A protein synthesis and nitric oxide-dependent presynaptic enhancement in persistent forms of long-term potentiation

Victoria P.A. Johnstone and Clarke R. Raymond

Department of Neuroscience, The John Curtin School of Medical Research & Eccles Institute of Neuroscience, The Australian National University, Canberra ACT 0200, Australia

Long-term potentiation (LTP) is an important process underlying learning and memory in the brain. At CA3–CA1 synapses in the hippocampus, three discrete forms of LTP (LTP1, 2, and 3) can be differentiated on the basis of maintenance and induction mechanisms. However, the relative roles of pre- and post-synaptic expression mechanisms in LTP1, 2, and 3 are unknown. Neurotransmitter release in the expression of LTP, 2, and 3 was measured via FM 1–43 destaining from CA3 terminals in hippocampal slices from male Wistar rats (7–8 wk). No difference in vesicle turnover rate was observed for LTP1 up to 160 min following induction by one train of theta-burst stimulation (ITBS). A presynaptic enhancement was found for LTP2 at 160 min after induction by 4TBS, and for LTP3 at both 80 and 160 min after induction by 8TBS. Inhibition of nitric oxide (NO) signaling blocked both LTP2 and LTP3 maintenance and the associated enhanced release. LTP2 maintenance and its presynaptic expression were dependent on protein synthesis, but not gene transcription. LTP3 maintenance was dependent on both translation and transcription, but like LTP2, the enhanced release only required translation. These data considerably strengthen the mechanistic separation of LTP1, 2, and 3, supporting a model of multiple, discrete forms of LTP at CA3–CA1 synapses rather than different temporal phases.

LTP expression

One aspect of LTP that continues to be debated concerns the relative contributions of pre- and post-synaptic changes to its expression (Bliss et al. 2003). There is a wealth of evidence for post-synaptic alterations that has been extensively reviewed elsewhere (Yuste and Bonhoeffer 2001; Malinow and Malenka 2002; Derkach et al. 2007). There is also a significant body of work supporting presynaptic changes in LTP expression (Lisman and Raghavachari 2006). Recent optical and quantal studies have revealed unequivocal presynaptic contributions to LTP expression at both early and late time points, but primarily when induced by “strong” protocols (Zakharenko et al. 2001; Sokolov et al. 2002; Stanton et al. 2005; Bayazitov et al. 2007; Enoki et al. 2009). Thus, enhanced presynaptic function could have a conditional involvement in LTP expression that may well contribute to discrepancies in the literature.

Other work has focussed on the retrograde signaling that must occur from the post-synaptic induction site to initiate a presynaptic enhancement. Several candidates for this so-called retrograde messenger have been proposed, with the properties of the gaseous neuromodulator nitric oxide (NO) making it a particularly attractive contender (Lancaster 1994). However, the role of NO in hippocampal LTP remains somewhat contentious (Shors and Matzel 1997; Regehr et al. 2009). While most studies indicate that NO augments transmission in the presynaptic cell and promotes synaptic plasticity (Böhme et al. 1991; O’Dell et al. 1991a; Arancio et al. 1995, 1996a; Son et al. 1996), others dispute its role in LTP, particularly in the context of learning (Bannerman et al. 1994a,b), or even whether it acts retrogradely (Ko and Kelly 1999).

LTP maintenance

New protein synthesis is required to maintain LTP beyond the lifetime of a typical synaptic protein and is also required for long-lasting memories (Abraham and Williams 2008). According to the LTP1, 2, 3 model both LTP2 and 3 are dependent on new protein synthesis, but only LTP3 requires transcription. LTP2 is proposed to involve translation from preexisting mRNA transcripts, perhaps in a synapse-specific manner (Otani et al. 1989; Raymond et al. 2000; Raymond 2007). This mechanistic differentiation has never been systematically tested under the constant experimental conditions of a single study. In addition, little is known about whether there is any overlap between these maintenance mechanisms and the expression mechanisms described above.
Results

LTP2 and LTP3 involve enhanced presynaptic function

It has previously been reported that induction of LTP either chemically or using specific patterns of electrical stimulation is associated with an increase in the rate of FM 1–43 release from CA3 terminals (Zakharenko et al. 2001; Stanton et al. 2005). However, none of these “forms” of LTP have been extensively characterized, and as such, it is unclear how they relate to one another or, more importantly, what the physiological significance of the finding is. In the present study presynaptic vesicle cycling was investigated in well-characterized, identifiable and mechanistically discrete forms of LTP induced by 1TBS, 4TBS, and 8TBS.

In control slices 1TBS resulted in weak LTP that decayed back to baseline ~180 min post-TBS, with a magnitude in the last 5 min of recording of 5% ± 1% (n = 5, Fig. 2A, below). Four trains of TBS resulted in LTP that had a larger initial magnitude to that induced by 1TBS and a significantly larger magnitude in the last 5 min of recording, measuring 27% ± 7% (n = 11, Fig. 2A, below). Eight trains of TBS induced LTP that had a similar initial magnitude to that produced induced by 4TBS, but a significantly larger final magnitude, measuring 83% ± 13% (n = 7) in the last 5 min of recording (Fig. 2A, below).

