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that viral infection of circulating cells, for example, monocytes, can succeed only if the virus prevents elimination of these cells by virus-specific CTLs. More work, however, will be required to identify the cell type supporting superinfection.

Although the biochemical and cell biological functions of US2, US3, US6, and US11 have been studied extensively (18), their role in viral pathogenesis had remained enigmatic. Analogous gene functions in murine CMV (MCMV) had been similarly found to be dispensable for both primary and persistent infection (10), although reduced viral titers have been reported for MCMV deleted for these genes (19). Thus, the reason all known CMVs dedicate multiple gene products to MHC-I down-regulation had remained elusive. Our current results now identify a critical role for these immunomodulators to enable superinfection of the CMV-positive host. Furthermore, these results suggest that the ability to superinfect is an evolutionary conserved function among CMVs and therefore might play an important role in the biology of these viruses. Superinfection could promote the maintenance of genetic diversity of CMV strains in a highly infected host population, which could provide an evolutionary advantage. However, there is another possibility. CMV is a large virus with thousands of potential T cell epitopes and therefore a high potential for CD8<sup>+</sup> T cell cross-reactivity (20). Indeed, in a study of pan-proteome HCMV T cell responses, 40% of HCMV seronegative subsets manifested one or more cross-reactive CD8<sup>+</sup> T cell responses to HCMV-encoded epitopes (5). As CMV recognition by cytotoxic T cells appears to effectively block primary CMV infection, individuals with cross-reactive CD8<sup>+</sup> T cell immunity might be resistant to CMV. Thus,

US2-11 function may be necessary to evade such responses and establish infection in this large population of individuals that might otherwise be CMV-resistant.

Our results also may explain why, so far, it has not been possible to develop a vaccine that efficiently protects humans from HCMV infection. Although antibody-mediated mucosal immunity might reduce the rate of superinfection (7, 21), once this layer of defense is breached, CMV-specific CTLs seem to be unable to prevent viral dissemination, due to MHC-I down-regulation by US2-11. Thus, although CMV-vaccines might be able to limit CMV viremia and associated morbidity, this MHC-I interference renders it unlikely that sterilizing protection against CMV infection is an achievable goal.

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22. We are grateful to P. Barry for providing the RhCMV-GFP and RhCMV-BAC and M. Messerle for plasmid ori6k-F5. The NIH Nonhuman Primate Reagent Resource Program provided the CD8 $\alpha$ -specific antibody cM-T807 used in this work (contracts AI040101 and RR016001), which was originally obtained from Centocor, Inc. We thank A. Townsend for help with the graphics. We thank D. Drummond, L. Coyne-Johnson, M. Spooner Lewis, C. Hughes, N. Whizin, M. Giday, J. Clock, J. Cook, J. Edgar, J. Dewane, and A. Legasse for technical assistance. This research was supported by the National Institutes of Health (RO1 AI059457 to K.F. and RO1 AI060392 to L.J.P.), the International AIDS Vaccine Initiative (to L.J.P.), the National Center for Research Resources (RR016025 and RR18107 to M.K.A.; RR00163 supporting the Oregon National Primate Research Center), the Ruth L. Kirschstein National Research Service Awards T32 AI007472 (to C.J.P.) and T32 HL007781 (to R.R.), the OHSU Tartar Trust fellowship (to C.J.P.), and the Achievement Reward for College Scientists (to R.R.). The CMV vector technology has been patented by Oregon Health and Science University (US2008/0199493A1) with L.J.P., J.A.N., and M.A.J. as coinventors. This patent has been licensed to the International AIDS Vaccine Initiative. The comparative genome sequencing data shown in fig. S5 were deposited in the Gene Expression Omnibus database (accession number GSE20308).

#### Supporting Online Material

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Materials and Methods  
Figs. S1 to S5  
Table S1  
References

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## Synchrony of Thalamocortical Inputs Maximizes Cortical Reliability

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Thalamic inputs strongly drive neurons in the primary visual cortex, even though these neurons constitute only ~5% of the synapses on layer 4 spiny stellate simple cells. We modeled the feedforward excitatory and inhibitory inputs to these cells based on in vivo recordings in cats, and we found that the reliability of spike transmission increased steeply between 20 and 40 synchronous thalamic inputs in a time window of 5 milliseconds, when the reliability per spike was most energetically efficient. The optimal range of synchronous inputs was influenced by the balance of background excitation and inhibition in the cortex, which could gate the flow of information into the cortex. Ensuring reliable transmission by spike synchrony in small populations of neurons may be a general principle of cortical function.

