relaxed. An object can be also be described by an imperfectly closed boundary together with constant texture or known structure.

Now, what is required for this? First, a possibility to group V1 cells without destroying their filtering properties, which is clearly provided by P&S. Second, the process must be cascadable, that is, groups of groups must be possible. This is a bit more problematic, because the number of *different* synchronized groups at a time is very limited. For the system proposed here, this is not a serious problem, because the processors keep their RF properties intact, so the relevant cells for a whole object may be synchronized into one group without confusion. For implementing, for example, a graph structure like the one used by Lades et al. (1993) it might constitute a serious limitation.

If such object descriptions develop in a self-organized way based on the input image it should be relatively easy to match them with stored models of objects, although the integration of flexible matching and associative memory remains an open problem. Anyway, the target article opens a couple of interesting new routes of investigation.

Authors' Response

Progress toward an understanding of cortical computation

W. A. Phillips^a and W. Singer^b

^aCenter for Cognitive and Computational Neuroscience, Department of Psychology, University of Stirling, FK9-4LA, Scotland, United Kingdom; wap@forth.stir.ac.uk www-psych.stir.ac.uk/~wap; ^bMax Planck Institute for Brain Research, D-60496, Frankfurt/Main, Germany; singer@mpihfrankfurt.mpg.de

Abstract: The additional data, perspectives, questions, and criticisms contributed by the commentaries strengthen our view that local cortical processors coordinate their activity with the context in which it occurs using contextual fields and synchronized population codes. We therefore predict that whereas the specialization of function has been the keynote of this century the coordination of function will be the keynote of the next.

We thank all of our commentators for producing thoughtful and insightful commentaries in the few weeks available. The commentaries come from five broad areas of expertise: neuroscience and neurology (13); computing, neural computing, and informatics (10); psychology (5); theoretical brain physics (2); and psychiatry (2). This diversity is a testament to the importance of trying to understand cortical computation. Anything common to such diverse sources of knowledge must reflect a deeper reality. We do not suppose, however, that all these perspectives can be boiled down to any single perspective, including that emphasized in the target article. Just as we suppose that cortical cells can cooperate so as to emphasize coherence while nevertheless maintaining their own distinctive contribution, so we suppose that different scientific perspectives can all shed light on a common underlying reality while nevertheless remaining distinct.

We first present a summary of the views taken on four central issues, and then discuss two of the most basic: the working hypothesis of common foundations, and the role of contextual coordination. The main body of the response is divided into nine sections. R1 to R5 discuss the five issues

Table R1. Views on four basic issues

Are common foundations for cortical computation likely?

- YES Amit, Bower, Bugmann, Eckhorn, Floreano, Gregson, Grossberg, Haase & Diniz, Iacoboni, König et al., McCollum, Morasso et al., Niebur & Usher, Palm & Wennekers, Silverstein & Schenkel, Smyth, Tononi & Edelman, Treves, Wang, Wright, and Würtz.
- NO ? Nunez, Stone.

Is the coordination of activity a fundamental goal?

- YES Amit, Bugmann, Eckhorn, Floreano, Gregson, Grossberg, Haase & Diniz, Iacoboni, König et al., McCollum, Morasso et al., Niebur & Usher, Nunez, Silverstein & Schenkel, Smyth, Stone, Tononi & Edelman, Wang, Wright, and Würtz.
- NO Treves.
- ? Bower, Palm & Wennekers.

Is the distinction between CFs and RFs useful?

- YES Bugmann, Eckhorn, Floreano, Grossberg, Iacoboni, König et al., McCollum, Morasso et al., Silverstein & Schenkel, Smyth, Stone, Tononi & Edelman, Wang, Wright, and Würtz.
- NO Niebur & Usher.
- Amit, Bower, Gregson, Haase & Diniz, Nunez, Palm & Wennekers, and Treves.

Does the cortex use synchronized population codes?

YES Bower, Bugmann, Eckhorn, Floreano, Gregson, Grossberg, Haase & Diniz, Iacoboni, König et al., McCollum, Niebur & Usher, Nunez, Palm & Wennekers, Silverstein & Schenkel, Tononi & Edelman, Wang, Wright, and Würtz.

NO Amit, Treves.

? Morasso et al., Smyth, and Stone.

listed in sections 6.1 to 6.5, and in the same order. R6 discusses possible additional foundations proposed by commentators. R7 discusses further issues raised by commentators. R8 takes up the invitation of one of the commentators to pursue the analogy between the scientific enterprise and cortical computation. Finally, R9 notes promising directions for future progress.

Table R1 classifies commentaries in relation to the views taken on four basic issues: (1) Are common foundations for cortical computation likely? (2) Is the coordination of activity a fundamental goal? (3) Is the distinction between contextual fields (CFs) and receptive fields (RFs) useful? (4) Does the cortex use synchronized population codes? Each commentary is classified as "yes," "no," or "agnostic" on each question. Most classifications are based on what is said explicitly, but in a few cases we inferred what seemed implied. Nevertheless, the classifications given are ours. There is no guarantee that commentators would agree.

On each of the four questions Table R1 classifies the majority of commentators as giving the same answers we

do. Few argue explicitly against the positions we take on these issues. They were not short of criticisms, of course, with the most prominent being as follows. **Amit** and **Treves** doubt any role for synchronization. **Bower** says that his biological models suggest a role for it, but a role that differs from ours. **Niebur & Usher** agree with us on the role for synchronization, but doubt CFs. **Palmer & Wennekers** think that synchronization is important, but criticize the computational studies for not modeling activity at the level of individual spikes. Some commentators see no role for information theory. There is thus no agreement between these criticisms, and much disagreement. Several commentaries call for more emphasis upon top-down processes. This is well within the spirit of our approach, and we agree.

All commentaries agree that it is worth searching for common foundations. This consensus does not show that view to be correct. There is no reason to suppose that those whose skeptical views were noted in sections 2.1, 2.2, and 2.3 are no longer skeptical. It is simply that their views are not represented here. It would help if they were, because their perspective suggests aspects of cognition that are unlikely to be directly comprehensible in terms of common foundations. Our search will be led seriously astray if it attempts to explain, without additions, aspects that require further capabilities. With respect to language, at least, the best supported hypothesis is that it does involve special capabilities (Pinker 1994). Whether these depend upon the common foundations, as our perspective suggests, remains to be seen.

Stone and **Nunez** both note that the hypothesis of common foundations is not well specified. We have not attempted any formal specification, simply using the word "common" to mean most or many, and not necessarily all. We do not see how this can be formalized. It is an empirical rather than a formal issue, so we keep our minds open as to what constitutes the range of cortical and cognitive processes to be understood under a common rubric. The more the better.

We now turn to views on the role of contextual coordination. We argued that functional specialization and contextual coordination together form two basic and mutually constraining principles of cortical organization (sect. 1.1). Functional specialization is undisputed. The issue is whether it is balanced against a need for coordination. Tononi & Edelman have argued elsewhere (Tononi et al. 1996), both formally and forcefully, that this is so, and von der Malsburg and Singer (1988) have shown how the emergence of global order from local cooperation is a very general property of physical systems composed of many distinct but interacting elements. The forms of contextual coordination emphasized in the target article include the selection of contextually relevant signals, their grouping into coherent subsets, and contextual disambiguation. Although most commentaries seem to agree with the need for such coordination, two raise important doubts. Treves argues for the value of dissonance and of letting cells be free rather than having them goose-stepping in synchrony. **Niebur & Usher** argue that as unexpected events can be both important for survival and highly salient, they constitute cases where our emphasis upon coherence is exactly the opposite of what is required.

Treves stressed the possibility that information transmission may be reduced rather than increased by synchrony. We will discuss evidence for synchrony as a cortical code in section R3.1. Here we focus upon conceptions of the goals of cortical processing. Underlying Treves's concern for squeezing as much information as possible out of cortical cells is the long and strong tradition that emphasizes the goal of recoding to reduce redundancy (sect. 1.2.1). If the goal is to transmit as much information as possible using a limited and noiseless channel, an optimal recoding will map activity into statistically independent variables. We agree that this perspective makes an important contribution to our understanding of cortical computation, but taken to the extreme it would lead to something worse than chaos. First, neither the external world nor the cortex are noiseless, so redundancy is necessary. More important, the transmission of as much information as possible seems to be a highly implausible goal for cortex. It is more plausible that it discovers what predicts what, including various aspects of reinforcement, and uses that to transmit information about just those variables that matter. This will emphasize signals that are coherently related, and thus reduce total information transmission.

Niebur & Usher note the sudden appearance of a predator as an example of the salience of "unexpected" events; they view this as evidence for the opposite of our emphasis upon coherence. We are well aware of the salience of abrupt changes. They were central to our studies of the detection of appearance and disappearance of single elements in random-dot arrays using psychophysical and physiological techniques (Phillips & Singer 1974; Singer & Phillips 1974). Gradients of change in both space and time are so crucial that they are central to RF specificity at all levels. This has long been clear, and it explains why contrast in either time or space (including higher feature spaces) is so noticeable. Does this imply that incoherent signals are more salient than coherent signals? Not at all. Consider a random-dot array in which two subsets of elements appear, one in some structured or familiar arrangement, the other at random. It is the structured or familiar subset that will be most salient, not the random one. The sudden appearance of a predator will produce activity across a wide array of cells in the visual cortex. Camouflage evolves so as to reduce the coherence of that activity, not so as to increase it. Thus, Niebur & Usher provide an excellent example that expresses at the intuitive and behavioral level our view that coherent patterns of activity, not incoherent ones, are most effective at the cellular level.

R1. Does the distinction between RFs and CFs have biological relevance?

R1.1. Niebur & Usher's doubts

Most commentaries agree that this distinction is useful. **Niebur & Usher**, however, argue explicitly that it is not. Their grounds for this are that lateral interactions in the cortex can be explained by excitatory and inhibitory interactions alone, making no use of gain-control mechanisms. **Smyth** discusses the same phenomena as Niebur & Usher, but, in contrast, concludes that such lateral interactions support the RF/CF distinction and should be included in the CFs. Who is right? In section R1.7 we will outline a scenario in which they could both be partly right. Here we emphasize two more general points. First, CFs are primarily distinguished by the effects they have, that is, modulatory and synchronizing, not by the source from which they come. They should not be identified with lateral interactions. Some long-range interactions may be concerned with computing RF features that require comparison with a large surround. RF features defined by comparing a center with a large surround are perfectly compatible with the contextual coordination of those features through CFs, however. Second, CFs are primarily distinguished by the effects they have, not by the mechanism producing those effects. Furthermore, although Niebur & Usher suggest a different mechanism for lateral interactions, they acknowledge elsewhere that voltage-dependent channels (e.g., NMDA) are common (Usher & Niebur 1996), and may play a role in "top-down attentional modulation." Thus, in our terms, they do assume a role for voltage-dependent CF mechanisms, suggesting that it is specifically intraregional voltage-dependent CFs that they doubt. If so, their view seems similar to that of Grossberg in this respect (sect. R1.7). The absence of such CFs is not yet clearly established. It cannot be established by showing that some longrange intraregional connections contribute to RFs, nor by showing that they may in part use mechanisms different from those we emphasized.

R1.2. How are RFs and CFs distinguished?

Given that this distinction is useful, how should it be drawn? At least five possible contrasts must be considered: (1) RFs have a driving effect (driving activity either up through excitation or down through inhibition); CFs are modulatory (producing either facilitative or suppressive gain-control); (2) RFs determine what information signals transmit; CFs help determine exactly when they are sent; (3) RFs determine what information signals transmit; CFs specify with which other signals they are to be grouped. (4) RFs determine what decision signals transmit whereas CFs help determine the confidence with which that decision is made; (5) RFs tend to be feedforward; CFs tend to be lateral and feedback.

Tononi & Edelman and Eckhorn cite further evidence that most of those contrasts are relevant. Our view is close to theirs but with a few differences of emphasis. Tononi & Edelman doubt the usefulness of distinguishing between decisions and confidence in those decisions. They rightly note the difficulty of making a categorical decision by distinguishing between active and inactive neurons. Our working assumption is that different outcomes of categorical decisions are represented by different groups of neurons with reciprocal inhibition between groups. The categorical decision could then be conveyed by the group transmitting more spikes or more synchronized ones, than the others, and the confidence by how much more. Tononi & Edelman put more emphasis on contrast 5 than we do. We propose that CFs are distinguished by the effects they have, not by the source from which they come. Nevertheless we agree that CFs are more likely to come from lateral and feedback sources. Eckhorn notes that there is little or no physiological evidence for suppressive effects between uncorrelated signals. This is a simple misunderstanding. The computational framework outlined in section 3 implies that CF connections will adapt so that they have no effect if they link streams between which activity is uncorrelated.

R1.3. Terminology for CFs

Given the similarities between our view of CFs and those of **Tononi & Edelman** and of **Eckhorn**, we explain briefly

why we use the terms we do, rather than those they propose. Calling the CFs reentrant connections puts the emphasis upon the source from which they come rather than upon the effects that they have. If it turns out that some bottom-up connections mediate contextual modulation, for example, to speed-up coordination, we could refer to them as feedforward CFs. Calling them feedforward reentrant connections would sound less coherent. Eckhorn calls the CFs association fields, as do Field et al. (1993). Although this has some appropriate connotations, it seems more appropriate to the retrieval of concepts. We therefore prefer to use the term associative connections to refer to the internal connections in attractor neural networks, such as those analyzed in detail by **Amit** (1989), Hopfield (1982), and many others.

R1.4. How could CFs produce synchronization?

It is noted by **Palm & Wennekers** that the computational theory outlined in section 3 does not show in detail how biological CFs could produce synchronization. Previous studies by many others have studied this in detail (sect. R1.9), however. An excellent example of the use of CFs in technological implementations is given here by **Wang.** His success in processing real images shows that CFs would be useful if they had biological reality. Further computational support for synchronizing CFs is provided by **Würtz** who notes that their use may overcome the problem of distinguishing multiple simultaneously active groups, which is a difficulty for the dynamic link architecture (DLA) (Lades et al. 1993). Thus, a system using CFs may be less flexible than the DLA, but closer to the biology.

R1.5. Is there a role for inter-regional feedback CFs?

There is an explicit call by Tononi & Edelman, König et al., Grossberg, and Bugmann for more emphasis upon top-down processes. We agree. Top-down sources for CFs are clearly shown in Figure 1c. Physiological evidence for such effects was cited in section 4, and their relevance to word perception was discussed in section 5.7.2. Nevertheless we understand why several commentators were misled about our views on this, because we did put more emphasis upon bottom-up stimulus-driven contextual coordination. This was not because we doubted the importance of topdown influences. Two classes of top-down effect must be distinguished, stimulus driven and strategically driven. There is clear physiological evidence for top-down contextual modulation that is stimulus driven (Zipser et al. 1996), and our computational studies have included such effects (e.g., Phillips et al. 1995a; 1995b). König et al. now provide evidence for strategic top-down coordination by showing that synchronization depends upon the current task and not just upon the current stimulus. We agree that such effects are important and could be mediated by CFs.

R1.6. Is there a role for internal sources of RF drive?

The flow of activity that arises from within the cortex itself is emphasized by **Tononi & Edelman**, and **Wright** suggests that intrinsic attractor dynamics be incorporated into the concept of a local-processor, and in such a way as to be seen as part of the RF. We see our approach as supporting both proposals. They rightly remind us that internal systems, for

example those concerned with plans and strategies, can provide a source of driving as well as of coordinating activity. Coordination of this internally generated activity is also crucial, however, because our thoughts and plans also need to be coherent (sect. R7.4).

R1.7. How are CFs related to long-range interactions in visual cortex?

The distinction between RFs and CFs is unnecessary, Niebur & Usher argue, because the effects of context can be explained without invoking a separation between the processing of context and content. It is not clear whether they are denying the functional distinction or proposing a different mechanism for it, however. Having denied the distinction, they then use it so that the sign of the effect of "context" depends upon the strength of the "input" (i.e., the RF input). If there were no difference between the processing of context and content then the sign of the effect of RF input should depend upon the strength of the context, and they do not claim that to be the case. It thus seems more likely that they are proposing a mechanism for the distinct effects of context. This interpretation is strengthened by their description of the physiological and psychophysical data to be explained as showing contextual modulation (Stemmler et al. 1995). The mechanism they propose is based upon the phenomenon of "stochastic resonance," in which noise enhances detection of weak signals by increasing the gain at weak signal strengths (Stemmler et al. 1995). Bezrukov & Vodyanov (1997) have now proved that stochastic resonance can occur in voltage-dependent ion channels, however, so the mechanism that Niebur & Usher propose may be more compatible than they thought with the one we emphasized. If so, that would enable them to preserve their account of the relevance of stochastic resonance without having to explain away evidence that long-range intra-regional connections activate voltagedependent channels (Hirsch & Gilbert 1991). Even that is unlikely to provide a complete account of long-range lateral interactions, however, because it is hard to see how enhancing the detectability of weak elements explains the figural grouping of high-contrast elements (e.g., as in Field et al. 1993).

Smyth, in clear contrast to Niebur & Usher, interprets the data on long-range interactions in V1 as evidence for CFs. Kastner et al. (1997) provide evidence for modulatory effects upon cells in visual cortex (V1). Their Figure 4a shows a cell with narrow orientation tuning. It is inhibited by a single central bar orthogonal to its preferred orientation. There is no sign of inhibition being produced by a surrounding array of parallel elements alone, but when they are presented along with an optimally oriented central bar they reduce firing rate from about 20 to about 5 spikes per sec above spontaneous. This is clear evidence for modulatory rather than driving effects, and Kastner et al. describe their findings in terms of effects upon "salience." This implies that they view the effects of the surround as not changing what is signaled, but as changing how effective that signal is at subsequent levels. This is in accord with what Smyth proposes.

