Optimal Information Transfer in the Cortex through Synchronization

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Abstract

In recent experimental work it has been shown that neuronal interactions are modulated by neuronal synchronization and that this modulation depends on phase shifts in neuronal oscillations. This result suggests that connections in a network can be shaped through synchronization. Here, we test and expand this hypothesis using a model network. We use transfer entropy, an information theoretical measure, to quantify the exchanged information. We show that transferred information depends on the phase relation of the signal, that the amount of exchanged information increases as a function of oscillations in the signal and that the speed of the information transfer increases as a function of synchronization. This implies that synchronization makes information transport more efficient. In summary, our results reinforce the hypothesis that synchronization modulates neuronal interactions and provide further evidence that gamma band synchronization has behavioral relevance.

Introduction

Gamma band synchronization has been found in many cortical areas and in a variety of tasks. It has been studied most extensively in the visual cortex of cats and monkeys [1–8]. Several authors have proposed that these synchronizations influence the interactions among neuronal groups [9,10], a hypothesis referred to as communication through coherence (CTC, [11]). In computational studies, it has been shown that entrainment enhances transmitted information between input and output spikes [12], that synchronization in the gamma frequency range increases the effective synaptic gain for the responses to an attended stimulus [13] and that the transmission time of responses of coupled oscillators depends on the phase difference in the stable synchronized state [14]. Also, several experimental studies have presented results supporting the CTC hypothesis [15–19].

In this study, we concentrate on the results shown by Womelsdorf et al. [17]. They explore the mutual influence of two groups of neurons as a function of their phase shift. These authors quantify the mutual influence of the multi unit activity (MUA) in the two groups as the Spearman rank correlation coefficient of the two MUA’s 60 Hz power. They show evidence that the correlation between the two groups of neurons varies as a function of the phase shift of the oscillations at 60 Hz. There is a specific phase shift at which the correlation between the two groups is highest. [17] conclude that the effective connectivity in a network can thus be maximized or minimized through synchronization of a specific phase relation, resulting in an effective interaction pattern.

While the results presented by [17] clearly support the CTC hypothesis, they leave some open questions. Is it only the 60 Hz power that depends on the 60 Hz phase? Do the MUA’s only correlate or is there mutual interaction between the two? Is this effect restricted to the gamma band or can it be generalized to other frequency bands? What is the influence of the total gamma power in the signal?

Here, to address these questions, we use a detailed biophysical model network with realistic spiking properties. A first advantage of using a model is that we can generate more data than in an experiment. This makes it possible to use an information theoretical measure for the mutual interaction instead of rank correlation. Many different interdependence measures such as mutual information, transfer information, nonlinear regression, phase synchronization and generalized synchronization have recently been proposed (see [20] and [21] for comparisons of the different methods). It has become evident that the appropriateness of each measure is determined by the data it is applied to. Thus, given our current data set, we opted to use transfer entropy (TE), introduced by [22]. The TE is an information theoretical measure that quantifies the statistical coherence between systems. It has the advantage that it does not only measure the coherence between two signals, but is able to distinguish between driving and responding elements and therefore between shared and transported information. This is called the directionality of the information flow. We measure the TE between the MUA of the two neuronal clusters, which allows us to study the interdependence of the spiking activity in each of them and not just the correlation of the spectral power in a specific frequency band, as was done in the experimental work. A further crucial advantage of the model is that we can change network parameters systematically and explore the dynamical range of the network.

The model we use in this study consists of integrate-and-fire neurons. One of two pools of excitatory neurons receives input
Author Summary

Different brain areas are involved in any cognitive task. This implies that information has to be transmitted between different brain areas. Recent experimental results suggest that synchronization plays a crucial role in information exchange between cortical areas. They show that synchronization is capable of rendering network connections effective or ineffective. We study this hypothesis using a neurodynamical model and present results suggesting that both phase and strength of neuronal oscillations in the gamma frequency band influence amount and speed of information transport. We conclude that neuronal synchronization is crucial for information transmission and therefore might even have behavioral relevance.

Methods

Experimental analysis

Womelsdorf et al. [17] analyzed four different data sets. The first data set consisted of measures from awake cats in area 17 [26], the second from awake cats in areas 18 and 21a [4], the third from awake monkeys in area V1 and the fourth from awake monkeys in area V4 [3,7]. In all four data sets they recorded multi unit activity (Poisson spike train) which it passes to a neighboring pool, connected by feedforward and feedback connections. Each pool of excitatory neurons is connected to a pool of inhibitory neurons, which generates oscillations in the gamma frequency band through a pyramidal-interneuron feedback loop [23]. Beta oscillations are obtained from the same network by parameter modification. Several methods have been proposed to generate beta oscillations [24,25]. Here, for the sake of simplicity, we modify the decay constants of the synapses. We show that the correlation as measured by the Spearman rank correlation coefficient depends on the phase relation in the gamma band. This result confirms the experimental finding of [17]. Secondly, after applying TE to measure the information exchange between two pools, we find that TE very similarly depends on the phase shift, i.e., that there is an optimal phase relation where the TE is maximal. Thirdly, we reveal such dependence also in the beta band. Fourthly, we demonstrate that the TE increases as a function of the power in the gamma band. Lastly, we show that the information exchange gets faster if the gamma band synchronization increases. In sum, we provide support for the CTC hypothesis and make the prediction that CTC is a general mechanism, not restricted to the gamma band.