Neurons can perform coincidence detection of synaptic inputs with a temporal integration window that depends on the time courses of the synaptic conductances and the intrinsic properties of the postsynaptic

neuron (1). Synchronous cortical inputs occur when there is a salient event in the sensory environment, such as the entrance of a moving object into a receptive field (2) or the deflection of a whisker in rodent (3). The precise

timing of action potentials has been shown to potentially provide information in addition to the spike rate (2, 4–7). For a population of presynaptic neurons to fire nearly simultaneously, however, requires resources to time spike initiation precisely, parallel anatomical pathways to carry the spikes, and energy costs for redundant spikes, which may outweigh the benefits of increased information rate (8, 9). We explored these issues in the projections from the lateral geniculate nucleus (LGN) to the primary visual cortex.

The question of efficient information transfer is particularly important for thalamocortical connections, because thalamic synaptic inputs,

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which are comparable in strength to cortical inputs, constitute only ~5% of the total synaptic input to cortical simple cells (10–12), but are nonetheless capable of reliably driving cortical neurons. To examine the relationship between synchrony and reliability, we performed computer simulations of a detailed biophysical model of a spiny stellate cell in layer 4 of area V1 (13) of the cat primary visual cortex (Fig. 1A). This cell received 300 synaptic inputs from the LGN, competing with 5500 other excitatory and inhibitory intracortical synapses, including feedforward inhibition. All synapses were stochastic, and the excitatory synapses included short-term history-dependent modulation of release probability. We used this model to quantify the number of synchronous synaptic inputs that maximizes the efficient transfer of information to the cortical cell, and we compared these predictions to experimental data obtained in vivo from anesthetized cats (14).

The pairwise correlation strength between two neurons is often used to estimate their degree of synchrony (15, 16). In the visual thalamocortical pathway, correlated spikes from two presynaptic LGN neurons strongly increase the probability of firing in a postsynaptic primary visual cortex (V1) simple cell (17–19) and ensure the transfer of information from the thalamus to the cortex, despite a low probability of synaptic transmission (3, 20). Our simulations support these results by showing that spike output reliability can be predicted by synchrony magnitude (SM), which is the number of thalamocortical synapses that are simultaneously driven by the same presynaptic thalamic spike train. Simultaneous recordings from more than two LGN neurons would be needed to measure the SM directly.

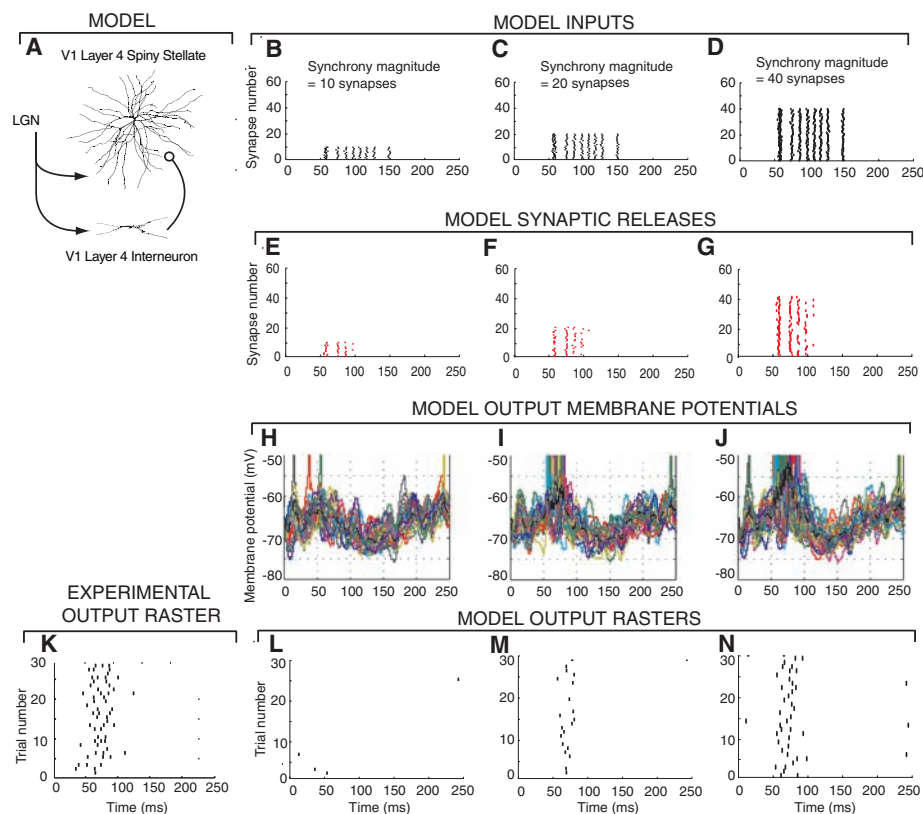
We compared the output of our model with in vivo recordings from Kara *et al.* (21), who simultaneously recorded spike trains from retinal ganglion cells, LGN relay cells, and cortical cells in the primary visual pathway of anesthetized cats. The visual stimulus was a drifting grating, which produced patterns of spikes without highly synchronous events, as occurs with flickering visual stimuli (2). The presynaptic spike times from the in vivo recordings of LGN cells were distributed across a varying number of input synapses in the model (Fig. 1, B to D), as well as across an inhibitory feedforward pathway leading to the spiny stellate cell. This pattern was transformed by the synapses into a sequence of neurotransmitter releases (Fig. 1, E to G) and integrated with the cortical background inputs to produce a fluctuating membrane potential (Fig. 1, H to J, and figs. S1 and S2). The average firing rate and the reliability (22) of the cortical cell spike pattern was then computed across 30 repeated 250-ms trials (Fig. 1, K to N), each using a different LGN cell spike pattern from the in vivo recordings (14).

