the binding of 4E-BP and eIF4E, allowing for 4E to bind eIF4G. This 4E–4G binding is critical for formation of the translation initiation complex eIF4F (Klann et al. 2004).

Disruption of the mTOR signaling cascade attenuates LTP expression generated by HFS (Tang et al. 2002). However, application of rapamycin at 1 μM, a concentration that blocks β-AR-induced phosphorylation of 4E-BP (Gelinas et al. 2007), did not block β-AR-mediated priming of LTP (Fig. 7A; fEPSPs were potentiated to 143% ± 6% 120 min after HFS). These results indicate a mechanistic divergence wherein ISO paired with HFS generates LTP that requires ERK and mTOR but not PKA (Gelinas and Nguyen 2005) and a metaplastic effect which requires PKA and β-AR but not mTOR.

GluA1 phosphorylation at S845 is persistently enhanced following β-AR stimulation

To examine the effects of β-AR activation on phosphorylation of AMPA receptor GluA1 subunits, we used phospho-site-specific antibodies and Western immunoblotting to test for potential changes in GluA1 phosphorylation at S845. In these experiments, CA1 minislices obtained from the same animal were either left untreated or exposed to a 10-min bath application of 1 μM ISO and then collected for analysis either immediately after ISO application, 60 min after ISO washout, or 120 min after ISO washout. ISO had no effect on total GluA1 levels but induced an approximately fourfold increase in GluA1 phosphorylation at S845 immediately after ISO application, 60 min after ISO washout, or 120 min after ISO washout. ISO had no effect on total GluA1 levels but induced an approximately fourfold increase in GluA1 phosphorylation at S845 immediately after ISO application (435% ± 44% of untreated control slices, n = 6, P < 0.05; Fig. 8). Consistent with the results of a previous study showing that β-AR activation induces an increase in GluA1 phosphorylation that persists for at least 1 h after ISO application (Vanhoose and Winder 2003), levels of S845 phosphorylated GluA1 were still significantly

---

**Figure 6.** ERK and PKA are required for β-AR-dependent metaplasticity. (A) U0126, a MEK inhibitor applied during β-AR activation blocked metaplasticity (open triangles). (B) Summary histogram comparing fEPSP slopes 120 min after HFS (n = 6). (C) HFS applied following β-AR activation induced L-LTP (black diamonds), which was blocked in the presence of KT5720 (a PKA inhibitor) overlapping with ISO application (open triangles). (D) Summary histogram for these experiments (n = 6). Sample traces were taken 10 min after commencement of baseline recordings and 120 min after HFS stimulation. Results in B and D represent means ± SEM; **P < 0.01.
GluA1 relative to the no-drug control significantly increased the levels of cell surface methodology, we found that ISO significantly increased the expression of GluA1-containing receptors at the cell surface. As we have previously shown that LTP-induced trafficking of some glutamate receptors is dependent on new protein synthesis (Williams et al. 2007), we tested whether the ISO-induced increase in surface GluA1 levels was translation-dependent. When ISO was applied in the presence of EME, there was a complete block of the ISO-triggered increase in cell surface GluA1 levels (0.82 ± 0.09, n = 10, P = 0.009). These data suggest that protein synthesis either promotes the expression of molecules required to traffic GluA1 to the cell surface or is required to maintain constitutively expressed GluA1 at the cell surface.

**Discussion**

Neuromodulators can modify the ability of synapses to undergo plasticity on timescales that extend well beyond normal synaptic transmission. Our results confirm previous studies showing that signals from neuromodulatory and neurotransmitter receptors can be integrated on extended timescales lasting at least 1 h (Christie and Abraham 1992; Huang et al. 1992; Cohen and Abraham 1996). In the present case, although application of a

![Figure 7.](image1.png)

**Figure 7.** mTOR is not required for metaplasiticy engaged through β-ARs. (A) Application of rapamycin did not inhibit the subsequent induction of LTP generated by HFS 1 h after ISO application (filled squares). (B) Summary histogram comparing fEPSP slopes 120 min after HFS. All sample traces were taken 10 min after commencement of baseline recording and 120 min after stimulation protocol. Results in B represent means ± SEM (n = 6).

