Temperature-Dependent Shift of Balance among the Components of Short-Term Plasticity in Hippocampal Synapses

Vitaly A. Klyachko and Charles F. Stevens
Howard Hughes Medical Institute and Molecular Neurobiology Laboratory, The Salk Institute, La Jolla, California 92037

Studies of short-term plasticity (STP) in the hippocampus, performed mostly at room temperature, have shown that small central synapses rapidly depress in response to high-frequency stimulation. This decrease in synaptic strength with synapse use places constraints on the use of STP as a dynamic filter for processing of natural high-frequency input. Here we report that, because of a strong but differential temperature dependence of STP components, the properties of STP in excitatory hippocampal synapses change dramatically with temperature. By separating the contributions of various STP processes during spike trains at different temperatures, we found a shift from dominating depression at 23°C to prevailing facilitation and augmentation at 33–38°C. This shift of balance among STP components resulted from a large increase in amplitudes of facilitation and augmentation ($Q_{10}$ = 2.6 and ~5.1, respectively) and little change in the amplitude of depression ($Q_{10}$ = 1.1) with temperature. These changes were accompanied by the accelerated decay of all three processes ($Q_{10}$ = 3.2, 6.6, and 2.1, respectively). The balance of STP components achieved at higher temperatures greatly improved the maintenance of synaptic strength during prolonged synaptic use and had a strong effect on the processing of natural spike trains: a variable mixture of facilitated and depressed responses at 23°C changed into a significantly more reproducible and depression-free filtering pattern at 33–38°C. This filtering pattern was highly conserved among cells, slices, and animals, and under various physiological conditions, arguing for its physiological significance. Therefore, the fine balance among STP components, achieved only at near body temperatures, is required for the robust function of STP as a dynamic filter during natural stimulation.

Key words: short-term synaptic plasticity; temperature dependence; natural stimulation patterns; synaptic filtering; information processing; hippocampus

Introduction

During ongoing activity, synaptic transmission is rapidly and dynamically modulated in a history-dependent manner. This short-term plasticity (STP) is thought to play a major role in information processing (Zucker and Regehr, 2002). Although properties of individual STP components have been studied extensively, the difficulty in analyzing the complex interplay of multiple forms of STP during the processing of realistic spike trains (Varela et al., 1997; Dobrunz and Stevens, 1999; Dittman et al., 2000; Ohliger-Frerking et al., 2003) hinders an understanding of their roles in synaptic computations at the majority of synapses. As a result, the function of STP has been characterized only in a few specific cases in which a single STP component dominates the synaptic response (Abbott et al., 1997; Tsodyks and Markram, 1997; Chung et al., 2002; Cook et al., 2003). An analysis introduced in studies of STP in the neuromuscular junction (Magleby, 1979; Magleby and Zengel, 1982; Kalkstein and Magleby, 2004), in which parameters of STP components were dissected from the decay phase of STP, can potentially circumvent this problem by separating contributions of overlapping components. Despite the importance of such an analysis for understanding STP function, it has not yet been attempted for CNS synapses. The goal of our study is to extend this approach for the analysis of STP components in small CNS synapses and to elucidate how the balance among these processes determines STP function during the processing of relevant spike trains.

Hippocampal CA3–CA1 synapses, used as a model system in our study, are one of the most well studied examples of CNS synapses expressing a mixture of multiple STP processes. Nevertheless, elucidating the role of STP in these synapses has been complicated by the fact that no strong correlation between the input spike pattern and the resulting synaptic response has been observed (Dobrunz and Stevens, 1999). The work cited above, however, as well as many other studies of hippocampal STP, has been performed at room temperature. A number of recent studies have shown that synaptic transmission can be affected dramatically by changes in temperature on both physiological and structural levels (Hardingham and Larkman, 1998; Dinkelacker et al., 2000; Taschenberger and von Gersdorff, 2000; Volgushev et al., 2000; Pyott and Rosenmund, 2002; Fernandez-Alfonso and...
Ryan, 2004; Roelandse and Matus, 2004; Micheva and Smith, 2005; Kushmerick et al., 2006). These findings suggest that contributions of different STP components, and the resulting computations performed by STP, may change significantly with temperature.

Here we have separated contributions of all major STP components expressed in excitatory hippocampal synapses to investigate how their roles in the processing of constant frequency and natural stimulation patterns change with temperature. Our data reveal a shift among STP components from dominating depression at 23°C toward prevailing facilitation and augmentation at 33–38°C. This shift greatly improves the maintenance of synaptic strength during high-frequency stimulation and has a strong effect on the processing of natural spike patterns. Our results indicate that a fine balance among STP components, achieved only at near physiological temperatures, is required for a robust and physiologically relevant STP function during natural stimulation.

Materials and Methods

Preparation. Transverse hippocampal slices (400 μm) were prepared from 14- to 25-d-old (for whole-cell recordings) or 21- to 35-d-old (for field potential recordings) Long–Evans rats as described previously (Pouille and Scanziani, 2001; Losonczy et al., 2002). In brief, slices were cut on a Leica Vibratome in ice-cold, modified ACSF containing (in mM): 85 NaCl, 75 sucrose, 2.5 KCl, 1.25 NaHPO4, 26 NaHCO3, 0.5 CaCl2, 4 MgCl2, 13 glucose, bubbled with 95% O2/5% CO2. Slices were incubated for ~1 h at 32–33°C and then held at the room temperature for 4–4 h before recordings in standard ACSF solution, of the same composition as above, but with 120 NaCl, 2 CaCl2, 1 MgCl2 (unless noted otherwise) and no sucrose. The CA3 region was surgically separated in each slice with incision to prevent recurrent excitation.

Electrophysiological recordings. Schaffer collaterals were stimulated (100 μs; 50–200 μA current injections) via a bipolar electrode placed in the stratum radiatum, and either whole-cell currents or extracellular field potentials were recorded using an Axopatch 200B amplifier. EPSCs were recorded from CA1 pyramidal cells at the holding potential of ~74 mV unless noted otherwise. Micropipettes (3–6 MΩ) were filled with solution containing (in mM): 130 K-gluc, 10 KCl, 9 NaCl, 10 HEPES, 5 EGTA, 2.5 MgATP, 0.3 Li2ATP; pH 7.25, 290–300 mOsm. Membrane holding potentials were corrected a posteriori for liquid junction potential (~14 mV). Access resistance was monitored continuously, and cells with unstable access resistance (>20% change) were excluded from analysis. Field EPSP (fEPSP) recordings were made in a submerged chamber with the recording electrode positioned in the stratum radiatum, two-thirds of the way from dentritic tree area, and filled with solution of the same composition as standard ACSF, but with 20 HEPES, 0 NaHPO4, 0 NaHCO3, pH adjusted to 7.25. All recordings were made in the presence of the GABA R antagonist picrotoxin (100 μM) or the more specific GABA R antagonist gabazine (10 μM) to block inhibitory responses and the NMDA receptor antagonist aminophosphonopentanoic acid (50 μM) to prevent possible long-term effects.