Since LTP is best classified on the basis of persistence the post-TBS data were fit with a double exponential decay function as described previously (Raymond et al. 2000; Raymond and Redman 2002). The mean time constant of decay of the slower exponential (τ) was used as a measure of LTP persistence. In control slices 1TBS resulted in LTP with a τ of 37 ± 15 min (n = 3, Fig. 2B, below). Four trains of TBS induced a more persistent form of LTP, with τ measuring 117 ± 14 min (n = 9, Fig. 2B, below). Eight trains of TBS produced LTP that was dramatically more persistent than that induced with 1TBS and 4TBS. We were unable to determine a τ value for 8TBS LTP in these experiments because all control LTP was essentially nondecremental over 2 h and could not be fit by exponential decay curves. For illustrative purposes the τ value obtained from a previous set of experiments (Raymond and Redman 2002) has been used (Fig. 2B, below). This 8TBS τ value is therefore an underestimate and is not used for statistical analysis. Based on these measurements and our previous data using 1TBS, 4TBS, and 8TBS, we classify these forms of LTP as LTP1, LTP2, and LTP3, respectively.

To determine whether presynaptic changes are associated with LTP1, 2, and 3, we simultaneously measured FM 1–43 release from potentiated CA3 terminals in the same slices as the LTP experiments above. FM 1–43 was loaded into terminals of CA3 neurons using a train of electrical stimulation applied to the Schaffer collateral pathway, following bath application of the dye (Fig. 1A). This method resulted in bright punctate staining that could be visualized using two-photon microscopy. The fluorescence of these puncta decreased rapidly following subsequent electrical stimulation, corresponding to diffusion of dye from synaptic vesicles (Fig. 1B). The kinetics of this destaining has previously been shown to strongly correlate with transmitter release at these synapses (Zakharenko et al. 2001).

Under baseline conditions, FM 1–43 destaining kinetics were consistent across each experimental group (baseline t1/2 1TBS = 156 ± 27 sec, n = 35 terminals, five slices; 4TBS = 122 ± 17 sec, n = 50 terminals, eight slices; 8TBS = 144 ± 11 sec, n = 124 terminals, 15 slices) (Fig. 2C–H). LTP1 was not associated with a change in FM 1–43 destaining at either 80 min (t1/2 LTP1 = 113 ± 16 sec, n = 50 terminals, seven slices) or 160 min post-induction (t1/2 LTP1 = 172 ± 32 sec, n = 45 terminals, five slices) (Fig. 2C,D). LTP2 was associated with enhanced exocytosis at 160 min after induction (t1/2 LTP2 = 58 ± 7 sec, n = 60 terminals, six slices, P < 0.05), but not at 80 min (t1/2 LTP2 = 95 ± 8 sec, n = 63 terminals, eight slices) (Fig. 2E,F). Finally, LTP3 was associated with enhanced exocytosis at both 80 min (t1/2 LTP3 = 59 ± 6 sec, n = 97 terminals, 14 slices, P < 0.05) and 160 min post-induction (t1/2 LTP3 = 63 ± 4 sec, n = 53 terminals, 10 slices, P < 0.05) (Fig. 2G,H). These findings demonstrate that short-lasting LTP1 is predominantly expressed post-synaptically, and that more persistent LTP2 and 3 recruit a presynaptic expression mechanism.

A single, stable population of terminals before and after LTP

It is important to verify that sampled puncta were primarily representative of CA3 terminals, and did not include large numbers of GABAergic inhibitory interneurons or astrocytes, both of which endocytose FM 1–43 (Hablitz et al. 2009; Li et al. 2009) and are frequently overlooked in hippocampal FM experiments. If this were the case, one might expect different populations of destaining kinetics to be apparent in our data, as both inhibitory interneurons and astrocytes exhibit heterogeneity in their release properties (McBain and Fisahn 2001; Moulder et al. 2007; Sakaba 2008). However, we found only one population of destaining kinetics, with an average basal t1/2 of 141 ± 9 (n = 215 boutons from 28 slices) (Fig. 3A). It was therefore concluded unlikely that release from astrocytes or inhibitory interneurons were appreciably affecting measurements of destaining from CA3 terminals. This is supported by evidence indicating that only 2.5% of terminals in the CA3 region are GABAergic (Hiscock et al. 2000). There was also no change in the mean number of release sites measured before and after LTP (basal: 7 ± 1, n = 209 puncta from 28 slices; LTP1: 9 ± 2, n = 44 puncta from five slices; LTP2: 9 ± 2, n = 71 puncta from eight slices; LTP3: 7 ± 3, n = 72 puncta from 10 slices) (Fig. 3B). This stability in the number of terminals loaded with FM 1–43 suggests that, under these conditions,
forms of LTP now identified at these synapses, or if it is specific for one set of effector mechanisms. In addition, evidence for the prevailing view that during LTP induction NO is released from the post-synaptic cell and acts on the presynaptic terminal to enhance transmitter release (for review, see Regehr et al. 2009) is largely based on indirect measurements of presynaptic function. We therefore determined whether inhibition of NO signaling differentially affects the magnitude and persistence of LTP1, 2, and 3 and if so, whether this correlates with direct measurements of exocytosis. Given that significant increases in exocytosis are observed at the 160-min time point for both LTP2 and 3, we confine this and all subsequent pharmacological analyses to this time point.

