Modulation of hippocampal long-term potentiation and long-term depression by corticosteroid receptor activation

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Long-term potentiation (LTP) and long-term depression (LTD) are lasting changes in synaptic efficacy which may underlie memory and learning processes. Hippocampal LTP has been shown to vary inversely with the degree of stress or adrenal activity, and adrenal corticosteroids (CORT) have been strongly implicated in this process. The present in vitro study was undertaken to further examine the role of glucocorticoids in LTP, and to extend this analysis to heterosynaptic LTD. Animals were injected twice daily (s.c.) for 2 days prior to sacrifice with either the corticosteroid biosynthesis inhibitor metyrapone (200 mg/kg/day) or CORT (20 mg/kg/day). Noninjected and saline-injected controls were also examined. CA1 population spike amplitudes and field EPSP slopes were assessed in response to activation of two separate stratum radiatum inputs. One input received tetanic stimulation and both inputs were monitored for 30 min following tetanization to determine long-term changes in synaptic efficacy (homosynaptic LTP of tetanized synapses; heterosynaptic LTD of non-tetanized synapses). Animals exhibiting low and high serum CORT titers exhibited significantly less LTP and LTD than did animals exhibiting moderate levels of CORT. In addition, in vitro perfusion of RU-28362, a Type II glucocorticoid receptor agonist, markedly reduced both LTP and LTD. There was no correlation between serum CORT and serum glucose titers, nor was there any correlation between serum glucose and either LTP or LTD. These results are consistent with recent reports of an inverted-U function for CORT levels versus primed-burst LTP (Bennett, Diamond, Fleshner, & Rose, 1991; Diamond, Bennett, Fleshner, & Rose, 1992) and further demonstrate that manipulation of corticosterone in adrenal-intact animals affects LTP and LTD in a similar fashion.

It is well established that the hippocampus and related brain structures are rich in specific corticosteroid receptors (CSRs) (Fuxe et al., 1985; McEwen, Wallach, & Magnus, 1974; McEwen, Weiss, & Schwartz, 1969; Reul & de Kloet, 1986). At least two types of cytosolic CSRs have been differentiated on the basis of their steroid binding characteristics: Type I (MR, mineralocorticoid/corticosterone) and Type II (GR, glucocorticoid) receptors. In the rat, Type I receptors exhibit the higher affinity for corticosterone (CORT, the naturally occurring endogenous glucocorticoid in the rodent) and are probably tonically activated even under conditions of low plasma titers of CORT. In contrast, Type II receptors exhibit a much lower affinity for CORT and, as such, appear to be activated mostly under conditions of high circulating CORT (e.g., diurnal peaks, stress; Dallman et al., 1987; de Kloet, Ratka, Reul, Sutanto, & van Eekelen, 1987; Reul, van den Bosch, & de Kloet, 1987). It appears that the Type I and Type II receptors are components of two functionally distinct systems that mediate differing metabolic and physiological responses within the hippocampus (see Joels & de Kloet, 1992a, 1992b; Schlatter & Dokas, 1989).

Because of the widely varying actions of corticosteroids on hippocampal function, and because of the prominent role of the hippocampus in memory and learning processes, a number of studies have been carried out in recent years to address the question of modulation of long-term potentiation (LTP) under varying conditions of stress or adrenal activity. Early studies demonstrated that LTP in the dentate gyrus is significantly diminished following adrenalectomy but can be restored by in vivo or in vitro administration of CORT (Dana et al., 1982; Dana & Martinez, 1984; Nowicky, Vardaris, & Teyler, 1983). In contrast, a number of investigators have demonstrated a significant impairment of LTP among animals exposed to CORT (Pavlides, Watanabe, & McEwen, 1993) or to a variety of stressful conditions, in comparison with unstressed controls (Diamond, Bennett, Stevens, Wilson, & Rose, 1990; Foy, Stanton, Levine, & Thompson, 1987; Shors, Seib, Levine, & Thompson, 1989). Similarly, several studies have indicated that surgical adrenalectomy (ADX) can prevent stress-related reductions in LTP (D-
Recently, in vivo studies of hippocampal region CA1 (Bennett, Diamond, Flesher, & Rose, 1991; Diamond, Bennett, Flesher, & Rose, 1992) have shown that the magnitude of primed-burst long-term potentiation (PB-LTP) varies with serum CORT levels according to an inverted-U shaped function, such that very low and very high CORT levels are associated with diminished PB-LTP, whereas midrange serum CORT titters (presumably sufficient to activate the Type I receptor only) are associated with a robust expression of PB-LTP. A recent report by Pavlides, Watanabe, Margarinos, and McEwen (1992) has indicated a strikingly similar pattern of LTP modulation by CORT in the dentate gyrus. In the latter study, surgically adrenalectomized (ADX) rats exhibited robust LTP in vivo following the administration of the Type I CSR agonist aldosterone, while exhibiting diminished LTP following administration of the Type II CSR agonist RU-28362 or vehicle (Pavlides et al., 1992).