Temperature measurement and control. Bath temperature in the recording chamber was monitored continuously with a bead thermistor probe positioned within ~5 mm from the slice. The probe provided a feedback readout for the automatic heater controller that continuously adjusted the temperature to a desired setting via an in-line solution heater positioned immediately before the recording chamber. For any temperature setting, slices were kept for 20 min at the desired temperature before the first control baseline measurements were started. With several additional minutes used to establish a baseline and ~2 min of control recordings at 0.1 Hz separating each train, the first actual recording was made after ~25 min. The 25 min waiting time was sufficient to establish stable recordings at all temperatures because synaptic responses were highly repeatable for the duration of recordings. To reduce variability associated with small changes in temperature during recordings (<0.5°C), parameters of STP components were measured multiple times in each slice and in random order.

Separation of STP components. To estimate the parameters of STP components, constant-frequency trains of 10–150 stimuli at 2–40 Hz were used. The amplitude and time course of each component were determined sequentially, starting with the slowest one, by selecting an appropriate time window, when that component dominated the synaptic response (see Fig. 2). For each train presentation, synaptic responses were normalized to an average of five control responses immediately preceding the train. Responses to four to six train presentations were averaged for each stimulation protocol. Parameters of each component were estimated using the monoeponential Levenberg-Marquardt least squares fitting algorithm in Origin and later used to compensate, where necessary, for the contributions of that component to the mixture with the faster STP processes (see Fig. 2). This approach is based on an analysis introduced in the studies of STP in the neuromuscular junction (NMJ) (Magleby and Zengel, 1976a, 1982; Kalkstein and Magleby, 2004), in which normalized synaptic response can be described as follows (see Fig. 2):

\[
\text{EPSP/EPSP}_0 = (1 + F)(1 + AUG)(1 - DEP). \tag{1}
\]

A multiplicative relationship between the components of potentiation and depression arises if these processes act independently to modulate the transmitter release, as has been suggested previously (Magleby and Zengel, 1976a; Magleby, 1979; Varela et al., 1999; Dittman et al., 2000; Zucker and Regehr, 2002; Abbott and Regehr, 2004; Kalkstein and Magleby, 2004). A multiplicative relationship between different forms of potentiation is suggested by previous studies in the NMJ (Magleby, 1979; Magleby and Zengel, 1982; Kalkstein and Magleby, 2004).

In this approach, parameters of augmentation (AUG) were determined in the time window of 5–30 s (10–60 s at 23°C) after the end of the train, when faster STP components, i.e., facilitation [τ ~ 150 ms (~500 ms at 23°C)] and depression [τ ~ 1.2 s (~2.5 s at 23°C)], had decayed completely. In some cases, when measurable amounts of a very slow component of depression (τ ~ 50–60 s) (Micheva and Wesseling, 1999) were observed, the decay of AUG was corrected for the very slow depression, assuming a multiplicative relationship between the components and taking into account that in the selected time window \( F = 0 \), \( \text{DEP} = 0 \) and calculating the amplitude of the very slow depression for each time point assuming monoeponential kinetics of decay. For the majority of stimulation paradigms used in this study, however, including all natural spike trains, the amplitude of the very slow depression was negligible and therefore did not contribute significantly to the temperature-dependent shift in synaptic response observed with all stimulus trains.

The kinetics of the rapid depression (DEP) was studied with a separate set of measurements by applying a single test pulse at different intervals [250 ms (500 ms at 23°C) to 6 s (5 s at 23°C)] after a 150-stimulus train at 10 or 40 Hz. Every time point was measured at least twice in each slice. Recovery from DEP was contaminated mostly by decaying facilitation and augmentation. To minimize the contribution from overlapping but rapidly decaying facilitation, the first time point was recorded at 250 ms at 33°C and at 500 ms at 23°C. Contributions from overlapping facilitation (determined with a paired-pulse protocol; see below) and augmentation were compensated using Equation 1 and assuming a multiplicative relationship among the components, as described above. In the selected time window, the contribution from the very slow component of depression (with \( \tau ~ 50–60 \) s) to the kinetics of the recovery from depression (\( \tau ~ 1–2 \) s) was negligible and was not corrected for.

Parameters of facilitation (\( F \)) were estimated with the commonly used paired-pulse protocol, which is insufficient to evoke other forms of STP (see Fig. 4) (Zucker and Regehr, 2002).

We also confirmed that the kinetics-based separation of STP components described above was consistent with a more direct approach, in which the entire data recorded after the stimulus train was fitted with Equation 1 directly. In these calculations we set \( F = 0 \), because facilitation had mostly decayed by the time the first data point for the recovery from depression was recorded, several hundred milliseconds after the train. This analysis produced results for the temperature-dependent changes in
depression and augmentation very similar to those observed with the first method, with \(Q_{10} = 1.1\) and 3.6 for the amplitudes and \(Q_{10} = 2.3\) and 6.0 for the time-constants of depression and augmentation, respectively.

Data analysis and presentation. Data were analyzed using software written in Matlab. Both peak amplitudes and initial slopes were analyzed and compared for consistency. Peak amplitudes were used for EPSCs, and initial slopes were used for fEPSPs. Peak amplitudes were measured relative to the baseline, which was calculated as an average of five points (1 ms) immediately preceding each spike. To account for the overlap of currents or potentials, which occurs mostly during natural stimulation at short interspike intervals (ISIs), the following method was applied to each set of responses to a stimulus train presentation and 6–8 preceding control measurements: first, a template of EPSC or fEPSP waveform was created by averaging responses separated by at least 100 ms from their neighbors and normalized to their peak values. For all ISIs \(<40\) ms, two previous responses were approximated by a template waveform scaled to their peaks, and the contributions to the current response were subtracted. Because of the rapid time course of EPSC and fEPSP decay, the contribution from the third preceding response during high-frequency epochs was indistinguishable from noise for all types of recordings. All data are presented as mean ± SEM. Statistical significance was evaluated using two-tailed \(t\) test.

For presentation only, the raw traces of spike train responses several seconds in duration (see Figs. 1, 5, 7) were down-sampled 4–10 times, and the baseline drifts were corrected either by using the method of moving average (2 and 10 Hz data) or by calculating an average for 1 ms intervals immediately preceding each spike and subtracting it from each ISI (40 Hz data).