Theoretical framework

We use a model with leaky integrate-and-fire (IF) dynamics, following [27]. Each IF unit charges up to its stationary value as long as its membrane potential stays below a threshold. The membrane potential $V(t)$ is given by:

$$I_{syn}(t) = I_{AMPA,ext} + I_{AMPA,rec} + I_{NMDA} + I_{GABA}.$$  (2)

The currents are defined as follows:

$$I_{AMPA,ext}(t) = g_{AMPA,ext}(V(t) - V_E) \sum_{j=1}^{N_{ext}} w_{j}^{AMPA,ext}(t)$$  (3)

$$I_{AMPA,rec}(t) = g_{AMPA,rec}(V(t) - V_E) \sum_{j=1}^{N_E} w_{j}^{AMPA,rec}(t)$$  (4)

$$I_{NMDA}(t) = \frac{g_{NMDA}(V(t) - V_F)}{1 + [Mg^{2+}] \exp(-0.062V(t))/3.57} \times \sum_{j=1}^{N_E} w_{j}^{NMDA}(t)$$  (5)

$$I_{GABA}(t) = g_{GABA}(V(t) - V_I) \sum_{j=1}^{N_I} w_{j}^{GABA}(t)$$  (6)

Figure 1. Spearman rank correlation coefficient. The rank correlation coefficient between the two MUA’s 60 Hz power is plotted as a function of their phase relation. The solid line indicates a cosine fit. Adapted from [17].

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The fractions of open channels are given by:

\[
\frac{ds_j^{AMPA,ext}(t)}{dt} = -\frac{s_j^{AMPA,ext}(t)}{\tau_{AMPA}} + \sum_k \delta(t - \tau_j^d)
\]

\[
\frac{ds_j^{AMPA,rec}(t)}{dt} = -\frac{s_j^{AMPA,rec}(t)}{\tau_{AMPA}} + \sum_k \delta(t - \tau_j^d)
\]

\[
\frac{ds_j^{NMDA}(t)}{dt} = -\frac{s_j^{NMDA}(t)}{\tau_{NMDA,decay}} + 2x_j(t)(1 - s_j^{NMDA}(t))
\]

\[
\frac{dx_j(t)}{dt} = -\frac{x_j(t)}{\tau_{NMDA,rise}} + \sum_k \delta(t - \tau_j^d)
\]

\[
\frac{ds_j^{GABA}(t)}{dt} = -\frac{s_j^{GABA}(t)}{\tau_{GABA}} + \sum_k \delta(t - \tau_j^d)
\]

\(\tau_{AMPA}\), \(\tau_{NMDA,decay}\) and \(\tau_{GABA}\) are the decay times and \(\tau_{NMDA,rise}\) is the rise time for the corresponding synapses. AMPA has a very short decay time (2 ms) while NMDA has a long one (100 ms) and the GABA decay time lies in-between (10 ms). The rise times of AMPA and GABA currents are neglected, as they are typically very short (<1 ms). The sums over \(k\) represent a sum over spikes formulated as \(\delta\)-peaks \(\delta(t)\) emitted by presynaptic neuron \(k\) at time \(\tau_j^d\). All input is generated via a Poisson process.

The equations are integrated using a fourth order Runge-Kutta method with a time step of 0.02 ms. The network is organized in pools. Neurons within a specific pool have stronger recurrent connections than neurons between the pools. The intention of this work is to study cortical neural interactions not limited to specific brain area. However, as our simulations needed to be directly comparable to [17], and have specific parameter sets, our network models two clusters of cortical neurons in visual cortex V4.

The network model consists of two parts (Fig. 2). In each part there are pools of excitatory and inhibitory neurons, with a total of 800 excitatory and 200 inhibitory neurons. The excitatory neurons are subdivided into a selective pool and a non-selective pool. The neurons in the selective pools (S, S') are the ones that receive input either from outside or from the connected selective pool. The non-selective neurons (NS, NS') simulate the surrounding brain areas. Each population of excitatory neurons is connected to a pool of inhibitory neurons (I, I'). This allows for generating oscillations in each population separately. The two parts of the network are connected via feedforward \(J_f\) and feedback \(J_k\) connections that project onto the selective pools. The external input \(v_{ext}\) is a Poisson spike train that projects to the selective pool \(S\) of the first part of the network. In addition to the recurrent connections, the network is exposed to an external current \(v_{ext}\), modeled as a Poisson spike train of 800 neurons, firing at 3 Hz. This models the spontaneous activity observed in the cerebral cortex. The network is fully connected.