**Figure 8.** β-AR activation drives protein synthesis-dependent membrane insertion of GluA1

Having shown that ISO increases phosphorylation of GluA1 at S845, we hypothesized that this may be associated with increased insertion or retention of GluA1 at the cell surface (Sun et al. 2005; Gao et al. 2006; Oh et al. 2006). To investigate this, we treated hippocampal slices with ISO (1 μM, 15 min) and isolated the cell surface proteins by applying the membrane impermeable biotin moiety, sulpho-NHS-SS-biotin, in conjunction with neutravidin pulldown assays (Williams et al. 2007). Using this methodology, we found that ISO significantly increased the levels of cell surface GluA1 relative to the no-drug control (1.47 ± 0.06, n = 11, P = 0.008), but this did not differ from ISO treatment alone (P = 0.677). These data confirm that ISO treatment increases the expression of GluA1-containing receptors at the cell surface. As we have previously shown that LTP-induced trafficking of some glutamate receptors is dependent on new protein synthesis (Williams et al. 2007), we tested whether the ISO-induced increase in surface GluA1 levels was translation-dependent. When ISO was applied in the presence of EME, there was a complete block of the ISO-triggered increase in cell surface GluA1 levels (0.82 ± 0.09, n = 10, P = 0.009). These data suggest that protein synthesis either promotes the expression of molecules required to traffic GluA1 to the cell surface or is required to maintain constitutively expressed GluA1 at the cell surface.

![Figure 8.](image2.png)

![Figure 8.](image3.png)
β-AR agonist to area CA1 of the hippocampus elicited only a very transient enhancement of synaptic strength, it substantially increased the magnitude and duration of LTP. Thus, a decremental LTP that does not require protein synthesis was converted to a translation-dependent, long-lasting LTP. The enhanced duration of LTP reflects the integration of diverse intracellular signals and provides a mechanism by which activation of neuromodulatory receptors can regulate synaptic plasticity subsequent to synaptic activity.

What are the mechanisms underlying β-AR-dependent metaplasticity? Metaplasticity induced by prior β-AR activation was immune to DPT, suggestive of a requirement for translation (Woo and Nguyen 2003). Inhibiting translation during β-AR activation but not during HFS, prevented LTP from being expressed. Importantly, shifting the application of ANI to overlap with HFS did not block the induction or expression of LTP, suggesting that previous activation of β-ARs is sufficient for priming transcriptional processes that once engaged, are no longer sensitive to protein synthesis arrest. Previous results indicate that isoproterenol applied during HFS induces LTP that is sensitive to translation inhibition (Gelinas and Nguyen 2005). Here, we have delayed HFS to 1 h after β-AR stimulation. In both our present and previous studies (Gelinas and Nguyen 2005), LTP was blocked only when protein synthesis inhibition coincided with β-AR activation. Thus, β-ARs appear to up-regulate translation necessary for subsequent expression of LTP, whereas HFS plays a role in induction of synaptic potentiation. Indeed, the induction of LTP primed by β-ARs was blocked when the NMDAR antagonist APV was applied during HFS.

In a broader perspective, as translation is required for LTP expression (Bradshaw et al. 2003; Gelinas and Nguyen 2005; Abraham and Williams 2006) and formation of long-term memories (Moncada and Viola 2007; Hernandez and Abel 2008), our data provide insight into the cellular mechanisms that may be required for associative memory processes that allow for incorporation of temporally separated components of an experience.