The view just outlined allows for the dependence of context effects on the strength of the central stimulus. The hypothetical activation function used in section 3.1 of the target article produces smaller effects of context as target strength increases, and in addition sect. 6.1 suggests that effects of context upon spike rate in cortex may reduce as target strength increases. Studies of context using visual evoked potentials show this to be so (Polat & Norcia 1996). Thus this aspect of "contrast-dependence" is catered for by the simple assumptions made in section 3, contrary to what **Smyth** may be taken as implying. The aspect that he intends to emphasize, however, is the change in *sign* of the context effects with changes in the strength of the central RF stimulus (personal communication with Smyth). We agree that this is not catered for in section 3.

For a broader view of long-range interactions it is crucial to distinguish several levels of processing. Two levels of analysis can be distinguished in the perception of textured stimuli such as those used in many of these experiments (Sagi 1995). At the lower level local features are computed, for example by anisotropic Gabor-like filters. At the higher level, further center-surround comparisons are used to compute gradients of change in the lower-level feature space (e.g., by isotropic Gabor-like filters with larger RFs than at the lower level). The model outlined by Sagi (1995) is concerned only with the basic feedforward RF architecture, so we need to add contextual coordination within levels and from the higher to the lower level. Detectors of distinct stimulus elements could therefore interact through modulatory CF effects at the lower level, but with their outputs being combined in RF computations at the higher level. At the higher level of analysis, however, long-range comparisons are used to compute gradients in feature space, and these are best thought of as being part of RF computation because information about the separate stimulus elements is lost (Sagi 1995). This is in keeping with the view expressed in section 1.3, except that it occurs in secondary rather than primary visual cortex.

This scenario may provide a reply to the second question that **Smyth** raises, that is, it shows how synchronization can have a role in input pre-processing. It allows for the dynamic sculpting of the exact extents of the effective RF centres and surrounds of cells in secondary visual cortex. These would otherwise be too rigid to reconcile, for example, the conflicting demands of segregation and integration (Braddick 1993). There is evidence that this involves topdown modulatory inputs (Salin & Bullier 1995), including those that are stimulus-driven (Zipser et al. 1996), as well as those that are task-dependent (Maunsell 1995). Zipser et al. (1996) found evidence for top-down contextual influences on V1 cells by showing that firing rate is increased by any of a variety of higher-level cues that made the features being signaled part of the figure rather than the ground. Synchronization of firing was not studied. Given the evidence for diffuse axonal bifurcation of the feedback connections (Salin & Bullier 1995), and given that pyramidal cells respond to synchronization over the relevant timescales (König et al. 1996), we predict that the outputs of V1 cells sensitive to different elements within the figure will tend to become more synchronized by the common top-down inputs that they receive.

An important difference between **Grossberg**'s perspective and the one we outlined is that in his models top-down and lateral connections have very different roles, with only the top-down being analogous to our CFs. He states his conviction that the function of horizontal intraregional connections is to complete missing information, as over blind spots and retinal veins, but cites no evidence that convinces us of this. It is hard to see how it could be the main part of the story, because lateral intraregional connections in V1 predominantly link cells with normal retinal input.

R1.8. What is the distribution of CF inputs?

Given the usefulness of distinguishing CFs from RFs, **Eckhorn** and **Morasso et al.** note that a major issue that arises is to map the distribution of facilitatory and suppressive CF inputs. We agree. Much relevant data concerning their distribution in V1 already exists (e.g., Kastner et al. 1997; Levitt & Lund 1996; Polat & Norcia 1996; Schmidt et al. 1997; Weliky et al. 1995; Zipser et al. 1996), but even there important issues await further clarification. In other regions there is much new territory to be explored, as Morasso et al. emphasize.

CFs are local in that each processor has its own particular set of direct CF inputs. All-to-all CF connectivity is neither possible nor desirable (**Wang**). If CF influences can be mediated through other processors, however, then each processor could be influenced indirectly by a much larger set. **Floreano**'s simulation of mediated long-range contextual influences clearly demonstrates this possibility. He shows how activity in distant parts of the network can influence RF learning given a chain of intermediate CF connections. An important constraint on this possibility, however, is that the mediating processors in the chain must themselves simultaneously receive RF input. This is in striking agreement with the capabilities and constraints observed by Polat and Sagi (1994b) in their psychophysical studies of perceptual learning in human vision.

R1.9. What synaptic receptor channels do CFs use?

Doubts are expressed by **Niebur & Usher** about whether NMDA-receptor channels can mediate synchrony at shorttime scales. NMDA channels have long been thought to have time constants of more than 150 msec, but it is now known that there are also fast components to the NMDA response with time constants closer to 15–50 msec (D'Angelo et al. 1990; Hestrin et al. 1990; Monyer et al. 1994). Although the fast components of the NMDA response are not well known, they have been used in models of STM (Jensen et al. 1996; Lisman & Idiart 1995). None of this implies that there are no other mechanisms for synchronization, and there is evidence for others (e.g., Cobb et al. 1995; Traub et al. 1996).¹

R2. Does the distinction between RFs and CFs (and/or synchronization) have psychological relevance?

In section 5 we argued that cognitive and neuropsychological evidence from a wide variety of paradigms supports our approach. Few of those arguments were questioned. Overall, commentaries did not subtract much from the psychological case presented, but, with one notable exception, neither did they add much. This suggests that a great deal remains to be done to foster facilitatory interactions between psychological and neurobiological studies of contextual coordination. The encouraging exception is provided by **Silverstein & Schenkel.** They have long used a variety of cognitive psychological paradigms to study perceptual grouping and other forms of contextual coordination in schizophrenia and conclude that it is selectively impaired in a distinct subset of patients. If so, studies of such states could provide new insights into the broader psychological significance of contextual coordination and of its pharmacological bases. Their suggestion will be considered closely in section R7.4.

R2.1. What is the role of top-down contextual control in cognition?

Internal sources of RF drive are dominant over external sources, Tononi & Edelman argue. This contrasts with the greater emphasis in the target article upon stimulus-driven effects. We agree that this is a major issue, and that the contextual modulation provided by strategic control may have much import for psychology. Perhaps the balance between internal and external sources varies, with the balance shifting too far toward external sources in some cases (e.g., attention deficit hyperactivity disorder, frontal lobe damage), and too far toward internal sources in others (e.g., psychotic delusions, dream states). They also argue for the importance of global context as provided by diffuse ascending systems. We agree, but emphasize that they do not carry semantic information and differ anatomically, pharmacologically, and functionally from mechanisms for local contextual coordination.

König et al. also emphasize the importance of internal strategic control variables, and show that they are amenable to combined electrophysiological and behavioral study. Their results indicate that such control uses similar CF synchronizing mechanisms as do stimulus-driven effects, and thus suggest promising new directions for future research. Section 1.4 in the target article notes several prior theories with similarities to our approach, including that of Grossberg. We did not specifically relate his work to these issues, however, and he has now done that better than we could have done. Instead, we put more emphasis upon relations to other theories that are closer to ours in spirit, and to the neurobiology in detail. Nevertheless, it is encouraging that Grossberg also sees important roles for synchronization and contextual modulation, for example in the topdown matching process of ART. Bugmann also makes a good case for top-down task-dependent contextual effects. We agree, but note that this does not imply the absence of bottom-up stimulus-dependent contextual effects. We await with interest for further developments in the use of CFs in computational models of strategic control.

R2.2. Are there modulatory interactions in word perception?

The relevance of contextual modulation to word perception is called into question by **Niebur & Usher** on the grounds that the effects of context and target are independent until additively combined at the final stage of response selection (Massaro 1989a; sect. R7.1.1). There are four replies to these doubts. First, Movellan and McClelland (1995) show how the data suggesting independence is compatible with interactions between streams even in the case where those interactions are mediated by driving excitatory and inhibitory connections. Second, even Massaro acknowledges an asymmetry in the effects of context and target, because it is only the target that determines the alternatives between

which choice is made. Third, Samuel (1996) shows that there are conditions where the signal detection measures used as evidence for independence do show effects of context upon signal detectability, even though these effects are small and fragile. Fourth, and most importantly, measures of signal detection are not equivalent to measures of information transmission. Changes in bias can have large effects on information transmission, so other measures are needed to test for the interactions that we propose. The appropriate conditional mutual information measures are presented in Smyth et al. (1996). We do not know of any studies applying these measures to context effects in word perception, but possible paradigms for doing so were outlined in section 5.7.2.

Iacoboni summarizes data showing that interhemispheric priming effects are mediated by the corpus callosum, as most theories would predict. This could reflect interhemispheric contextual modulation, but more direct evidence is required. In particular, measures that distinguish modulatory from driving effects (e.g., Smyth et al. 1996) would have to be used in order to show that the interhemispheric priming is indeed modulatory. Until then, a prima facie case for the hypothesis can be made on the grounds that response selection is determined by the target, with cross-hemispheric context predominantly influencing the RT.

R.3. Does the cortex use synchronized population codes?

R3.1. The doubts of Amit and Treves

Both Treves and Amit doubt that synchronization is used. This question can only be resolved experimentally. Spurious synchronization in Amit's models in no way detracts from the many observations of stimulus-dependent synchronization in real animals. Treves's use of information measures may well help analyze this data, but it is not just the transmission of information in general that must be measured, but information about the particular stimulus relations that affect synchronization. Stimulation paradigms have to be used that require response selection and grouping, and evidence has to be obtained that the result of these operations is signaled by changes in synchronization rather than by changes in discharge rate. Such evidence is available. Whether neurons with spatially segregated RFs are activated with a single continuous contour or with two different stimuli is reflected by the degree of synchronization rather than by rate changes in the cat retina and LGN (Neuenschwander et al. 1996), cat area 17 (Engel et al. 1991; Freiwald et al. 1995; Gray et al. 1989), for groups of neurons distributed across area 17 of the two hemispheres (Engel et al. 1991; Munk et al. 1996), or across areas 17 and PMLS (Engel et al. 1991), in area MT of awake monkeys (Kreiter & Singer 1996), and in primary auditory cortex (deCharms & Merzenich 1996). In amblyopia, a developmental disorder associated with deficits in contextual grouping, the only detectable abnormality in primary visual cortex was drastically reduced synchronization among responses conveyed by the amblyopic eye. There were no differences in discharge rates between neurons driven by the normal or the amblyopic eye even when these responses were evoked by gratings whose spatial frequency was so high that they could no longer be resolved by the cats when

R3.2. Should external (stimulus-locked) and internal synchronization be distinguished?

Stimulus-locked and internal synchronization are distinguished by Tononi & Edelman, Eckhorn, and Wright. We agree. In both cases synchronization is exploited for binding. Psychophysical evidence suggests that stimuluslocked synchronization is used for figure-ground segmentation if figure elements are presented with temporal offset relative to background elements. The temporal precision of this segmentation is better than 8 msec and the temporal information is provided by the magnocellular pathway (Leonards et al. 1996; Leonards & Singer 1997; but see Kiper et al. 1996 for a different result). This grouping by external timing can be overridden by internal grouping, however. If the figure is defined by textual coherence of its elements, detection is not impeded by false temporal conjunctions. If two different figures are defined, one by temporal cues and the other by texture cues, there is rivalry and the figure defined by the more salient cues is perceived.

R3.3. What are the time-scales of synchronization?

Questions concerning the relevant time scales of synchronization are raised by **Palm & Wennekers** and **Nunez**. One of the main advantages of synchronization is that it can operate on a much faster time scale than selecting responses by increasing their rate – provided that it is possible to synchronize responses rapidly. If responses are selected by rate increases, a cell has to integrate over a sufficiently large number of incoming EPSPs before emitting a response in order to assume that the response is actually generated by the selected input and not by accidentally arriving EPSPs or nonselected inputs. The required duration of this safety interval depends on average firing level and can be set by modulating the membrane potential (i.e., the distance to threshold and the membrane time constants). Assuming response frequencies of cortical neurons in the range of 50 Hz and postulating summation over at least 4-5 EPSPs before reacing a decision amounts to transmission times per processing step of at least 80 to 100 msec. This is far too slow. Input selection by spike synchronization is much faster because it does not rely on temporal summation. Coincident EPSPs can reach firing level within a few msec, thus reducing transmission times to near the synaptic delay. The speed of selection is thus primarily constrained by the time required to obtain synchrony. Contrary to earlier models, which generated synchrony by coupling harmonic oscillators, more recent models based on spiking neurons (i.e., relaxation "oscillators") indicate that synchrony can be obtained very rapidly (Wang; Somers & Kopell 1993). This is supported by experimental evidence. In the retina, responses to coherent stimuli are

In the cortex, it is much more difficult to determine experimentally the time required for the alignment of spikes by cross-correlation analysis because the windows must have a minimal duration to obtain sufficient entries, and because response onset is not well defined when moving stimuli are used. The data suggest that newly induced oscillatory responses synchronize right away once oscillatory patterning develops (Singer & Gray 1995). As cortical activity has a complex oscillatory structure (Arieli et al. 1996) and as oscillatory modulation of the membrane potential shifts spike timing, feature specific synchronization may occur very rapidly. The rate at which different assemblies can be organized successively without merging depends critically on the integration time constants of cortical neurons. Here views diverge, ranging from estimate of less than a millisecond (König et al. 1995; Softky 1995) to tens of milliseconds (Shadlen & Newsome 1994). Data on echo location and spatial location of sound sources show that neurons are capable of very precise conicidence detection (Yan & Suga 1996). Whether this is also true in the cortex remains to be seen, but recent evidence on action potentials backpropagating into dendrites (Stuart & Sakmann 1994) and other regenerative events in dendrites (Connors & Gutnick 1990) may lead to a drastic change in views on the timescale of dendritic integration.

R3.4. What is the role of oscillation?

Questions are raised by McCollum, and Haase & Diniz concerning the relation between oscillation and synchronization. Experimental observations indicate a close relation between oscillation and the precision of synchrony (König et al. 1995). Also, there is evidence that cortical neurons engage in synchronous oscillations in the beta and gamma frequency range across visual, association, somatosensory, and motor areas when the animal focuses its attention in the preparation of a visuomotor reaction; this synchronization increases in a task-specific way during execution of the task but collapses upon completion (Roelfsema et al. 1997). This suggests that oscillatory modulation helps synchronize responses, facilitating handshaking among neurons that need to be bound together for the execution of the task. Whether this frame-setting oscillatory modulation of membrane potential is due to oscillatory input from inhibitory or excitatory sources is not crucial for the principle and requires experimental testing.

Oscillatory patterning may be advantageous for rapid synchronization of discharges because it allows for temporal shifting of responses to synaptic input in both directions. The duration of the possible shift intervals is determined by oscillation frequency, and in the range of about a half-cycle. An assembly defined by synchronized discharges need not necessarily extend over many oscillation cycles. In some cases it may be sufficient to organize a single volley of synchronized discharges and have a new assembly at the next processing stage as proposed by Abeles (1991). In that case different assemblies can be organized on successive cycles. However, oscillatory patterning can also serve to generate the same synchronous volleys over several cycles, thereby refreshing assemblies that need to be stabilized for a while. The advantage over stabilizing assemblies by simple rate increases would be that it reduces the possibility of getting false conjunctions if several assemblies need to be maintained within the same array of neurons, as false bindings are avoided if unrelated assemblies do not fire in synchrony. This does not require fixed phase shifts, as it can be achieved by interactions that modulate the regularity of the respective oscillations.

R4. What forms of learning occur within the cortex?

R4.1. Can the abstract learning rules discover higher-order variables?

There is a recognition by **Stone** of the potential of using context to guide learning to variables that are statistically related to variables of significance to the organism, including, but not limited to, reinforcement. We agree that this can include temporal as well as spatial structure. The algorithms that use temporal constraints to discover statistical structure (e.g., Becker 1996; de Sa & Ballard 1997; Stone 1996a; 1996b) are sufficiently similar to those outlined in section 3 that ways in which the two constraints can be used together merit further study.

Floreano shows that the learning rules outlined in section 3.3 can discover nonlinear transforms. The example used is stereo depth. He studies the possibility that this is guided by contextual input from other cues to depth. As there are usually multiple cues it may be that those that are either genetically specified or more easily learned guide the discovery of others. Floreano assumes that different depth cues are processed in different streams, and this is also plausible because different cues are not always in agreement (e.g., depth of a picture plane versus depth in the picture). The success of Floreano's simulation in sometimes discovering stereo depth using such an internal teacher is therefore encouraging. It did not do so reliably, however, and although his suggestion as to why that was so may be part of the story, the capabilities and limitations of this whole class of learning algorithms remain unclear.

R4.2. Does unsupervised learning in the cortex discover higher-order variables?

In section 6.4 we asked whether there is any evidence that self-organization in the cortex can discover nonlinear variables such as XOR. No such evidence was offered in the commentaries, nor have we yet found any from other sources. The continued failure of such evidence to appear suggests that reliable discovery of such nonlinear variables may not be a fundamental capability of cortex.

R4.3. Is there a floating threshold for LTP, and if so what are its determinants?

Sections 3.3 and 6.4 note converging evidence for learning rules of the BCM/ABS type (Artola & Singer 1993; Bienenstock et al. 1982; Hancock et al. 1991a). A central feature of these learning rules is a non-monotonic dependence of synaptic change upon post-synaptic activity with a threshold (that may be movable) below which synapses are weakened (LTD) and above which they are strengthened (LTP). **Grossberg** notes that some of his learning rules have a similar form, and **Stone** also notes their importance. A major goal for the neurobiology of learning is therefore to

find out whether the threshold for LTP does move and if so what it depends upon.²

R4.4. Does SMA modulate sensorimotor learning in PMdc?

Neuroimaging evidence is outlined by **Iacoboni** showing that activity in the supplementary motor area (SMA) provides a contextual input that guides the learning of a sensorimotor mapping from the rostral sector of dorsal premotor cortex (PMdr) to the caudal sector of dorsal premotor cortex (PMdc). We agree that this may be so but other explanations of the results are possible. One way forward may be to show that activity in both PMdr and PMdc is necessary for learning, but that activity in SMA is not, even though it has a guiding effect when present. Experiments of this sort are now under way (Iacoboni, personal communication).

R5. To what extent do these processes provide a representation of the external world?