Natural spike patterns. The natural stimulation patterns used here represent timings of action potential firing recorded in vivo from the hippocampal place cells of awake, freely moving rats generously provided by Drs. A. A. Fenton and R. U. Muller (Fenton and Muller, 1998). Six different patterns of 128 stimuli were used. Natural stimulation patterns were presented multiple times to the same cell or slice, separated by 2–3 min control sections at 0.1–0.2 Hz. Each subset of data was normalized to an average of four to six control responses immediately preceding each train. Spikes with ISIs \(<10\) ms were treated as a single stimulus, because the delay between the action potential firing and the peak of postsynaptic currents (potentials) prevented resolution of individual synaptic responses at shorter ISIs.

Results

Temperature-dependent shift from depression to potentiation

Our basic findings are illustrated in Figure 1A, in which we compare fEPSPs evoked in the CA1 area of the hippocampus by a series of constant-frequency trains at 2–40 Hz at room temperature (23°C) and at near physiological temperature (33°C). In agreement with previous studies (Dobrunz and Stevens, 1997; Brager et al., 2002; Sun et al., 2005), the synaptic response at room temperature consisted of a small initial increase in amplitude that was rapidly overcome by accumulating depression (\(n = 9–16\) slices) (Fig. 1A, black traces). The relative contribution of apparent depression (estimated as a difference between the maximal response and the minimal response during the train, normalized by the maximal response) increased with the frequency of stimulation from 10 ± 3% at 2 Hz to 45 ± 2% at 10 Hz, reaching 96 ± 4% at 40 Hz (Figs. 1A, 3C). In contrast, high-frequency stimulation at 33°C evoked a large increase in synaptic strength that persisted throughout the train at all frequencies tested (Fig. 1A, red traces). No significant depression was evident at frequencies \(<10\) Hz, and even at 40 Hz, slowly accumulating depression (32 ± 4%, \(n = 16\)) was mostly countered by synaptic potentiation (Fig. 1A, right panel) (compare with 96% of depression at 23°C). These data show that the balance among STP components shifts with temperature from the dominating depression at room temperature toward potentiation at near physiological temperatures.

Because fEPSP recordings represent a combined response from many cells, we sought to confirm these findings in individual CA1 neurons with the whole-cell recordings. Using minimal stimulation, we first confirmed the previous observations that different forms of STP coexist in putative single CA3–CA1 synapses (supplemental Fig. 1, available at www.jneurosci.org as supplemental material) (Dobrunz and Stevens, 1997). Next, we used multifiber stimulation at different temperatures to evaluate the temperature-dependent changes in STP during high-frequency trains. Figure 1B shows an example of whole-cell currents evoked in the same neuron by a 40 Hz, 150-stimulus train at three different temperatures: 23, 33, and 38°C. As in fEPSP recordings, the initial increase in the EPSC amplitude at 23°C was dominated by accumulating depression, which caused synaptic responses to decline rapidly (\(n = 11\) cells) (Fig. 1B, black traces).
(fEPSP and EPSC train responses are also shown on an expanded time scale in supplemental Fig. 2, available at www.jneurosci.org as supplemental material). At 33 and 38°C, however, responses showed large increases in amplitude that persisted throughout the train (Fig. 1B, red traces). Again, accumulation of apparent depression during trains proceeded much faster at room temperature, increasing with frequency of stimulation from 12 ± 5% at 2 Hz to 29 ± 6% at 10 Hz, and reaching 75 ± 4% at 40 Hz (but only 17 ± 4% at 33°C; p < 10\(^{-12}\)) [supplemental Fig. 3A, available at www.jneurosci.org as supplemental material (n = 7–11 cells; two to five train presentations per cell)].

These data show that the results reported by whole-cell and fEPSP recordings are basically similar, with the fEPSPs representing averaged whole-cell responses from multiple cells [the somewhat larger extent of apparent depression observed in fEPSP recordings may reflect the age difference of the animals used for whole-cell and fEPSP recordings (14–25 vs 21–35 d old, respectively) (Bolshakov and Siegelbaum, 1995; Hsia et al., 1998; Dekay et al., 2006)]. Therefore, the temperature-dependent shift among STP components is a robust effect that occurs both in single cells as well as in large populations of cells. In the experiments described below, we further confirmed that results reported by whole-cell and fEPSP recordings are equivalent for all of our major findings. Because this study required experiments with multiple train presentations and hours of continuous recordings, the experiments described below were typically performed using the more stable fEPSP recordings, with the key observations (data in Figs. 1, 3, 7, 8, and 9) being confirmed by whole-cell recordings.

As seen in Figure 1B, synaptic responses at 33 and 38°C were qualitatively similar, with only a slight difference in the magnitude of increase during trains. Thus, most of the shift from depression to potentiation occurred in the range of 23–33°C. A detailed analysis of the STP temperature dependence presented below confirmed this observation by showing that parameters of all STP components changed mostly in the 23–33°C temperature range but did not change significantly between 33 and 38°C (see Fig. 6). Because maintaining healthy slices at 38°C for prolonged periods of time is difficult as a result of oxygen escape from the solution at this temperature, we performed most of the high-temperature recordings at 33°C. The above observations gave us confidence that recordings made at 33°C accurately reflected properties of synaptic transmission at body temperature. As an additional control, however, we also confirmed the key observations at 38°C (Figs. 1, 3–8).

Postsynaptic recordings may reflect not only presynaptic changes attributable to STP, but they may also reflect several postsynaptic effects, such as receptor saturation and desensitization. We tested whether these processes could play an appreciable role in the observed shift in synaptic responses, such as receptor saturation and desensitization. We tested whether these processes could play an appreciable role in the observed shift in synaptic responses. Receptor desensitization and receptor saturation were reduced with cyclothiazide (CTZ; 100 \(\mu\)M) (Diamond and Jahr, 1995) and a commonly used low-affinity AMPA antagonist, \(\gamma\)-D-glutamylglycine (\(\gamma\)-DGG; 1–2 mM) (Liu et al., 1999; Wadiche and Jahr, 2001), respectively. The design of these experiments was similar to that used in several previous studies (Neher and Sakaba, 2001; Scheuss et al., 2002; Taschenberger et al., 2002). Although application of CTZ and \(\gamma\)-DGG increased the half-width of EPSCs and reduced their amplitude, as expected from previous reports (Diamond and Jahr, 1995; Hjelmstad et al., 1999; Liu et al., 1999), we found no significant effects of CTZ and \(\gamma\)-DGG on the pattern of synaptic responses to high-frequency trains or on the shift in synaptic responses with temperature (n = 7) (supplemental Fig. 4, available at www.jneurosci.org as supplemental material). These results show that under the conditions of our experiments, receptor desensitization and receptor saturation do not play a significant role in the observed effects. Thus, postsynaptic recordings can be reliably used to analyze changes in synaptic responses caused by STP.