Gamma oscillations in a network with excitatory and inhibitory neurons are generated through a pyramidal-interneuronal feedback loop [23,28]. Pyramidal neurons excite interneurons and inter-neurons in turn send inhibition back on pyramidal cells. The population frequency is determined by the sum of excitatory and inhibitory lags. The recurrent excitatory connections tend to decrease the oscillation frequency (as compared to only excitatory-inhibitory and excitatory-inhibitory-excitatory connections) as they tend to prolong the positive phase in each cycle. In our network we can therefore generate and control the oscillations in the gamma frequency band by adjusting the AMPA and NMDA conductances. For example, increasing the \(g_{AMPA}\) and decreasing \(g_{NMDA}\) shifts the balance in the network towards excitation (AMPA) and slow inhibition (GABA) and thus increases the gamma frequency band oscillations. The conductances in our network are varied according to the following rule: \(g_{NMDA} = g_{NMDA}(1 - \delta)\) and \(g_{AMPA} = g_{AMPA}(1 + 10\delta)\). Throughout the paper, we will refer to the parameter \(\delta\) as the \(g_{AMPA}/g_{NMDA}\) modification ratio.

The factor 10 stems from the fact that near the firing threshold, the ratio of NMDA:AMPA components becomes 10 in terms of charge entry, as stated in [27]. Therefore, in order not to change the spontaneous state, a decrease in \(g_{NMDA}\) is compensated by a tenfold increase in \(g_{AMPA}\). All recurrent conductances (both inhibitory and excitatory) are changed according to these rules.

By adjusting the synaptic decay constants, the oscillation frequency can be shifted into the beta band. The crucial parameter is \(\tau_{GABA}\). An increase of \(\tau_{GABA}\) slows down the rhythm of the pyramidal-interneuronal loop and will therefore yield an oscillation at a lower frequency. To generate oscillations in the beta range (around 20 Hz), we use \(\tau_{AMPA} = 1.5\) ms and \(\tau_{GABA} = 38\) ms. To generate phase shifts in the gamma oscillations between the two parts of the network, we introduce a delay. The
delay is set bidirectionally in the feedforward and feedback connections of the selective pools. Each spike emitted in $S$ arrives at $S'$ after $\Delta t$ and vice versa. This lag in spike transmission generates a phase lag in the oscillations. A delay of, e.g., 4 ms yields a phase shift of about 90 in a 60 Hz oscillation. The actual value of the mean phase shift is not crucial to the obtained results. All trials are initiated with a period of 400 ms in which no stimulus is present, followed by a period of 5500 ms composed of the presentation of the stimulus, followed by 100 ms in which no stimulus is present. Each simulation consists of 100 trials. All parameter values are listed in Table 1.

**Analysis**

**Multi unit activity.** From our spiking simulations we calculate the multi unit activity (MUA) to analyze our simulations, in order to be able to compare our results directly with the experiments. To simulate the MUA, we randomly chose 10 neurons in each of the selective pools. This point process data is converted to a time series by binning the spikes in windows of 5 ms. The binning window is shifted in steps of 1 ms. The time series is then normalized to zero mean and unit variance. We use the multitaper method [29,30] to calculate the spectral power of our normalized time series to estimate power spectrum and transfer entropy. Normalization is applied to rule out the possible influence of rate changes.

**Power spectrum and phase estimation.** We use the multitaper method [29,30] to calculate the spectral power of our data. The signal in each time window (1000 ms) is multiplied with a set of Slepian data tapers. The tapered signal is then Fourier transformed, according to:

$$\tilde{x}_k(v) = \sum_{t=1}^{N} w_k(t)x(t)e^{-2\pi itv}$$  \hspace{1cm} (12)

where $w_k(t)$ are $K$ orthogonal taper functions, $x(t)$ is the time series of our signal, and $N$ is the number of elements in each time window. The power spectrum is then the squared amplitude of $\tilde{x}_k(v)$, averaged over the $K$ tapers. We used $K = 4$ tapers in our study. The cross spectrum $(S_{xy})$ between two signals $x_k(v)$ and $y_k(v)$ (averaged over $K$ tapers) is given by:

$$S_{xy} = \frac{1}{K} \sum_{k=1}^{K} \tilde{x}_k(v)\tilde{y}_k(v).$$  \hspace{1cm} (13)

The phase relation between two signals $x_k(v)$ and $y_k(v)$ is defined as the angle of the cross spectrum. We use this method for phase estimation to be able to compare directly to the experimental results.