One of our central suggestions is that local processors could lay foundations for representation and meaning by discovering commonalities across diverse data sets. This is because CFs provide a means whereby local processors can select the relevant information. Morasso et al., Stone, Tononi & Edelman, and Wright all supported this suggestion and no one argued against it. Amit, Eckhorn, and McCollum all overlooked this suggestion, however, calling for a notion of relevance as though we had not addressed the issue. We must thus reiterate our suggestion, which is essentially a simple one. If local processors receive only RF input then we can see no way in which they could distinguish the relevant from the irrelevant. If they also receive CF input, then they can do so by, for example, selectively transmitting only the RF information that is statistically related to the context. Thus, in clear contrast to what McCollum assumes, this does involve information loss, and crucially so.

In section 6.5 we asked whether the processors we propose could help form proper intentional representations of the external world. We suggested that while they might contribute they could not be sufficient. We have so far not been very precise as to what "external input" means. In relation to the simulations it simply means external to the network being simulated. For most local processors in the cortex this is analogous to input from another cortical subsystem, and not to input from a world external to the system as a whole. To make this distinction the system must have some notion of "itself." **Wright**'s insightful suggestion that the ability to discover coherence in multiple sources of activity could apply to sources from within the organism may be relevant here, but philosophical clarification of the issues involved would also be helpful.

R6. Additional computational foundations suggested by commentators

We proposed functional specialization and contextual coordination as two fundamental and mutually constraining foundations of cortical computation, expecting thereby to provoke proposals of additional or alternative foundations. Few others were forthcoming. **Grossberg** refers us to his neural network models. **Tononi & Edelman** suggest a perspective from which the predominant information flow is between sources internal to the cortex, with the main role of external input being to modulate that flow. We agree that the distinction between internal and external sources is important, and that the balance between them is a topic of fundamental importance (sect. R2.1). It is clear that at the level of local circuits Tononi & Edelman emphasize just the same two fundamental principles as we do.

In keeping with an emphasis upon internal sources, Wright notes that local processors may be organized so that they are autonomously active and with an intrinsic dynamic. In this case they cannot be adequately described only in terms of a transfer function. We agree that in this case such sources should be seen as being included within the total set of primary RF inputs, and that developments of the kind he suggests are well worthwhile. Haase & Diniz and Stone note that structure in time is as important as structure in space, so an ability to process temporal sequence is also crucial. This seems compatible with our emphasis upon both coherence and synchrony. Indeed, as Haase & Diniz note, precision in temporal sequencing may be well served by the use of synchronized population codes (Abeles 1991). Overall, functional specialization and contextual coordination still appear most prominent as fundamental principles of cortical computation.

R7. Further issues raised by commentators

R7.1. What mathematical formalisms are useful?

We take it for granted that if there are common foundations then it must be possible to describe them in abstract terms that are independent of any specific content.

R7.1.1. Is information theory useful? We can ask, echoing Amit's questions concerning synchronization "Is information theory necessary to an understanding of cortical function?" and "Is it sufficient?" To both we answer with an unequivocal "No." McCollum identified our methods with information theory and neural network simulations. Given our backgrounds in experimental psychology and neurophysiology, and the allocation of only 20% of the target article to computational studies we do not see that as being accurate. Nevertheless, it is justified to ask "Is information theory useful to an understanding of cortical function?" and "Are other formalisms useful?" To both we answer with an unequivocal "Yes." Of the commentators, Stone, Tononi & Edelman, Treves, and Wright also find information theory useful, but Gregson, Eckhorn, Nunez, and Mc-Collum do not. These differences may be due to a misunderstanding of the uses proposed for information theory, so we will try again to make them clear.

Three uses are exemplified in the target article and commentaries. First, there is its use for job specification, that is, to provide a conception of the essential information processing operations performed by local cortical circuits. In clear contrast to **Grossberg**'s perception of the target article, its central concern is with general principles. In using information theory to formulate a conception of the goals of cortical computation we are in a long (e.g., Attneave 1954; Barlow 1959) and still strong (e.g., Intrator & Cooper 1995a; Olshausen & Field 1996) tradition. The goals of information compression, data reduction, and sparsification are relevant to the concerns rightly raised by **Gregson** and by **Eckhorn** in relation to metabolic constraints. In accord with the work of **Stone** and of **Tononi & Edelman**, a central aim of the target article is to extend these conceptions to include the challenging task of information *selection* (sects. 1.2.1, R5).

Second, and also in accord with the work of **Stone**, we have used the information theoretic job specification to derive a learning algorithm that changes connection strengths to better approximate the goal, given the statistical structure within the input received. Our description of the goal as maximizing the objective function specified (sect. 3.2) may mislead some commentators, for example, Amit. There is no reason to suppose that in any realistic situation perfect optimization is either necessary or possible. All that is required for the learning to be useful is that it more closely approximates the system to such a state. Although it may not be necessary to use information theory to understand learning, should synaptic plasticity be found to have a dynamic threshold for LTP that moves as specified by one of the abstractly derived learning rules (sect. R4.3), then many will wish to understand better how that rule was derived.

Third, information theory can be used for data analysis, for example, to measure information transmitted, as emphasized by **Treves.** We agree that such measures are useful (sect. R3.1). In addition, we use information theory to tackle a problem that no commentator discussed, that is, to provide an adequate specification of what should count as "context." There was agreement that context is important, but this means little unless we can give an adequate account of what can count as context.

Consider a psychophysical experiment in which there are two cues to texture segregation and that both influence performance, although subjects are asked to base their judgments on just one of them. We could call the specified cue the target, and the other the "context," thereby hoping to provide evidence for the relevance of context. Though common, this conception of context is wholly inadequate. On that conception "target" and "context" are distinguished simply by what experimenter and subject call them. It may be that both cues contribute in essentially the same way to the observed responses, so that dividing them into target and context is arbitrary.

This is exactly what is implied by Massaro's (1989a) fuzzy logic model of perception. Perceptual channels are independent until they all contribute in essentially the same way to response decision. In such a system there is no need to distinguish target from context. Either an input variable contributes to response or it does not. The same can also apply to the cellular level; either an input effects output or it does not. If it does, it is part of the cell's RF, and if not, it isn't. From this viewpoint there is no need to distinguish RF influences from CF influences (Gilbert, personal communication). Thus, in contrast to what Amit assumes, the case for "context" does have to be made. Our use of information theory to specify what can count as context is developed further in Smyth et al. (1996), and is being used at Stirling to analyze the interaction of cues to texture segregation by human subjects (thus addressing Gregson's doubts about the empirical validity of the transfer function used in the computational theory). We hope to put the study of contextual coordination on a sounder basis than is provided either by showing that variables other than those that the experimenter calls the target can affect subjects' responses, or by showing that cellular activity is affected by more than what the experimenter calls the classical receptive field.

R7.1.2. What other formalisms are useful? We do not expect to squeeze all of cortical function into a single mathematical formalism. We agree with **Gregson**, **Nunez**, **McCollum**, and **Morasso et al.** that others are needed, and non-linear dynamic systems analysis in particular. They complement rather than supplant the uses just listed for information theory, however. The challenge for such formalisms is to give rise to revealing experimental paradigms and to make their findings more comprehensible than they would otherwise have been. Having tried to do that for what are perhaps some of the more basic and intrinsically comprehensible aspects of cortical function, we know how big a challenge that is.

R7.2. What is the role of modeling?

None of the uses listed in section R7.1.1 for information theory can be described as "modeling." Simulations are outlined in section 3.4 but they are not models of biological systems. As Bower indicates, they are "proof of concept" simulations, being designed to explore the abstract goals, transfer functions, and learning rules. Würtz also understood this, and thought our focus on simple computational concepts a distinct advantage. Nevertheless, Palm & Wennekers's attempt to interpret the simulations as models of spiking neurons, and then criticize them for not being so. Their attempt to interpret the simple demonstration whose results are shown in Figure 10 of the target article shows that they misunderstood it in several ways.³ Palm & Wennekers ask questions that are not pertinent to any of the uses made of the computational theory, but they could be asked of a biological model, or more pertinently still of the biological system itself (sect. R3.3).

Bower continues to argue for the relevance of detailed biological models. We agree that they are useful, but their limitations as a primary focus for research are well revealed by Bower's commentary. Necessary responses to the question posed in his title are "What details?" and "To what is their relevance being assessed?" If they matter, they are not details. If we were trying to discover how birds stay up in the air it would be counterproductive to try to include all details of bird structure and physiology. The "details" that matter are those essential to aerodynamic lift. Bower implies that the details that matter in the case of cortical computation are those in his models. Grossberg also describes his models as being biologically realistic, but they are not the same as Bower's and are used to support different conclusions. The most convincing way to test the relevance of a theory to biology is to test it on real organisms, not on models that try to replicate them. We therefore direct our efforts towards seeking a conceptual understanding of cortical computation that can be tested and developed through investigations of real brains and real people, as exemplified here by König et al. and by Silverstein & Schenkel. As the role of computational theory is still so widely and deeply misunderstood Table R2 compares it

Table R2. Relations between theory and biology

	Bird flight	Cortical computation
Goal ¹	Stay up	Coordinate activity ²
Strategy	Use aerodynamic lift	Use contextual information
Formalization	Aerodynamic equa- tions for lift, drag, moment, etc.	Transfer functions and information theoretic objective functions, etc.
Mechanism	Wings ³	Contextual fields ³
Testing the strategy	Wings in wind tunnels ^{4,5}	Neural net "simula- tions" ^{4,5}
Biological testing	Do bird's wings have the required aerofoil section?	Do cortical cells re- ceive contextual input?
	Is air velocity adequate?	Does it affect output as required?

1. Only one goal is considered in each case. Both bird flight and cortical computation have other goals but separate goals are best considered separately.

2. The goals of cortical computation are far less obvious than those of flight. Discovering what they are is part of the problem we have to solve.

3. Wings and CFs are neither necessary nor sufficient for their goals.

4. The theories are best tested using versions of the mechanism that are as simple as possible yet consistent with achieving the goal in accordance with the theory.

5. The mechanisms tested are real. The wing in the wind tunnel generates lift. The "simulated" net computes. Thinking of them as models of something else does not contribute anything to their role in testing the strategy.

with the role of aerodynamics in the study of flight, in the hope that that will clarify our views on this issue.

R7.3. What is the role of CFs and/or synchronization in motor control?

The possible relevance of CFs and synchronization to motor control is discussed by Iacoboni, McCollum, and Morasso et al. The selection of particular motor responses raises combinatorial problems similar to those raised by scene segmentation. The number of different constellations of muscle contractions needed for the execution of different movements is very large. As the same muscles are used in different combinations for different movements, the commands for their contraction have to be bound in ever different constellations. Representing every possible movement by a command unit that distributes its output to the appropriate constellation of motor neurons leads to the same combinatorial explosion as representing every distinguishable object by a pontifical cell. The analogy with scene segmentation and object representation is obvious if one equates motor units with elementary features and a motor program with an object. This predicts that one should observe oscillations and synchronization during the initial selection phases of motor programming and a sequence of successively structured synchronized assemblies of the type ABCD etc. during execution. The findings of Murthy and Fetz (1996) are compatible with such a view.

R7.4. Is contextual coordination deficient in schizophrenia?

Contextual coordination, Silverstein & Schenkel suggest, may be impaired in particular schizophrenic states. Prima facie, this has an intuitive appeal. If there are specialized mechanisms for contextual coordination, fragmentation will result from their impairment. If these mechanisms are widely distributed across cortex, disorganized perceptions, thoughts, and actions are all possible consequences. Many studies of cognition in schizophrenia can be seen as evidence for such disorganization. Silverstein & Schenkel's suggestion is further strengthened by evidence for underactivity of NMDA-receptor channels in schizophrenic states (Olney & Farber 1995) and for the psychotomimetic effects of ketamine (Krystal et al. 1994), which blocks NMDA-channels. This is in keeping with our hypothesis that contextual coordination is achieved via CFs that exert gain control, for example, via NMDA-channels. It thus becomes of paramount importance to understand the basic pharmacology of coordinating gain-control channels, because that may advance our understanding of schizophrenia in a way that directly links molecular mechanisms with their cognitive consequences.

Attempts to understand schizophrenia have been plagued by finding a task on which patients are impaired, and then building a grand explanatory model on that (Ronan O'Carroll, personal communication). A particular problem that arises here is that the "contextual coordination" that is impaired in schizophrenia may have little or nothing to do with the "contextual coordination" for which the target article presents evidence. We therefore first consider studies of perceptual grouping in schizophrenia, then note unresolved issues that arise.

Silverstein et al. (1996a) review many studies of perceptual grouping in schizophrenia beginning with those of Cox and Leventhal (1978) and Place and Gilmore (1980), and present further evidence using the Banks and Prinzmetal (1976) paradigm to study the effect of visual grouping on visual search. All these studies suggest that in severe schizophrenic states perception is more fragmented than normal, with reduced effects of processes that group the stimulus elements into larger units. The evidence suggests that both stimulus-induced and top-down attentional processes may be involved in this impairment. The impairment is indicative of schizophrenia (Knight 1992; Silverstein et al. 1992), and correlates with the severity of other psychotic symptoms (Silverstein et al. 1996c). Using Navon's (1977) terminology, these patients have a tendency to see the trees before the forest, whereas control subjects have a tendency to see the forest before the trees. The evidence cannot be dismissed as being due to a general impairment of performance for at least two reasons: (1) The altered processes involve a reversal in the relative difficulty of conditions as compared with controls (Silverstein et al. 1996a); (2) This patient group performs better in tasks where it is advantageous to see the trees, not the forest (Place & Gilmore 1980; Rabinowicz et al. 1996). These perceptual disorders thus fit well with those to be expected, given impairments to the processes of contextual coordination hypothesized in the target article, however several unresolved issues then arise.

1. This patient group shows no deficits in grouping under conditions where there are either strong stimulus cues to

grouping (Knight 1992) or a strong push to allocate attention to global properties (Silverstein et al. 1996a). The hypothesis must therefore be expressed in terms of a reduced *tendency* to use grouping, and this needs further conceptual clarification and empirical testing. Gestalt grouping processes may not be at all impaired, either because they are genetically specified, or because if they are acquired or refined by visual input then the input correlations upon which they depend are so overwhelmingly present in natural input that they are learned to asymptotic levels even by an impaired learning mechanism. The acquisition of new assemblies at higher levels is not based upon such strong statistical structure, however, so impairments in acquiring and using those may be associated with impaired binding and reduced synchronization both within and between regions.

2. As **Silverstein & Schenkel** note, the evidence suggests impairments of perceptual learning as well as of perceptual processing, and this is in keeping with other perspectives on this disorder (e.g., Gray et al. 1991). An unresolved issue that arises here is whether it is only CF learning (i.e., assembly formation) that is impaired and, if so, how this can be reconciled with the hypothesis that CFs guide RF learning.

3. Some of the top-down attentional effects on perceptual grouping that have been shown to be impaired in schizophrenia clearly involve episodic and working memory processes (Silverstein et al. 1996b). The framework as presented in the target article would therefore have to be extended to show how it is related to higher level processes such as attention and episodic and working memory, if it is to be applicable to such findings. The value of such an extension is also emphasized by **Bugmann, König et al.**, and **Tononi & Edelman** (sect. R3).

4. It is easy to see how an impairment in contextual coordination could produce symptoms of disorganization, but it may at first seem less easy to see how it could account for the positive symptoms, such as hearing voices. One approach would be to seek a separate explanation for these positive symptoms. Alternatively, we could speculate that they may reflect: (1) reduced modulatory control by external input of the flow of activity arising from internal thoughts; and (2) reduced knowledge of the source of internal speech and percepts because of reduced interregional binding.⁴ The pharmacological evidence encourages the search for an integrated explanation because drug-induced psychoses that reduce NMDA activity mimic both positive and negative symptoms (Krystal et al. 1994).

Silverstein & Schenkel's hypothesis thus survives close scrutiny well and, as they argue, it opens important directions for further work. One possibility may be to study the effect of drug-induced psychoses on perceptual learning and inter-regional synchrony using local field potentials. More broadly, this hypothesis should encourage physiological, psychological, and neuroimaging studies of contextual coordination in schizophrenic and drug-induced psychotic states using both human and animal subjects.

R8. Science as a search for coherence

It is suggested by **Treves** that if the commentaries "turn out not to be fully sychronized" (by which we assume he means not in full agreement) then they will contribute to the population response but by means other than synchronizing it. We agree that it may be both revealing and entertaining to draw an analogy between the outputs of local cortical processors and scientific writings. Each paper written tries to convey something that is both distinctive and relevant. Relevance does not imply agreement, however. Different members of a group can have opposing effects at a higher level of analysis. The primary code used by *BBS* to signal mutual relevance is simultaneous publication of target articles and associated commentaries, that is, grouping through synchronous transmission. This is not because synchrony is necessary to grouping, but because it makes it a lot easier for readers to process them as a whole.

This analogy is also useful for another reason. By seeing science as a continuation of the search for coherence, by means beyond the reach of any algorithm possessed by mammals in general, we are reminded that local processors should be seen as moving toward that goal, not as reaching it (sect. R7.1.1).

R9. Promising directions for future progress

Specialization of function has been the keynote of this century. Coordination of function may be the keynote of the next. This will require more attention to contextual interactions, synchronized population codes, and fast dynamics. We believe that future research must emphasize relational codes rather than focusing only upon response properties of single cells. The relevant relations can be detected only by analyzing simultaneously recorded activity. We shall have to look at the outputs of any processing level in the same comprehensive way as it is looked at by the subsequent processing stages to which it projects. This requires analysis of mutual dependencies among the responses of a large number of neurons, and hence the study of ensemble dynamics. There is thus a logical progression from early studies of the topology of brain functions, to the functional analysis of individual neurons located in different compartments, and now to the attempt to understand the spatio-temporal patterns of coordinated neuronal activity. In conclusion, then, our prediction is that the temporal organization and coordination of brain activity is as sophisticated and subtle as is its topological organization, the former being an emergent property of the latter.