Although LTP has been reported to vary with serum CORT titters, there is still little actual evidence that glucocorticoids (alone) are involved in the various reported effects. That is, many of the studies have utilized surgical ADX or stress (or both) to produce variations in serum CORT, without providing sufficient control of ADX- and stress-induced variations in adrenal catecholamines, adrenal opioids, and ACTH. The latter factors have also been implicated in the modulation of LTP (Gold, Delanoy, & Merrin, 1984; Shors, Levine, & Thompson, 1990a, 1990b) and memory and learning processes (de Wied, 1989; Gold & van Buskirk, 1975, 1976), and they should be considered as possible experimental confounds (or cofactors; cf. Axelrod & Reisine, 1984) in at least some of the glucocorticoid studies cited above.

In addition, most of the conclusions to date regarding LTP have been based on extracellular measures of population spike amplitude only, and more direct measures of altered synaptic efficacy by assessment of the field EPSP have been largely neglected. Moreover, although a homosynaptic primed-burst long-term depression (PB-LTD) was recently observed in very high CORT animals (Diamond, Bennett, et al., 1992), neither this nor other forms of synaptic plasticity such as heterosynaptic LTD have been systematically examined. Thus, a number of questions regarding the modulation of LTP and LTD by stress or ADX remain unresolved.

In the present experiments, adrenal-intact adult rats were administered either the corticosteroid biosynthesis inhibitor metyrapone or corticosterone as a means of directly manipulating serum CORT levels in vivo. LTP and heterosynaptic LTD were then assessed in vitro, utilizing measures of both the population spike and the field EPSP to assess changes in synaptic efficacy and gross hippocampal function. (preliminary results of this study have appeared in abstract form: Kerr, Huggett, & Abraham, 1993). Consistent with recent reports (Bennett et al., 1991; Diamond, Bennett, et al., 1992; Pavlides et al., 1992), our results indicate a strong inverted-U shaped relationship between serum CORT level and the magnitude of LTP. In addition, modulation of heterosynaptic LTD also conforms to an inverted-U shaped function. For both LTP and LTD, alterations in the slope of the field EPSP lend support to the hypothesis that the changes observed are due to direct effects on synaptic efficacy, as opposed to modulation of local inhibition or neuronal excitability.

METHOD

Hippocampal slices (400-μm transverse sections) were prepared from 3- to 4-month-old male Sprague-Dawley rats and were maintained fully submerged in a standard brain slice chamber. The slices were constantly perfused with an artificial cerebrospinal fluid consisting of 124.0 mM NaCl, 3.2 mM KCl, 1.25 mM NaH2PO4, 26.0 mM NaHCO3, 2.0 mM CaCl2, 2.0 mM MgCl2, 10.0 mM glucose, and 0.5 μM picrotoxin (33°-34°C, saturated with 95% CO2/5% O2). Picrotoxin has been shown to enhance the induction of LTD in the hippocampus (Abraham & Wickens, 1991; Kerr & Abraham, 1993) and was included here for that reason. In order to reduce potential hyperexcitability and the spread of seizure-like activity during tetanization, region CA3 was removed by a manual knife cut during hippocampal dissection (Figure 1). This procedure, previously employed by Abraham and Wickens (1991) among others, does not adversely affect the evoked responses normally observed in vitro.