**Transfer entropy.** In Ref. [17] the mutual influence between two neuronal groups is quantified as the Spearman rank correlation coefficient of spectral power. The Spearman rank correlation is a non-parametric measure of correlation, which makes no assumptions about normality or linearity of the data. However, it is a symmetric measure and therefore fails to measure directionality of the flow of information. Thus, to overcome this limitation, here we use TE [22], which enables us to distinguish between shared and transported information. TE measures the deviation from the following generalized Markov property:

$$p(x_{t+1}|x_t^k) = p(x_{t+1}|x_t^k, y_t^l),$$  \hspace{1cm} (14)

where $p$ is the transition probability and $k$ and $l$ are the dimensions of the delay vectors. $x_t$ and $y_t$ are the time series of the signal. We write $x_t$ and $y_t$ instead of $x(t)$ and $y(t)$, respectively, for better readability. If the deviation is small, then $Y$ has no relevance for the transmission probability of $X$. The incorrectness of this assumption can be quantified by the Kullback entropy

$$T_{y\rightarrow x} = \sum_{t} p(x_{t+1}^k, y_t^l) \log_2 \frac{p(x_{t+1}^k|x_t^k, y_t^l)}{p(x_{t+1}^k|x_t^k)}.$$

In other words, transfer entropy represents the information about a future observation of variable $x_t$ obtained from the simultaneous observation of past values of both $x_t$ and $y_t$, after discarding the information about the future of $x_t$ obtained from the past of $x_t$ alone [20]. For computational reasons, we set $k = l = 1$. Conditional probabilities required in equation 15 are calculated from the joint probabilities. We approximate the joint probabilities by coarse-graining the continuous state space at resolution $r$ and using the histograms of the embedding vector (naive histogram technique [31]). When the available data is limited (number of samples $N < 1000$) and the coupling between the time series is small, TE suffers from a finite sample effect, in particular for small resolution ($r < 0.05$), which makes the assessment of the significance of the obtained values difficult [31]. However, for all our simulations $N > 1000$ and $r > 0.05$, so we can assume that the finite sample issue affects our results to a negligible extent. We calculate the TE between the MUA in the two neuronal pools.

**Results**

First we describe how the mean phase shift between pools of neurons is set by the delay in the feedforward and feedback connections. We then show that the correlation between the gamma power in the two pools depends on the phase relation in the gamma band. We demonstrate that TE has a very similar

| Parameter | Value | Parameter | Value |
|-----------|-------|-----------|-------|
| $C_m$ (excitatory) | 0.5 nS | $V_m$ | −70 mV |
| $C_m$ (inhibitory) | 0.2 nS | $V_{out}$ | −55 mV |
| $\Delta$PA (excitatory) | 2.08 nS | $\Delta$NE | −50 mV |
| $\Delta$PA (inhibitory) | 1.62 nS | $w_+$ | 1.5 |
| $\Delta$PA (excitatory) | 0.104 nS | $w_-$ | 1.0 |
| $\Delta$PA (inhibitory) | 0.081 nS | $\tau$ | 0.5 ms$^{-1}$ |
| $\gamma$ABA (excitatory) | 1.287 nS | $\gamma$ABA | 250 Hz |
| $\gamma$ABA (inhibitory) | 1.002 nS | $\gamma$ABA | 2.4 kHz |
| $\gamma$NMDA (excitatory) | 0.327 nS | $\gamma$NMDA | 2 ms |
| $\gamma$NMDA (inhibitory) | 0.258 nS | $\gamma$NMDA (beta osc.) | 1.5 ms |
| $\tau$ (excitatory) | 25 nS | $\tau$ABA | 10 ms |
| $\tau$ (inhibitory) | 20 nS | $\tau$ABA (beta osc.) | 38 ms |
| $\tau$ | 1.8 | $\tau$MDA (delay) | 100 ms |
| $\delta$ | 0.6 | $\tau$MDA (τm) | 2 ms |
| $\tau_{ref}$ (excitatory) | 800 | $\tau_{ref}$ | 2 ms |
| $\tau_{ref}$ (inhibitory) | 800 | $\tau_{ref}$ | 1 ms |
| $\tau_{delay}$ | 200 | $\Delta t$ | 4 ms |
| $\tau_{delay}$ | 0 mV | feedback/feedforward ratio | 1/3 |
| $\tau_{delay}$ | −70 mV | }
dependence on the phase shift and that TE increases as a function of gamma power. Finally, we reveal that if gamma power is high, information flow as measured by TE commences earlier.

Delay–phase relation

Raster plots for 20 neurons from each neuronal pool are shown in (Fig. 3). The power spectrum of the MUA in our network shows a clear peak in the gamma band (Fig. 4), in accordance with the experimental results. Therefore, the introduced delay sets the phase shift for oscillations in the gamma band. The delay, however, sets only the mean phase shift, but the phase shifts fluctuate over time. Thus, even for a fixed delay they show a broad distribution around this mean phase shift. This distribution is shown in Fig. 5a. The mean phase in this specific simulation is $91.4^\circ$. This, however, is just an example, as the mean phase shift in the simulations can be set to any value by adjusting the delays accordingly. The phases are similarly widely distributed as in the experimental results by [17], shown in Fig. 5b.

Dependence of correlations on phase

The phase shifts at 60 Hz between the two pools show a broad range of phases. We determine the phase shift in each time window of 500 ms. Then we calculate the correlation between the two pools for this time window by calculating the Spearman rank coefficient for the 60 Hz power in the two pools. The obtained correlation can now be sorted into different bins for the different phase shifts. We find that the correlation of the gamma band power between the two pools depends on the mean phase shift in the gamma band. Fig. 6 shows the rank correlation plotted against the phase shifts. The correlation is highest for the bin containing the mean phase shift and drops as it moves away from the mean. This confirms the experimental results of [17].

