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Coupling of Total Hemoglobin Concentration, Oxygenation, and Neural Activity in Rat Somatosensory Cortex

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pretation of the resulting maps. The present report sis was limited to frequencies above 40 Hz. represents the first systematic study with sufficient Another potential problem is that the linear fits re-

modynamic signals, simultaneous recordings of neu- activity. Since the hemodynamic response is measured ral activity, and an event-related stimulus paradigm, as a change from baseline, the response to the absence we demonstrate that (1) there is a strongly nonlinear of a stimulus (or zero stimulus amplitude) should be zero, relationship between electrophysiological measures and thus the curve would be expected to go through the of neuronal activity and the hemodynamic response, origin. (2) the hemodynamic response continues to grow be- The present report represents the first systematic yond the saturation of electrical activity, and (3) the study with sufficient statistical power to quantitatively initial increase in deoxyhemoglobin that precedes an characterize the nonlinear relationship between changes increase in blood volume is counterbalanced by an in blood oxygen content and the neural spiking and synapequal initial decrease in oxyhemoglobin. tic activity by employing high signal-to-noise ratio (SNR)

Neural activity changes in brain tissue are coupled to Results changes in blood flow, blood volume, and blood oxygenation, collectively referred to as the hemodynamic re- Spectroscopic optical measurements of total hemoglothem functional magnetic resonance imaging (fMRI), in- et al., 2003) were performed simultaneously with electrofer neural activity from hemodynamic changes (Kwong physiological recordings in rat somatosensory (barrel) et al., 1992; Ogawa et al., 1992). However, the nature of cortex. The barrel cortex in the rat is a well-studied the coupling between the fMRI signal and brain electrical example of topographic mapping, where each one of activity is still under debate (Arthurs and Boniface, 2002; the large facial vibrissae (whiskers) is mapped onto a zen, 2001). Many fMRI studies have relied on a linear der Loos, 1970). Tactile whisker stimulation produces coupling model, where the hemodynamic signal is as- neural and hemodynamic responses that localize to corsumed to be proportional to a measure of neural activity. responding barrels (Masino et al., 1993). This system is this would greatly simplify the analysis and interpreta- cortical activations. tion of fMRI data (Heeger and Ress, 2002). Indeed, a Single deflections of a single whisker produced a reli-

number of recent studies have provided evidence in support of a linear coupling model, showing that hemodynamic signals correlate linearly with synchronized synaptic activity (Arthurs et al., 2000; Brinker et al., 1999; Ngai et al., 1999) as well as with neural firing rates (Smith Charlestown, Massachusetts 02129 et al., 2002). Across-species comparison of single-cell activity in monkeys and human fMRI data also yielded 2Program in Biophysics Harvard Medical School a linear relationship (Heeger et al., 2000; Rees et al., Boston, Massachusetts 02115 2000). However, these studies have generally had insuf- ³ Institute for Psychology of the Hungarian Academy ficient statistical power to detect subtle departures from of Sciences linearity due to measurement noise, insufficient number Budapest 1068 of data points, or a narrow range of stimulus intensities. Hungary For instance, only two conditions were used in Smith et al. (2002). Ngai et al. (1999), on the other hand, used a much larger number of data points (n 34). However, Summary the measurement variance makes it difficult to distinguish between linear and nonlinear models. In the report Recent advances in brain imaging techniques, includ- by Logothetis et al. (2001), the dynamic range of the ing functional magnetic resonance imaging (fMRI), of- stimulus intensities may have been too narrow to reveal fer great promise for noninvasive mapping of brain a nonlinear relationship. Furthermore, in order to reduce function. However, the indirect nature of the imaging artifacts in the electrophysiological recordings introsignals to the underlying neural activity limits the inter- duced by simultaneous fMRI measurements, the analy-

statistical power to quantitatively characterize the re- ported in several studies (Arthurs et al., 2000; Logothetis lationship between changes in blood oxygen content et al., 2001; Ngai et al., 1999) suggest a significant posiand the neural spiking and synaptic activity. tive or negative change in the hemodynamic signal cor-Using two-dimensional optical measurements of he- responding to the baseline level of electrophysiological

hemodynamic measurements with rich data sampling Introduction over a wide range of neural response amplitudes.

bin concentration and hemoglobin oxygenation (Dunn specific cortical area, called a barrel (Woolsey and Van **If a linear relationship were a satisfactory approximation, perfectly suited for studying small and spatially localized**

able hemodynamic signal with a spatial extent compara- *Correspondence: adevor@nmr.mgh.harvard.edu ble to that reported in previous studies (Peterson et

(A) Ratio images of activation for each of the 6 filters (wavelengths et al., 1993). indicated above) were calculated by dividing the response (aver- MUA and LFP signals integrated over the time window aged 1.5–2.5 s following the stimulus) by the baseline image (aver-
aged from 7 s preceding the stimulus). 990 trials were averaged for