CA1 field potentials, recorded from stratum pyramidale and stratum radiatum, were assessed in response to monosynaptic activation of separate Schaffer collateral/commissural inputs. Monopolar electrical microstimulation with 25-μm teflon-coated steel wires, applied on opposite sides of the recording site (see Figure 1), yielded robust synaptic responses. These microstimulation procedures have been shown to reliably yield greater than 95% separation of the activated converging afferent inputs, as evidenced by summation tests of the individual slopes of the fiber volleys (Kerr & Abraham, 1993). One input served as the conditioning pathway, the other (test) pathway received no concurrent stimulation, and recordings continued for at least 30 min posttetanization (Figure 3). Population spike amplitudes (in millivolts, defined as the height of the spike from peak negativity to a point of intersection with a tangent line along the rising face of the positive wave; see Figure 1 legend) and field EPSP initial slopes (millivolts/millisecond) were measured off line, using ASYST-based software written in our lab, and were expressed as percent change from baseline values. Slices not exhibiting stable baseline potentials for at least 10 min prior to conditioning were discarded from the study. Slices were not discarded for reasons of failure to exhibit LTP following conditioning. Student t tests and two-way analyses of variance (ANOVAs) were performed with repeated measures designs for the 10-, 15-, and 30-min posttetanization time points in order to assess group differences following the various treatments. Earlier time points were excluded from these analyses in order to avoid con-
Serum CORT and Glucose Titers

Radioimmunoassay of serum CORT revealed that it differed for the four groups assessed in Experiment 1 (Figure 2A; final ns indicated for each group below.). Saline-injected and noninjected controls exhibited low to moderate levels of serum CORT, respectively (mean ± SEM: SAL, 3.8 ± 0.7, n = 6; INT, 7.1 ± 3.8, n = 5). CORT-injected animals exhibited elevated serum CORT titers (25.4 ± 9.3, n = 4) consistent with those values observed in acutely or chronically stressed animals (Eldridge, Brodish, Klute, & Landfield, 1989; Foy et al., 1985; Sapolsky, Krey, & McEwen, 1985) respectively. Noninjected and saline-injected animals were included to control for stress effects due to handling and rehousing.

RESULTS

Figure 1. Top: Schematic diagram of hippocampal slice. Area CA1 was removed during dissection to reduce hyperexcitability. Extracellular recording electrodes were positioned in stratum pyramidale and stratum radiatum of area CA1; insets show typical evoked responses observed at these sites. Stimulation sites are indicated by the black dots superimposed over the Schaffer collateral-commissural fibers in stratum radiatum, on opposite sides of the recording sites. Bottom: Representative waveforms (population spike above, field EPSP below) illustrating methods of waveform analysis. Spike amplitudes (millivolts) were defined as the height of a line extending from the peak negativity to a tangent across the rising face of the positive wave (vertical dashed line). Field EPSP slopes (millivolts/millisecond) were defined as the rate of change along the most linear portion of the negative-going face of the EPSP.

tamination of the long-term changes with the more dramatic short-term depression normally arising after strong tetanization. Statistical significance was determined at a confidence level of p < .05.

Blood Collection, CORT, and Glucose Assays

Following anesthetization (ether vapor) and rapid decapitation, trunk bloods were collected from each animal. Samples were maintained on ice and allowed to clot for approximately 3 h; then they were centrifuged at 800 × g for 10 min to yield whole-cell and platelet-free serum. Serum corticosterone was assayed with the use of a commercially prepared radioimmunoassay (RIA) highly specific for rat corticosterone (Coat-a-Count RIA Kit; Diagnostic Products Corp., Los Angeles, CA). Minimum detectable levels for this assay are less than 1 μg/dl, and nonspecific binding is negligible (≤0.6%). Under our assay conditions, parallelism tests indicated no interference by metyrapone or any other endogenous serum components, and although not assessed during the present study, intrasand interassay coefficients of variation for this assay generally vary between 4.0%-4.3% and 4.8%-5.8%, respectively.

Because CORT can affect peripheral glucose titers, and because brain glucose levels have been shown to correlate with certain types of learning (Gold, 1986; Lee, Graham, & Gold, 1988), we also assessed serum glucose levels in all animals used in Experiment 1 of the present study. Glucose titers were determined with the use of an automatic analyzer and a commercially prepared diagnostic kit (Glucose HK kit, Roche Inc.) utilizing a standard two-step enzymatic reaction (hexokinase and glucose-6-phosphate dehydrogenase) yielding NADH quantified by measurement of absorbance at 340 nm.