ACKNOWLEDGMENTS

We thank Peter Hancock, Jim Kay, Steve Silverstein, Leslie Smith, and Darragh Smyth for comments on a draft of this response.

NOTES

1. Indeed, if NMDA receptor activation is combined with oscillatory modulation of the membrane potential, latency of responses is shifted as a function of oscillation phase and output phase is locked to input phase (Volgushev et al., submitted). This can synchronize discharges within a single oscillation cycle with high temporal precision because of the voltage-dependent gating of the MG^{2+} -block, which allows for very rapid activation and inactivation of NMDA-receptor mediated input.

2. Evidence is becoming available that it is not the absolute level of postsynaptic depolarization or the absolute level of the postsynaptic Ca^{2+} -surge alone that determines whether a particular synapse undergoes LTP or LTD. The actual state of the synapse, that is, the actual release probability with which it operates (Volgushev et al. 1997) and the rate with which it is activated during induction of the modification also matter (Hansel et al., in preparation). Thus synaptic changes are affected by information stored locally at each synapse, and this makes complex

learning rules such as those of Phillips et al. (1995b) and Kay et al. (in press) less implausible.

3. The simulations assume four levels of organization: cells, units, local processors, and layers. They operate at the level of units, but assume them to be composed of stochastic binary elements. The simulation shown in Figure 10 is intended to be analogous to the perception of an ambiguous figure, such as the Rubin vase, where alternative perceptions emerge successively, even though the external stimulus does not change. Thus, in contrast to what **Palm & Wennekers** assumed, the input cannot represent direct input from the external stimulus. Instead, the processing layer to which the simulation relates is assumed to be some layers distant from the external stimulus. Its inputs from the previous layer fluctuate because processing is stochastic. Palm & Wennekers suggest that the simulation in Figure 10 would work in the same way if the graded response units were replaced by single binary units. This is not so. If outputs are restricted to their extremal values, the CFs have no effect, in which case the outputs in at each iteration in Figure 10 would simply be identical to the inputs. We should therefore have made in clear in section 3.1 that the apparent insensitivity of performance to such a replacement refers only to the learning, with which most of the simulations were concerned. The two sets of 3 imes 3 units with internal coherence in the study of grouping shown in Figure 10 represent the two alternative percepts. CFs were therefore positive within and negative between these two alternative groupings. The background elements received no CF input. The first six iterations show the simultaneous emergence from the background of all elements in one of the two alternative groupings. Thus, contrary to what Palm & Wennekers conclude, this demonstrates the role of internal grouping processes, because the coherent sets of elements are distinguished from the background only by their CFs. To show the successive emergence of alternative percepts activity in the units was stopped and restarted after six simulations. The cortex is not short of mechanisms that would have such an effect, so there is no need for either it or a simulation to become stuck in the stationary state to which Palm & Wennekers refer. The choice of six iterations was arbitrary, but by varying it in an appropriate way periodic or other temporal structure could easily be introduced to the simulation. That would not change the effects with which it was concerned, however.

4. The latter possibility was suggested to us by Frith's (1992) interpretation of psychotic hallucinations as being due to a failure of self-monitoring. Words activated within the articulatory-loop can arise from either internal or external sources. Their source may therefore be unknown if they are not bound to activity in the other cortical modules from which they arise.

References

Letters "a" and "r" appearing before authors' initials refer to target article and response, respectively.

- Abeles, M. (1982) Local cortical circuits: Studies of brain function, vol 6. Springer-Verlag. [GP, aWAP, DLW]
- (1991) Corticonics: Neural circuits of cerebral cortex. Cambridge University Press. [arWAP]
- Abeles, M., Bergman, H., Margalit, E. & Vaadia, E. (1993b) Spatiotemporal firing patterns in the frontal cortex of behaving monkeys. *Journal of Neurophysiology* 70:1629–38. [aWAP]
- Abeles, M., Vaadia, E., Bergman, H., Prut, Y., Haalman, I. & Slovin, H. (1993a) Dynamics of neuronal interactions in the frontal cortex of behaving monkeys. *Concepts in Neuroscience* 4:131–58. [aWAP]

Able, K. P. & Bingham, V. P. (1987) The development of orientation and navigation behavior in birds. *Quarterly Review of Biology* 62:1– 29. [aWAP]

- Aertsen, A. M. H. J., Gerstein, G. L., Habib, M. K. & Palm, G. (1989) Dynamics of neuronal firing correlation: Modulation of "effective connectivity." *Journal* of Neurophysiology, vol. 61, 5:900–17. [GP]
- Allman, J., Miezen, F. & McGuinness, E. (1985) Stimulus specific responses from beyond the classical receptive field: Neurophysiological mechanisms

for local-global comparisons in visual neurons. Annual Review of Neuroscience 8:407-30. [aWAP]

- Amari, S. (1983) Field theory of self-organizing neural nets. *IEEE Transactions* on Systems, Man, and Cybernetics SMC-13:741–48. [PM]
- Amit, D. J. (1989) Modelling brain function: The world of attractor neural networks. Cambridge University Press. [RAMG, arWAP]
- (1995) The Hebbian paradigm reintegrated: Local reverberations as internal representations. *Behavioral and Brain Sciences* 18:617–57. [JJW]
- Amit, D. J. & Brunel, N. (submitted) Dynamics of a recurrent network of spiking neurons before and following learning. (See e.g. www.fiz.huji.ac.il/staff/acc/faculty/damita.) [DJA]
- Andrew, C. & Pfurtscheller, G. (1996) Event-related coherence as a tool for studying dynamic interaction of brain regions. *Electroencephalography and Clinical Neurophysiology* 98:144–48. [PLN]
- Arieli, A., Sterkin, A., Grinvald, A. & Aertsen, A. (1996) Dynamics of ongoing activity: Explanation of the large variability in evoked cortical responses. *Science* 273:1868–71. [rWAP]
- Armstrong-James, M., Welker, E. & Callahan, C. A (1993) The contribution of NMDA and non-NMDA receptors to fast and slow transmission of sensory information in the rat SI barrel cortex. *Journal of Neuroscience* 16:480– 87. [aWAP]
- Artola, A., Brocher, S. & Singer, W. (1990) Different voltage-dependent thresholds for the induction of long-term depression and long-term potentiation in slices of the rat visual cortex. *Nature* 347:69–72. [aWAP]
- Artola, A. & Singer, W. (1993) Long-term depression of of excitatory synaptic transmission and its relationship to long-term potentiation. *Trends in the Neurosciences* 15:218–26. [SG, arWAP]
- Ascher, P., Bregestrovski, P. & Nowak, L. (1988) N-methyl-D-aspartate activated channels of mouse central neurones in magnesium-free solutions. *Journal of Physiology* 339:207–26. [aWAP]
- Atick, J. J. (1992) Could information theory provide an ecological theory of sensory processing? *Network* 3:213–51. [aWAP]
- Atick, J. J. & Redlich, A. N. (1990) Predicting ganglion and simple cell receptive field organizations. *International Journal of Neural Systems* 1:305. [aWAP]
- (1993) Convergent algorithm for sensory receptive field development. *Neural Computation* 5:45–60. [aWAP]
- Attneave, F. (1954) Informational aspects of visual perception. Psychological Review 61:183–93. [arWAP]
- Baddeley, R. J. & Hancock. P. J. B. (1991) A statistical analysis of natural images matches psychophysically derived orientation tuning curves. *Proceedings of the Royal Society of London B* 246:219–23. [aWAP]
- Bair, W., Koch, C, Newsome, W., Britten, K. & Niebur, E. (1994) Power spectrum analysis of bursting cells in area MT in the behaving monkey. *Journal of Neuroscience* 14:870–92. [aWAP]
- Baldi, B. & Hornik, K. (1989) Neural networks and principal component analysis: Learning from examples without local minima. *Neural Networks* 2:5358. [aWAP]
- Banks, W. P. & Prinzmetal, W. (1976) Configural effects in visual information processing. *Perception & Psychophysics* 19:361–67. [rWAP]
- Barlow, H. (1996) Intraneuronal information-processing, directional selectivity and memory for spatiotemporal sequences. *Network: Computation in neural* systems 7(2):251–59. [JVS]
- Barlow, H. B. (1959) Sensory mechanisms, the reduction of redundancy, and intelligence. In: *The mechanisation of thought processes*. Her Majesty's Stationery Office. [arWAP]
 - (1961) Possible principles underlying the transformations of sensory messages. In *Sensory communication*, ed. W. A. Rosenblith. MIT Press. [aWAP]
 - (1972) Single units and sensation: A neuron doctrine for perceptual psychology? *Perception* 1:371–94. [aWAP]
- (1989) Unsupervised learning. *Neural Computation* 1:295–311. [aWAP]
 (1993) The biological role of neocortex. In: *Information processing in the cortex*, ed. A. Aertsen & V. Braitenberg. Springer-Verlag. [aWAP]
- Barlow, H. B. & Foldiak, P. (1989) Adaptation and decorrelation in the cortex. In: *The computing neuron*, ed. R. Durbin, C. Miall & G. Mitchison. Addison-Wesley. [RE, aWAP]
- Becker, S. (1993) Learning to categorize objects using temporal coherence. In: Advances in neural information processing systems 5. Morgan Kaufmann. [aWAP]
- (1996) Mutual information maximization: Models of cortical self-organization. *Network* 7:7–31. [arWAP, JVS]
- Becker, S. & Hinton, G. E. (1992). A self-organizing neural network that discovers surfaces in random-dot stereograms. *Nature* 355:161–63. [DF, aWAP]
- Bentin, S. (1992) Phonological awareness, reading, and reading acquisition: A survey and appraisal of current research. In: Orthography, phonology, morphology, and meaning, ed. R. Frost & L. Katz. Elsevier. [aWAP]
- Bentin, S., Hammer, R. & Cahan, S. (1991) The effects of aging and first grade

schooling on the development of phonological awareness. *Psychological Science* 2:271–74. [aWAP]

- Bernander, O., Douglas, R. J., Martin, K. A. C. & Koch, C. (1991) Synaptic background activity influences spatiotemporal integration in single pyramidal cells. *Proceedings of the National Academy of Sciences USA* 88:11569– 73. [aWAP]
- Bernander, O., Koch, C. & Usher, M. (1994) The effect of synchronized inputs at the single neuron level. *Neural Computation* 6:622-41. [DJA]
- Bernstein, N. (1967) *The co-ordination and regulation of movements.* Pergamon Press. [GM]
- Bezrukov, B. M. & Vodyanov, I. (1997) Stochastic resonance in non-dynamical systems without response thresholds. *Nature* 385:319–12. [rWAP]
- Bialek, W., Ricke, F., Vansteveninck, R. R. D. & Warland, D. (1991) Reading a neural code. Science 252:1854–57. [JMB]
- Bialek, W. & Rieke, F. (1992) Reliability of information transmission in spiking neurons. *Trends in Neurosciences* 15: 428–34. [VGH]
- Biederman, I. (1972) Perceiving real-world scenes. Science 177:77–80. [aWAP] (1987) Recognition-by-component: A theory of human image understanding. Psychological Review 94:115–47. [DLW]
- Bienenstock, E. (1995) A model of neocortex. Network 6:179-224. [aWAP]

Bienenstock, E., Cooper L. N. & Munro P. W. (1982) Theory for the development of neuron selectivity: Orientation specificity and binocular interaction in visual cortex. *Journal of Neuroscience* 2:32–48. [arWAP]

- Blakemore, C. & Tobin, E. A. (1972) Lateral inhibition between orientation detectors in the cat's visual cortex. *Experimental Brain Research* 15:439– 40. [aWAP]
- Bower, J. M. (1995) Reverse engineering the nervous system: An in vivo, in vitro, and in computo approach to understanding the mammalian olfactory system. In: An introduction to neural and electronic networks, second edition, ed. S. Zornetzer, J. Davis and C. Lau. Academic Press. [JMB]
- Braddick, O. (1993) Segmentation versus integration in visual motion processing. *Trends in the Neurosciences* 16:263–68. [rWAP]
- Braitenberg, V. (1978) Cortical architectonics: General and areal. In: Architectonics of the cerebral cortex, ed. M. A. B. Brazier & H. Petsch. Raven. [aWAP]
- Braitenberg, V. & Schüz, A. (1991) Anatomy of the cortex. Springer-Verlag. [aWAP]
- Bressler, S. L. (1995) Large scale cortical networks and cognition. Brain Research Reviews 20:288–304. [PLN]
- Brousse, O. & Smolensky, P. (1989) Virtual memories and massive generalization in connectionist combinatorial learning. In: *Proceedings of the 11th Annual Conference of the Cognitive Science Society*. Erlbaum. [aWAP]
- Bruce, V. (1988) Recognising faces. Erlbaum. [aWAP] Bruner, J. S. (1972) Beyond the information given. Norton. [GT]
- Bugmann, G. & Taylor, J. G. (1993) A model for latencies in the visual system. In: Proceedings of the third conference on artificial neural networks
- (ICANN'93, Amsterdam), ed. S. Gielen & B. Kappen. Springer-Verlag. [GB]
- (1994a) Role of short-term memory in neural information propagation. In: Extended abstract book of the International Symposium on Dynamics of Neural Processing, Washington. [GB]
- (1994b) A top-down model for neuronal synchronization. Research Report NRG-94-02, School of Computing, University of Plymouth. [GB]
- Bullier, J., Munk, M. H. J. & Nowak, L. G. (1992) Synchronization of neuronal firing in areas V1 and V2 of the monkey. *Society of Neuorsciences Abstracts* 18:11.7. [aWAP]
- Bullock, T. H., McClune, M. C., Achimowicz, J. Z., Iragui-Madoz, V. J., Duckrow, R. B. & Spencer, S. S. (1995) EEG coherence has structure in the millimeter domain: Subdural and hippocampal recordings from epileptic patients. *Electroencephalography and Clinical Neurophysiology* 95:161– 77. [PLN]
- Byrne, R. W. & Whiten, A. (1992) Cognitive evolution in primates: Evidence from tactical deception. *Man* 27:609–27. [aWAP]
- Calev, A., Venables, P. H. & Monk, A. F. (1983) Evidence for distinct verbal memory pathologies in severely and mildly disturbed schizophrenics. *Schizophrenia Bulletin* 9:247–64. [SMS]
- Campbell, R. N. & Olson, D. R. (1990) Children's thinking. In: Understanding children: Essays in honour of Margaret Donaldson, ed. R. Grieve & M. Hughes. Blackwell. [aWAP]
- Carpenter, G. A. (1989) Neural network models for pattern recognition and associative memory. *Neural Networks* 2:243–57. [aWAP]
- Carpenter, G. A. & Grossberg, S. (1991) Pattern recognition by self-organizing neural networks. MIT Press. [SG]
- (1993) Normal and amnesic learning, recognition, and memory by a neural model of cortico-hippocampal interactions. *Trends in Neurosciences* 16:131– 37. [SG]
- Castelfranco, A.M., Robertson, L. T. & McCollum, G. (1993) Detail, proportion,

and foci among face receptive fields of climbing fiber responses in the cat cerebellum. *Somatosensory and Motor Research* 11:27–46. [GM]

- Cattell, J. M. (1886) The time taken up by the cerebral operations. *Mind* 11:377–92. [aWAP]
- Chevalier-Skolinkoff, S. (1983) Sensori-motor development in orangutans and other primates. *Journal of Human Evolution* 12:545–46. [aWAP]
- Chiang, C., von Stein, A. & König, P. (1996) Synchronous activity between primary visual and sensorimotor cortex in awake behaving cat. Society for Neuroscience Abstracts 22:255.2. [PK]
- Cirelli, C., Pompeiano, M. & Tononi, G. (1996) Neuronal gene expression in the waking state: A role for the Locus Coeruleus. *Science* 274:1211–15. [GT]
- Clark, A. & Thornton, C. (1997) Trading spaces: Computation, representation and the limits of uninformed learning. *Behavioral and Brain Sciences* 20(1) 57–90. [aWAP]
- Clothiaux, E. E., Cooper, L. N. & Bear, M. F. (1991) Synaptic plasticity in visual cortex: Comparison of theory with experiment. *Journal of Neurophysiology* 66: 1785–1804. [aWAP]
- Cobb, S. R., Buhl, E. H., Halasy, K., Paulsen, O. & Somogyi, P. (1995) Synchronization of neuronal activity in hippocampus by individual GABAergic interneurons. *Nature* 378:75–78. [rWAP]
- Cohen, J. D. & Servan-Schreiber, D. (1992) Context, cortex, and dopamine: A connectionist approach to behavior and biology in schizophrenia. *Psychological Review* 99:45–77. [SMS]
- Connors, B. W. & Gutnick, M. J. (1990) Intrinsic firing patterns of diverse neocortical neurons. Trends in the Neurosciences 13:99–104. [rWAP]
- Cox, M. D. & Leventhal, D. N. (1978) A multivariate analysis and modification of a preattentive perceptual dysfunction in schizophrenia. *Journal of Nervous and Mental Disease* 166:709–18. [rWAP]

Creutzfeldt, O. D. (1978) The neocortical link: Thoughts on the generality of structure and function of the neocortex. In: Architectonics of the cerebral cortex, ed. M. A. B. Brazier & H. Petsche. Raven Press. [JVS]

- Crick, F. (1988) What mad pursuit. Penguin. [aWAP]
- (1989) The recent excitement about neural networks. *Nature* 337:129– 32. [DF]
- Crick, F. & Koch, C. (1990) Towards a neurobiological theory of consciousness. Seminars in the Neurosciences 2:263–75. [aWAP]

Damasio, A. R. (1989) The brain binds entities and events by multiregional activity from convergence zones. *Neural Computation* 1:123–32. [aWAP]