Dependence of TE on phase

We apply TE to the same data as in the previous section. However, we measure the TE between the MUA in the two pools...
and not only the spectral power at 60 Hz, as was done in the experiment. We find that the TE depends strongly on the phase relation in the gamma band between the spiking activities of the two groups of neurons. It is highest for the mean phase between the two signals and drops as it moves away from the mean. This is consistent with our results for correlation. The phase dependence is illustrated in Fig. 7. TE is plotted as a function of the mean phase shift. The solid line represents TE from the first to the second pool (forward) and the dashed line TE from the second to the first one (backward). Forward TE is stronger than backward TE, implying that TE correctly detects the causal dependence of the second neuronal pool on the first one. Forward TE is stronger than backward TE even if the feedforward and feedback connections are symmetrical (not shown). The stronger the feedforward and the weaker the feedback connections, the bigger the difference in the TE for the two directions, as shown in Fig. 8. We plot the relative difference in the TE, calculated as \( \frac{\text{TE}_{\text{f}} - \text{TE}_{\text{b}}}{\text{TE}_{\text{b}}} \). The feedback/feedforward ratio is defined as \( J_{\text{f}}/J_{\text{b}} \). We use a feedback/feedforward ratio of 1/3 in the baseline simulations. To make sure the phase dependence is not only a by-product of changes in spectral power, we sorted the trials according to their spectral power in the gamma frequency band and calculated the phase dependence both for trials with power below and above the median. In both cases, the phase dependence is very similar (not shown).

**Different frequency bands**

Another result we obtain is that the phase dependence of information transport is not restricted to the gamma band. We find that even in simulations with a network oscillating strongly in

![Figure 5. Phase distribution.](https://doi.org/10.1371/journal.pcbi.1000934.g005)

![Figure 6. Spearman rank correlation coefficient.](https://doi.org/10.1371/journal.pcbi.1000934.g006)

![Figure 7. TE as a function of phase shifts and directionality.](https://doi.org/10.1371/journal.pcbi.1000934.g007)
the TE for a network oscillating in the gamma band. The trials are sorted according to their phase relation in the gamma band. Fig. 9b shows the same network but with the trials now sorted according to their phase relation in the beta band. The phase dependence curve becomes a lot flatter and the optimal phase for maximal TE is much less pronounced. Fig. 9c shows the TE for a network oscillating in the beta band with trials sorted according to the phase relation in the gamma band. And Fig. 9d shows the TE for a network oscillating in the beta band with trials sorted according to the phase relation in the beta band. It becomes clear that it is the phase of the dominating frequency band that is responsible for high or low TE. We therefore conclude that it is not only the gamma band that has the ability to shape effective network connections via the phase, but that it is a general mechanism, observable in different frequency bands.

Dependence of TE on gamma band power (without parameter modification)

We further find that TE depends on the spectral power in the gamma band (30–85 Hz). For a fixed parameter set, we first sort all the trials according to their power in the gamma band into bins. In each of these bins, we measure the TE for the mean phase relation. The TE as a function of the power in the gamma band is plotted in Fig. 10. We find that the TE increases as a function of power. Note, however, that instead of sorting the trials according to their gamma band power for a fixed parameter set, we can also vary the parameters in the network. This allows us to vary the power over a wider range and the effect becomes clearer (see the next section).

Figure 9. TE as a function of the mean phase shift in the gamma and beta band. The network oscillates strongly in one frequency band (either beta or gamma). The trials are sorted according to their phase shift either in the beta or gamma band. (a) Network oscillating in the gamma band. The trials are sorted according to their phase shift in the gamma band. (b) Network oscillating in the gamma band. The trials are sorted according to their phase shift in the beta band. (c) Network oscillating in the beta band. The trials are sorted according to their phase shift in the gamma band. (d) Network oscillating in the beta band. The trials are sorted according to their phase shift in the beta band. In all four graphics, $\frac{g_{\text{AMPA}}}{g_{\text{NMDA}}}$ modification ratio $\approx 0.12$, averaged over 100 trials.

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Dependence of TE on gamma band power (with parameter modification)

In the previous section we have shown how TE depends on power in the gamma band for a fixed parameter set. Now we explicitly vary the amount of gamma power and study the TE dependence. Gamma band oscillations in a network of excitatory and inhibitory integrate-and-fire neurons appear when excitation is faster than inhibition [28]. Thus, we made the network oscillate by increasing AMPA conductance and decrease NMDA conductance. This change was applied to both excitatory and inhibitory neurons. In our simulations, we vary the $g_{\text{AMPA}}/g_{\text{NMDA}}$ modification ratio from 0 to 0.12, which results in a gamma band that contains from 10 to 65% of power. If we sort the data according to its shift as in the previous section, we find that, for the different $g_{\text{AMPA}}/g_{\text{NMDA}}$ modification ratios, the TE shows a similar dependence on the phases. However, if the $g_{\text{AMPA}}/g_{\text{NMDA}}$ modification ratio is too low, the phase measurement is not reliable any more and the curve gets flat, consistent with the case of random phase distribution. Fig. 11 shows the TE as a function of phase shifts and gamma oscillations. If the oscillations are strong in the gamma band ($g_{\text{AMPA}}/g_{\text{NMDA}}$ modification ratio = 0.12), there is a clear phase shift between the two groups of neurons and the phase dependence curve is clearly bell shaped. If the oscillations are too low, there is no meaningful phase shift and the curve becomes flat ($g_{\text{AMPA}}/g_{\text{NMDA}}$ modification ratio = 0.02). Averaged over 100 trials. doi:10.1371/journal.pcbi.1000934.g011