Experiment 1

Twenty-four rats (average body weight, 328 g; range, 310-365 g) were divided into four groups: noninjected intact controls (INT, n = 8), saline-injected controls (SAL, n = 6), and two groups consisting of animals injected with either the corticosteroid biosynthesis inhibitor metyrapone (Sigma Chemical Co., St. Louis; MET, 200 mg/kg/day, s.c., n = 5) or corticosterone (Sigma Chemical Co.; CORT, 20 mg/kg/day, s.c., n = 5). All animals were weighed and rehoused in isolation. Experimental subjects and saline-injected controls were each injected twice daily (two half-doses), once at 9:00 a.m. and again at 4:00 p.m. for 2 days prior to sacrifice, and once again on the morning of Day 3, approximately 2 h prior to sacrifice. This method was adopted to ensure slow release, yielding relatively constant levels of metyrapone and CORT throughout the treatment period and to ensure that any CORT-related genomic events were at equilibrium well before the in vitro studies. These dosages are within ranges previously reported to produce either relatively CORT-free, pharmacologically adrenalec-tomized animals (cf. Murison, Overmeier, Hellhammer, & Carmona, 1989; Stein & Sapolsky, 1988), or stress-equivalent high-CORT animals (cf. Sapolsky, Krey, & McEwen, 1985) respectively. Noninjected and saline-injected animals were included to control for stress effects due to handling and rehousing.

Experiment 2

To assess possible direct effects of metyrapone itself (i.e., effects on synaptic plasticity not related to inhibition of CORT synthesis), an additional group of animals was injected with both metyrapone and CORT (MET + CORT, n = 6, dosages as above). In addition, to directly assess the role of the Type II glucocorticoid receptor in the modulation of LTP and LTD, a highly specific Type II receptor agonist, RU-28362, was administered to slices by constant infusion (10 μM in artificial cerebrospinal fluid). These experiments were carried out in metyrapone-injected animals (MET + RU, n = 2) as well as noninjected controls (INT + RU, n = 4).
and are summarized as follows (glucose, mean millimoles/l ± SEM; SAL, 10.65 ± 0.34, n = 6; INT, 9.68 ± 0.20, n = 5; MET, 10.36 ± 0.10, n = 5; CORT, 11.18 ± 1.17, n = 4). These data are shown in Figure 2B. Regression analyses indicated no significant correlation between serum CORT and serum glucose levels.

**LTP and LTD: Experiment 1**

LTP and LTD were assessed in several slices from each animal, and group means were based on the total number of stable slices per group (total slices: SAL, n = 16; INT, n = 17; MET, n = 14; CORT, n = 15). All data summarized below are uniformly expressed as mean percent change from the baseline level (±SEM), at 30 min post-tetanization. Significant differences in both LTP and LTD were observed between conditions (Figure 3).

Saline-injected (low serum CORT) and CORT-injected (high serum CORT) animals exhibited only modest increases in population spike amplitude homosynaptically (SAL, 58.6% ± 6.8%; CORT, 57.1% ± 13.8%) and decreases heterosynaptically (SAL, −24.9% ± 4.6%; CORT, −26.9% ± 9.4%) following homosynaptic conditioning. Intact (noninjected) and metyrapone-injected animals (midrange serum CORT levels) exhibited greater LTP and LTD of the spike following tetanization (INT, LTP, 91.7% ± 10.6%; LTD, −31.7% ± 8.3%; MET, LTP, 102.1% ± 11.4%; LTD, −55.0% ± 6.3%). An analysis of variance (ANOVA) indicated a significant main effect of pharmacological treatment (LTP: F = 5.68, p < .001; LTD: F = 3.97, p < .002), and post hoc t tests revealed significant differences between the MET and SAL groups (p ≤ .01, both LTP and LTD) and the MET and CORT groups (p ≤ .025, both LTP and LTD).

EPSP slope measurements confirmed that the pattern of changes observed for population spikes were related to changes in synaptic efficacy. Saline-injected and CORT-injected animals exhibited small percent changes in EPSP slope (SAL, LTP, 5.8% ± 3.0%; LTD, −1.8% ± 1.9%; CORT, LTP, 1.4% ± 5.5%; LTD, −2.4% ± 13.8%) which were not significantly different between groups. Noninjected and metyrapone-injected animals generally exhibited considerably larger EPSP slope changes both homosynaptically and heterosynaptically than the saline- and CORT-injected groups (INT, LTP, 58.6% ± 6.8%; LTD, −7.3% ± 2.6%; MET, LTP, 17.1% ± 2.3%; LTD, −8.1% ± 2.5%). An ANOVA indicated a significant main effect across conditions for LTP of the EPSP (F = 2.56, p < .03), but not for heterosynaptic LTD of the EPSP (F = 0.94, p > .05). Post hoc analyses revealed significant differences between the MET and SAL groups (p = .01) and the MET and CORT groups (p < .03) for LTP of the EPSP.