What roles do PKA and ERK play in the metaplasticity processes engaged by β-AR activation? Stimulation of β-ARs initiates ERK and mTOR signaling cascades, which independently converge at regulation of the translation factor eIF4E during β-AR-dependent LTP (Gelinas et al. 2007). ERK has been implicated in translational control of synaptic plasticity (Kelleher et al. 2004; Gelinas and Nguyen 2005; Gobert et al. 2008) and memory processes (Kelly et al. 2003; Duvarti et al. 2005; for review, see Sweatt 2004; Giovannini 2006). ERK regulates protein synthesis through a Mnk1-mediated phosphorylation of eIF4E, which has been associated with increased rates of translation (Panja et al. 2009). Interestingly, mTOR inhibition did not block the β-AR-induced metaplasticity observed in the current study. Previous results from our lab determined that mTOR is required for LTP induced by pairing ISO application with HFS (Gelinas and Nguyen 2005). This suggests a divergence in mechanisms supporting these types of plasticity. For metaplasticity, ERK signaling may generate a long-lasting (at least 1 h) state of primed translation that does not require mTOR. Functionally, this may relate to the saliency of the signals; the contiguous protocol may represent a “strong” signal that initiates immediate, long-term changes in synaptic strength through multiple signaling cascades coregulating translation. Metaplastic processes, on the other hand, must by nature be highly labile to allow for incorporation of signals over time, and therefore may not require the same magnitude of translation regulation observed with “strong” stimuli.

Emetine and anisomycin inhibit translation by blocking the 48S ribosome, whereas rapamycin blocks translation by inhibiting phosphorylation of 4E-BP, precluding assembly of the preinitiation complex at the 5′ caps of target mRNAs. Interestingly, our β-AR-dependent metaplasticity did not require activation of mTOR, as it was not blocked by rapamycin. This suggests that cap-independent translation, involving internal ribosomal entry sites (IRESs) (reviewed by Klann et al. 2004), may be recruited by activation of β-ARs 1 h prior to LTP induction. Some dendritic mRNA species that are known to contain IRESs and that may be translated in a cap-independent manner include Arc (activity-regulated cytoskeletal protein), CaMK2-α, and MAP2 (microtubule-associated protein-2) (Pinkstaff et al. 2001). Further research is needed to directly assess the involvement of cap-independent translation, and of these specific proteins, in β-AR-induced metaplasticity of LTP.

Activation of β-ARs can increase phosphorylation of AMPARs in a PKA-dependent manner (Man et al. 2007). The increased surface expression of GluA1 observed in the present study suggests one mechanisms of β-AR metaplasticity, namely, trafficking of GluA1-containing AMPARs to the cell surface through a PKA-dependent phosphorylation of S845 and protein synthesis. Recent work suggests that PKA phosphorylation of GluA1 S845 regulates AMPAR surface expression by both increasing receptor insertion and decreasing internalization (Man et al. 2007). Phosphorylation of S845 can also enhance the subsequent activity-dependent insertion of GluA1 by mobilizing AMPARs to extrasynaptic locations (Oh et al. 2006). The role of translation in this process is less clear, but a similar effect has been reported following LTP in the dentate gyrus in vivo, although for subunits other than GluR1 (Williams et al. 2007). While the identities of the synthesized proteins remain to be determined, they could include molecular chaperones important for the trafficking process, as the total level of GluA1 itself is not changed. Regardless, the localization of GluR5 extrasynaptically is an attractive metaplasticity mechanism, as this positions them for activity-dependent lateral movement into the synapse, thereby enhancing the subsequent LTP. This mechanism could help explain the previously reported increase in the induction of LTP following ISO priming (Cohen et al. 1999).

Monitoring baseline responses to application of ISO alone revealed a transient increase in potentiation which returned to
Hippocampal slice preparation
Male C57BL/6 mice (8–13 wk) (Charles River, Montréal, Québec, Canada) were used for all experiments. Animals were housed and cared for at the University of Alberta using guidelines approved by the Canadian Council on Animal Care. After cervical dislocation and decapitation, transverse hippocampal slices (400 μm thick) were prepared as described by Nguyen and Kandel (1997). Slices were maintained in an interface chamber at 28°C and perfused 1–2 ml/min with artificial CSF (aCSF) composed of the following (in mM): 124 NaCl, 4.4 KCl, 1.3 MgSO₄, 1.0 NaH₂PO₄, 26.2 NaHCO₃, 2.5 CaCl₂, and 10 glucose, aerated with 95% O₂ and 5% CO₂. Slices recovered for 60 min before experiments commenced.