Inhibition of nitric oxide synthase (NOS) with N·ω-nitro-L-arginine methyl ester (L-NAME, 100 μM) had no significant effect on either the magnitude at 160 min post-TBS (control = 5 ± 1%, n = 5; L-NAME = 10 ± 8%, n = 4) or persistence of LTP1 (control τ = 37 ± 15 min, n = 3; L-NAME = 72 ± 10 min, n = 3) (Fig. 4A). Likewise, no effect of the membrane impermeable NO scavenger 2-(4-carboxyphenyl)−4,4,5,5-tetramethylimidazoline-1-oxyl-3-oxide (cPTIO, 40 μM) was observed for LTP1 in terms of persistence (cPTIO τ = 80 ± 9 min, n = 4) or magnitude (2.5 ± 8%, n = 4) at 160 min post-TBS (Fig. 5A). Neither drug had any effect on normal synaptic transmission when applied in the absence of TBS (Figs. 4A, 5A). Unsurprisingly, exocytotic rates measured at 160 min after TBS were also unaffected by inhibition of NO signaling by either L-NAME (control τ1/2 = 172 ± 32 sec; L-NAME = 140 ± 4.5 sec, n = 75 terminals from four slices) (Fig. 4B) or cPTIO (132 ± 4 sec, n = 140 terminals from four slices) (Fig. 5B).

In contrast, inhibition of NO signaling significantly reduced LTP2 persistence (control τ = 117 ± 14 min, n = 9; L-NAME = 59 ± 16 min, n = 3; cPTIO = 73 ± 9 min, n = 4; P < 0.05) and magnitude (control = 27 ± 7%, n = 11; L-NAME = 10 ± 5%, n = 4; cPTIO = 7 ± 8%, n = 4; P < 0.05) (Figs. 4C, 5C). Both inhibitors also abolished the enhanced exocytosis associated with LTP2 measured 160 min post-TBS (control τ1/2 = 58 ± 7 sec; L-NAME = 136 ± 10 sec, n = 20 terminals from four slices; cPTIO = 147 ± 9 sec, n = 38 terminals from four slices) (Figs. 4D, 5D).

Inhibition of NO signaling with L-NAME and cPTIO qualitatively reduced LTP τ values in comparison with previously published data: control τ = 230 min (from Raymond and Redman 2002) (L-NAME = 69 ± 8 min, n = 4; cPTIO = 77 ± 10 min, n = 3); however, statistical analysis of τ values for LTP3 was not possible in this case. Both inhibitors significantly reduced LTP magnitude at 160 min post-8TBS (control = 83 ± 13%, n = 7; L-NAME = 1 ± 6%, n = 4; cPTIO = 34 ± 18%, n = 3; P < 0.05) (Figs. 4E, 5E).

induction of LTP1, 2, and 3 does not “unsilence” presynaptically inactive synapses (Voronin et al. 2004).

LTP2 and 3 are NO dependent

Several studies have shown that inhibition of NO signaling in CA1 prevents the induction of LTP (Böhme et al. 1991; O’Dell et al. 1991a; Schuman and Madison 1991; Haley et al. 1992). However, it is not clear whether NO signaling is important for all

Figure 2. (A) Mean percent change of field excitatory post-synaptic potential (fEPSP) slope from slices receiving 1TBS (n = 4), 4TBS (n = 8), 8TBS (n = 5) or no TBS (n = 4). LTP was induced (arrow) after a 20-min baseline period and recordings continued for 160 min post-TBS. (B) Decay of LTP1, 2, and 3. The mean time constant of decay, τ, was calculated for LTP induced with 1TBS and 4TBS. A τ value could not be calculated for LTP induced by 8TBS due to the robust nature of the LTP; therefore, a value has been reproduced from Raymond and Redman (2002). (C) LTP1 is not associated with an enhancement of FM 1–43 release from CA3 terminals. Mean ± SEM time course of FM 1–43 destaining prior to LTP induction (baseline), 80 min and 160 min post-induction. (D) Summary histogram showing average half time of decay (t(1/2)) of FM 1–43 fluorescence for the baseline, 80 min and 160 min unload. (E) FM 1–43 destaining time course shows LTP2 is associated with an enhancement of release 160 min post-induction. (F) Summary histogram of average t(1/2) in LTP2 experiments. (*) P < 0.05. (G) FM 1–43 destaining time course shows LTP3 is associated with enhanced release at 80 min and 160 min post-TBS. (H) Summary histogram of average t(1/2) in LTP3 experiments. (*) P < 0.05.