- D'Angelo, E., Rossi, P. & Garthwaite, J. (1990) Dual-component NMDA receptor currents at a single central synapse. *Nature* 346:467–69. [rWAP]
- Das, A. & Gilbert, C. D. (1995) Long-range horizontal connections and their role in cortical reorganization revealed by optical recording of cat primary visual cortex. *Nature* 375:780–84. [aWAP]
- deCharms, R. C. & Merzenich, M. M. (1996) Primary cortical representation of sounds by the coordination of action-potential timing. *Nature* 381:610– 13. [rWAP]
- De Loach, J. (1987) Rapid change in the symbolic functioning of very young children. Science 238:1556–57. [aWAP]
- Der, R. & Smyth, D. (in press) Local online learning of coherent information. Neural Networks. [aWAP]
- de Sa, V. (1994a) Unsupervised classification learning from cross-modal environmental structure. PhD thesis, University of Rochester, NY. [aWAP]
 (1994b) Learning classification with unlabeled data. In: Advances in neural information processing systems 6. Morgan Kaufmann. [aWAP]
- de Sa, V. & Ballard, D. H. (1997) Category learning through multi-modality sensing. *Neural Computation* (in press). [arWAP]
- Douglas, R. J. & Martin, K. A. C. (1990) Neocortex. In: *The synaptic organization of the brain*, ed. G. M. Shepherd. Oxford University Press. [aWAP]
- Douglas, R. J. & Martin, K. A. C. (1991a) A functional microcircuit for cat visual cortex. *Journal of Physiology* 440:735–69. [PK]
 - (1991b) Opening the grey box. *Trends in the Neurosciences* 14:286–93. [aWAP]
- (1994) The canonical microcircuit: A co-operative neuronal network for neocortex. In: *Structural and functional organization of the neocortex*, ed. B. Albowitz, K. Albus, U. Kuhnt, H.-Ch. Nothdurft & P. Wahle. Springer-Verlag. [JVS]
- Dudek, S. M. & Bear, M. F. (1992) Homosynaptic long-term depression in area CA1 of hippocampus and the effects on NMDA receptor blockade. *Proceedings of the National Academy of Sciences USA* 89:4363– 67. [aWAP]
- Durgin, F. H. (1995) Contingent aftereffects of texture density: Perceptual learning and contingency. PhD thesis, Department of Psychology, University of Virginia. [aWAP]
- Ebdon, M. (1993) Is the cerebral neocortex a uniform cognitive architecture? Mind and Language 8(3):369–403. [JVS]
- Eckhorn, H. (1997, in press) Neural mechanisms of scene segmentation. *Journal of Artificial Neural Networks*. [RE]

Eckhorn, R. (1994) Oscillatory and non-oscillatory synchronizations in the visual cortex and their possible roles in associations of visual features. *Progress in Brain Research* 102:405–26. [GP]

Eckhorn, R., Bauer, R., Jordan, W., Brosch, M., Kruse, W., Munk, M. & Reitboeck, H. J. (1988) Coherent oscillations: A mechanism of feature linking in the visual cortex? *Biological Cybernetics* 60:121–130. [RE, DLW]

Eckhorn, R., Dicke, P., Arndt, M., & Reirboeck, H. (1991a) Flexible linking of visual features by stimulus-related synchronizations of model neurons. In: *Induced rhythms in the brain*, ed. E. Basar & T. H. Bullock. Birkhauser. [aWAP]

Eckhorn, R., Frien, A., Bauer, R., Woelbern, T. & Kehr, H. (1993) High frequency (60–90 Hz) oscillations in primary visual cortex of awake monkey. *NeuroReport* 4:243–46. [RE]

Eckhorn, R. & Gruesser, O. -J., Kroeller, J., Pellnitz, K. & Poepel, B. (1976) Efficiency of different neural codes. *Biological Cybernetics* 22:40–60. [RE]

Eckhorn, R., Reitboeck, H. J., Arndt, M. & Dicke, P. (1990) Feature linking among distributed assemblies. *Neural Computation* 2:293-306. [RE]

Eckhorn, R., Schanze, T. Brosch, M., Salem, W. & Bauer, R (1991b) Stimulusspecific synchronizations in cat visual cortex: Multiple microelectrode and correlation studies from several cortical areas. In: *Induced rhythms in the brain*, ed. E. Basar & T. H. Bullock. Birkhauser. [aWAP]

Edelman, G. M. (1978) Group selection and phasic re-entrant signalling: A theory of higher brain function. In: *The mindful brain*, ed. G. M. Edelman & V. B. Mountcastle. MIT Press. [aWAP, GT]

(1987) Neural Darwinism: The theory of neuronal group selection. Basic Books. [PK, GT]

- (1989) The remembered present: Biological theory of consciousness. Basic Books. [aWAP]
- (1993) Neural Darwinism: Selection and re-entrant signalling in higher brain function. *Neuron* 10:1–20. [GT]
- Edelman, G. M. & Mountcastle, V. B. (1978) *The mindful brain.* MIT Press. [aWAP]
- Ellis, A. W. & Young, A. W. (1988) *Human cognitive neuropsychology.* Erlbaum. [aWAP]

Engel, A. K., König, P., Kreiter, A. K., Schillen, T. B. & Singer, W. (1992) Temporal coding in the visual cortex: New vistas on integration in the nervous system. *Trends in the Neurosciences* 15:218–26. [aWAP]

Engel, A. K., König, P. Kreiter, A. K. & Singer, W. (1991a) Interhemispheric synchronization of oscillatory neuronal responses in cat visual cortex. *Science* 252:1177–79. [MI, arWAP]

Engel, A. K., König, P. & Singer, W. (1991b) Direct physiological evidence for scene segmentation by temporal coding. *Proceedings of the National Academy of Sciences USA* 88:9136–40. [GB, arWAP]

Engel, A. K., Kreiter, A. K., König, P. & Singer, W. (1991c) Synchronization of oscillatory neuronal response between striate and extrastriate visual cortical areas of the cat. *Proceedings of the National Academy of Sciences USA* 88:6048–52. [arWAP]

Felleman, D. J. & Van Essen, D. C. (1991) Distributed hierarchical processing in the primate cerebral cortex. *Cerebral Cortex* 1:1–47. [aWAP]

Field, D. J., Hayes, A. & Hess, R. F. (1993) Contour integration by the human visual system: Evidence for a local "association" field. *Vision Research* 33:173–79. [SG, arWAP]

Finkel, L. H. & Edelman, G. M. (1989) The integration of distributed cortical systems by reentry: A computer simulation of interactive functionally segregated visual areas. *Journal of Neuroscience* 9:3188–208. [aWAP]

Fishman, M. C. & Michael, C. R. (1973) Integration of auditory information in the cat's visual cortex. *Vision Research* 13:1415–19. [aWAP]

Floreano, D., Phillips, W. A. & Kay J. (1995) A computational theory of learning visual features via contextual guidance. *Perception* 24(suppl):22. [aWAP]

Fodor, J. A. & Pylyshyn, Z. W. (1988) Connectionism and cognitive architecture: A critical analysis. *Cognition* 28:3–71. [aWAP]

Foldiak, P. (1990) Forming sparse representations by local anti-Hebbian learning. Biological Cybernetics 64:165–70. [aWAP]

(1991) Learning invariance from transformation sequences. *Neural Computation* 3(2):194–200. [JVS]

Fox, K. & Daw, N. (1992) A model for the action of NMDA conductances in the visual cortex. *Neural Computation* 4:59–83. [aWAP]

- Fox, K., Sato, H. & Daw, N. (1990) The effect of varying stimulus intensity on NMDA-receptor activity in cat visual cortex. *Journal of Neurophysiology* 64:1413–28. [aWAP]
- Francis, G., Grossberg, S. & Mingolla, E. (1994) Cortical dynamics of feature binding and reset: Control of visual persistence. *Vision Research* 34:1089– 1104. [SG]

Freiwald, W. A., Kreiter, A. K. & Singer, W. (1995) Stimulus dependent intercolumnar synchronization of single unit responses in cat area 17. *NeuroReport* 6:2348–52. [rWAP]

Frien, A., Eckhorn, R., Bauer, R., Woelbern, T. & Kehr, H. (1994) Stimulus-

specific fast oscillations at zero phase between visual areas V1 and V2 of awake monkey. *NeuroReport* 5:2273–77. [RE]

- Frien, A., Eckhorn, R. & Reitboeck, H. J. (1996) Fast oscillations in V1 of awake monkey. Society for Neuroscience Abstracts 22:255.5. [RE]
- Fries, P., Roelfsma, P. R., Engel, A. K., König, P. & Singer, W. (1996) Synchronized gamma frequency oscillations correlate with perception during binocular rivalry in awake squinting cats. *Society of Neuroscience Abstracts* 22(1):117.3 [arWAP]

Friston, K. J., Tononi, G., Reeke, G. N., Jr., Sporns, O. & Edelman, G. M. (1994) Value-dependent selection in the brain: Simulation in a synthetic neural model. *Neuroscience* 59:229–43. [GT]

Frith, C. D. (1992) The Cognitive neuropsychology of schizophrenia. Erlbaum. [rWAP]

Gallistel, C. R. (1995) The replacement of general-purpose theories with adaptive specializations. In: *The cognitive neurosciences*, ed. M. S. Gazzaniga. MIT Press. [aWAP]

- Geman, S., Bienenstock, E. & Doursat, R. (1992) Neural networks and the bias/variance dilemna. *Neural Computation* 4:1-58. [GT]
- Georgopoulos, A. P. (1990) Neural coding of the direction of reaching and a comparison with saccadic eye-movements. *Cold Spring Harbour Symposium on Quantitative Biology* 55:849–59. [aWAP]
- Gerstner, W. (1995) Time structure of the activity in neural network models. *Physical Review E* 51(6):738–58. [GP]
- Ghose, G. M. & Freeman, R. D. (1992) Oscillatory discharge in the visual system: Does it have a functional role? *Journal of Neurophysiology* 68:1558– 74. [VGH]
- Giffins, R. (1985) Canonical analysis: A review with applications in ecology (Biomathematics 12). Springer-Verlag. [aWAP]
- Gilbert, C. D. (1992) Horizontal integration and cortical dynamics. *Neuron* 9:1– 13. [aWAP]
- (1995) Dynamic properties of adult visual cortex. In: *The cognitive neurosciences*, ed. M. S. Gazzaniga. MIT Press. [aWAP]
- Gilbert, C. D. & Wiesel, T. N. (1983) Clustered intrinsic connections in cat visual cortex. *Journal of Neuroscience* 3:1116–33. [aWAP]

(1989) Columnar specificity of intrinsic horizontal and cortico-cortical connections in cat visual cortex. *Journal of Neuroscience* 9:2432– 42. [aWAP]

(1990) The influence of contextual stimuli on the orientation selectivity of cells in primary visual cortex of the cat. *Vision Research* 11:1689–1701. [aWAP]

- Gluck, M. A. & Rumelhart, D. E. (1990) Neuroscience and connectionist theory. Erlbaum. [aWAP]
- Goebel, R. (1993) Perceiving complex visual scenes: An oscillator neural network model that integrates selective attention, perceptual organisation, and invariant recognition. In: Advances in neural information processing systems 5, ed. S. J. Hanson, J. D. Cowan & C. L. Giles. Morgan Kaufmann. [aWAP]
- Golomb, D., Hertz, J., Panzeri, S., Treves, A. & Richmond, B. J. (1997) How well can we estimate mutual information from limited samples of neuronal responses? *Neural Computation* 9:649–65. [AT]
- Goodall, W. C. (1994) Neuropsychological studies of reading and writing. PhD thesis, University of Stirling, Scotland, UK. [aWAP]
- Goodall, W. C. & Phillips, W. A. (1994) Three routes from print to sound: Evidence from a case of acquired dyslexia. *Cognitive Neuropsychology* 12:113–47. [aWAP]
- Gove, A., Grossberg, S. & Mingolla, E. (1995) Brightness perception, illusory contours, and corticogeniculate feedback. *Visual Neuroscience* 12:1027– 52. [SG]
- Gray, C. M., König, P., Engel, A. K. & Singer, W. (1989) Oscillatory responses in cat visual cortex exhibit inter-columnar synchronization which reflects global stimulus properties. *Nature* 338:334–37. [JMB, arWAP, DLW]
- Gray, C. M. & McCormick, D. A. (1996) Chattering cells superficial pyramidal neurons contributing to the generation of synchronous oscillations in the visual cortex. *Science* 274:109–13. [JJW]

Gray, C. M. & Singer, W. (1989) Stimulus-specific neuronal oscillations in orientation columns of cat visual cortex. *Proceedings of the National Academy of Sciences USA* 86:1698–1702. [aWAP]

- Gray, C. M. & Viana Di Prisco, G. (1993) Properties of stimulus-dependent rhythmic activity of visual cortical neurons in the alert cat. *Society of Neuroscience Abstracts* 19:359.8. [aWAP]
- Gregson, R. A. M. (1988) Nonlinear psychophysical dynamics. Erlbaum Associates. [RAMG]

(1992) *n-Dimensional nonlinear psychophysics*. Erlbaum Associates. [RAMG] (1993) Learning in the context of nonlinear psychophysics: The Gamma Zak

Embedding. British Journal of Mathematical and Statistical Psychology 46:31-48. [RAMG]

(1995) Cascades and fields in perceptual psychophysics. World Scientific. [RAMG]

Gregson, R. A. M. & Harvey, J. P. (1992) Similarities of low-dimensional chaotic auditory attractor sequences to quasirandom noise. *Perception and Psychophysics* 51:267–78. [RAMG]

Grossberg, S. (1968) Some nonlinear networks capable of learning a spatial pattern of arbitrary complexity. Proceedings of the National Academy of Sciences 60:758–65. [SG]

- (1971) Pavlovian pattern learning by nonlinear neural networks. Proceedings of the National Academy of Sciences 68:828–31. [SG]
- (1982) Studies of mind and brain. Reidel. [SG]
- (1987) The adaptive brain, vol. II. Elsevier. [SG]

 (1993) Self-organizing neural models of categorization, inference and synchrony. *Behavioral and Brain Sciences* 16:460–61. [aWAP]
 (1995) The attentive brain. *American Scientist* 83:438–49. [SG]

- Grossberg, S., Boardman, L. & Cohen, M. (1997a) Neural dynamics of variablerate speech categorization. *Journal of Experimental Psychology: Human Perception and Performance* 23:481–503. [SG]
- Grossberg, S. & Mingolla, E. (1985) Neural dynamics of perceptual grouping: Textures, boundaries, and emergent segmentations. *Perception and Psychophysics* 38:141–71. [SG]

Grossberg, S., Mingolla, E. & Ross, W. (1997b) Visual brain and visual perception: How does the cortex do perceptual grouping? *Trends in Neurosciences* 20:106–111. [SG]

Grossberg, S. & Somers, D. (1991) Synchronized oscillations during cooperative feature linking in a cortical model of visual perception. *Neural Networks* 4:453–66. [SG, aWAP]

Guettler, A., Eckhorn, R., Juergens, E. & Frien, A. (1997) Neural correlation contrast in visual cortex of monkey changes with stimulus contrast across an object-background-border. In: *From membrane to mind*. Thieme. [RE]

Gur, M. & Akri, V. (1992) Isoluminant stimuli may not expose the full contribution of color to visual functioning: Spatial contrast sensitivity measurements indicate interaction betwen color and luminance processing. *Vision Research* 32:1253–62. [aWAP]

Hamming, R. W. (1980) Coding and information theory. Prentice-Hall. [aWAP]

Hancock, P. J. B., Smith, L. S. & Phillips W. A. (1991a) A biologically supported error-correcting learning rule. *Neural Computation* 3:201–12. [arWAP] (1991b) A biologically supported error-correcting learning rule. In: *Proceedings*

of the International Conference on Artificial Neural Networks, ed O. Simula. Elsevier. [arWAP]

- Hansel, C., Artola, A. & Singer, W. (in preparation) Relation between dendritic Ca²⁺ levels and the polarity of synaptic long-term modifications in rat visual cortex neurons. [rWAP]
- Harter, M. R. (1967) Excitability cycles and cortical scanning: A review of two hypotheses of central intermittency in perception. *Psychological Bulletin* 68:47–58. [VGH]

Hebb, D. O. (1949) The organization of behaviour. Wiley. [PK, GP, aWAP]

Heeger, D. J. (1993) Modeling single cell direction selectivity with normalized, half-squared, linear operators. *Journal of Neurophysiology* 70(5):1885– 98. [DS]

Hestrin, S., Sah, P. & Nicoll, R. A. (1990) Mechanisms generating the time course of dual component excitatory synaptic currents recorded in hippocampal slices. *Neuron* 5:247–53. [rWAP]

Hinton, G. E., McClelland, J. L. & Rumelhart, D. E. (1986) Distributed representations. In: *Parallel distributed processing. Explorations in the microstucture of cognition*, vol. 1, ed. D. E. Rumelhart. & J. L. McClelland. MIT Press. [aWAP]

Hirsch, J. A. & Gilbert, C. D. (1991) Synaptic physiology of horizontal connections in cat's visual cortex. *Journal of Neuroscience* 11:1800– 9. [arWAP]

 (1993) Long-term changes in synaptic strength along specific intrinsic pathways in the cat visual cortex. *Journal of Physiology* 461:247– 62. [aWAP]

Hopfield, J. J. (1982) Neural networks and physical systems with emergent collective computational capabilities. *Proceedings of the National Academy* of Sciences USA 79:2554–58. [arWAP]

Horgan, J. (1995) From complexity to perplexity. *Scientific American* June:74–79. [aWAP]

Hotelling, H. (1936) Relations between two sets of variables. *Biometrika* 28:321– 77. [aWAP]

Huang, Y-Y., Colino, A., Selig, D. K. & Malenka, R. C. (1992) The influence of prior synaptic activity on the induction of long-term potentiation. *Science* 255:730–33. [aWAP]

Hummel, J. E. & Biederman, I. (1992) Dynamic binding in a neural network for shape recognition. *Psychological Review* 99:480–517. [aWAP]

Hummel, J. E. & Holyoak, K. J. (1993) Distributing structure over time. Behavioral and Brain Sciences 16:464. [aWAP] Humphreys, G. W. & Riddoch, M. J. (1987) To see but not to see: A case study of visual agnosia. Erlbaum. [aWAP]