Electrophysiology
fEPSPs were recorded with a glass microelectrode filled with aCSF (resistances, 2–3 MΩ) and positioned in the stratum radiatum of area CA1. fEPSPs were elicited by applying LFS (5 Hz, 3 min) 10 min before application of the NMDAR antagonist overlapping with HFS. This suggests that activation of β-ARs appears to engage a long-lasting (1 h) priming mechanism that facilitates the activity-dependent mobilization of AMPARs to synaptic locations required for LTP. Synaptic incorporation of AMPARs appears to be an NMDA-receptor-dependent event as application of the NMDAR antagonist overlapping with HFS blocked the induction of LTP. Thus, the role of HFS in this protocol may be to drive the synaptic incorporation of AMPARs through Ca²⁺-dependent events, which may stabilize AMPARs synthetically by regulating receptor anchoring (Borgdorff and Choquet 2002; Tardin et al. 2003).

Our results provide novel evidence for a β-AR-dependent facilitation of subsequent induction of LTP on timescales requiring metaplastic mechanisms. Because persistent LTP requires de novo protein synthesis, β-ARs may engage translation to gate the future establishment of long-lasting plasticity. As β-ARs also couple to PKA and ERK-dependent signaling cascades, this provides a biochemical mechanism for the metaplastic enhancement of LTP generated by prior activation of β-ARs (Fig. 10). Here, we established that β-AR-dependent metaplasticity recruits PKA and ERK activation as LTP was attenuated by inhibition of either PKA or ERK during ISO application. Inhibition of translation with EME or ANI blocked the expression of LTP but only when inhibitors were applied overlapping with ISO and well before LTP induction. The effects of translational inhibition appear to affect stabilization of GluA1 membrane insertion, which was increased following stimulation of β-AR, but reduced in the presence of EME. Indeed, activation of β-ARs increased GluA1 phosphorylation at site S845, which facilitates membrane insertion of GluA1 (Man et al. 2007). Our data suggest a metaplastic mechanism whereby stimulation of β-ARs primes synaptic incorporation of GluA1 through phosphorylation and translation-dependent stabilization of recently inserted receptors.

In summary, our data characterize the biochemical signaling mechanisms by which activation of β-ARs can enhance the future persistence of LTP. Metastability mechanisms engaged by neuro-modulators, by enhancing translation regulation, could facilitate the encoding of information presented on timescales much longer than normal synaptic transmission. Our data add the β-AR to a growing number of neuromodulatory receptors that when activated alter the threshold for future changes in synaptic strength. Under circumstances that require attention or elicit arousal, global levels of NA increase (Berridge and Waterhouse 2003). Release of NA lowers the threshold for subsequent synaptic change by priming translational processes necessary for long-term synaptic modifications (Gelinas and Nguyen 2007). Our results suggest that the priming effect of β-AR activation was limited to 1 h, despite the persistence of GluA1 phosphorylation for at least 2 h. As shifting the application of anisomycin beyond ISO application failed to block the subsequent induction of LTP, this suggests a time-limited role for translational regulation in priming of future synaptic events. In the course of everyday events, experience is best framed in the second to minute to hourly time frame (Dudai 2009). Therefore, an effective way for updating of a previously established memory would be to encode information that presents within a limited (≤1 h) time window relative to the plasticity-inducing stimulus. This also raises the possibility that NA acting at β-ARs obviates the need for ongoing attentional processes to facilitate association of new items into the previously established memory as primed plasticity mechanisms would lower the threshold for subsequent changes in synaptic efficacy supporting memory updating. This would be an efficient method for ensuring that information that may have value is properly encoded in association with the initial priming event.

Materials and Methods

Hippocampal slice preparation
Male C57BL/6 mice (8–13 wk) (Charles River, Montréal, Québec, Canada) were used for all experiments. Animals were housed and cared for at the University of Alberta using guidelines approved by the Canadian Council on Animal Care. After cervical dislocation and decapitation, transverse hippocampal slices (400 μm thick) were prepared as described by Nguyen and Kandel (1997). Slices were maintained in an interface chamber at 28°C and perfused 1–2 ml/min with artificial CSF (aCSF) composed of the following (in mM): 124 NaCl, 4.4 KCl, 1.3 MgSO₄, 1.0 NaH₂PO₄, 26.2 NaHCO₃, 2.5 CaCl₂, and 10 glucose, aerated with 95% O₂ and 5% CO₂. Slices recovered for 60 min before experiments commenced.