Figure 3. (A) Frequency histogram of half-time of intensity decay (t(1/2)) of individual puncta suggests that measurements are taken from a single population of synaptic vesicles. Data not well fit by a Gaussian distribution (P < 0.05, D’Agostino and Pearson omnibus normality test). (B) No difference in the number of puncta analyzed per field before LTP induction (basal) and 160 min post-induction of LTP1, LTP2, and LTP3 indicates there is no “unsilencing” of synapses post-LTP (P = 0.88, analysis of variance [ANOVA]).
Inhibition of NO signaling also prevented the enhanced exocytosis associated with LTP1 at 160 min post-TBS (control $\tau_{1/2} = 63 \pm 4$ sec; L-NAME $= 146 \pm 4$ sec, $n = 25$ terminals from four slices; cPTIO $= 138 \pm 13$ sec, $n = 44$ terminals from three slices) (Figs. 4F, 5F).

Although in some cases cPTIO and L-NAME appear to enhance the initial induction of LTP, a significant increase in the maximum EPSP amplitude was measured only for LTP2 in the presence of L-NAME (control $= 86 \pm 18\%$, $n = 11$; L-NAME $= 132 \pm 21\%$, $n = 4$). Given that we find no significant effect in all other cases and that other reports indicate that neither drug affects peak EPSP magnitude, nor various other post-synaptic properties of these neurons (Zhu and Erdemli 1995; Ko and Kelly 1999), we conclude that the observed effect on LTP2 is not specifically drug related and that inhibition of NO signaling does not generally enhance LTP induction.

The data presented here demonstrate that only the more robust LTP2 and LTP3 require both activation of NO and diffusion of NO in the extracellular space, and that NO is necessary to trigger the enhanced exocytosis associated with more persistent forms of LTP.

LTP2 and LTP3 are protein synthesis dependent

Until now LTP1, 2, and 3 have been differentiated on the basis of distinct induction and maintenance mechanisms (Raymond 2007; Reymann and Frey 2007). With regard to maintenance, LTP1 is proposed to involve post-translational modifications of existing proteins and glutamate receptor trafficking (equivalent to early LTP) (Lovinger et al. 1987; Malenka et al. 1989; O’Dell et al. 1991b; Shi et al. 1999). LTP2 is dependent on protein synthesis but not gene transcription (Raymond et al. 2000). Finally, LTP3 is translation and transcription dependent, equivalent to traditional late LTP (Nguyen et al. 1994; Kandel 2001; Reymann and Frey 2007). To confirm this aspect of the model under constant experimental conditions, either the translation inhibitor anisomycin (ANI, 20 $\mu$M) or the transcription inhibitor Actinomycin D (Act D, 40 $\mu$M) were both applied for 45 min beginning 20 min prior to LTP induction. Importantly, both drugs do not interfere with synaptic function by actions other than the inhibition of translation and transcription (Linden 1996; Shitaka et al. 1996; Wiegert et al. 2009) and exposure to Act D and ANI had no effect on normal synaptic transmission or FM 1–43 release in the absence of LTP induction (Figs. 6A, 7A).

Inhibition of translation with ANI or transcription with Act D had no effect on LTP1 persistence (control $\tau = 37 \pm 15$ min, $n = 3$; ANI $= 68 \pm 29$ min, $n = 4$; Act D $= 51 \pm 16$ min, $n = 4$) or magnitude at 160 min post-TBS (control $= 5 \pm 1\%$, $n = 5$; ANI $= 1 \pm 2\%$, $n = 4$; Act D $= 1 \pm 1\%$, $n = 4$) (Figs. 6A, 7A). This is consistent with this form of LTP depending only on post-translational modifications. Inhibition of translation with ANI significantly reduced LTP2 magnitude at 160 min post-4TBS (control $= 27 \pm 7\%$, $n = 11$; ANI $= 9 \pm 4\%$, $n = 6$; $P < 0.05$) and enhanced its

**Figure 4.** LTP2 and LTP3 and associated enhanced release are dependent on NO synthesis. (A) Mean percent change in fEPSP slope in controls and slices incubated with L-NAME (100 $\mu$M) during the 1TBS induction protocol. Application of L-NAME did not affect LTP1 persistence. Data for nonetanolized L-NAME incubated slices are also plotted. (Inset) Representative fEPSPs for the period immediately prior to LTP induction (black) and recording termination (gray). (Top) Control, (bottom) L-NAME. Scale bars $= 1$ mV, 10 msec. (B) Histogram of mean FM 1–43 destaining kinetics ($t_{1/2}$) in LTP1 experiments showing L-NAME did not affect exocytotic rate 160 min post-1TBS. (C) L-NAME reduced LTP2 persistence and (D) eliminated enhanced release at 160 min post-4TBS. (E) L-NAME reduced LTP3 persistence and (F) eliminated enhanced release 160 min post-8TBS. (*) $P < 0.05$.