**Separation of STP components**

The complex response to constant frequency stimulation, shown in Figure 1, resulted from the interplay of at least three major forms of STP: facilitation (F), augmentation (AUG), and depression (DEP). During the train (shown on an expanded timescale), all components were present simultaneously, and their contributions were difficult to identify. Because of different time courses of decay, however, various components became unmasked at different times after the end of stimulation. Facilitation, which decayed first with a time course of approximately hundreds of milliseconds, was studied separately with a paired-pulse protocol. The second most rapidly decaying process, recovery from DEP, dominated the synaptic response almost immediately after the end of the train (250 ms to 6 s (500 ms to 8 s at 23°C)). Finally, ~5 s (~10 s at 23°C) after the end of the train, AUG remained the last major STP component, because contributions from rapidly decaying F and DEP became insignificant. We took advantage of this time-based separation of STP components and traced their contributions backward, starting with the slower one. Parameters of the slower components were used in turn to compensate, where necessary, for their contributions to the mixture with the faster components, assuming a multiplicative relationship among the components of potentiation and depression (see Materials and Methods).

![Figure 2](https://via.placeholder.com/150)

**Figure 2.** Separation of STP components. Complex response to constant-frequency stimulation (representative fEPSP response to 20 Hz, 150 stimuli is shown) resulted from the interplay of at least three major forms of STP: facilitation (F), augmentation (AUG), and depression (DEP). The design of these experiments was similar to that used in several previous studies (Neher and Sakaba, 2001; Scheuss et al., 2002; Taschenberger et al., 2002). Although application of CTZ and γ-DGG increased the half-width of EPSCs and reduced their amplitude, as expected from previous reports (Diamond and Jahr, 1995; Hjelmstad et al., 1999; Liu et al., 1999), we found no significant effects of CTZ and γ-DGG on the pattern of synaptic responses to high-frequency trains or on the shift in synaptic responses with temperature (n = 7) (supplemental Fig. 4, available at www.jneurosci.org as supplemental material). These results show that under the conditions of our experiments, receptor desensitization and receptor saturation do not play a significant role in the observed effects. Thus, postsynaptic recordings can be reliably used to analyze changes in synaptic responses caused by STP.
slower components could be used in turn to compensate, where necessary, for their contributions to the mixture with the faster components, an approach pioneered by Magleby and Zengel (Magleby, 1979; Magleby and Zengel, 1982). To compensate for contributions from overlapping processes, we assumed that components of potentiation and depression interact multiplicatively, as had been suggested in previous studies of STP (Magleby, 1979; Magleby and Zengel, 1982; Tsodyks and Markram, 1997; Varela et al., 1997; Markram et al., 1998; Hempel et al., 2000; Okatan and Grossberg, 2000; Brager et al., 2002; Kalkstein and Magleby, 2004) (see Materials and Methods). Because the detailed mechanisms of interactions among STP components are poorly understood and might be more complex than assumed in our analysis, we also determined the extent to which this compensation affected our conclusions (see Discussion). Below, we describe the temperature dependence of STP components, separated as described above but presented in a different order for clarity.

Recovery from depression is much faster at warmer temperatures

Rapid synaptic depression is commonly interpreted as the depletion of the readily releasable pool (RRP) of vesicles, and therefore the time course of this component is determined by the RRP refilling rate (Zucker and Regehr, 2002; Abbott and Regehr, 2004). We compared the kinetics of depression at 23 and 33°C by applying test pulses at different intervals [250 ms to 6 s (or 500 ms to 8 s at 23°C) after 150-stimulus trains at 10 or 40 Hz (Fig. 3A (responses to 40 Hz trains are shown)). During this time window, changes in synaptic strength were governed mainly by recovery from depression (Fig. 2) because facilitation had mostly decayed, whereas longer lasting augmentation changed relatively little; however, contributions from other STP components, mainly decaying augmentation, were evident at 33°C (see additional test pulses at 6–9 s) (Fig. 3A). To isolate depression, then, we corrected synaptic responses for contributions from other STP processes by approximating each component with a monoexponential fit and assuming a multiplicative relationship among the components of potentiation and depression (see Materials and Methods).

This analysis revealed a much faster recovery from depression at warmer temperatures than at room temperature (τ = 2.5 ± 0.4 s at 23°C; τ = 1.18 ± 0.16 at 33°C; n = 10, 8; p < 0.01). Perhaps surprisingly, we also found that depression accumulated to a similar extent at room and near body temperatures (0.79 ± 0.03 and 0.84 ± 0.04, respectively; p > 0.49) (Fig. 3B). This finding may seem inconsistent with the observation that the amount of apparent depression during trains was significantly reduced at warmer temperatures (Fig. 3C). These results can be understood if we consider a situation in which depression accumulates linearly so that the amount of depression at the end of the train is proportional to the product of stimulus frequency, the amount of depression per stimulus, and the time constant of recovery. Because we found that the time constant for recovery is decreased approximately twofold at the higher temperature, the amount of depression per stimulus must increase at warmer temperatures to produce a similar extent of depression at the end of the train. If the mechanism of depression is the depletion of the RRP, this increase in depression per impulse is expected. The fact that at warmer temperatures components of potentiation compensate for depression to maintain elevated levels of transmitter release during constant frequency stimulation (see below) means that the release probability must be higher during trains at 33°C, and thus the average depression per impulse is also increased. Thus, the amplitude of apparent depression during trains reflects the interaction of several STP processes rather than synaptic depression alone. This analysis, also confirmed by whole-cell recordings (supplemental Fig. 3, available at www.jneurosci.org as supplemental material), emphasizes the need for separation of STP components to correctly estimate their contributions to synaptic response.

Both facilitation and augmentation increase strongly with temperature and decay much faster at body temperature

Two major components of STP that increase synaptic strength have been identified in hippocampal synapses: facilitation, with a time course on the order of hundreds of milliseconds, and augmentation, with a time course on the order of a few seconds (Magleby and Zengel, 1976b; Stevens and Wesseling, 1999). We first isolated facilitation with a commonly used paired-pulse protocol that was insufficient to evoke other STP components (Fig. 4) (Zucker and Regehr, 2002). Pairs of pulses at 10–1000 ms intervals revealed a large increase in the amplitude of facilitation with temperature (from 0.26 ± 0.03 at 23°C to 0.70 ± 0.05 at 33°C; extrapolated to t = 0; p < 10−5; n = 12, 16; two to five recordings per slice) (Fig. 4A, B) and an approximately threefold acceleration in the time course of decay (from 465 ± 79 ms to 143 ± 14 ms) (Fig. 4A, C). Both amplitude and time course of facilitation were essentially the same at 33 and 38°C (F38 = 0.66 ± 0.08; τ38 = 163 ± 21 ms; n = 5; p > 0.3 for both).