Electrophysiology
fEPSPs were recorded with a glass microelectrode filled with aCSF (resistances, 2–3 MΩ) and positioned in the stratum radiatum of area CA1. fEPSPs were elicited by applying LFS (5 Hz, 3 min) 10 min before application of the NMDAR antagonist overlapping with HFS. This suggests that activation of β-ARs appears to engage a long-lasting (1 h) priming mechanism that facilitates the activity-dependent mobilization of AMPARs to synaptic locations required for LTP. Synaptic incorporation of AMPARs appears to be an NMDA-receptor-dependent event as application of the NMDAR antagonist overlapping with HFS blocked the induction of LTP. Thus, the role of HFS in this protocol may be to drive the synaptic incorporation of AMPARs through Ca²⁺-dependent events, which may stabilize AMPARs synthetically by regulating receptor anchoring (Borgdorff and Choquet 2002; Tardin et al. 2003).

Our results provide novel evidence for a β-AR-dependent facilitation of subsequent induction of LTP on timescales requiring metaplastic mechanisms. Because persistent LTP requires de novo protein synthesis, β-ARs may engage translation to gate the future establishment of long-lasting plasticity. As β-ARs also couple to PKA and ERK-dependent signaling cascades, this provides a biochemical mechanism for the metaplastic enhancement of LTP generated by prior activation of β-ARs (Fig. 10). Here, we established that β-AR-dependent metaplasticity recruits PKA and ERK activation as LTP was attenuated by inhibition of either PKA or ERK during ISO application. Inhibition of translation with EME or ANI blocked the expression of LTP but only when inhibitors were applied overlapping with ISO and well before LTP induction. The effects of translational inhibition appear to affect stabilization of GluA1 membrane insertion, which was increased following stimulation of β-AR, but reduced in the presence of EME. Indeed, activation of β-ARs increased GluA1 phosphorylation at site S845, which facilitates membrane insertion of GluA1 (Man et al. 2007). Our data suggest a metaplastic mechanism whereby stimulation of β-ARs primes synaptic incorporation of GluA1 through phosphorylation and translation-dependent stabilization of recently inserted receptors.

In summary, our data characterize the biochemical signaling mechanisms by which activation of β-ARs can enhance the future persistence of LTP. Metastability mechanisms engaged by neuro-modulators, by enhancing translation regulation, could facilitate the encoding of information presented on timescales much longer than normal synaptic transmission. Our data add the β-AR to a growing number of neuromodulatory receptors that when activated alter the threshold for future changes in synaptic strength. Under circumstances that require attention or elicit arousal, global levels of NA increase (Berridge and Waterhouse 2003). Release of NA lowers the threshold for subsequent synaptic change by priming translational processes necessary for long-term synaptic modifications (Gelinas and Nguyen 2007). Our results suggest that the priming effect of β-AR activation was limited to 1 h, despite the persistence of GluA1 phosphorylation for at least 2 h. As shifting the application of anisomycin beyond ISO application failed to block the subsequent induction of LTP, this suggests a time-limited role for translational regulation in priming of future synaptic events. In the course of everyday events, experience is best framed in the second to minute to hourly time frame (Dudai 2009). Therefore, an effective way for updating of a previously established memory would be to encode information that presents within a limited (≤1 h) time window relative to the plasticity-inducing stimulus. This also raises the possibility that NA acting at β-ARs obviates the need for ongoing attentional processes to facilitate association of new items into the previously established memory as primed plasticity mechanisms would lower the threshold for subsequent changes in synaptic efficacy supporting memory updating. This would be an efficient method for ensuring that information that may have value is properly encoded in association with the initial priming event.

Figure 10. Hypothetical model for primed AMPAR mobility following β-AR activation. β-ARs facilitate AMPAR insertion through PKA-dependent phosphorylation of GluA intracellular C-terminal domains and translation of plasticity related proteins (PRPs), which may include transmembrane AMPAR regulatory proteins (TARPs) that regulate AMPAR trafficking. PKA phosphorylation of S845 on GluA1 and newly synthesized PRPs increase AMPAR expression at extrasynaptic sites. HFS may drive lateral mobilization of AMPARs into the synapse resulting in enhanced synaptic strength.