Output reliability was a highly nonlinear function of SM, rising steeply from 20 to a maximum at 40 synapses (Fig. 2A), more rapidly than the output firing rate (Fig. 2C). The reliability-per-SM (RPSM) function, defined by dividing the reliability by the SM, reached an optimal synchrony magnitude (OSM) at approximately 30 synapses (Fig. 2B, dashed line). The reliability-per-spike (RPS) ratio, a measure of the reliability increase for each additional output spike computed by dividing each reliability output by its corresponding firing rate, also peaked at approximately 30 synchronous synapses (Fig. 2D). Similar results were obtained when the synaptic inputs were generated from groups of five synchronous inputs corresponding to single LGN afferents with each group from a different recorded LGN input (fig. S2).

In experimental recordings from thalamic and cortical cells, the trial-to-trial spike-count variability was low (21, 23), indicating that

the spike rates of these cells conveyed important information about the input stimulus. Using spike trains from LGN recordings as inputs to the model cell (21), we also found low spike-count variability: The Fano factor (FF), defined as the sample variance divided by sample mean, achieved a minimum of 0.2 to 0.4 in the range of 20 to 80 synchronous synapses (fig. S4).

The quantitative analysis of output reliability can be used to predict the number of synchronous inputs that drive cortical cell responses during in vivo behavioral experiments (Fig. 3). We plotted data from in vivo recordings of V1 cells supplied by Kara *et al.* (21) against the transfer function of input synchrony as a function of output reliability, as determined by our model, to infer synchrony magnitude of the cells in vivo. Each of the four neurons (from different animals) predicted input synchrony in the range of 20 to 60 synchronous synapses (Fig. 3). The reliability and



**Fig. 1.** Varying the synchrony of input synapses affects output reliability and firing rates. (A) Morphology of a reconstructed V1 layer 4 spiny stellate cell that was modeled with 748 compartments, thalamocortical (TC) synapses with a release probability of  $P = 0.2$  and short-term plasticity (13, 14), and a feedforward inhibitory interneuron that receives input from LGN and projects to the spiny stellate cell. (B to D) Rastergrams of 60 thalamocortical synaptic inputs into the model cell for one trial. An input spike train obtained from in vivo LGN recordings (21) (referred to as event times) was repeated on a number of input synapses (SM). The input events were time jittered on average by 1 ms and had 1 out of 10 spikes randomly deleted. (E to G) Synaptic release rastergram for inputs (B to D). (H to J) Superimposed spiny stellate membrane potentials from 30 trials. (K) Cortical cell output spike trains from experimental in vivo recordings (21). (L to N) Rastergrams of outputs from 30 trials of inputs based on different LGN spike trains from the in vivo data.