**Figure 5.** LTP2 and LTP3 and associated enhanced release are dependent on extracellular NO signaling. (A) Mean percent change in fEPSP slope in controls and slices incubated with cPTIO (40 $\mu$M) during the 1TBS induction protocol. Application of cPTIO did not affect LTP1 persistence. Data for nonetanolized cPTIO incubated slices are also plotted. (Inset) Representative fEPSPs for the period immediately prior to LTP induction (black) and recording termination (gray). (Top) Control, (bottom) cPTIO. Scale bars $= 1$ mV, 10 msec. (B) Histogram of mean FM 1–43 destaining kinetics ($t_{1/2}$) showing cPTIO did not affect LTP1 exocytotic rate 160 min post-1TBS. (C) cPTIO reduced LTP2 persistence and (D) eliminated enhanced release at 160 min post-4TBS. (E) cPTIO reduced LTP3 persistence and (F) eliminated enhanced release 160 min post-8TBS. (*) $P < 0.05$. 
mechanisms, our data suggest an intriguing possibility that enhanced exocytosis is also protein synthesis dependent. As expected, neither Act D nor ANI had any effect on LTP1 exocytotic rate measured at 160 min post-TBS (control \(t_{1/2} = 172 \pm 32\) sec; ANI = 136 \(\pm\) 5 sec, \(n = 52\) terminals from four slices; Act D = 134 \(\pm\) 3 sec, \(n = 88\) terminals from four slices) (Figs. 6B, 7B).

Together with the LTP1 decay data this supports the idea that LTP1 is expressed post-synaptically and independently of protein synthesis.

In concordance with LTP2 decay data ANI prevented the increase in exocytotic rate that was observed in controls 160 min post-LTP2 induction, whereas Act D had no significant effect (control \(t_{1/2} = 58 \pm 7\) sec; ANI = 126 \(\pm\) 10 sec, \(n = 91\) terminals from six slices; Act D = 56 \(\pm\) 5 sec, \(n = 47\) terminals from five slices) (Figs. 6D, 7D).

The presynaptic component of LTP is protein synthesis dependent

Although the requirement for protein synthesis in long-lasting LTP is generally associated with post-synaptic expression
Intriguingly, while the enhanced exocytic rate associated with LTP3 was also prevented by ANI, it remained unaffected by inhibition of transcription with Act D (control $t_{1/2} = 63 \pm 4$ sec; ANI = 128 ± 16 sec, $n = 36$ terminals from four slices; Act D = 67 ± 4 sec, $n = 53$ terminals from five slices) (Figs. 6F, 7F).

These data demonstrate that the enhanced presynaptic function associated with persistent LTP only depends upon translation and does not require gene transcription.

Discussion

It is increasingly accepted that LTP is not a unitary phenomenon and that even within one population of synapses different forms of LTP can coexist. Our findings confirm and elaborate upon the LTP1, 2, and 3 classification model at hippocampal CA3–CA1 synapses. LTP1, 2, and 3 were initially separated on the basis of persistence, but in recent times the differentiation has been expanded to include discrete Ca$^{2+}$ signaling mechanisms (Raymond and Redman 2002, 2006) and different requirements for post-synaptic action potentials during induction (Raymond 2008). We now show that the recruitment of a presynaptic expression mechanism further distinguishes LTP2 and LTP3 from LTP1. Using a more precise categorization of LTP with well-characterized and discrete mechanisms, our results elaborate on previous studies that showed persistent forms of LTP are associated with presynaptic functional changes (Huang and Kandel 1994; Zakharenko et al. 2001, 2003; Huang et al. 2005; Stanton et al. 2005; Ahmed and Siegelbaum 2009).

Role of new protein synthesis in LTP2 and LTP3

We have now confirmed a central feature of the LTP1, 2, and 3 model by showing that the maintenance of LTP3 is both transcription and translation dependent, whereas LTP2 maintenance requires only translation and LTP1 is protein synthesis independent. Protein synthesis-dependent LTP (historically known as late LTP) has been extensively characterized in the hippocampus, generally involving CREB-mediated transcription of key plasticity-related gene products (Abraham 2003; Pittenger and Kandel 2003; Benito and Barco 2010). In this case, mRNA can be trafficked to the dendrites where it is locally translated (Steward and Schuman 2003), or translated at the soma and the newly synthesized proteins trafficked to the dendrites to be utilized by “tagged” synapses (Frey and Frey 2008). On the other hand, a form of LTP requiring transcription-independent protein synthesis was first demonstrated in the dentate gyrus in vivo (Otani et al. 1989) and was later also identified in CA1 in vitro (Raymond et al. 2000; Gelinas and Nguyen 2005). In these studies LTP persistence was dependent on local protein synthesis, presumably from a constitutive pool of dendritic mRNA, triggered by activation of either Group 1 metabotropic glutamate receptors or β-adrenergic receptors.