Augmentation, a longer-lasting component of potentiation, was evoked by high-frequency stimulation and could be isolated and studied a few seconds after the end of the stimulus train, when both facilitation and depression had decayed completely (Fig. 5A). To compare parameters of augmentation at different temperatures, we applied test pulses at 0.2 Hz starting 5 s (or 10 s at 23°C) after the end of a 150-stimulus, 10 Hz train (Fig. 5A), to
allow for complete refilling of the RRP ($\tau \sim 2.5$ s at $23^\circ$C and $\tau \sim 1.2$ s at $33^\circ$C). This particular stimulation protocol was chosen because it was insufficient to evoke measurable amounts of the very slow component of depression ($n = 9$ slices; data not shown); therefore, in the selected time window, the reported changes in synaptic strength were associated with pure augmentation. We found that the temperature dependence of augmentation was qualitatively similar to that of facilitation (Fig. 5B–D): augmentation underwent a fivefold increase in amplitude (from $0.76 \pm 0.08$ to $3.9 \pm 0.3$; $n = 13, 18$; extrapolated to $t = 0; p < 10^{-7}$) and decayed approximately seven times faster ($37 \pm 3$ vs $5.6 \pm 0.5$ s; $p < 10^{-5}$) as temperature increased from 23 to $33^\circ$C. As with facilitation, parameters of augmentation did not change significantly between 33 and $38^\circ$C ($p > 0.37$ for both; $n = 18, 4$) (Fig. 5C, D). Together, these data indicate that both components of potentiation exhibit strong temperature dependence, with a large increase in amplitude and a much faster decay at higher temperatures. In contrast, depression changes little in amplitude but decays twice as fast at higher temperatures (Table 1).

**Amplitudes and rates of STP components have different transition points in their temperature dependence**

To determine the range of temperatures at which the transition in properties of STP components occurred, we measured the amplitude and time constant of each process at several intermediate temperatures between 23 and $38^\circ$C (Fig. 6). In agreement with previous studies suggesting common underlying mechanisms of facilitation and augmentation (Zucker and Regehr, 2002), these processes underwent qualitatively similar temperature-dependent transitions in both amplitudes and rates. Interestingly, we also found that for both components of enhancement, the transition in amplitudes occurred several degrees higher ($2-4^\circ$C) than the transition in rates, suggesting that different temperature-dependent mechanisms may underlie these changes. The transition point for the rate of depression was shifted $3^\circ$C toward warmer temperatures relative to that of facilitation and augmentation, and no significant transition was observed for the amplitude of depression (Fig. 6). For all STP processes, changes in the amplitudes and rates were independent of the direction of the temperature change in the transition range, indicating that no significant hysteretic effects were observed in the temperature dependence of STP components (supplemental Fig. 5, available at www.jneurosci.org as supplemental material).

**Pharmacological and mechanistic separation of STP components**

In addition to the kinetics-based method applied above, several other approaches have been used to isolate various STP processes in the NMJ and other preparations. We therefore tested whether these methods are applicable to the hippocampal slice preparation to confirm our observation of the temperature-dependent shift in balance among STP components.

Because elevation in the residual calcium is commonly thought to underlie both facilitation and augmentation, chelating the residual calcium with EGTA-AM should eliminate both components of enhancement (Swandulla et al., 1991; Regehr and Delaney, 1994) and be expected to reveal pure depression. We found, however, that although application of EGTA-AM (250 $\mu$M) completely blocked paired-pulse facilitation after $\sim 30$ min of incubation, it was not sufficient to block augmentation even after $4$ h of incubation ($n = 6$) (supplemental Fig. 6, available at www.jneurosci.org as supplemental material). The effects of EGTA-AM on augmentation were nevertheless apparent as it progressively slowed down the increase in synaptic response during trains and prolonged the decay of augmentation with the incubation time, reaching saturation after $\sim 2$ h. Therefore, the approach based on the application of EGTA-AM could not be used in our preparation to separate STP components.

Previous studies suggest that augmentation can be isolated by lowering extracellular calcium to decrease the release probability and minimize depression (Dittman and Regehr, 1998). We reduced $[Ca^2+]_o$ to 0.75 mM and increased $[Mg^{2+}]_o$ to 2.25 mM and compared the responses to 150-stimulus trains at 23 and $33^\circ$C. Depression was greatly reduced at both temperatures (supplemental Fig. 7A, available at www.jneurosci.org as supplemental material), as expected from previous studies, whereas the temperature dependence of augmentation remained qualitatively similar to that found above with the kinetic separation of STP components (supplemental Fig. 7, B and C, available at www.jneurosci.org as supplemental material). The differences in the $Q_{10}$ values for the amplitudes and rates of augmentation at 0.75 and 2 mM $[Ca^2+]_o$ may reflect the $Ca^{2+}$ dependence of the processes underlying augmentation (Khanashvili et al., 1995; Blaustein and Lederer, 1999), which are also likely to be temperature dependent (see Discussion). This approach confirmed our findings of the strong temperature dependence of augmentation with a large increase in amplitude and a several-fold acceleration of decay.

Augmentation is also known to be selectively enhanced in the NMJ by $Ba^{2+}$ application (Zengel and Magleby, 1977; Kalkstein and Magleby, 2004), and this property of $Ba^{2+}$ can potentially be used as another approach to dissect STP components. We found, however, that $Ba^{2+}$ application had very different effects in hip-

**Table 1. Amplitudes and rates of STP components have different temperature dependence**

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<tr>
<th>STP process</th>
<th>$Q_{10}$ Amplitude</th>
<th>$Q_{10}$ Time course (acceleration)</th>
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<td>Facilitation</td>
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Figure 4. Facilitation increased in amplitude and decayed much faster at $33 - 38^\circ$C. A. Facilitation was measured at $23^\circ$C (□) and $33^\circ$C (●) with a paired-pulse protocol at 10 – 1000 ms intervals. Inset, Representative paired-pulse fEPSPs at 23 and $33^\circ$C. B, C. Extrapolated amplitude (B) and the time course of decay (C) of facilitation at 23, 33, and $38^\circ$C. Facilitation was more than two times larger and decayed approximately three times faster at body temperature. Parameters of facilitation did not vary significantly between 33 and $38^\circ$C. **$p < 0.01$; ***$p < 0.001$. 

**Q10 values for the amplitudes and rates of STP components at 0.75 and 2 mM $[Ca^2+]_o$:**

- **Facilitation:** $Q_{10} = 2.6$
- **Augmentation:** $Q_{10} = 5.1$
- **Depression:** $Q_{10} = 1.1$

Table 1. Amplitudes and rates of STP components have different temperature dependence

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pocampal synapses, where it produced strong and long-lasting depression during constant-frequency stimulation (2–40 Hz) (supplemental Fig. 8A, available at www.jneurosci.org as supplemental material) and during application of natural stimulus patterns (supplemental Fig. 8B, available at www.jneurosci.org as supplemental material). These effects were observed in a wide range of Ba²⁺ concentrations (0.6–2 mM). Therefore, the action of Ba²⁺ in the hippocampal synapses is very different from that observed in the NMJ and thus cannot be used to selectively increase augmentation in our preparation. Together, our analysis of different approaches to separation of STP components indicates that, for hippocampal synapses, the kinetics-based method is the most complete approach that allowed dissection of all STP processes.