Humphreys, G. W., Troscianko, T., Riddoch, M. J., Boucart, M. Donnely, N. & Harding, G. F. A. (1992) Covert processing in different visual recognition systems. In: *The neuropsychology of consciousness*, ed. A. D. Milner & M. D. Rugg. Academic Press. [aWAP]

Iacoboni, M., Rayman, J. & Zaidel, E. (1996a) Left brain says yes, right brain says no: Normative duality in the split brain. In: *Toward a science of consciousness: The first Tucson discussions and debates*, eds. S. R. Hameroff, A. W. Kaszniak, A. C. Scott. MIT Press. [MI]

(in preparation) Lateralized lexical decision with unilateral and bilateral presentations: Evidence from the split brain. [MI]

Iacoboni, M., Woods, R. P., Lenzi, G. L. & Mazziotta, J. C. (1997) Merging of oculomotor and somatomotor space coding in the human right precentral gyrus. *Brain* 120(3). [MI]

Iacoboni, M., Woods, R. P. & Mazziotta, J. C. (1996b) Brain-behavior relationships: Evidence from practice effects in spatial stimulus-response compatibility. *Journal of Neurophysiology* 76:321–31. [MI]

Iacoboni, M. & Zaidel, E. (1996) Hemispheric independence in word recognition: Evidence from unilateral and bilateral presentations. *Brain and Language* 53:121-40. [MI]

Ingber, L. (1985) Statistical mechanics of neocortical interactions: Stability and duration of the 7+/- rule of short-term memory capacity. *Physical Review* A., 31:1183–86. [PLN]

(1995) Statistical mechanics of multiple scales of neocortical interactions. In: *Neocortical dynamics and human EEG rhythms*, ed. P. L. Nunez. Oxford University Press. [PLN]

Intrator, N. & Cooper, L. N. (1995a) Information theory and visual plasticity. In: *The handbook of brain theory and neural networks*, ed. M. A. Arbib. MIT Press. [arWAP]

(1995b) BCM theory of visual cortical plasticity. In: *The handbook of brain theory and neural networks*, ed. M. A. Arbib. MIT Press. [arWAP]

Ishai, A & Sagi, D. (1995) Common mechanisms of visual perception and imagery. *Science* 268:1772–74. [aWAP]

Jakobson, R. (1964) Towards a linguistic typology of aphasic impairments. In: Disorders of language. Cyba Foundation Symposium, ed. A. V. S. de Reuck & M. O'Connor. Little, Brown & Company. [VGH]

James, W. (1890) The Principles of Psychology. Reprint. Dover. [GT]

Jensen, O., Idiart, M. A. P. & Lisman, J. E. (1996) Physiologically realistic formation of autoassociative memory in networks with theta/gamma oscillations: Role of fast NMDA channels. *Learning and Memory* 3:243–56. [rWAP] Jerison, H. J. (1973) *Evolution of the brain and intelligence*. Academic

Press. [aWAP]

Jirsa, V. K. & Haken, H. (1996) Field theory of electromagnetic brain activity. *Physical Review Letters* 77: 960–63. [PLN]

Johnston, J. C. & McClelland, J. L. (1973) Visual factors in word perception. Perception and Psychophysics 14:365–70. [aWAP]

Juergens, E. & Eckhorn, R. (1997) Parallel processing by a homogenous group of coupled model neurons can enhance, reduce and generate signal correlations. *Biological Cybernetics* 76:217–27. [RE]

Kaas, J. H. (1995) The reorganization of sensory and motor maps in adult mammals. In: *The cognitive neurosciences*, ed. M. S. Gazzaniga. MIT Press. [aWAP]

Kanevsky, D. (1989) A multiple source, or, is a striped apple more than a striped orange? *Behavioral and Brain Sciences* 12:767–69. [aWAP]

Kapadia, M. K., Ito, M., Gilbert, C. D. & Westheimer, G. (1995) Improvement in visual sensitivity by changes in local context: Parallel studies in human observers and in V1 of alert monkeys. *Neuron* 15:843–56. [SG, aWAP]

Karni, A. & Sagi, D. (1991) Where practice makes perfect in texture discrimination: Evidence for primary visual cortex plasticity. *Proceedings of* the National Academy of Sciences USA 88:4966–70. [aWAP]

Kastner, S., Nothdurft, H. -C. & Pigarev, I. N. (1997) Neuronal correlates of pop-out in cat striate cortex. *Vision Research* 37(4):371–76. [DS]

Katznelson, R. D. (1981) Normal modes of the brain: Neuroanatomic basis and a physiologic theoretical model. In: *Electric fields of the brain: The neurophysics of EEG*, ed. P. L. Nunez. Oxford University Press. [PLN]

Kay, J. (1992) Feature discovery under contextual supervision using mutual information. Proceedings of the International Joint Conference on Neural Networks Book 4:79–84. Baltimore. [aWAP]

Kay, J., Floreano, D. & Phillips, W. A. (in press) Contextually guided unsupervised learning using local multivariate binary processors. *Neural Networks*. [aWAP]

Kay, J. & Phillips, W. A. (1994) Activation functions, computational goals and learning rules for local processors with contextual guidance (Technical Report CCCN-15). Centre for Cognitive and Computational Neuroscience, University of Stirling, Scotland, UK. [aWAP]

Kay, J. & Phillips, W. A. (1997) Activation functions, computational goals and learning rules for local processors with contextual guidance. *Neural Computation* 9:763–78. [arWAP]

Kiper, D. C., Gegenfurtner, K. R. & Movshon, J. A. (1996) Cortical oscillatory responses do not affect visual segmentation. *Vision Research* 36(4):539– 44. [rWAP]

Kirkwood, A., Rioult, M. G. & Bear, M. F. (1996) Experience-dependent modification of synaptic plasticity in visual cortex. *Nature* 381:526– 28. [aWAP]

Kisvardy, Z. F., Martin, K. A. C., Freund, T. F., Magloczky, Z., Whitteridge, D. & Somogyi, P. (1986) Synaptic targets of HRP-filled layer III pyramidal cells in the cat striate cortex. *Experimental Brain Research* 64:541–52. [aWAP]

Knierem, J. J. & Van Essen, D. C. (1992) Neuronal responses to static texture patterns in area V1 of the alert macaque monkey. *Journal of Neurophysiology* 67:961–80. [aWAP]

Knight, R. A. (1992) Specifying cognitive deficiencies in poor premorbid schizophrenics. In: *Progress in experimental personality and psychopathology research: Vol. 15*, ed. E. F. Walker, R. H. Dworking & B. A. Cornblatt. Springer.

Knight, R. A. & Silverstein, S. M. (In press) The role of cognitive psychology in guiding research on cognitive deficits in schizophrenia. In: *Experimental psychopathology and the pathogenesis of schizophrenia*, ed. M. F. Lenzenweger & R. H. Dworkin. APA Press. [SMS]

Kohonen, T. (1982) Self-organizing formation of topologically correct feature maps. *Biological Cybernetics* 43:59–69. [PM]

König, P., Engel, A. K., Löwel, S. & Singer, W. (1993) Squint affects synchronization of oscillatory response in cat visual cortex. *European Journal* of Neuroscience 5:501–8. [GB, aWAP]

König, P., Engel, A. K., Roelfsema, P. R. & Singer, W. (1996a) Coincidence detection or temporal integration. The role of the cortical neuron revisited. *TINS* 19:130–37. [PK]

 König, P., Engel, A. K. & Singer, W. (1995) Relation between oscillatory activity and long-range synchronization in cat visual cortex. *Proceedings of the National Academy of Sciences USA* 92:290–94. [arWAP]
 (1996b) Integrator or coincidence detector? The role of the cortical neuron

revisited. Trends in the Neurosciences 19:130–37. [rWAP]

Kottmann, H., Eckhorn, R., Woelbern, T., Frien, A. & Bauer, R. (1996) Synchronized fast oscillations in striate cortex correlate with perception of grating orientation in a binocular rivalry task in monkey. In: *Brain and evolution*, ed. N. Elsner & H. -U. Schnitzler. Thieme. [RE]

Kreiter, A. K. & Singer, W. (1992) Oscillatory neuronal responses in the visual cortex of the awake macaque monkey. *European Journal of Neuroscience* 4:369–75. [aWAP]

(1994) Global stimulus arrangement determines synchronization of neuronal activity in the awake macaque monkey. *Supplement European Journal of Neuroscience* 7:153. [aWAP]

(1996) Stimulus-dependent synchronization of neuronal responses in the visual cortex of the awake macaque monkey. *Journal of Neuroscience* 16:2381–96. [DJA, arWAP]

Kröse, B. J. A. & Julesz, B. (1989) The control and speed of shifts of attention. Vision Research 29:1607–9. [aWAP]

Krüger, J. & Becker, J. D. (1991) Recognizing the visual stimulus from neuronal discharges. Trends in Neurosciences 14:282–86. [VGH]

Krueger, L. E. & Shapiro, R. G. (1979) Letter detection with rapid serial presentation: Evidence against word superiority at feature extraction. *Journal of Experimental Psychology: Human Perception and Performance* 5:657–73. [aWAP]

Kruse, W. & Eckhorn, R. (1996) Inhibition of sustained gamma oscillations (35– 80 Hz) by fast transient responses in cat visual cortex. *Proceedings of the National Academy of Sciences USA* 93:6112–17. [RE]

Krystal, J. H. et al (1994) Subanesthetic effects of the noncompetitive NMDA antagonist ketamine in humans. Archives of General Psychiatry 51:199– 214. [rWAP]

Kuramoto, Y. (1984) Chemical oscillations, waves and turbulence. Springer. [GP]

Lades, M., Vorbrügen, J. C., Buhmann, J., Lange, J., von der Malsburg, C., Würtz, R. P. & Konen, W. (1993) Distortion invariant recognition in the dynamic link architecture. *IEEE Transactions on Computers* 42:300– 11. [arWAP, RPW]

Langdon, R. B. & Sur, M. (1990) Components of field potentials evoked by white matter stimulation in isolated slices of primary visual cortex: Spatial distributions and synaptic order. *Journal of Neurophysiology* 64:1484– 1501. [PK]

Lashley, K. (1951) The problem of serial order in behavior. In: Cerebral Mechanisms in Behavior. The Hixon Symposium, ed. L. A. Jeffress. Wiley. [VGH]

Laurent, G. (1996) Dynamical representation of odors by oscillating and evolving neural assemblies. *Trends in Neurosciences* 19:489–96. [VGH]

Leonards, U. & Singer, W. (1997) Interactions between segmentation mechanisms with differential sensitivity for color and luminance contrast. Submitted to *Vision Research*. [rWAP] Leonards, U., Singer, W. & Fahle, M. (1996) The influence of temporal phase differences on texture segmentation. *Vision Research* 36:2689–97. [arWAP]

Levitt, J. B. & Lund, J. S. (1996) Contrast dependence of modulatory surround effects in macaque striate neurons. *Investigative Ophthalmology & Visual Science* 37(3):S482. [DS]

- Li, Z. & Atick, J. J. (1994) Efficient stereo coding in the multiscale representation. *Network* 5:157–74. [aWAP]
- Linsker, R. (1988) Self-organization in a perceptual network. Computer 21:105– 17. [aWAP, JVS]
- Lisman, J. E. & Idiart, M. A. P. (1995) Storage of 7+/-2 short-term memories in oscillatory subcycles. *Science* 267:1512–14. [rWAP]
- Llinás, R. & Sasaki, K. (1989) The functional organization of the olivo-cerebellar system as examined by multiple Purkinje cell recordings. *European Journal* of Neuroscience 1:587–602. [GM]

Llinas, R. R., Grace, A. A. & Yarom, Y. (1991) In vitro neurons in cortical layer 4 exhibit intrinsic oscillatory activity in the 10 to 50 Hz frequency range. *Proceedings of the National Academy of Sciences USA* 88:897–901. [JJW]

Lorentó de No, R. (1949) Cerebral cortex: Architecture, intracortical conections, motor projections. In: *Physiology of the nervous system*, ed. J. F. Fulton. Oxford University Press. [aWAP]

Löwel, S. & Singer, W. (1992) Selection of intrinsic horizontal connections in the visual cortex by correlated neuronal activity. *Science* 255:209–12. [aWAP]

Lumer, E. D., Edelman, G. M. & Tononi, G. (1997) Neural dynamics in a model of the thalamocortical system. I. Layers. loops, and the emergence of fast synchronous oscillations. *Cerebral Cortex* 7:207–27. [GT]

MacLennan, B. (1997) Field computation in motor control. In: Self-organization, computational maps, and motor control, ed. P. Morasso & V. Sanguineti. Elsevier. [PM]

- Macphail, E. M. (1987) The comparative psychology of intelligence. *Behavioral and Brain Sciences* 10:645–95. [aWAP]
- Mainen, Z. F. & Sejnowski, T. J. (1995) Reliability of spike timing in neocortical neurons. *Science* 268:1503–06. [PK]

Marr, D. (1970) A theory for cerebral neocortex. Proceedings of the Royal Society of London (B) 176:161–234. [JVS] (1982) Vision. Freeman. [aWAP]

Martin, K. A. C. (1988) The Wellcome Prize lecture: From single cells to simple circuits in the cerbral cortex. *Quarterly Journal of Experimental Physiology* 73:637–702. [aWAP]

Martinetz, T. & Schulten, K. (1994) Topology representing networks. Neural Networks 7:507–22. [PM]

Massaro, D. W. (1979) Letter information and orthographic context in word perception. Journal of Experimental Psychology: Human Perception and Performance 5:595–609. [aWAP]

(1989a) Testing between the TRACE model and the fuzzy logic model of speech perception. *Cognitive Psychology* 21:398–21. [arWAP]

(1989b) Multiple book review of Speech perception by ear and eye: A paradigm for psychological inquiry. Behavioral and Brain Sciences 12:741–94. [aWAP]

 Massaro, D. W. & Cohen, M. M. (1983) Phonological constraints in speech perception. *Perception and Psychophysics* 34:338–48. [aWAP]
 (1991) Integration versus interactive activation: The joint influence of stimulus

and context in perception. Cognitive Psychology 23:558-614. [aWAP]

Massaro, D. W. & Friedman, D. (1990) Models of integration given multiple sources of information. *Psychological Review* 97:225–52. [aWAP]

Matelli, M., Luppino, G. & Rizzolatti, G. (1991) Architecture of superior and mesial area 6 and the adjacent cingulate cortex in the macaque monkey. *Journal of Comparative Neurology* 311:445–62. [MI]

Maunsell, J. H. R. (1995) The brain's visual world: Representations of visual targets in cerebral cortex. *Science* 270:764–68. [rWAP]

- Mayford, M., Wang, J. Kandel, E. R. & O'Dell, T. J. (in press) CaMKII regulates the frequency response function of hippocampal synapses for the production of both LTD and LTP. *Cell.* [aWAP]
- McCarthy, R. A. & Warrington, E. K. (1990) Cognitive neuropsychology. A clinical introduction. Academic Press. [aWAP]

McClelland, J. L. (1978) Perception and masking of wholes and parts. Journal of Experimental Psychology: Human Perception and Performance 4:210– 23. [aWAP]

(1991) Stochastic interactive processes and the effect of context on perception. *Cognitive Psychology* 23:1–44. [aWAP]

McClelland, J. L. & Elman J. L. (1986) The TRACE model of speech perception. *Cognitive Psychology* 18:1–86. [aWAP]

McClelland, J. L., McNaughton, B. L. & O'Reilly, R. C. (1995) Why there are complementary learning systems in the hippocampus and neocortex: Insights from the successes and failures of connectionist models of learning and memory. *Psychological Review* 102:419–507. [aWAP]

McClelland, J. L. & Rumelhart, D. E. (1981) An interactive activation model of context effects in letter perception: 1. An account of basic findings. *Psychological Review* 88:375–407. [aWAP]

- McClelland, J. L., Rumelhart, D. E. & Hinton, G. E. (1986) The appeal of parallel distributed processing: In: *Parallel distributed processing: Explorations in the microstucture of cognition*, vol. 1, ed. D. E. Rumelhart & J. L. McClelland. MIT Press. [aWAP]
- McClurkin, J. W., Optican, L. M., Richmond, B. J. & Gawne, T. J. (1991) Concurrent processing and complexity of temporally encoded messages in visual perception. *Science* 253:675–77. [VGH]
- McCollum, G. (1994) Navigating a set of discrete regions in body position space. Journal of Theoretical Biology 167:263–71. [GM]
- (1997) Dynamics of cerebellar neurons. *International Journal of Theoretical Physics* (in press). [GM]
- McCollum, G., Holroyd, C. & Castelfranco, A. M. (1995) Forms of early walking. *Journal of Theoretical Biology* 176:373–90. [GM]
- McGuire, B. A., Gilbert, C. D., Rivlin, P. K. & Wiesel, T. N. (1991) Targets of horizontal connections in macaque primary visual cortex. *Journal of Comparative Neurology* 305:370–92. [aWAP]

Michalski, A., Gerstein, G. L., Czarkowska, J. & Tarnecki, R. (1983) Interactions between cat striate cortex neurons. *Experimental Brain Research* 51:97– 107. [aWAP]

- Milner, P. M. (1974) A model for visual shape recognition. *Psychological Review* 81:521–35. [aWAP, DLW]
- Mizobe, K., Polat, U., Kasamatsu, T. & Norcia, A. M. (1996) Lateral masking reveals facilitation and suppression from the same single cells in cat area 17. *Investigative Ophthalmology & Visual Science* 37(3):S483. [DS]

Mohr, B., Pulvermuller, F., Rayman, J. & Zaidel, E. (1994) Interhemispheric cooperation during lexical processing is mediated by the corpus callosum: Evidence from the split-brain. *Neuroscience Letters* 181:17–21. [MI]

Monyer, H., Burnashev, N., Laurie, D. J., Sakmann, B. & Seeburg, P. H. (1994) Developmental and regional expression in the rat brain and functional properties of four NMDA receptors. *Neuron* 12:529–40. [rWAP]