In this study, both Act D and ANI were applied using concentrations and application durations that are known to induce substantial inhibition of transcription and translation. Here we applied the transcription inhibitor Act D for 45 min, beginning 20 min preinduction, which is similar to other studies (Huber et al. 2000; Kelleher et al. 2004; Leveson et al. 2004) and is sufficient to achieve substantial transcription reduction (Perry and Kelley 1970). The translation inhibitor ANI was also applied for a similar duration to that used in other studies (Stanton and Survey 1984; Huang and Kandel 1994; Hardingham et al. 1999; Gelinas and Nguyen 2005) and at a concentration known to block protein synthesis by >80% (Stanton and Survey 1984). In addition to its role in protein synthesis inhibition, ANI is also known to activate the p38/MAPK pathway under some conditions (Shifrin and Anderson 1999), and can elicit synaptic depression at specific frequencies (Xiong et al. 2006). We conclude it unlikely that these or other nonspecific effects of ANI and Act D are significant in our experiments, as inclusion of both drugs in the bath did not appreciably affect the magnitude of the fEPSP (Figs. 6A, 7A).

Until now there has been little to link the protein synthesis requirements of LTP2 and LTP3 to a presynaptic expression mechanism. Our data show that in the absence of translation the enhanced presynaptic function associated with the maintenance of both LTP2 and LTP3 is eliminated. Interestingly, although LTP3 as measured via fEPSPs is significantly impaired by Act D, the presynaptic expression component remains intact. This suggests that LTP3 maintenance also involves a significant post-synaptic expression mechanism. The potentiation remaining in the absence of transcription likely reflects the presynaptic component, whereas the post-synaptic expression mechanism, which presumably underlies the greater persistence of LTP3 over LTP2, appears to be supported by new transcription. It should be noted however, that since the CA3 cell bodies containing the presynaptic transcriptional machinery are removed during our hippocampal slicing protocol, it remains possible that presynaptic transcription may contribute to LTP persistence in vivo. Indeed, changes in the levels of several synapse-related transcripts have been observed in area CA3 following LTP induction at Schaffer collateral synapses (for review, see Abraham and Williams 2003).

Previous studies using FM 1–43 or synaptophysin–Hourin-expressing mice suggest that L-type voltage-dependent Ca$^{2+}$ channel (L-VDCC) activation is necessary for LTP involving a presynaptic component (Zakharenko et al. 2001; Bayazitov et al. 2007). Certainly, with respect to LTP3 our data are consistent with this since LTP3 is dependent on somatic Ca$^{2+}$ signaling via L-VDCCs (Raymond and Redman 2006). However, our model suggests that L-VDCCs are not obligatory for activating a presynaptic LTP expression mechanism since LTP2, which is independent of L-VDCCs (Raymond and Redman 2002, 2006), also involves a presynaptic enhancement. Rather, our data suggest that L-VDCCs are upstream of a crucial common effector of presynaptic change—protein synthesis—and that different signaling pathways (e.g., internal Ca$^{2+}$ release via IP$_3$ receptors) activated by weaker stimulation protocols may also suffice.

The translation required for the presynaptic component of both forms of LTP could either occur post-synaptically, or perhaps more interestingly presynaptically in response to NO. Local post-synaptic protein synthesis has been extensively characterized (Steward and Levy 1982; Sutton and Schuman 2006; Bramham and Wells 2007) and is often associated with LTP. There is also mounting evidence to support a role for translation in presynaptic terminals (for review, see Akins et al. 2009). Local presynaptic protein synthesis has been established in both immature and mature rat hippocampal cultures (Sebeo et al. 2009), in forms of LTP and long-term depression in Xenopus nerve–muscle cultures (Zhang and Poo 2002), in corticostriatal fibers (Yin et al. 2006), and in hippocampal mossy fiber–CA3 synapses (Huang and Hsu 2004).

Onset of presynaptic expression

Another interesting aspect of the current findings is the difference in onset timing of the presynaptic expression. While enhanced presynaptic function was only apparent for LTP2 at the 160 min post-TBS unload, enhanced release was also observed for LTP3 at 80 min post-induction. Other studies have reported a slow onset for the presynaptic expression associated with more persistent forms of LTP (Bayazitov et al. 2007) as is also observed here; however, this is the first demonstration that different forms of LTP may recruit changes in presynaptic expression at different times. Based on these results it seems likely that 8TBS stimulation may more
rapidly trigger the translation that is required for enhanced release. Indeed, the additional dendritic Ca\(^{2+}\) signal evoked by 8TBS over 4TBS (Raymond and Redman 2006) could more effectively activate Ca\(^{2+}\)-dependent dendritic translation (Atkins et al. 2004). Such temporal coordination of expression mechanisms may act to more effectively enhance synaptic strength, as the earlier onset of presynaptic changes reinforces the post-synaptic signaling events. Further investigations using methods with higher temporal resolution are necessary to address these issues.