**Much better maintenance of synaptic strength during prolonged use at near body temperatures**

Recent experiments in hippocampal cell cultures have shown that, at body temperature, endocytosis is sufficiently fast to compensate for exocytosis at stimulus rates of up to ~10 Hz so that synaptic strength is maintained during continued synapse use; however, at room temperature, RRP depletion occurs at frequencies above ~2 Hz (Fernandez-Alfonso and Ryan, 2004). To examine how a shift of balance among STP components affects the maintenance of synaptic strength at near body temperature, we recorded a set of responses to high-frequency trains (10 and 40 Hz) of increasing duration (25–600 stimuli) [Fig. 7 (only trains up to 300 stimuli are shown for clarity)]. As with the results reported for hippocampal cell cultures, synaptic strength was maintained much better at near body temperatures. At least 600 stimuli at 10 Hz (but only ~15 at 23°C) (Fig. 1) and ~300 stimuli at 40 Hz (but only ~4 at 23°C) could be transmitted at near body temperatures without depressing below control levels (Fig. 7). As shown in Figure 1, with the same frequency of stimulation, whole-cell currents exhibit somewhat less apparent depression and a better maintenance of synaptic transmission than do fEPSPs. Nevertheless, percentage of depressed EPSC responses during high-frequency trains declined ~12-fold at warmer temperatures (n = 18; data not shown).

Together, this set of experiments with constant-frequency stimulation has shown that STP components exhibit strong temperature dependence with a wide range of Q₁₀ factors and that the balance among the components achieved at near body temperatures results in a much better maintenance of synaptic strength during high-frequency trains.

**Implications for processing of natural spike trains**

Although the use of constant-frequency trains provides an informative tool for assessing the roles of different STP components in synaptic function, these spike trains are a poor representation of the input that hippocampal synapses would experience in vivo. Therefore, we studied synaptic responses to more physiologically relevant spike patterns recorded in vivo from the hippocampal place cells of behaving rodents (Fenton and Muller, 1998). These patterns, qualitatively like those exhibited by all hippocampal place cells, consist of high-frequency discharges that are also called bursts (Abbott and Regehr, 2004) (average duration of 12.6 ± 0.9 action potentials; range, 3–30 action potentials; and average action potential frequency of 28.6 ± 1.2 Hz; range, 4–62 Hz; data from 69 bursts), separated by periods of low activity (Fig. 8A, top). Although parameters of individual bursts vary widely among different cells within each hippocampal subfield as well as among ensembles of cells in CA1 and CA3 (Fenton and Muller, 1998; Leutgeb et al., 2004), the firing patterns used here provide a typical structure and a range of ISIs, burst durations, and other statistical properties of the spike patterns that hippocampal synapses may encounter naturally in the circuit. Below we will refer to these spike patterns as "natural" to indicate their origin.

To use natural spike trains recorded at 38°C for experiments performed at room temperature, a previous study (Dobrunz and Stevens, 1999) scaled all ISIs by an empirical factor of 3. The choice of a factor 3, intended to compensate for the differences in the characteristic times of various STP components, corresponded to Q₁₀ ~2, similar to that reported by other investigators. As described above, however, we found that the Q₁₀ values for different STP components varied significantly (from ~1 to 600 stimuli at 10 Hz (but only ~15 at 23°C) (Fig. 1) and ~300 stimuli at 40 Hz (but only ~4 at 23°C) could be transmitted at near body temperatures without depressing below control levels (Fig. 7). As shown in Figure 1, with the same frequency of stimulation, whole-cell currents exhibit somewhat less apparent depression and a better maintenance of synaptic transmission than do fEPSPs. Nevertheless, percentage of depressed EPSC responses during high-frequency trains declined ~12-fold at warmer temperatures (n = 18; data not shown).

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and so the choice of a single scaling factor is problematic. Therefore, we recorded responses at room temperature to unscaled natural input patterns (as is commonly done with the constant-frequency trains) as well as to patterns scaled by a factor of 3 to facilitate comparison with the previous study (Dobrunz and Stevens, 1999).

Figure 8A shows an example of fEPSP recordings from the same slice at 23 and 33°C and from a different slice at 38°C in response to the natural spike pattern shown above (unscaled at both temperatures). As reported previously (Dobrunz and Stevens, 1999; Dekay et al., 2006), synaptic responses at room temperature underwent large and rapid changes in amplitude that consisted of a variable mixture of facilitation and depression (minimum, 0.48 ± 0.08; maximum, 2.04 ± 0.15; n = 12) (Fig. 8A, top panel). The distribution of changes in synaptic strength during the train had no evident structure, and the ratio of depressed to potentiated responses varied widely from slice to slice (the relative amount of depressed responses varied from 9 to 70% among different slices; n = 12). This pattern of rapid alteration between potentiation and depression was independent of the ISI scaling because it was also observed during application of natural spike patterns scaled by a factor of 3 (n = 8; data not shown, but see Fig. 8C).

As expected from the strong temperature dependence of STP components, the response to natural stimulation underwent large changes with heating to 33–38°C (Fig. 8A, middle and bottom panels). Although the absolute amplitude of changes was similar to that at 23°C (2.52 ± 0.17 at 23°C vs 2.57 ± 0.12 at 33°C; p > 0.8; n = 12, 35), changes in synaptic strength were “shifted
correlation analysis to compare responses to natural stimulation at 23, 33, and 38°C (Fig. 8C). Pairs of averaged responses to the same natural spike train from different slices were plotted point-by-point versus each other, and the correlation coefficient (R) was determined with linear regression. The high similarity of responses corresponded to correlation coefficient values close to 1. As in experiments with constant-frequency stimulation (Fig. 1), synaptic responses were highly correlated in the temperature range of 33–38°C (Fig. 8C) (R_{33–38C} = 0.85 ± 0.02; 16 randomly selected pairs), but only a weak (and negative) correlation was found between responses at 23 and 33°C (R_{23–33C} = −0.55 ± 0.04; 22 randomly selected pairs). These data confirm our results obtained with constant-frequency stimulation that most of the temperature-dependent shift in synaptic response occurs in the 23–33°C range.