Timing

Finally, we are interested in whether the gamma band oscillations also have an influence on the speed of information exchange, on top of the increased amount of information exchange. To do this, we measure the time required until the stimulus presentation to the first pool leads to an increase in TE towards the second pool. We find that the onset of TE increase is significantly earlier when there is a lot of power in the gamma band. While for a $g_{\text{AMPA}}/g_{\text{NMDA}}$ modification ratio of 0.02 it requires 28 ms to reach 50% of the average TE, for a $g_{\text{AMPA}}/g_{\text{NMDA}}$ modification ratio of 0.12 it takes only 17 ms. The onset of information flow is clearly faster for higher levels of gamma band power (Fig. 13). This increase in speed is a further demonstration of how gamma oscillations increase network performance and shows how a network can be made more competitive.

Figure 10. TE of trials sorted by power in the gamma band. Network parameters are kept fixed. The TE increases as a function of gamma power. $g_{\text{AMPA}}/g_{\text{NMDA}}$ modification ratio = 0.12, averaged over 100 trials. Error bars indicate 95% confidence intervals. doi:10.1371/journal.pcbi.1000934.g010

Figure 11. TE as a function of phase shifts and gamma oscillations. If the oscillations are strong in the gamma band ($g_{\text{AMPA}}/g_{\text{NMDA}}$ modification ratio = 0.12), there is a clear phase shift between the two groups of neurons and the phase dependence curve is clearly bell shaped. If the oscillations are too low, there is no meaningful phase shift and the curve becomes flat ($g_{\text{AMPA}}/g_{\text{NMDA}}$ modification ratio = 0.02). Averaged over 100 trials. doi:10.1371/journal.pcbi.1000934.g011

Figure 12. Mean TE as a function of gamma frequency band power. We plot the TE for six different $g_{\text{AMPA}}/g_{\text{NMDA}}$ modification ratios (solid line). A higher $g_{\text{AMPA}}/g_{\text{NMDA}}$ modification ratio causes the network to oscillate in the gamma frequency range and thus increases the power in the gamma frequency band (dashed line). Error bars indicate 95% confidence intervals; averaged over 100 trials. doi:10.1371/journal.pcbi.1000934.g012
Communication through coherence

It has been hypothesized that interactions among neuronal groups depend on neuronal synchronization. Recent results show that gamma band oscillations and especially the phase relation in the gamma band can modify the strength of correlations in a network and therefore influence the effectiveness of connections in it [17]. These effects could be used as a mechanism to connect and disconnect areas in a network without altering the physical connections. Here, using a model network of IF neurons, we intend to test this hypothesis. We demonstrate that also in a model network, the correlation between two areas depends on the phase shift in the gamma band between these two areas. Our modeling approach enables us to use an information theoretical measure, as it allows us to generate as much data as needed for such a measure. Thus, we use transfer entropy, which has the advantage of being able to distinguish driving and responding elements in a network. We show that also for TE there is an optimal phase shift between two neuronal groups, where TE is highest. We study this phase dependence in different bands (beta and gamma). Our results demonstrate that, in a network with strong beta oscillations, TE depends on the phase shift in the beta band similarly to the way TE depends on the phase shift in the gamma band in a network with strong gamma oscillations. The ability to shape network connections seems therefore not to depend on the frequency range and seems to be a general mechanism. This confirms recent experimental results that have pointed out the importance of beta synchrony in functional integration [32,33]. We also study how TE depends on power in a specific frequency band. We do this here for the gamma band. For a fix set of parameters, we sort the trials in a simulation according to their power in the gamma band. We find that within a simulation, the trials with high gamma power have a high TE. Then we modify the parameters and vary the gamma power over a wider range. Again, we find that TE increases as a function of gamma power. Finally, we reveal that it is not only the amount of exchanged information that increases but also the speed: The higher the power in the gamma band, the earlier the onset of the information flow.