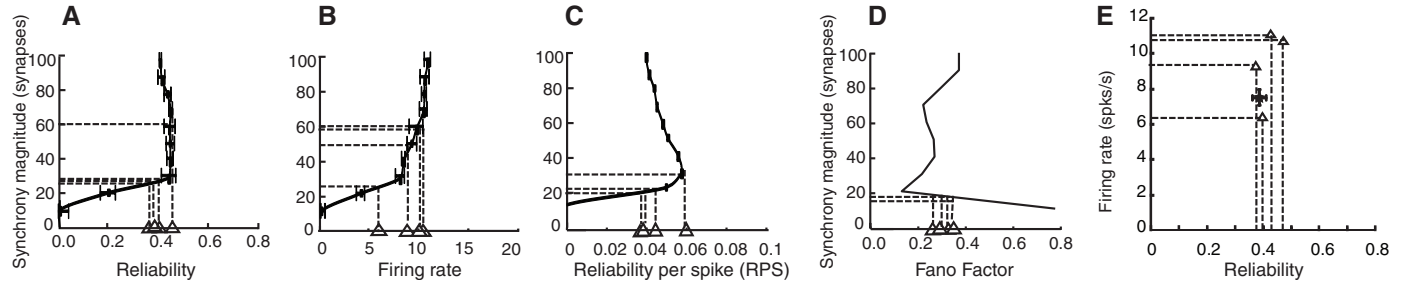
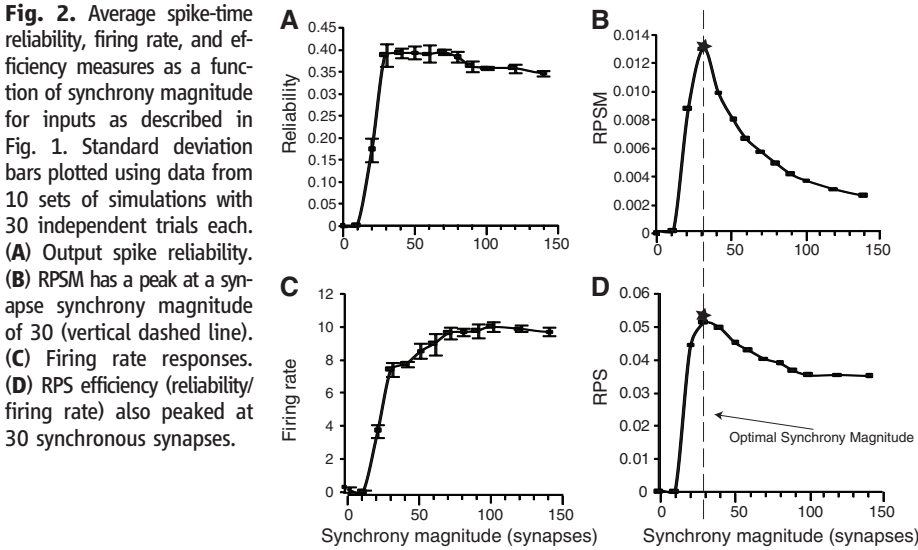
firing rate at the OSM from our model (Fig. 2, stars), using recorded thalamic input spike trains, fall within the observed ranges of the experimental values (Fig. 3E).

These results were robust to increasing the jitter and varying the strengths of thalamic inputs and feedforward inhibitory synapses (Fig.

4, A and B, and fig. S4). However, the OSM depended on the balance between background excitatory (*E*) and inhibitory (*I*) inputs to the 5500 intracortical synapses. When the integrated inputs were balanced (total average excitatory input equal to the total average inhibitory input), the neuron became highly sensitive to correlated

fluctuations of the membrane potential, which affected the reliability (23) as well as the gain of the input spike rate to output spike rate curve (24). We varied the cortical background rate and the ratio of inhibitory to excitatory input rates,  $\beta = I/E$ , to assess their influence on the reliability of the synchronous inputs to LGN synapses (Fig. 4C and fig. S3). Background excitatory inputs above 1 spike/s depressed the reliability response by directly competing with the LGN excitatory inputs and introducing high levels of spurious output firing. Increasing the background inhibition also increased the OSM and was therefore a potential mechanism for setting the threshold for synchrony detection. The sensitivity of the cortex to synchronous inputs can be varied over a wide range by regulating  $\beta$ .