Drugs
ISO, a specific β-AR agonist, (R-[–]-isoproterenol [+] -bitartarate, 1 μM; Sigma) was prepared fresh daily in distilled water as a 2 mM
stock solution. The β-AR antagonist (+)-propranolol hydrochloride (PROP; 50 μM; Research Biochemicals) was also prepared daily in distilled water as a 50-mM stock solution. The NMDAR antagonist, APV (DL-2-amino-5-phosphonopentanoic acid, 50 μM; Sigma) was prepared in distilled water as a 50-mM stock solution. An mTOR inhibitor, rapamycin (1 μM; Sigma) and a MEK inhibitor, U0126 (20 μM; Sigma), were prepared as 40-mM stock solutions in DMSO and 20 mM in distilled water, respectively. APV, propranolol, ANI, EME, and rapamycin were applied 30 min before ISO application, throughout ISO application, and for 10 min after ISO application. All drug experiments were performed under dimmed light conditions due to drug photosensitivity. Drug experiments were interleaved with drug-free controls.

Data analysis

The initial slope of the fEPSP was measured as an index of synaptic strength (Johnston and Wu 1995). The average “baseline” slope values were acquired over a period of 20 min before experimental protocols and all fEPSP initial slope measurements were expressed as a percentage of the baseline. fEPSP slopes were measured at 120 min post-stimulation for comparisons of LTP. Student’s t-test was used for statistical comparisons of mean fEPSP slopes between two groups, with a significance level of P < 0.05. All values shown are means ± SEM, with n = number of slices.

Preparation of cell surface fractions

Hippocampal cell-surface proteins were isolated by reversible tagging with a membrane-impermeable biotin moiety and precipitated by addition of Neuravidin conjugated to agarose beads using a method based on our established protocol (Williams et al. 2007), with modifications. Slices (400 μm) were bathed in aCSF (2 h/28 °C) with constant perfusion, prior to treatment (15 min). Groups included no drug (n = 7), 1 μM ISO (n = 11), 1 μM ISO + HFS (n = 11), and 1 μM ISO + 20 μM EME (n = 10). After washout (1 h), slices were removed to ice-cold aCSF (5 min) followed by incubation with 2.5 mg/ml sulfo-NHS-SS-biotin (Pierce, 29200; 2 h, 4 °C) and, following three washes in TBST, blots were incubated for 2–4 h at room temperature in the appropriate HRP-conjugated secondary antibody. Immunoreactive bands were visualized using enhanced chemiluminescence and a cooled CCD camera-based image acquisition and analysis system (Quantity One; Bio-Rad) was used to acquire images of the blots and perform densitometric analysis. All blots were reprobed with an antituBulin antibody and the optical density values for each band of interest was normalized to the density values obtained for tubulin in the same lane to control for potential variations in loading. These values were then normalized to levels for the same protein measured in samples from untreated control slices. Friedman repeated-measures ANOVA on ranks followed by Student-Newman-Keuls tests for multiple pair-wise comparisons were used to determine statistical significance.

Phosphorylation status immunoblotting

To examine the effects of β-AR activation on phosphorylation of AMPA receptor GluA1 subunits and ERK activation, we use CA1 “minislices” that were prepared by removing the dentate gyrus, CA3 region, and subiculum from freshly cut, 400 μm-thick hippocampal slices. CA1 minislices were then maintained in interface-type slice chambers (typically three to six slices/chamber) perfused (2 mL/min) with warm (28°C), oxygenated aCSF, and allowed to recover for at least 2 h. Slices were then either left in aCSF (untreated controls) or exposed to a 10 min bath application of ISO (1 μM) and collected for analysis either immediately after ISO application or at different time points following washout of ISO with drug-free aCSF. In all groups slices were snap-frozen by rapidly transferring them into prechilled microcentrifuge tubes (Kontes Glass Company) and plunging the tube into a bed of crushed dry ice.