**Nitric oxide**

Our studies have also identified a role for NO signaling in the persistence of LTP2 and 3, and the associated enhanced release. While a number of studies have reported a dependence of LTP persistence on NO signaling (Schuman and Madison 1991; Haley et al. 1992; Arancio et al. 1996a,b; Ko and Kelly 1999; Bon and Garthwaite 2001), this is the first study in which unequivocal NO-dependent effects on both LTP and presynaptic function have been directly measured. One previous study observed a partial effect of NO antagonism on FM 1–43 kinetics, but no effect on LTP (Stanton et al. 2005). The general scheme for NO action in hippocampal LTP is that Ca\(^{2+}\) influx via NMDA receptors activates NOS, which synthesizes NO from l-arginine (Boehning and Snyder 2003; Garthwaite 2008). Once synthesized, NO diffuses out of the post-synaptic cell and acts on soluble guanylyl cyclase in the presynaptic neuron, switching on cGMP-dependent protein kinases. Our results are consistent with this hypothesis, showing that NO production and diffusion into the extracellular space is necessary for LTP-associated presynaptic enhancement. Alternatively, NO may act on astrocytes or other intermediary cells that then initiate signaling to the presynaptic neuron. Indeed, there is increasing evidence to support a role for astrocytes in the regulation of LTP in culture and slice preparations (Filosa et al. 2009; Bélair et al. 2010; Henneberger et al. 2010). Although it is possible that NO is acting post-synaptically to prolong LTP (Ko and Kelly 1999) it seems unlikely in our experiments given the concomitant inhibition of the LTP-associated presynaptic enhancement by NO antagonists. If this were the case NO would have to be involved in the release of a second retrograde messenger.

Nevertheless, it is interesting to speculate whether NO is upstream of or downstream from the protein synthesis that we now show is necessary for presynaptic LTP expression. NO production per se could depend upon local dendritic protein synthesis that is known to be important for some forms of LTP (Raymond et al. 2000; Gelin and Nguyen 2005), although evidence so far suggests increased NOS expression only at the very late stages of LTP maintenance (Abraham and Williams 2003). Alternatively, a NO–cGMP–MAP kinase pathway has been implicated in synaptic plasticity and memory (Izumi et al. 2008; Ota et al. 2008) and MAPK/ERK signaling is a well-known activator of local protein synthesis in dendrites (Sutton and Schuman 2005). Since it is unlikely that NO is acting post-synaptically in our experiments, it is possible that this cascade is activated presynaptically to trigger local protein synthesis in or near synaptic terminals. An alternate possibility is that NO signaling acts in parallel with protein synthesis and that both pathways therefore act synergistically to trigger enhanced exocytosis.

**Concluding remarks**

The LTP1, 2, and 3 model is considerably strengthened by the addition of our current findings. It is now abundantly clear that CA3–CA1 synapses can support at least three mechanistically discrete forms of LTP. LTP1 is dependent entirely on local, post-synaptic induction, expression, and maintenance mechanisms. LTP2 induction involves dendritic Ca\(^{2+}\) signaling, modulated by backpropagating action potentials, and both pre- and post-synaptic expression mechanisms maintained by local dendritic and/or presynaptic protein synthesis. Finally, LTP3 involves cell-wide mechanisms, including somatic Ca\(^{2+}\) signaling during induction, a presynaptic expression component dependent on protein synthesis, and a post-synaptic expression mechanism supported by gene transcription. There is evidence for a similar LTP repertoire at the perforant path input to the dentate gyrus (Abraham and Otani 1991; Reymann and Frey 2007) and whether such variants exist in other synaptic pathways outside the hippocampus will be an interesting question for future studies.

Another important question is whether each form of LTP subserves a different function in learning and memory processing. We contend that many of the controversies in CA1 LTP research have arisen from the coexistence of LTP1, 2, and 3, and that appreciation of their individual characteristics and underlying mechanisms should assist in the design and interpretation of future investigations.