Morasso, P. (1981) Spatial control of arm movements. *Experimental Brain Research* 42:223–27. [PM]

Morasso, P. & Sanguineti, V. (1996) How the brain can discover the existence of external egocentric space. *Neurocomputing* 12:289–310. [PM]

Morton, J. (1969) Interaction of information in word recognition. *Psychological Review* 76:165–78. [aWAP]

Movellan, J. R. & McClelland, J. L. (1995) Stochastic interactive processing, channel separability, and optimal perceptual inference: An examination of Morton's law (Technical Report PDP.CNS.95.4). Department of Psychology, Carnegie Mellon University, Pittsburgh, PA. [arWAP]

- Mozer, M. C., Zemel, R. S., Behrmann, M. & Williams, C. K. I. (1992) Learning to segment images using dynamic feature binding. *Neural Computation* 4:650–65. [aWAP]
- Mumford, D. (1992) On the computational architecture of the neocortex: 2. The role of cortico-cortical loops. *Biological Cybernetics* 66:241–51. [aWAP]

Munk, M. H. J., Nowak, L. G., Chouvet, G., Nelson, J. I. & Bullier, J. (1992) The structural basis of cortical synchronization. *European Journal of Neuroscience Supplement* 5:21. [aWAP]

Munk, M. H. J., Nowak, L. G., Nelson, J. I. & Bullier, J. (1995) Structural basis of cortical synchronization. 2. Effects of cortical-lesions. *Journal of Neurophysiology* 74:2401–14. [GB]

Munk, M. H. J., Roelfsema, P. R., K(nig, P., Engel, A. K. & Singer, W. (1996) Role of reticular activation in the modulation of intracortical synchronization. *Science* 272:271–74. [rWAP]

Murthy, V. N. & Fetz, E. E. (1992) Coherent 25- to 35- Hz oscillations in the sensorimotor cortex of awake behaving monkeys. *Proceedings of the National Academy of Sciences (USA)* 89:5670–74. [VGH]

(1996) Synchronization of neurons during local field potential oscillations in sensorimotor cortex of awake monkeys. *Journal of Neurophysiology* 76:3968–82. [rWAP]

Nakayama, K. & Mackeben, M. (1989) Sustained and transient components of focal visual attention. *Vision Research* 31:1221–36. [aWAP]

Navon, D. (1977) Forest before trees: The precedence of global features in visual perception. *Cognitive Psychology* 9:353–83. [rWAP]

Nelson, J. I. (1995) Binding in the visual system. In: *The handbook of brain theory and neural networks*, ed. M. A. Arbib. MIT Press. [aWAP]

Nelson, J. I. & Frost, B. J. (1978) Orientation-selective inhibition from beyond the classic visual receptive field. *Brain Research* 139:359–65. [aWAP]

Neuenschwander, S., Engel, A. K., König, P., Singer, W. & Varela, F. J. (1996) Long-range synchronization of oscillatory light responses in the cat retina and lateral geniculate nucleus. *Nature* 379:728–33. [arWAP]

Neven, H. & Aertsen, A. Rate coherence and event coherence in the visual cortex: A neuronal model of object recognition. *Biological Cybernetics* 67: 309–22. [aWAP]

Nunez, P. L. (1974) The brain wave equation: A model for the EEG. *Math Biosciences* 21:279–97. [PLN]

(1981) *Electric fields of the brain: The neurophysics of EEG.* Oxford University Press. [PLN]

References/Phillips & Singer: Cortical computation

(1989) Generation of human EEG rhythms by a combination of long and short-range neocortical interactions. *Brain Topography* 1:199–215. [PLN] (1995a) *Neocortical dynamics and human EEG rhythms*. Oxford University

(1995a) Neocortical dynamics and human EEG rhythms. Oxford University Press. [PLN]
(1995b) Multicole proceeding dynamics apparimental EEC measures and

(1995b) Multiscale neocortical dynamics, experimental EEG measures, and global facilitation of local cell assemblies. *Behavioral and Brain Sciences* 19:305–306. [PLN]

Nunez, P. L., Silberstein, R. B., Carpentar,, M. R., Srinivasan, R., Wijesinghe, R. S., Tucker, D. M., Cadusch, P. J. & Wu, X. (1997) EEG coherency II: Comparisons of multiple measures of EEG coherency differences between resting and cognitive states. *Electroencephalography and Clinical Neurophysiology*, submitted. [PLN]

Olney, J. W. & Farber, N. B. (1995) Glutamate receptor dysfunction and schizophrenia. Archives of General Psychiatry 52:998-1007. [SMS]

- Olshausen, B. A., Anderson, C. H. & van Essen, D. C. (1993) A neurobiological model of visual attention and invariant pattern recognition based on dynamic routing of information. *Journal of Neuroscience* 13(11):4700– 19. [RPW]
- (1995) A multiscale dynamic routing circuit for framing size- and positioninvariant object representations. *Journal of Computational Neuroscience* 2:45–62. [RPW]
- Olshausen, B. & Field, D. J. (1996) Emergence of simple-cell receptive field properties by learning a sparse code for natural images. *Nature* 318:607– 09. [rWAP]

Palm, G. (1982) Neural assemblies. An alternative approach to artificial intelligence. Springer. [GP]

(1993) On the internal structure of cell assemblies. In: *Brain theory*, ed. A. Aertsen. Elsevier Science Publishers. [GP]

- Palmer, S. E. (1975) The effects of contextual scenes on the identification of objects. *Memory and Cognition* 3: 519–26. [aWAP]
- Pandya, D. N. & Yeterian, E. H. (1985) Architecture and connections of cortical association areas. In: *Cerebral cortex*, ed. A. Peters & E. G. Jones. Plenum Press. [GB]

Panzeri, S., Biella, G., Rolls, E. T., Skaggs, W. E. & Treves, A. (1996) Speed, noise, information and the graded nature of neuronal responses. *Network* 7:365–70. [AT]

Pece, A. E. C. (1992) Redundancy reduction of a Gabor representation: A possible computational role for feedback from primary visual cortex to lateral geniculate nucleus. In: *Artificial neural networks 2*, ed. I. Aleksander & J. Taylor. Elsevier. [aWAP]

- Phillips, W. A. (1971) Does familiarity affect transfer from an iconic to a short-term memory? *Perception and Psychophysics* 10:153–57. [aWAP]
- (1997) Theories of cortical computation. In: *Cognitive neuroscience*, ed M. D. Rugg. University College Press. [aWAP]

Phillips, W. A. & Goodall, W. C. (1994) Lexical writing can be non-semantic and fluent without practice. *Cognitive Neuropsychology* 12:149–74. [aWAP]

Phillips, W. A., Hay, I. M., & Smith, L. S. (1993) Lexicality and pronunciation in a simulated neural net. *British Journal of Mathematical and Statistical Psychology* 46:193–205. [aWAP]

Phillips, W. A., Kay, J., & Smyth, D. (1995a) How local cortical processors that maximise coherent variation could lay foundations for representation proper. In: *Neural computation and psychology*, ed. L. S. Smith & P. J. B. Hancock. Springer-Verlag. [arWAP]

Phillips, W. A., Kay, J. & Smyth, D. (1995b) The discovery of structure by multistream networks of local processors with contextual guidance. *Network* 6:225–46. [arWAP]

Phillips, W. A. & Singer, W. (1974) Function and interaction of on and off transients in vision: 1. Psychophysics. *Experimental Brain Research* 19:493– 506. [arWAP]

Piaget, J. (1954) The construction of reality by the child. Basic Books. [aWAP]

Picard, N. & Strick, P. L. (1996) Motor areas of the medial wall: A review of their location and functional activation. *Cerebral Cortex* 6:342–53. [MI]

Pinker, S. (1994) *The language of instinct*. Penguin. [rWAP]

Place, E. J. S. & Gilmore, G. C. (1980) Perceptual organization in schizophrenia. Journal of Abnormal Psychology 89:409–18. [rWAP]

Plaut, D. C. & McClelland, J. L. (1993) Generalization with componential attractors: Word and nonword reading in an attractor network. In: *Proceedings of the 15th Annual Conference of the Cognitive Science Society*. Erlbaum. [aWAP]

- Poggio, T., Fahle, M. & Edelman S. (1992) Fast perceptual learning in visual hyperacuity. *Science* 256:1018–20. [aWAP]
- Polat, U. & Norcia, A. M. (1996) Neurophysiological evidence for contrast dependent long-range facilitation and suppression in the human visual cortex. *Vision Research* 36(14):2099–2109. [DS]

Polat, U. & Sagi, D. (1993) Lateral interactions between spatial channels: Suppression and faciltation revealed by lateral masking experiments. *Vision Research* 33:993–99. [aWAP]

- (1994a) The architecture of perceptual spatial interactions. *Vision Research* 34:73–78. [aWAP]
- (1994b) Spatial interactions in human vision: From near to far via experiencedependent cascades of connections. *Proceedings of the National Academy of Sciences USA* 91:1206–9. [arWAP]
- Poppel, E. (1970) Excitability cycles in central intermittency. *Psychologische Forschung* 34:1–9. [VGH]
- (1978) Time perception. In: Handbook of sensory physiology. Vol. VIII: Perception, ed. R. Held, H. W, Leibowitz & H. -L. Teuber. Springer. [VGH]
- (1994) Temporal mechanisms in perception. International Review of Neurobiology 37:185–202. [VGH]
- Posner, M. I. & Rothbart, M. K. (1994) Constructing neuronal theories of mind. In: Large-scale neuronal theories of brain function, ed. C Kock & J. L. Davis. MIT Press. [aWAP]
- Proctor, R. W. & Dutta, A. (1993) Do the same stimulus-response relations influence choice reactions initially and after practice? *Journal of Experimental Psychology: Learning, Memory, & Cognition* 19:922– 30. [MI]
- Protopapas, A. & Bower, J. M. (1994) Sensitivity in the response of piriform cortex pyramidal cells to fluctuations in synaptic timing. In: *Computation in neurons and neural systems*, ed. F. Eeckman. Kluwer Press. [JMB]
- Purves, D., Riddle, D. R. & LaMantia, A.-S. (1992) Iterated patterns of brain circuitry (or how the cortex gets its spots). *Trends in the Neurosciences* 15:362–68. [aWAP]
- Rabinowicz, E. F., Opler, L. A., Owen, D. R. & Knight, R. A. (1996) The dot enumeration perceptual organization task (DEPOT): Evidence for a shortterm memory deficit in schizophrenia. *Journal of Abnormal Psychology* 105:336–48. [rWAP]
- Rakic, P. & Singer, W. (1988) Neurobiology of neocortex. Wiley. [aWAP]
- Redlich, A. N. (1993) Redundancy reduction as a strategy for unsupervised learning. *Neural Computation* 5:289–304. [aWAP]
- Reeke, G. N, Jr., Finkel, L. H., Sporns, O. & Edelman, G. M. (1990) Synthetic neural modelling: A multilevel approach to the analysis of brain complexity. In: *Signal and sense: Local and global order in perceptual maps*, ed. G. M. Edelman, W. E. Gall & W. M. Cowan. Wiley. [aWAP, GT]
- Reicher, G. M. (1969) Perceptual recognition as a function of meaningfulness of stimulus material. *Journal of Experimental Psychology* 81:274–80. [aWAP]
- Richman, H. B. & Simon, H. A. (1989) Context effects in letter perception: Comparison of two theories. *Psychological Review* 96:417-32. [aWAP]
- Robinson, P. A. Rennie, C. J. & Wright, J. J. (1997a) Propagation and stability of waves of electrical activity in the cerebral cortex. *Physical Review E.* (in press) [JJW]
- Robinson, P. A., Wright, J. J. & Rennie, C. J. (1997b) Synchronous oscillation in the cerebral cortex. *Physical Review E.* (submitted). [JJW]
- Rockel, A. J., Hiorns, R. W. & Powell, T. P. S. (1980) The basic uniformity in structure of the neocortex. *Brain* 103:221–44. [aWAP]
- Rockland, K. & Lund, J. S. (1983) Intrinsic laminar lattice connections in primate visual cortex. *Journal of Comparative Neurology* 216:303– 18. [aWAP]
- Roelfsma, P. R., Engel, A. K., König, P. & Singer, W. (1994a) Oscillations and synchrony in the visual cortex: Evidence for their functional relevance. In: Oscillatory event-related brain dynamics, ed. C. Pantev. Plenum. [arWAP]
- (1997) Visuomotor integration is associated with zero time-lag synchronization among cortical areas. *Nature* 385:157–61. [rWAP]
- Roelfsma, P. R., König, P., Engel, A. K., Sireteanu, R. & Singer, W. (1994b) Reduced synchronization in the visual cortex of cats with strabismic amblyopia. *European Journal of Neuroscience* 6:1645–55. [GB, arWAP]
- Rolls, E. T., Treves, A. & Tovee, M. J. (1997) The representational capacity of the distributed encoding of information provided by populations of neurons in the primate temporal visual cortex. *Experimental Brain Research* 114:149–62. [AT]
- Rucci, M., Tononi, G. & Edelman, G. M. (1997) Registration of neural maps through value-dependent learning: Modeling the alignment of auditory and visual maps in the bran owl's optic tectum. *Journal of Neuroscience* 17:334– 52. [GT]
- Ruhnau, E. (1995) Time-gestalt and the observer. In: *Conscious experience*, ed. T. Metzinger. Imprint Academic Schöningh. [RAMG]
- Rumelhart, D. E., Hinton, G. E. & Williams, R. J. (1986) Learning internal representations by back-propagating errors. *Nature* 323:533–36. [DF, aWAP]
- Rumelhart, D. E. & McClelland, J. L. (1982) An interactive activation model of context effects in letter perception: 2. The contextual enhancement effect and some tests and extensions of the model. *Psychological Review* 89:60– 84. [aWAP]
- (1986) Parallel distributed processing: Explorations in the microstucture of cognition, vol. 1. MIT Press. [aWAP]

- Sagi, D. (1995) The psychophysics of texture segmentation. In: *Early vision and beyond*, ed. T. V. Papathomas et al. MIT Press. [rWAP]
- Salin, P. & Bullier, J. (1995) Corticocortical connections in the visual system: Structure and function. *Physiological Reviews* 75:107–54. [rWAP]
- Samuel, A. G. (1981) Phonemic restoration: Insights from a new methodology. Journal of Experimental Psychology: General 110:474–94. [aWAP] Does lexical information influence the perceptual restoration of phonemes?
- Journal of Experimental Psychology: General 125:28–51. [arWAP] Sanes, J. N. & Donoghue, J. P. (1993) Oscillations in local field potentials of the
- primate motor cortex during voluntary movement. Proceedings of the National Academy of Sciences (USA) 90:4470–74. [VGH]
- Sanger, T. D. (1994) Theoretical considerations for the analysis of population coding in motor cortex. *Neural Computation* 6:29–37. [PM]
- Sanguineti, V., Morasso, P. & Frisone, F. (1997) Cortical maps of sensorimotor spaces. In: Self-organization, computational maps, and motor control, ed. P. Morasso and V. Sanguineti. Elsevier. [PM]
- Schillen, T. B. & König, P. (1994) Binding by temporal structure in multiple feature domains of an oscillatory neuronal network. *Biological Cybernetics* 70:397–405. [aWAP]
- Schlaggar, B. L. & O'Leary, D. D. M. (1991) Potential of visual cortex to develop an array of functional units unique to somatosensory cortex. *Science* 252:1556-60. [aWAP]
- Schmidhuber, J. & Prelinger, D. (1993) Discovering predictable classifications. Neural Computation 5:625–35. [aWAP]
- Schmidt, K. E., Löwel, S., Goebel, R. & Singer, W. (1997) The perceptual grouping criterion of colinearity is reflected by anisotropies of connections in primary visual cortex. *European Journal of Neuroscience* 9:1083– 89. [aWAP]
- Schwartz, C. & Bolz, J. (1991) Functional specificity of the long-range horizontal connections in cat visual cortex: A cross-correlation study. *Journal of Neuroscience* 11:2995–3007. [aWAP]
- Schwartz, E. L. (1977) Spatial mapping in the primate sensory projection: Analytic structure and relevance to perception. *Biological Cybernetics* 25:181–94. [GM]
- Sejnowski, T. J., Koch, C. & Churchland, P. S. (1988) Computational neuroscience. Science 241:1299–1306. [aWAP]
- Shadlen, M. N. & Newsome, W. T. (1994) Noise, neural codes and cortical organization. *Current Biology* 4:569–79. [rWAP]
- Shallice, T. (1988) From neuropsychology to mental structure. Cambridge University Press. [aWAP]
- Shallice, T. (1991) Précis of From neuropsychology to mental structure. Behavioral and Brain Sciences 14:429–69. [aWAP]
- Shastri, L. & Ajjanagadde, V. (1993) From simple associations to systematic reasoning: A connectionist representation of rules, variables and dynamic bindings using temporal synchrony. *Behavioral and Brain Sciences* 16:417– 94. [aWAP]
- Shepherd, G. M., ed. (1990) *The synaptic organization of the brain*. Oxford University Press. [aWAP]
- Shepherd, G. M. & Koch, C. (1990) Introduction to synaptic circuits. In: *The synaptic organization of the brain*, ed. G. M. Shepherd. Oxford University Press. [aWAP]
- Silberstein, R. B. (1995) Neuromodulation of neocortical dynamics. In: Neocortical dynamics and human EEG rhythms, ed. P. L. Nunez. Oxford University Press. [PLN]
- Sillito, A. M., Jones, H. E., Gerstein, G. L. & West, D. C. (1994) Feature-linked synchronization of thalamic relay cell firing induced by feedback from the visual cortex. *Nature* 369:479–82. [GB, SG, aWAP]
- Silverstein, S. M., Bakshi, S., Chapman, R. M. & Nowlis, G. (1996c) Reduced plasticity of the visual information processing system in schizophrenia. Evidence from the processing of visual patterns (submitted). [SMS]
- Silverstein, S. M., Knight, R. A., Schwarzkopf, S. B., West, L. L., Osborn, L. M. & Kamin, D. (1996a) Configural and context effects in perceptual organization in schizophrenia. *Journal of Abnormal Psychology* 105:410– 20. [SMS]
- Silverstein, S. M., Matteson, S. & Knight, R. A. (1996b) Reduced top-down influence in auditory perceptual organization in schizophrenia. *Journal of Abnormal Psychology* 105:663–67. [SMS]
- Silverstein, S. M., Raulin, M. R., Pristach, E. A. & Pomerantz, J. R. (1992) Perceptual organization and schizotypy. *Journal of Abnormal Psychology* 101:265–70. [rWAP]
- Silverstein, S. M., Schenkel, L. S., Light, G. A. & Schwarzkopf, S. B. (1996d) Clinical features associated with perceptual organization dysfunction in schizophrenia (submitted). [SMS]
- Simpson, J. I., Wylie, D. R. & De Zeeuw, C. I. (1996) On climbing fiber signals and their consequence(s). *Behavioral and Brain Sciences* 19:380–94. [GM]
- Singer, W. (1987) Activity-dependent self-organization of synaptic connections as a substrate of learning. In: *The neural and molecular bases of learning*, ed. J.-P. Changeux & M. Konishi. Wiley. [aWAP]