Our results support the CTC hypothesis. If the effective connections in a network are to be influenced by the phase lock in a specific frequency band between two areas, it is important that it not only affects the coherence between them, but also the throughput of information in a specific direction. By measuring TE instead of the Spearman rank coefficient, we extend the work of [17]. Our result is also more general, as we use the rates to measure TE and not only the 60 Hz power. Our study of different frequency bands is a further extension. We provide evidence that the CTC mechanism is not restricted to the gamma band, but also functions in different frequency bands. In addition, our modeling approach also enables us to study how the information transport depends on the total power in a specific frequency band. Our finding that TE increases as a function of power suggests that both the phase and the power in a specific frequency band are important to shape effective connections in a network. The phase dependence of information transmission is not only a byproduct of the power dependence, as we find the same phase dependence both in trials with high gamma power as well as in trials with low gamma power. Furthermore, we demonstrate that the onset of information exchange depends on the power, which contributes to effectively shaping the connections in a network. We have already shown in the context of attention that gamma power increases the network effect of an attentional bias and that it makes the network more efficient [34]. Here, we can confirm this finding and put it in a more general context, independent of attention.

Implications for visual information transmission

As we are modeling results from visual cortical areas, we can assume that the neuronal clusters in the model transmit largely visual information. Several recent studies have contributed to the understanding of visual information transmission. These studies suggest that LFP power gradually increases as a function of stimulus contrast and gamma band LFP power increases differentially, that is, to a higher extent with respect to the baseline than relative to either higher or lower bands [35]. For the highest stimulus contrast, these authors report a clear peak in the gamma frequency band. In other words, the contrast dependence of the LFP is different in different frequency bands and the LFP power spectrum changes shape depending on contrast, with a peak in the gamma band emerging at high contrast. [36] studied the encoding of naturalistic sensory stimuli in LFPs and spikes. They found that the most informative LFP frequency ranges were 1–8 Hz and 60–100 Hz. They showed that the LFP in the 60–100 Hz high gamma band showed little noise correlation during visual stimulation but showed the highest observed signal correlation across all LFP frequencies. The high gamma band had the highest proportional power increase during visual stimulation. These experimental results are supported by the modeling work of [37]. These authors showed that their modeling network encoded static input spike rates into gamma-range oscillations generated by inhibitory-excitatory neural interactions. In sum, these reports indicate that the gamma frequency range is the one used most frequently to encode visual information in the visual cortex and that visual information is encoded by the power of gamma range oscillations. These observations, together with our result that gamma oscillations increase both the overall amount and the speed of information indicates that information about preferred stimuli is treated preferentially and, in consequence, that cortical modules mostly exchange information about their preferred stimuli.
In sum, we provide results to support the CTC hypothesis, we show evidence that CTC is a general mechanism independent of a specific frequency band and show that not only the phase but also the power is important to effectively shape the flow of information in a network.

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Author Contributions

Conceived and designed the experiments: AB. Performed the experiments: AB. Analyzed the data: AB. Wrote the paper: AB.