**Materials and Methods**

**Slice preparation and electrophysiology**

Experiments were performed in accordance with the Australian National University Animal Experimentation Ethics Committee guidelines. Male Wistar rats (6–8 wk) were anesthetized with isoflurane via inhalation, decapitated, and the brains submerged in ice-cold dissecting solution (mM: 124 NaCl, 3.2 KCl, 1.25 NaH\(_2\)PO\(_4\), 26 NaHCO\(_3\), 0.5 CaCl\(_2\), 7 MgCl\(_2\) and 10 D-glucose, equilibrated with 95% O\(_2\)/5% CO\(_2\)). Transverse hippocampal slices (400 µm) were prepared and area CA3 was removed to reduce potential hyperexcitability. Slices were immediately transferred to a holding solution as above, except the Ca\(^{2+}\), Mg\(^{2+}\) and D-glucose concentrations were adjusted to 2.5 mM, 1.3 mM, and 25 mM, respectively. Slices were maintained at 34°C for at least 30–40 min before recording.

Slices were perfused in a continuous flow (≈2 mL min\(^{-1}\)) of recording solution (as per holding solution) at 32–34°C. Field excitatory postsynaptic potentials (fEPSPs) were recorded from stratum radiatum in area CA1 using glass microelectrodes (2–5 M\(_\Omega\)) filled with recording solution. Baseline synaptic responses were evoked by stimulation of the Schaffer collateral/commissural axons at 0.033 Hz (0.1-msec pulse width) with a Teflon-insulated tungsten bipolar electrode placed 100–200 µm away from the recording pipette. The stimulation intensity was adjusted to evoke fEPSPs of approximately two-thirds maximal amplitude. Only fEPSPs between 1 mV and 4 mV in amplitude were accepted. LTP was induced by TBS, consisting of trains of 10 × 100 Hz bursts (five pulses/burst) with a 200-msec interburst interval, at the test pulse intensity. When multiple trains were delivered (i.e., 4TBS and 8TBS) the intertrain interval was 30 sec.

**FM 1–43 loading and unloading**

After recording stable Schaffer collateral-evoked fEPSPs for at least 20 min synaptic vesicles were loaded by delivering a train of 1200 stimuli (at 10 Hz) in the presence of FM 1–43 (5 µM, Biotium Inc.) and D-APV (100 µM, Tocris) (Fig. 1A). The dye application continued for 1 min past the duration of the loading stimuli to ensure completion of endocytosis and dye uptake. FM 1–43 was then washed out in the presence of ADVASEP-7 (0.5 mM, Biotium) for 20 min to remove any extracellularly bound dye (Kay et al. 1999). Destaining was achieved by stimulating at 1 Hz for 6 min in the presence of ADVASEP-7 and D-APV. Results from preliminary experiments demonstrated that this protocol maximally unloaded the dye (data not shown). In addition, the rate of destaining was both Ca\(^{2+}\)- and temperature-dependent in a manner consistent with it being a valid measure of exocytosis (data not shown; Zakharenko et al. 2001). Importantly, three bouts of

**Presynaptic enhancement in persistent LTP**

We contend that many of the controversies in CA1 LTP research have arisen from the coexistence of LTP1, 2, and 3, and that appreciation of their individual characteristics and underlying mechanisms should assist in the design and interpretation of future investigations.
staining and destaining were performed within one slice (base-line, 80-min and 160-min post-LTP induction) allowing for intra-slice comparisons of the effects of LTP on exocytotic rate at time points following LTP1, 2, and 3 induction.

Two-photon imaging
Fluorescence of labeled release sites was visualized using a Zeiss LSM 510 two-photon laser-scanning microscope with a water immersion objective (Achroplan 40 × 0.75 W). Two-photon excitation was achieved with a Ti:Sapphire laser tuned to 840 nm. All fields imaged were typically 10–20 μm deep, and were 40–80 μm away from the stimulating electrode. A series of 5–10 images (512 × 512) of 0.15 μm/pixel in the x–y axes) were taken at 1–2 μm intervals in the z-plane. These z-stacks were repeated every 30 sec, beginning 1 min prior to and during the unloading stimulation. In offline analyses, maximal z-projections were generated from each image series to create one image per 30-sec time point. Circular regions of interest were defined around the center of brightly stained punctate fluorescent spots, and 2–69 terminals and three background regions were measured at each time point. Only puncta that satisfied four predetermined criteria were included in analysis. These were: (1) diameter of 0.5–1 μm, (2) approximately circular shape, (3) minimal x–y movement, and (4) activity-dependent destaining that was well fit by a first order exponential decay function. A fluorescence time course was generated by subtracting the average background fluorescence at each time point then normalizing the fluorescence of each punctum at each time point to the average of the pre-stimulus values. Photobleaching was corrected for by normalizing to the average background fluorescence at the corresponding time point throughout the unloading stimulation. The half-time of decay of fluorescence intensity during unloading (t1/2) was calculated for each punctum from single exponential decay curves fitted to the 6 min of stimulus-induced destaining.

Data analysis
Two-tailed unpaired Student’s t-tests were used to determine statistical significance at the 95% confidence level for all analysis (unless otherwise stated). All data represented as mean ± SEM.

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A protein synthesis and nitric oxide-dependent presynaptic enhancement in persistent forms of long-term potentiation

Victoria P.A. Johnstone and Clarke R. Raymond

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