Linear and polynomial regressions were performed to compare mean level of LTP or LTD (both spike and EPSP) for each animal versus serum CORT and serum glucose. Second-order polynomial regressions revealed strong correlations between serum CORT titer and magnitude of LTP or LTD for the homosynaptic spike, heterosynaptic spike, and heterosynaptic EPSP.
CORTICOSTERONE, LTP, AND LTD

CORT

METYRAPONE

baseline LTP 30 min

baseline LTP 30 min

Figure 3. Top: Population spikes and field EPSPs recorded from representative slices obtained from CORT-treated and metyrapone-treated animals. Each waveform represents the average of four to five individual, consecutive waveforms during the 1 min prior to tetanization (baseline) or at 30 min posttetanization (long-term potentiation, LTP). Double-spiking following tetanization is routinely observed in picrotoxin (see Method section). Calibration bars apply to all waveforms shown. Bottom: LTP and long-term depression (LTD) for the four groups in Experiment 1. Values shown are population spike amplitude (mean percent change from baseline ± SEM). Conditioning stimulation (see Method section) was administered at time = zero (arrow). Dashed line represents baseline levels measured prior to conditioning stimulation. Filled circles, CORT-treated animals; open circles, intact (noninjected) animals; filled triangles, metyrapone-treated animals; open triangles, saline-injected animals.

In addition, regression analyses were performed to establish curves of best fit describing group mean population spike and EPSP slope changes versus group mean serum CORT titers (Figures 4 and 5). Again, second-order polynomial functions were found to provide good descriptions of plasticity versus hormone level for each of the four conditions (spike LTP, \( r = 0.987 \); spike LTD, \( r = 0.960 \); EPSP LTP, \( r = 0.850 \); EPSP LTD, \( r = 0.956 \)). No significant correlations were observed between any of the measures of synaptic plasticity and serum glucose levels.

As shown in Figures 4 and 5, the CORT dependency of both LTP and LTD appears to be well described by inverted-U shaped functions. Accordingly, maximal synaptic plasticity, evidenced by LTP or LTD of either the population spike or EPSP slope, was evident at serum CORT levels of approximately 14–15 \( \mu \)g/dl. At lower or higher levels of serum CORT, plasticity (as assessed by any of these four measures) was markedly reduced.

**LTP and LTD: Experiment 2**

Additional experiments were conducted in order to assess the possible role of side effects of metyrapone on the modulation of synaptic plasticity seen in Experiment 1, and to determine whether plasticity could also be modulated by in vitro perfusion with the Type II glucocorticoid receptor agonist RU-28362. Animals injected with both metyrapone and CORT (n = 17 slices) exhibited reductions in LTP and LTD comparable to those levels seen in animals injected with CORT alone (spike LTP, 47.0% ±
Figure 4. Long-term potentiation (LTP) of the spike and field EPSP for all groups (both Experiments 1 and 2). Bar graphs (top) indicate mean percent change from the baseline level ± SEM; line graphs (below) illustrate mean percent change from baseline ± SEM for the four groups in Experiment 1 versus mean serum CORT titer for each group. Second-order regressions (lines of best fit) have been superimposed, illustrating inverted-U shaped modulation functions for CORT versus LTP.

10.5%; spike LTD, -22.5% ± 4.0%; EPSP LTP, 2.7% ± 2.1%; EPSP LTD, -4.7% ± 1.7%). Comparison of the MET + CORT group versus animals injected with metyrapone alone revealed significant differences (unpaired t tests, p ≤ .001) for all measures except LTD of the EPSP.

Slices taken from MET-injected animals and subsequently exposed to RU-28362 (n = 7 slices) exhibited significant reductions in LTP (p < .01) and LTD (p < .001) of the spike in comparison with MET-injected animals alone. Similarly, noninjected controls exhibited marked reductions in synaptic plasticity following in vitro perfusion with RU-28362 (n = 8 slices). Student t tests revealed significant differences (p < .01) between this group and the MET group on all measures except LTD of the EPSP.