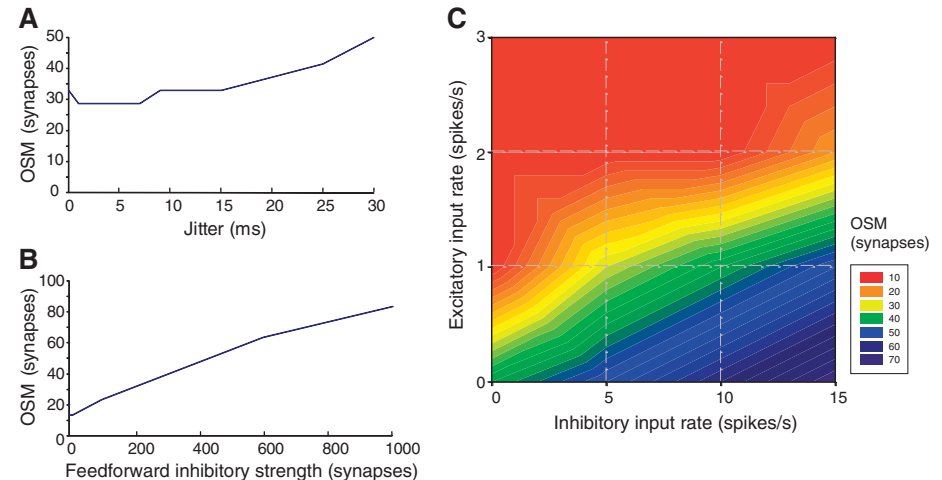
Spiking due to input synchrony may be a way to ensure that important events are registered by the spiny stellate neurons in the cortex, regardless of asynchronously arriving spikes from ongoing cortical computation. This prediction could be tested with intracellular recordings in vivo by using a dynamic clamp (25) to inject somatic conductances reflecting events in the presence of in vivo background noise.



**Fig. 3.** Predicted input synchrony ranges for in vivo recordings. Graphs from Fig. 2 were inverted to make synchrony magnitude the dependent variable. In each plot, four triangles are positioned along the x axis (input value), corresponding to in vivo experimentally measured values from four separate animals (21), and the inferred output synchrony magnitude is shown as a horizontal dashed line from the first intersection on

each curve. Predictions were made based on (A) reliability, (B) firing rates, (C) RPS, and (D) FF (taken from fig. S4C). (E) The predicted firing rate and reliability at SM = 30 (solid circle with error bars indicating SD) is plotted along with the measured values from the four sets of recordings (triangles). Despite the small sample of cells with enough trials, the estimated reliabilities cluster around 0.4.

**Fig. 4.** Effect of input jitter, inhibitory interneuron strength, and the balance of inhibitory and excitatory background inputs on the predicted OSM. (A) The jitter of the input event signals was varied from 0 to 30 ms. The default used in Figs. 1 to 3 used jitter = 1 ms. (B) The number of synapses from the feedforward inhibitory interneurons was varied from 0 to 1000 synapses. The default used in Figs. 1 to 3 was 200 synapses. (C) The Poisson-distributed presynaptic spike trains for the 4500 excitatory (glutamate) and 1000 inhibitory ( $\gamma$ -aminobutyric acid) intracortical synapses were covaried from 1 to 3 spikes/s excitatory background inputs and 1 to 15 spikes/s inhibitory background inputs. The default in Figs. 1 to 3 was 1 excitatory spike/s and 5 inhibitory spikes/s.



Moving visual inputs give rise to synchronous spikes in retinal ganglion cells (26). A single retinal ganglion cell can drive four or more LGN cells (27), which can make one to eight synapses each (28), with a spiny stellate cell in V1 with overlapping receptive fields (17). Thus, assuming an average of four synapses for each LGN axon and an OSM of 20 to 40, as few as 5 to 10 LGN cells could effectively drive a cortical neuron. This prediction could be tested by studying the effects of modulating the contrast of visual stimuli on the firing rates and reliability of spike timing in cortical cells (21, 29). Cortical feedback to the thalamus could also regulate the degree of synchrony among thalamocortical cells.

We have quantified the number of synchronous thalamic spikes needed to reliably report a major sensory event to cortical neurons (19), which is consistent with previous physiological experiments that found correlated firing with 1-ms precision in visual cortex (4, 17) and mouse barrel cortex (3). Spike synchrony, through converging anatomical pathways, enhances the information transfer rate and speeds up processing (30, 31).

The output spike pattern of a layer 4 neuron is thus determined by the temporal pattern, as well as the rate, of the synchronous thalamic inputs according to the history-dependent dynamics of its synapses acting coherently within 6 to 8 ms. The same analytical methods used here could be applied to study reliability and

connectivity in other types of neurons. Spike synchrony, observed throughout the cortex, may also have a more general function in ensuring information transmission between cortical areas (32).

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References

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