For Western blot analysis the slices were homogenized in ice-cold buffer containing 50 mM Tris-HCl, 50 mM NaF, 10 mM EGTA, 10 mM EDTA, 80 μM sodium molybdate, 5 mM sodium pyrophosphate, 1 mM sodium orthovanadate, 1 mM phenylmethylsulfonyl fluoride, 0.01% Triton X-100, and 4 mM para-nitrophenylphosphate that contained both protease inhibitors (Protease Inhibitors Complete, Roche Molecular Biochemicals) and protein phosphatase inhibitors (Phosphatase Inhibitor Cocktails I and II; Sigma-Aldrich). Aliquots from each of the samples were removed to determine protein concentration using the Bradford assay and samples were then boiled for 5 min after the addition of 2× loading buffer. Equal amounts of total protein (20 μg/lane) from each of the samples were resolved on 12% SDS-PAGE gels, transferred onto nitrocellulose membranes, and then blocked for 1 h in Tris-buffered saline containing 0.05% Tween-20 (TBST) and 4% fat-free dry milk. Primary antibody incubations were done overnight (at 4 °C) and, following three washes in TBST, blots were incubated for 2–4 h at room temperature in the appropriate HRP-conjugated secondary antibody. Immunoreactive bands were visualized using enhanced chemiluminescence and a cooled CCD camera-based image acquisition and analysis system (Quantity One; Bio-Rad) was used to acquire images of the blots and perform densitometric analysis. All blots were reprobed with an antituBulin antibody and the optical density values for each band of interest was normalized to the density values obtained for tubulin in the same lane to control for potential variations in loading. These values were then normalized to levels for the same protein measured in samples from untreated control slices. Friedman repeated-measures ANOVA on ranks followed by Student-Newman-Keuls tests for multiple pair-wise comparisons were used to determine statistical significance.

References

Abel T, Nguyen PV, Barad M, Deuel TA, Kandel ER, Bourtchouladze R. 1997. Genetic demonstration of a role for PKA in the late phase of LTP and in hippocampus-based long-term memory. Cell 88: 615–626.

Abraham WC. 2008. Metaplasticity—tuning synapses and networks for plasticity. Nat Rev Neurosci 9: 387–399.

Abraham WC, Bear MF. 1996. Metaplasticity: The plasticity of synaptic plasticity. Trends Neurosci 19: 126–130.

Abraham WC, Williams JM. 2008. LTP maintenance and its protein synthesis-dependence. Neuron 59: 260–268.

Alberini CM. 2008. The role of protein synthesis during the labile phases of memory: Revisiting the skepticism. Neuron 59: 234–246.
Williams JM, Mason-Parker SE, Abraham WC, Tate WP. 1998. Biphasic changes in the levels of N-methyl-D-aspartate receptor-2 subunits correlate with the induction and persistence of long-term potentiation. *Mol Brain Res* **60**: 21–27.

Williams JM, Guévremont D, Mason-Parker SE, Luxmanan C, Tate WP, Abraham WC. 2007. Differential trafficking of AMPA and NMDA receptors during long-term potentiation in awake adult animals. *J Neurosci* **27**: 14171–14178.

Winder DG, Martin KC, Muzzio IA, Rohrer D, Chruscinski A, Kobilka B, Kandel ER. 1999. ERK plays a regulatory role in induction of LTP by theta frequency stimulation and its modulation by beta-adrenergic receptors. *Neuron* **24**: 715–726.

Woo NH, Nguyen PV. 2003. Protein synthesis is required for synaptic immunity to depotentiation. *J Neurosci* **23**: 1125–1132.

Received August 5, 2010; accepted in revised form September 28, 2010.
'Silent' priming of translation-dependent LTP by β-adrenergic receptors involves phosphorylation and recruitment of AMPA receptors

Gustavo Tenorio, Steven A. Connor, Diane Guévremont, et al.

Learn. Mem. 2010, 17:
Access the most recent version at doi:10.1101/lm.1974510

References
This article cites 65 articles, 27 of which can be accessed free at:
http://learnmem.cshlp.org/content/17/12/627.full.html#ref-list-1

License

Email Alerting Service
Receive free email alerts when new articles cite this article - sign up in the box at the top right corner of the article or click here.