Posttetanic Potentiation and Homosynaptic Depression

In addition to CORT-dependent differences in the induction of LTP and LTD, significant differences were also apparent between groups from 1 to 5 min posttetanization (Figure 3). Homosynaptic posttetanic potentiation (PTP) and heterosynaptic posttetanic depression (PTD) were reduced in both the SAL and the CORT groups relative to MET-injected animals. An ANOVA indicated a significant treatment effect for both LTP and LTD of the spike, as well as LTP of the EPSP, and post hoc comparisons revealed significant differences between the SAL and MET (p < .05) and CORT and MET (p < .05) conditions. Again, the pattern of change across the four groups conformed to an inverted-U shaped function for CORT level versus degree of synaptic plasticity.

In both Experiments 1 and 2, a number of slices from the low CORT, high CORT, and RU-28362 conditions not only failed to exhibit spike LTP homosynaptically following tetanization, but in some cases even exhibited a degree of lasting homosynaptic depression. No slices from the intermediate CORT conditions (INT and MET) failed to exhibit LTP (Table 1). EPSP slope changes, while relatively small and more variable, showed the same trend.

DISCUSSION

The results of the present study confirm a prominent role of corticosteroids in the modulation of synaptic plasticity. Using adrenal-intact animals, and thereby avoiding many of the experimental confounds associated with
stress or surgical ADX, we have shown that direct in vivo manipulations of serum CORT can affect both long-term potentiation as well as heterosynaptic long-term depression. Animals with either low or high serum CORT levels exhibited significantly less LTP and LTD than did animals with midrange CORT titers. A consistent inverted-U shaped function for CORT versus LTP or LTD was observed in vitro for both the population spike and the field EPSP. The fact that EPSP changes mirrored those occurring for the population spike suggests that corticosteroids modulate plasticity directly at the level of the synapse. The modulation of synaptic plasticity by our in vivo CORT manipulations most likely occurred as a direct result of CSR activation, as opposed to indirect CORT effects. First, selective activation of the Type II glucocorticoid receptor by in vitro application of RU-28362 produced significant reductions in LTP and LTD. Second, although serum glucose levels have been shown to correlate with memory processes in an inverted-U relationship (Gold & van Buskirk, 1975; Lee et al., 1988), in the present study serum glucose levels did not vary between groups and no relationship was observed between serum CORT and glucose, or between serum glucose and either LTP or LTD.

These findings are similar to those of recent electrophysiological studies conducted by Bennett et al. (1991) and Diamond, Bennett, et al. (1992) in hippocampal region CA1, and by Pavlides et al. (1992) in the dentate gyrus, in which inverted-U shaped functions were observed for CORT (or specific CSR activation) versus LTP. In the Table 1

| Percent of Slices Either Failing to Exhibit Long-Term Potentiation or Exhibiting Homosynaptic Depression of the Population Spike |
|---------------------------------------------------------------|
| MET | MET | INT |
| SAL | INT | MET | CORT | CORT | RU | RU |
| No LTP | 7 | 0 | 0 | 14 | 6 | 0 | 0 |
| (100-115%) | | | | | | | |
| Homosynaptic LTD | 0 | 0 | 0 | 14 | 19 | 14 | 44 |
| (<100%) | | | | | | | |

Note—LTP, long-term potentiation; LTD, long-term depression; SAL, saline-injected controls; INT, noninjected intact controls; MET, injected with metyrapone; CORT, injected with corticosterone; RU, Type II receptor agonist RU-28362, administered in MET or INT animals.
been associated with lasting elevations of adrenocorticosteroid receptor activity that is likely to ensure a steady release and presumably constant inhibition of CORT synthesis for the 2 days prior to sacrifice. However, chronic metyrapone treatment has significantly reduced serum CORT levels in this group of animals. The reduced serum CORT levels in this group of animals while unexpected, proved fortuitous in that the robust expression of LTP and LTD seen in the MET-injected animals is due to a side effect of the metyrapone itself. However, animals injected with both MET and RU-28362 produced reductions in both forms of LTP/LTD consistent with previous reports demonstrating (1) a dose-dependent suppression of PB-LTP by the Type II glucocorticoid stereo-steroid receptor agonist dexamethasone in urethane-anesthetized ADX rats (presumably exhibiting high ACTH in vivo) (Diamond, Branch, Bennett, Fleschner, & Rose, 1992), and (2) a Type II receptor dependent suppression of LTP in the dentate gyrus of ADX rats (Pavlices et al., 1992). We also considered the possibility that the robust expression of LTP and LTD seen in the MET-injected group was due to a side effect of the metyrapone itself. However, animals injected with both MET and CORT exhibited significant reductions in both forms of synaptic plasticity suggesting either that MET does not exert non-CORT related modulatory effects on LTP/LTD, or that the co-administration of CORT is sufficient to override or reverse this modulation.