- (1990) Search for coherence: A basic principle of cortical self-organization. Concepts in Neuroscience 1:1–26. [VGH, aWAP]
- (1993) Synchronization of cortical activity and its putative role in information processing and learning. *Annual Review of Physiology* 55:349–74. [VGH, PLN, aWAP]
- (1994a) Coherence as an organizing principle of cortical functions. International Review of Neurobiology 37:153–83. [aWAP]
- (1994b) Time as a coding space in neocortical processing: A hypothesis. In: Temporal coding in the brain, ed. G. Buzsáki, R. Llinás, W. Singer, A. Berthoz & Y. Christen. Springer. [VGH]
- (1995) Development and plasticity of cortical processing architectures. *Science* 270:758–64. [aWAP]
- Singer, W. & Artola, A. (1994) Plasticity of the mature cortex. In: *Cellular and molecular mechanisms underlying higher neural functions*, ed. A. I. Selverston & P. Ascher. Wiley. [aWAP]
- Singer, W. & Gray, C. M. (1995) Visual feature integration and the temporal correlation hypothesis. *Annual Review of Neuroscience* 18:555–86. [aWAP, JJW]
- Singer, W. & Phillips, W. A. (1974) Function and interaction of on and off transients in vision: 2. Neurophysiology. *Experimental Brain Research* 19:507–21. [arWAP]
- Smyth, D. (1994) Simulations of networks of local processors with contextual guidance. M.Sc. thesis, Centre for Cognitive and Computational Neuroscience, University of Stirling, Scotland, UK. [aWAP]
- Smyth, D. & Der, R. (1995) Learning to bind. Poster presented at the Symposium on Phenomena and Architectures of Cognitive Dynamics, Universität Leipzig, June. [aWAP]
- Smyth, D., Kay, J. & Phillips, W. A. (1994) Discovery of high-order functions in multi-stream, multi-stage nets without external supervision. Paper presented to the Neural Computation and Psychology Workshop, Centre for Cognitive and Computational Neuroscience, University of Stirling, Scotland, UK. [aWAP]
- Smyth, D., Phillips, W. A. & Kay, J. (1996) Measures for investigating the contextual modulation of information transmission. *Network*. 7:307– 16. [aWAP]
- Softky, W. R. (1995) Commentary: Simple codes versus efficient codes. Current Biology 5(2):239–47. [rWAP]
- Somers, D. & Kopell, N. (1993) Rapid synchronization through fast threshold modulation. *Biological Cybernetics* 68:393–407. [rWAP, DLW]
- Soutif, M. (1970) Vibrations, propagation, diffusion. Dunod. [VGH]
- Spinelli, D. N., Starr, A. & Barrett, T. W. (1968) Auditory specificity in unit recordings from cat's visual cortex. *Experimental Neurology* 22:75– 84. [aWAP]
- Sporns, O., Gally, J. A., Reeke, G. N., Jr. & Edelman, G. M. (1989) Reentrant signaling among simulated neuronal groups leads to coherency in their oscillatory activity. *Proceedings of the National Academy of Sciences USA* 86:7265–69. [aWAP]
- Sporns, O., Tononi, G. & Edelman, G. M. (1991) Modeling perceptual grouping and figure-ground segregation by means of active re-entrant connections. *Proceedings of the National Academy of Sciences USA* 88:129–33. [aWAP, GT, DLW]
- Squire, L. R. (1992) Memory and the hippocampus: A synthesis from findings with rats, monkeys, and humans. *Psychological Review* 99:195– 231. [aWAP]
- Stemmler, M., Usher, M. & Niebur, E. (1995) Lateral interactions in primary visual cortex: A model bridging physiology and psychophsyics. *Science* 269:1877–80. [DS]
- Steriade, M., Curro-Dossi, R. & Contreras, D. (1993) Electrophysiological properties of intralaminar thalamocortical cells discharging rhythmic approx.
 40 Hz spike bursts at approx. 1000 Hz during waking and rapid eye movement sleep. *Neuroscience* 56:1–9. [JJW]
- Stone, J. V. (1996a) A canonical microfunction for learning perceptual invariances. *Perception* 25(2):207–20. [JVS]
 - (1996b) Learning perceptually salient visual parameters through spatiotemporal smoothness constraints. *Neural Computation*. [aWAP, JVS]
- Stone, J. & Bray, A. (1995) A learning rule for extracting spatio-temporal invariances. *Network* 6:429–36. [aWAP]
- Stryker, M. P. (1988) Group report. Principles of cortical self-organization. In: Neurobiology of neocortex, ed. P. Rakic & W. Singer. Wiley. [aWAP]
- Stuart, G. J. & Sakmann, B. (1994) Action propaganda of somatic action potentials into neocortical pyramidal cell dendrites. *Nature* 367:69– 72. [rWAP]
- Sur, M., Garraghty, P. & Roe, A. (1988) Experimentally induced visual projections into auditory thalamus and cortex. *Science* 242:1437– 41. [aWAP]
- Swindale, N. V. (1990) Is the cerebral cortex modular? Trends in the Neurosciences 12:487–92. [aWAP]

- Szenatgoathai, J. (1978) The neuron network of the cerebral cortex: A functional approach. *Proceedings of the Royal Society London* (B) 201:219–48. [JVS]
- Tass, P. & Haken, H. (1996) Synchronized oscillations in the visual cortex a synergetic model. *Biological Cybernetics* 74:31–39. [JJW]
- Terman, D. & Wang, D. L. (1995) Global competition and local cooperation in a network of neural oscillators. *Physica D* 81:148–76. [DLW]
- Theunissen, F. & Miller, J. P. (1995) Temporal encoding in nervous systems: A rigorous definition. *Journal of Computional Neuroscience* 2:149– 62. [VGH]
- Thorpe, S. J., O'Regan, K. & Pouget, A. (1989) Humans fail on XOR pattern classification problems. In: *Neural networks from models to applications*, ed. L. Personnaz & G. Dreyfus. IDSET. [aWAP]
- Tiitinen, H., Sinkkonen, J., Reinikainen, K. Alho, K., Lavikaainen, J. & Näätänen, R. (1993) Selective attention enhances the auditory 40-Hz transient response in humans. *Nature* 364:59–60. [aWAP]
- Tononi, G., Sporns, O. & Edelman, G. M. (1992a) The problem of neuronal integration: Induced rhythms and short-term correlations. In: Induced rhythms in the brain, ed. E. Basar & T. H. Bullock. Birkhauser. [aWAP]
- Tononi, G., Sporns, O. & Edelman, G. M. (1992b) Reentry and the problem of integrating multiple cortical areas: Simulation of dynamic integration in the visual system. *Cerebral Cortex* 2:310–35. [aWAP, GT]
- Tononi, G., Sporns, O. & Edelman, G. M. (1994) A measure for brain complexity: Relating functional segregation and integration in the nervous system. *Proceedings of the National Academy of Sciences USA* 91:5033– 37. [aWAP, GT]
- Tononi, G., Sporns, O. & Edelman, G. M. (1996) A measure for selective matching of signals by the brain. *Proceedings of the National Academy of Sciences USA* 93:3422–27. [aWAP, GT]
- Tooby, J. & Cosmides, L. (1995) Mapping the evolved functional organization of mind and brain. In: *The cognitive neurosciences*, ed. M. S. Gazzaniga. MIT Press. [aWAP]
- Tovee, M. J. & Rolls, E. T. (1992a) Oscillatory activity is not evident in the primate temporal visual cortex with static stimuli. *NeuroReport* 3:369– 72. [VGH]
- (1992b) The functional nature of neuronal oscillations. *Trends in the Neurosciences* 15:387. [aWAP]
- Toyama, K., Kimura, M. & Tanaka, K. (1981) Organization of cat visual cortex as investigated by cross-correlation techniques. *Journal of Neurophysiology* 46:202–14. [aWAP]
- Traub, R. D., Whittington, M. A., Stanford, I. M. & Jeffreys, J. G. R. (1996) A mechanism for generation of long-range synchronous fast oscillations in the cortex. *Nature* 383:621–24. [rWAP, JJW]
- Treves, A. & Panzeri, S. (1995) The upward bias in measures of information derived from limited data samples. *Neural Computation* 7:399–407. [AT]
- Treves, A., Skaggs, W. E. & Barnes, C. A. (1996) How much of the hippocampus can be explained by functional constraints? *Hippocampus* 6:666–74. [AT]
- Troscianko, T. (1994) Contribution of colour to the motion aftereffect and motion perception. *Perception* 23:1221–31. [aWAP]
- Troscianko, T., Davidoff, J., Humphreys, G., Landis, T., Fahle, M., Greenlee, M., Brugger, P. & Phillips, W. A. (1996) Human colour discrimination based on a non-parvocellular pathway. *Current Biology* 6:200–10. [aWAP]
- Troscianko, T., Landis, T. & Phillips, W. A. (1993) Chromatic discrimination in cerebral achromotopsia: Additional evidence favouring magno-based perception, and a neural-net model. *Perception* 22(suppl):8-9. [aWAP]
- Troscianko, T., Prince, C., Fahle, M. & Regard, M. (1995) The uses of hightemporal-frequency chromatic information in visual perception. *Perception* 24(suppl.):59. [aWAP]
- Ts'o, D., Gilbert, C. & Weisel, T. N. (1986) Relationship between horizontal interactions and functional architecture in cat striate cortex as revealed by cross-correlation analysis. *Journal of Neuroscience* 6:1160–70. [aWAP]
- Ullman, S. (1994) Sequence seeking and counterstreams: A model for bidirectional information flow in the cortex. In: *Large-scale neuronal theories of brain function*, ed. C Koch & J. L. Davis. MIT Press. [aWAP]
- Usher, M. & Niebur, E. (1996) Modeling the temporal dynamics of It neurons in visual search: A mechanism for top-down selective attention. *Journal of Cognitive Neuroscience* 8:311–27. [rWAP]
- Van Essen, D. C., Anderson, C. H. & Olshausen, B. A. (1994) Dynamic routing strategies in sensory, motor and cognitive processing. In: *Large-scale neuronal theories of brain function*, ed. C. Koch & J. L. Davis. MIT Press. [aWAP]
- Van Essen, D. C. & Maunsell, J. H. R. (1983) Hierarchical organization and functional streams in the visual cortex. *TINS* 6:370–75. [PK]
- van Rotterdam, A., Lopes da Silva, F. H., van der Ende, J., Viergever, M. A. & Hermans, A. J. (1982) A model of the spatial-temporal characteristics of the alpha rhythm. *Bulletin of Mathematical Biology* 44:283–305. [PLN]
- Volgushev, M., Chistiakova, M. & Singer, W. (submitted) Modification of discharge patterns of neocortical neurons by induced oscillations of the membrane potential. [rWAP]

Volgushev, M., Voronin, L. L., Chistiakova, M. & Singer, W. (1997) Relations between long-term synaptic modifications and paired-pulse interactions in the rat neocortex. *European Journal of Neuroscience* (in press). [rWAP]

- von der Malsburg, C. (1981) The correlation theory of brain function (Internal report 81-2.). Department of Neurobiology, Max-Planck-Institute for Biophysical Chemistry, Gottingen, Germany. [aWAP, DLW] (1998) Bettern recognition by labled graph matching. Neural Networks 1:141
- (1988) Pattern recognition by labled graph matching. *Neural Networks* 1:141–48. [aWAP]
- von der Malsburg, C. & Schneider, W. (1986) A neural cocktail-party processor. Biological Cybernetics 54:29–40. [aWAP]
- von der Malsburg, C. & Singer, W. (1988) Principles of cortical network organization. In: *Neurobiology of neocortex*, ed. P. Rakic & W. Singer. Wiley. [rWAP]
- Von Stein, A., Chiang, C. & König, P. (1996) Expectancy-driven synchronization between primary visual cortex and parietal cortex in cats. *Society for Neuroscience Abstracts* 22:255.3. [PK]
- Wang, D., Buhmann, J. & von der Malsburg, C. (1990) Pattern segmentation in associative memory. *Neural Computation* 2:94–106. [aWAP]
- Wang, D. L. (1993a) Modeling global synchrony in the visual cortex by locally coupled neural oscillators. *Proceedings of the Fifteenth Annual Conference* of the Cognitive Science Society 1058–63. [DLW]
- (1993b) Pattern recognition: Neural networks in perspective. *IEEE Expert* 8:52–60, August. [DLW]
- (1995) Emergent synchrony in locally coupled neural oscillators. *IEEE Transactions on Neural Networks* 6:941–48. [DLW]
- Wang, D. L. & Terman, D. (1995) Locally excitatory globally inhibitory oscillator networks. *IEEE Transactions on Neural Networks* 6:283–86. [DLW]
 (1997) Image segmentation based on oscillatory correlation. *Neural*
- Computation 9:805–36. [DLW] Warren, R. M. (1970) Perceptual restoration of missing speech sounds. Science 167:392–93. [aWAP]
- Weisstein, N. & Harris, C. S. (1974) Visual detection of line segments: An object-superiority effect. *Science* 186:752–55. [aWAP]
- Weliky, M., Kandler, K., Fitzpatrick, D. & Katz, L. C. (1995) Patterns of excitation and inhibition evoked by horizontal connections in visual cortex share a common relationship to oreintation columns. *Neuron* 15:541– 52. [DS]
- Wenneckers, T. & Palm, G. (1997) On the relation between neural modelling and experimental neuroscience. *Theory in Biosciences* (in press). [GP]
- Wever, R. (1965) Pendulum vs. relaxation oscillation. In: Circadian clocks. Proceedings of the Feldafing Summer School, ed. J. Aschoff. North Holland. [VGH]
- White, E. L. (1989) Cortical circuits: Synaptic organization of the cerebral cortex: Structure, function and theory. Birkhauser. [aWAP]
- Wilson, H. R. & Cowan, J. D. (1973) A mathematical theory of the functional

dynamics of cortical and thalamic nervous tissue. *Kybernetik* 13:55-80. [PLN]

- Wilson, M. A. & Bower, J. M. (1988) A computer simulation of olfactory cortex with functional implications for storage and retrieval of olfactory information. In: *Neural information processing systems*, ed. D. Anderson. AIP Press. [JMB]
 - (1991) A computer simulation of oscillatory behavior in primary visual cerebral cortex. *Neural Computation* 3:498–509. [JMB]
 - (1992) Cortical oscillations and temporal interactions in a computer simulation of piriform cortex. *Journal of Neurophysiology* 67:981–95. [JMB]
- Wimmer, H. & Perner, J. (1983) Beliefs about beliefs: Representing and constraining function of wrong beliefs in children's understanding of deception. *Cognition* 13:103–28. [aWAP]
- Wright, J. J. (1997) EEG simulation: Variation of spectral envelope, pulse synchrony and approx. 40 Hz oscillation. *Biological Cybernetics* (in press). [JJW]
- Wright, J. J. & Liley, D. T. J. (1996) Dynamics of the brain at global and microscopic scales: Neural networks and the EEG. *Behavioral and Brain Sciences* 19:285–295. [PLN, JJW]
- Wurtz, R. P. (1995) Multilayer dynamic link networks for establishing image point correspondences and visual object recognition, volume 41 of Reihe Physik. Verlag Harri Deutsch. [RPW]
- (1997) Object recognition robust under translations, deformations and changes in background. *IEEE Transactions on Pattern Recognition and Machine Intelligence* (in press). [RPW]
- Yamaguchi, Y. & Shimizu, H. (1994) Pattern-recognition with figure-ground separation by generation of coherent oscillations. *Neural Networks* 3:49– 63. [aWAP]
- Yan, J. & Suga, N. (1996) Corticofugal modulation of time-domain processing of biosonar information in bats. *Science* 273:1100–03. [rWAP]
- Young, M. P., Tanaka, K. & Yamane, S. (1992) On oscillating neuronal responses in the visual cortex of the monkey. *Journal of Neurophysiology* 67:1464– 74. [VGH, aWAP]
- Zaitchik, D. (1990) When representations conflict with reality. *Cognition* 35:41– 68. [aWAP]
- Zemel, R. S. & Hinton, G. E. (1991) Discovering viewpoint-invariant relationships that characterize objects. In: Advances in neural information processing systems 3:299–305. Morgan Kaufmann. [aWAP]
- Zhadin, M. N. (1984) Rhythmic processes in cerebral cortex. Journal of Theoretical Biology 108:565–95. [PLN]
- Zipser, D. & Andersen, R. A. (1988) A back-propagation programmed network that simulates response properties of a subset of posterior parietal neurons. *Nature* 331:679–84. [DF]
- Zipser, K., Lamme, V. A. F. & Schiller, P. H. (1996) Contextual modulation in primary visual cortex. *The Journal of Neuroscience* 15:7376–89. [rWAP]