References

1. Gray CM, König P, Engel AK, Singer W (1989) Oscillatory responses in cat visual cortex exhibit inter-columnar synchronization which reflects global stimulus properties. Nature 338: 334–337.
2. Maldonado PE, Friedman-Hill S, Gray CM (2000) Dynamics of striate cortical activity in the alert macaque: II. Fast time scale synchronization. Cerb Cortex 10: 1117–1131.
3. Fries P, Reynolds JD, Rorie AE, Desimone R (2001) Modulation of oscillatory neuronal synchronization by selective visual attention. Science 291: 1560–1563.
4. Fries P, Schröder JH, Roeloffs PR, Singer W, Engel AK (2002) Oscillatory neuronal synchronization in primary visual cortex as a correlate of stimulus selection. J Neurosci 22: 3739–3754.
5. Bichot NP, Rossi AF, Desimone R (2003) Parallel and serial neural mechanisms for visual search in macaque area V4. Science 300: 529–534.
6. Taylor K, Mandon S, Freiwald WA, Kreiter AK (2005) Coherent oscillatory activity in monkey area V4 predicts successful allocation of attention. Cerb Cortex 15: 1424–1437.
7. Womelsdorf T, Fries P, Mira PP, Desimone R (2006) Gamma-band synchronization in visual cortex predicts speed of change detection. Nature 439: 733–736.
8. Fries P, Womelsdorf T, Oostenveld R, Desimone R (2008) The effects of visual stimulation and selective visual attention on rhythmic neuronal synchronization in macaque area V4. J Neurosci 28: 4823–4835.
9. Varela F, Lachaux JP, Rodriguez E, Martinerie J (2001) The brainweb: phase synchronization and large-scale integration. Nat Rev Neurosci 2: 229–239.
10. Salinas E, Sejnowski TJ (2001) Correlated neuronal activity and the flow of neural information. Nat Rev Neurosci 2: 539–550.
11. Fries P (2005) A mechanism for cognitive dynamics: neuronal communication through neuronal coherence. Trends Cogn Sci 9: 474–480.
12. Tiesinga PH, Fellous JM, Jose JV, Sejnowski TJ (2002) Information transfer in entorhinal cortical networks. Network Comput Neural Syst 13: 41–66.
13. Zeiter M, Fries P, Gielen S (2008) Biased competition through variations in amplitude of gamma-oscillations. J Comput Neurosci 25: 89–107.
14. Tass PA (2004) Transmission of stimulus-locked responses in two coupled phase oscillators. Phys Rev E Stat Nonlin Soft Matter Phys 69: 051909.
15. Buschman TJ, Miller EK (2007) Top-down versus bottom-up control of attention in the prefrontal and posterior parietal cortices. Science 315: 1060–1062.
16. Salesmann YB, Pigarev IN, Vidyasagar TR (2007) Neural mechanisms of visual attention: how top-down feedback highlights relevant locations. Science 316: 1612–1615.
17. Womelsdorf T, Schoffelen JM, Oostenveld R, Singer W, Desimone R et al. (2007) Modulation of neuronal interactions through neuronal synchronization. Science 316: 1609–1612.
18. Pesaran B, Nelson MJ, Andersen RA (2008) Free choice activates a decision circuit between frontal and parietal cortex. Nature 453: 486–489.
19. Gregorios GG, Gots SJ, Zhou H, Desimone R (2009) High-frequency, long-range coupling between prefrontal and visual cortex during attention. Science 324: 1207–1210.
20. Ishiguro K, Otsu N, Lungarella M, Kuniyoshi Y (2008) Comparison of nonlinear Granger causality extensions for low-dimensional systems. Phys Rev E Stat Nonlin Soft Matter Phys 77: 036217.
21. Wendling F, Ansari-Ad K, Bartolomei F, Sengadji I (2009) From EEG signals to brain connectivity: a model-based evaluation of interdependence measures. J Neurosci Methods 183: 9–18.
22. Schreiber T (2000) Measuring information transfer. Phys Rev Lett 85: 461–464.
23. Traub RD, Jefferys JG, Whittington MA (1997) Simulation of gamma rhythms in networks of interneurons and pyramidal cells. J Comput Neurosci 4: 141–150.
24. Kopell N, Ermentrout GB, Whittington MA, Traub RD (2000) Gamma rhythms and beta rhythms have different synchronization properties. Proc Natl Acad Sci USA 97: 1676–1672.
25. Roopun AK, Kramer MA, Carracedo LM, Kaiser M, Davies CH, et al. (2008) Temporal Interactions between Cortical Rhythms. Front Neurosci 2: 145–154.
26. Fries P, Roeloffs PR, Engel AK, König P, Singer W (1997) Synchronization of oscillatory responses in visual cortex correlates with perception in interocular rivalry. Proc Natl Acad Sci USA 94: 12699–12704.
27. Brunel N, Wang XJ (2001) Effects of neuromodulation in a cortical network model of object working memory dominated by recurrent inhibition. J Comput Neurosci 11: 63–85.
28. Brunel N, Wang XJ (2003) What determines the frequency of fast network oscillations with irregular neuronal discharges? I. Synaptic dynamics and excitation-inhibition balance. J Neurophysiol 90: 415–430.
29. Percival DB, Walden AT (1993) Spectral Analysis for Physical Applications: Multitaper and Conventional Univariate Techniques Cambridge University Press.
30. Mitra PP, Pesaran B (1999) Analysis of dynamic brain imaging data. Biophys J 76: 691–708.
31. Lungarella M, Ishiguro K, Kuniyoshi Y, Otsu N (2007) Methods for quantifying the causal structure of bivariate time series. Int J Bifurcat Chaos 17: 903–921.
32. Brovelli A, Ding M, Ledberg A, Chen Y, Nakamura R, et al. (2004) Beta oscillations in a large-scale sensorimotor cortical network: directional influences revealed by Granger causality. Proc Natl Acad Sci USA 101: 9849–9854.
33. Houweling S, Beek PJ, Daffertshofer A (2010) Spectral changes of interhemispheric crosstalk during movement instabilities. Cerb Cortex Advance Access: doi:10.1093/cercor/bhp088.
34. Lungarella M, Ishiguro K, Kuniyoshi Y, Otsu N (2007) Methods for quantifying the causal structure of bivariate time series. Int J Bifurcat Chaos 17: 903–921.
35. Brovelli A, Ding M, Ledberg A, Chen Y, Nakamura R, et al. (2004) Beta oscillations in a large-scale sensorimotor cortical network: directional influences revealed by Granger causality. Proc Natl Acad Sci USA 101: 9849–9854.
36. Belitski A, Gretton A, Magri C, Murayama Y, Montemurro MA, et al. (2008) The neuronal basis of attention: Rate versus synchronization modulation. J Neurosci 28: 7679–7686.
37. Heurie JA, Shapley R (2005) LFP power spectra in V1 cortex: the graded effect of stimulus contrast. J Neurophysiol 94: 479–490.
38. Belitski A, Gretton A, Magri C, Murayama Y, Montemurro MA, et al. (2008) Low-frequency local field potentials and spikes in primary visual cortex convey independent visual information. J Neurosci 28: 5696–5709.
39. Mazziotti A, Panzeri S, Logothetis NK, Brunel N (2005) Encoding of naturalistic stimuli by local field potential spectra in networks of excitatory and inhibitory neurons. PLoS Comput Biol 4: e1000239. doi:10.1371/journal.pcbi.1000239.