Corticosteroid receptor up- and down-regulation have been demonstrated following various long-term manipulations (Reul et al., 1987), but it does not seem likely that the group differences seen in this study are due to individual differences in receptor capacity. In general, CSR up-regulation in young adult rats does not become apparent before 3–7 days of ADX (Eldridge, Fleenor, Kerr, Campbell, & Landfield, 1989), and significant down-regulation of CSRs has not been observed earlier than 1 week or longer following exogenous CORT administration (Meaney, Sapolsky, & McEwen, 1985) or chronic stress (Eldridge, Brodish, et al., 1989). Our manipulations took place over a period of 2 days (sacrifice on the morning of the 3rd day), and presumably sufficient time had not elapsed for MR or GR autoregulation. The finding that RU-28362 produced reductions in LTP/LTD suggests that Type II receptor occupancy facili-
tates synaptic plasticity, but that Type II receptor occupancy is inhibitory. This idea is consistent with the findings of Joels and de Kloet that Type I and Type II receptor activation produces opposite effects on cell excitability. Type I receptor activation produces increases in cell excitability, as is evidenced by measures of spike accommodation and suppression of 5-HT induced hyperpolarizations (Joels & de Kloet, 1990; Joels, Hesen, & de Kloet, 1991), whereas Type II activation produces consistent decreases in cell excitability, which are apparently due to prolongation of the Ca\(^{2+}\)-dependent afterhyperpolarization (AHP\(_c\); Joels & de Kloet, 1989, 1990; Kerr, Campbell, Hao, & Landfield, 1989).

Specific mechanisms underlying the modulation of LTP/LTD by CORT have not been identified. However, because of the fact that the Type I and Type II receptors exert opposing actions on CA1 AHPs, Bennett et al. (1991) and Diamond, Bennett, et al. (1992) have suggested that the group differences in PB-LTP that they observed may be related to differences in the effectiveness of the primed burst conditioning. That is, CORT enhancement of the AHP\(_c\) that follows the priming pulse may have shunted or offset the depolarization normally produced during the high-frequency burst delivered afterward. The same reasoning may also apply in our use of theta-burst conditioning in the present study. Thus, by differentially regulating cell excitability, CORT could be indirectly affecting the voltage-dependent activation of NMDA receptor/channels, which are known to be important for both LTP and heterosynaptic LTD (Abraham & Wickens, 1991; Christie & Abraham, 1992). As selective Type II CSR activation has been shown to modulate Ca\(^{2+}\) influx through both N- and L-type Ca\(^{2+}\) channels (Kerr, Campbell, Thibault, & Landfield, 1992), the hypothesis that corticosteroids modulate LTP and LTD through a Ca\(^{2+}\)-dependent process seems attractive and certainly merits further attention. For example, if that is true, then it is possible that the application of specific Ca\(^{2+}\) channel blockers might prevent CORT-related reductions in synaptic plasticity.

Regardless of the precise mechanisms underlying these effects, the present findings support the contention that glucocorticoids alone can modulate hippocampal synaptic plasticity, and that this modulation may depend on the degree of Type I or Type II CSR activation. Our data are consistent with recent reports demonstrating an inverted-U shaped function for CORT versus LTP in the dentate gyrus (Pavlides et al., 1992) and area CA1 (Bennett et al., 1991; Diamond, Bennett, et al., 1992), and they further demonstrate a similar pattern of corticosteroid modulation of heterosynaptic LTD. Collectively, these findings may be relevant to previous reports of LTP variation during the circadian rhythm (Dana & Martinez, 1984) or following ADX versus acute stress (Shors et al., 1990a), and may further relate to recent behavioral evidence of differential roles of the hippocampal MR and GR receptor systems in memory and learning (Bodnoff, Humphreys, Rose, & Meaney, 1992; Oitzl & de Kloet, 1992a, 1992b). When considered together with the inverted-U shaped learning functions previously demonstrated for peripheral epinephrine (Gold & van Buskirk, 1975) and central glucose (Lee et al., 1988), the relatively slow and lingering actions of corticosteroids may well act in concert with the more immediate adrenal catecholamines to facilitate the acquisition and consolidation of information in a stressful environment.

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