The pattern of synaptic response at 33°C is conserved across a wide range of calcium concentrations

The physiological extracellular calcium concentration ([Ca\(^{2+}\)]_o) in the brain is known to be ~1.2 mM (Jones and Keep, 1988; Brumberg et al., 2000) rather than 2 mM, as commonly used in laboratory experiments. Therefore, we attempted to mimic physiologically relevant conditions more closely by recording responses to natural spike trains in 1 mM [Ca\(^{2+}\)]_o (and 2 mM [Mg\(^{2+}\)]_o). Synaptic responses had the same pattern in 1 mM [Ca\(^{2+}\)]_o and 2 mM [Ca\(^{2+}\)]_o (Fig. 8B,D) (R = 0.88 ± 0.01; 30 randomly selected pairs), whereas the average increase during bursts was significantly larger in 1 mM [Ca\(^{2+}\)]_o, as compared to natural stimulation at 23, 33, and 38°C (Fig. 8C). This trend continued at even lower extracellular calcium (0.6–0.7 mM [Ca\(^{2+}\)]_o and 2.4 mM [Mg\(^{2+}\)]_o), with the preservation of the synaptic response structure (R = 0.81 ± 0.02; 24 randomly selected pairs) and an even larger average increase during bursts, reaching 7.2 ± 1.6 (p < 0.05; n = 4; data not shown). These data demonstrate that the robust pattern of synaptic response, one that becomes evident at near body temperature because of a precise balance of STP components, is highly conserved across different experimental conditions, including not only a range of temperatures (33–38°C) but also at least a threefold range of calcium concentrations (0.6–2 mM).

Responses to natural stimulation are significantly more reproducible at 33–38°C than at 23°C

To further test the robustness of excitatory synaptic responses to natural stimulation at different temperatures, we compared the reproducibility of whole-cell and fEPSP recordings within single cells or slices as well as across different cells, slices, and animals for the same natural spike train. To facilitate this comparison, we plotted the data from Figure 8A versus stimulus number rather than time, so that closely spaced points during the bursts could be compared more easily (Fig. 9A). As with the findings reported in

The shift in synaptic responses to natural spike patterns occurs mostly over the temperature range of 23–33°C

To estimate the range of temperatures at which synaptic responses underwent the transition described above, we used a...
previous studies (Dobrunz and Stevens, 1999; Dekay et al., 2006), responses to natural spike trains at room temperature were highly conserved within the same slice for both unscaled and scaled spike patterns \( [R = 0.82 \pm 0.02 \text{ (unscaled); } R = 0.85 \pm 0.03 \text{ (scaled)}; n = 12, 7; 3 \text{ different patterns tested}] \), but were significantly less reproducible among different slices \( [R = 0.63 \pm 0.06 \text{ (unscaled); } R = 0.65 \pm 0.05 \text{ (scaled)}; p < 0.01 \text{ for both}; n = 30, 21 \text{ pairs}] \) (Fig. 9D, left panels). Similar results were obtained with whole-cell recordings \( [R = 0.58 \pm 0.03 \text{ within cells}; n = 12; \text{ and } R = 0.38 \pm 0.05 \text{ across cells}; n = 16 \text{ random pairs}] \), and there was no strong correlation between EPSC and fEPSP responses (Fig. 9, compare A, B) \( [R = 0.13 \pm 0.07; n = 15 \text{ pairs}; p > 0.47 \text{ compared with noncorrelated responses}] \). Notice that there is a higher response correlation in fEPSP recordings than in EPSC recordings. This likely reflects the high intrinsic variability of whole-cell responses, which is significantly reduced in field potential recordings because of the averaging across many cells. The high variability of responses at room temperature is illustrated in Figure 9C (left panel): three whole-cell and three fEPSP recordings at 23°C, all from different animals, are superimposed in the same graph and bear little resemblance to each other. Therefore, as has been reported previously (Dobrunz and Stevens, 1999), at room temperature, synaptic responses to natural stimulation vary significantly among different cells, slices, and animals.

In contrast, at near body temperature, both whole-cell and fEPSP recordings were highly conserved within single cells \( [R = 0.75 \pm 0.02; n = 36; 5 \text{ patterns tested}] \) or single slices \( [R = 0.90 \pm 0.01; n = 35; 6 \text{ patterns tested}] \) as well as across different cells \( [R = 0.81 \pm 0.02; n = 36], \text{ slices, and animals (} R = 0.91 \pm 0.01; n = 35 \text{ slices from 14 animals; 6 patterns tested}] \) (Fig. 9A–D, right panels). The striking robustness of synaptic response at higher temperatures is emphasized in Figure 9C (right panel), where three whole-cell recordings and three fEPSP recordings, all from different animals, are superimposed on the same graph and are virtually the same (EPSC and fEPSP responses were highly correlated, with \( R = 0.81 \pm 0.01; 60 \text{ random pairs; five spike patterns} \)). Thus, the selective and predictable synaptic response observed at near physiological temperatures is highly conserved among different cells, slices, and animals, and this conservation of response properties argues for its physiological significance.

**Discussion**

In this work, we have isolated and analyzed the contributions of various STP components expressed in excitatory hippocampal synapses during constant frequency and natural stimulation at different temperatures. Our data have shown that, because of a strong but differential temperature dependence, the balance among STP components shifts from dominating depression at room temperature to facilitation- and augmentation-driven potentiation at body temperature. This shift results in a greatly improved maintenance of synaptic strength during prolonged stimulation and qualitatively changes the synaptic response to natural spike trains. The highly conserved filtering pattern observed during natural stimulation at higher temperatures suggests that the fine balance among STP components, achieved only at near physiological temperatures, is required for the robust and physiologically relevant STP function during the processing of realistic spike trains.

Elucidating the functional roles of STP components has been complicated by the presence of multiple overlapping processes during synaptic activity. Here we took advantage of the large difference in the decay kinetics of various STP components to separate their contributions to synaptic modulation. This approach is based on two considerations. (1) Different STP components can be studied during appropriate time windows, during which a particular process dominates changes in synaptic response (Fig. 2); and (2) contributions from the overlapping processes can be corrected for assuming a multiplicative relationship among the components. The validity of the first assumption is supported by our additional analysis, in which the entire data, rather than different time windows, are fitted with Equation 1.
yielding essentially the same results (see Materials and Methods). However, because the relationships among STP components are not completely understood and probably more complex than assumed in our analysis, it is important to determine the extent to which the second assumption and the associated correction for the overlapping STP components affect our conclusions. Examination of the data reveals that only in the case of depression are the estimates for the temperature-dependent effects significantly altered by the correction (Fig. 3). The observed accelerated recovery from depression with temperature is significant, however, in both cases ($p < 0.01$), and our estimate for the $Q_{10}$ value of this process ($2.1 \pm 0.5$) fits well with the previously published values ($Q_{10} \approx 2–3$) (Dinkelacker et al., 2000; Pyott and Rosenmund, 2002; Fernandez-Alfonso and Ryan, 2004; Micheva and Smith, 2005; Kushmerick et al., 2006). For all other STP components, the correction is either not required or the amplitudes of temperature-dependent effects are significantly larger than the changes introduced by the correction. Thus, the assumption of multiplicative relationships among STP components does not appreciably affect our conclusions.

The approach applied here to analyze the balance among STP components at different temperatures revealed a diminishing role of depression in the direct modulation of synaptic response at higher temperatures and an increasing dominance of facilitation and augmentation (Fig. 1). Our experiments do not allow us to elucidate the cellular biological mechanisms of these temperature-dependent changes; nevertheless, our current understanding of the mechanisms underlying STP processes suggests that the decreasing role of depression originates from accelerated vesicle retrieval and a much faster replenishment of the RRP at higher temperatures (Pyott and Rosenmund, 2002; Fernandez-Alfonso and Ryan, 2004). In the case of facilitation and augmentation, the strong temperature dependence of the amplitudes of these processes may arise, in part, from a larger Ca$^{2+}$ influx through Ca$^{2+}$ channels that has been shown to increase with temperature several-fold (Taylor, 1988; McAllister-Williams and Kelly, 1995). Hippocampal neurons also express high levels of the temperature-gated vanilloid receptors (Toth et al., 2005) that are known to affect Ca$^{2+}$ homeostasis and may have an effect on the amplitudes of facilitation and augmentation by altering basal Ca$^{2+}$ levels (Guatteo et al., 2005). In addition to a large increase in amplitude with temperature, both components of enhancement exhibited a several-fold acceleration of decay (Figs. 4, 5). These temperature-dependent changes may result from the acceleration of calcium-cleaning mechanisms, such as the activity of Na$^{+}$/Ca$^{2+}$ exchanger, which increases with temperature approximately threefold ($Q_{10}$) (Powell et al., 1993; Khananshvili et al., 1995). The Ca$^{2+}$ dependence of the Na$^{+}$/Ca$^{2+}$ exchange rate also varies with temperature (Khananshvili et al., 1995; Blaustein and Lederer, 1999) and may contribute to the differences in the $Q_{10}$ values of the two components because they are typically observed at different intracellular Ca$^{2+}$ levels (Delaney and Tank, 1994).

The temperature-dependent shift among STP components occurred in parallel with a greatly improved maintenance of synaptic strength during continuous synaptic use. At least 10 times as many spikes could be transmitted at higher temperatures without depressing below control levels (Figs. 1, 7). Considering a weak temperature dependence of the basal release probability in hippocampal synapses (Pyott and Rosenmund, 2002), our data suggest that better maintenance of synaptic strength is likely a consequence of the fine balance among STP components achieved at higher temperatures. These results are in agreement with the recent findings in hippocampal cell cultures (Pyott and Rosenmund, 2002; Fernandez-Alfonso and Ryan, 2004), and may be a widespread synaptic property, because a similar effect of temperature has been also found in such structurally and physiologically different types of synapses, as the calyx of Held (Taschenberger and von Gersdorff, 2000) and end-bulb synapses in the avian nucleus magnocellularis (Brenowitz et al., 1998).

The most striking effect of temperature observed here is the large change in the processing of natural spike trains with heating from 23 to 33–38°C. A combination of several effects can account for these temperature-dependent changes. First, cooling to 23°C has been shown to affect several important synaptic properties, including spike generation (Stevens and Zador, 1998; Volgushev et al., 2000), RRP size, and kinetics of exocytosis and endocytosis (Pyott and Rosenmund, 2002; Fernandez-Alfonso and Ryan, 2004; Micheva and Smith, 2005; Kushmerick et al., 2006), as well as the progressive degradation of dendritic spine structure and motility with cooling time (Roelands and Matsus, 2004). The variability in potentiation and depression, observed during natural stimulation at 23°C, may reflect, in part, the different extent to which these structural and physiological changes have taken place, because the duration of cooling varies in preparations from the same as well as different animals. Indeed, we found consistent (although not statistically significant) changes in the amplitudes of all three STP components with cooling time (~45, 15, and 7% increase in the amplitudes of facilitation, augmentation, and depression, respectively). These changes had an impact on the overall structure of synaptic responses to natural stimulation at room temperature as they shifted the relative number of facilitated and depressed responses during the trains (from 49% of depressed responses to 34%). However, prolonged cooling had little effect on natural spike train responses at 33–38°C, because depression was always masked during natural stimulation at these temperatures, and the overall structure of synaptic responses remained largely unaffected. In addition to the effect of different cooling times, we also found an overall larger variability in the parameters of STP components at 23°C than at 33°C. This included both the amplitude and time constant of facilitation (~60 and ~50% increase in coefficient of variation, respectively) and the amplitude of augmentation (~25% increase). Thus, the reduced variability in the parameters of STP components also contributes to the higher reproducibility of natural spike train responses at warmer temperatures. Finally, another source of variability of STP at room temperature is suggested by the finding that the ratio of facilitation and depression observed during natural spike trains at 23°C changed dramatically with the animals’ age (postnatal days 12–35) (Dekay et al., 2006). As a result, the responses to natural stimulation were poorly correlated among animals of different ages. Here we found that this was not the case at 33–38°C, where no depression was observed at any time during natural spike trains independently of the animals’ age, in a comparable range of postnatal days 14–35. Moreover, the animals’ age had no significant effect on response correlation at higher temperatures: the correlation was similarly strong for animals of the same and different ages ($p > 0.6$). Thus the improved reproducibility of responses at higher temperatures may also reflect the developmental changes in expression of facilitation and depression at room temperature that introduce much less variability at higher temperatures, where apparent depression during natural spike trains is insignificant.
An important consequence of this temperature-dependent shift is the appearance of a physiologically relevant pattern of synaptic response to natural stimulation, obscured at room temperature by dominating depression. Unlike responses at 23°C, this pattern can be predicted from the structure of the input spike train alone: at higher temperatures, STP acts to amplify synaptic transmission selectively during spike bursts (Fig. 8). The physiological significance of this burst-amplification pattern is suggested by the finding that it is highly conserved across different cells, slices, and animals and under various physiological conditions. Its possible functional role is further emphasized by the fact that high-frequency spike bursts signal hippocampal place fields that carry information about the animal’s position in the environment (O’Keefe and Dostrovsky, 1971). Therefore, at higher temperatures, STP in excitatory hippocampal synapses may act to amplify the specific information associated with the place field of the input neuron.

In summary, we found that components of STP undergo a temperature-dependent shift of balance that results in a much better maintenance of synaptic strength at higher temperatures and reveals a highly conserved and predictable pattern of synaptic response to natural stimulation.

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