Responses for each of the 6 filters are superimposed. The stimulus ures 4C and 4D). onset is denoted by an arrow. Legend: 560 nm (dark blue), 570 nm In order to address the laminar patterns of electro-

al., 1998). Figure 1A shows ratio images obtained by
dividing the maximal optical response image by the
baseline image for the largest of the 9 stimulus ampli-
tudes at each of the 6 wavelengths. The observed differ-
ence **of total hemoglobin (HbT). It also reflects blood volume to peak responses of Hb, HbO, and HbT averaged from** under the assumption of constant hematocrit. On the other hand, imaging at 610 nm creates a map highly biased toward Hb (Malonek and Grinvald, 1996). Figure 1B shows the averaged time course for each of the **stimulus amplitudes measured from the region of inter- function of stimulus intensity. As a result, the relationest (ROI) outlined by a square in Figure 1A. As expected, the amplitude of the response increased with stimulus ure 5C). The hemodynamic signals also increased with**

signal changes at all wavelengths to a model, taking higher amplitudes (Figure 5D). As a result, the relation-

into account the respective absorption spectra (see Experimental Procedures). Figure 2 shows percent change maps for Hb, HbO, and HbT calculated from the spectral data in Figure 1. An initial increase in Hb (the "initial dip") was observed shortly following stimulus onset. The initial dip was originally reported by Grinvald and collaborators (Frostig et al., 1990) and since then has been observed by other groups (Thompson et al., 2003). In contrast to previous studies (Vanzetta and Grinvald, 2001), the initial increase in Hb was always balanced by an equal decrease in HbO, so that the blood volume (HbT) remained unchanged during the time of the initial dip. Blood volume started to increase about 600 ms following the stimulus, probably reflecting increased blood flow. As was first pointed out by Fox and Raichle (1986), an excessive delivery of fresh blood, measured in our study as an increase in blood volume, leads to an increase in HbO and decrease in Hb, HbO peaking before Hb (Figure 2).

Electrophysiological recordings of multiple unit activity (MUA) and local field potential (LFP) were performed simultaneously with optical measurements from the center of the barrel (Figure 3). LFP measures a weighted sum of transmembrane currents due to synaptic and dendritic activity (Eccles, 1951; Plonsey and Heppner, 1967), whereas MUA measures population spiking activity (Legatt et al., 1980; Mitzdorf, 1987). Both MUA and LFP showed a fast transient response followed by a delayed response with lower amplitude. This biphasic response has been observed in previous studies (Moore and Nelson, 1998) and has been attributed to the de-Figure 1. Multiwavelength Imaging of Intrinsic Signals layed activation of NMDA receptors (Armstrong-James

of activation (Σ MUA and Σ LFP) increased as the stimulus aged 1.5–2.5 s following the stimulus) by the baseline image (aver-
aged from 7 s preceding the stimulus). 990 trials were averaged for
each stimulus amplitudes and approached saturation at high
from the baseline. Scale b **lated by averaging the signal from the ROI (square in A, first panel). transient and the delayed responses components (Fig-**

light green), 580 nm (purple), 590 nm (light blue), 600 nm (red), 610 physiological activity, MUA and LFP were measured us-
Ing a linear array multi-electrode (see Experimental im (dark green). Vertical scale bar equals 0. **Procedures). Current source density (CSD) analysis was**

 MUA and Σ **LFP MUA,** Σ **LFP, Hb, HbO, and HbT** are averaged from 5 animals, and the error bars represent the intersubject standard error. **EMUA** (Figure 5A) **LFP (Figure 5B) behaved in a similar way as a** MUA and Σ LFP was close to linear (Fig**intensity from stimulus amplitude 1 to 9. increasing stimulus intensity. However, in contrast to Quantitative estimates of the concentrations of Hb, the electrophysiological parameters, there was no indi-HbO, and HbT were obtained by fitting the observed cation of saturation of the hemodynamic response at**

Figure 2. Spatial-Temporal Evolution of HbO, Hb, and HbT

Each image represents an individual frame (average of 990 trials). Time between consecutive images is 200 ms. (B) is a continuation of the time series shown in (A). The signal for Hb and HbO is expressed in percent change relative to its own baseline concentration (40 and 60 μM, **respectively). HbT was calculated as a sum of Hb and HbO. Scale bar equals 500 m.**

ship between the observed electrophysiological mea- Discussion sures and hemodynamic signals is nonlinear (Figure 5E).

$$
f = ax^b, \hspace{1cm} (1)
$$

relation corresponds to $b = 1$. However, the value $b =$ **shown above the plots, indicating a significant nonline- moglobin. arity. A number of improvements in experimental design**

The observed relationship between electrophysiologi- Using simultaneous multiwavelength optical and eleccal measures and the hemodynamic response ampli- trophysiological recordings, we have demonstrated that tude can be well approximated by a power-law function (1) there is a strongly nonlinear relationship between electrophysiological measures of neuronal activity and *f ax* **the hemodynamic response, (2) the hemodynamic re-** *^b***, (1) sponse continues to grow beyond the saturation of elec**as shown in Figure 5E (solid lines). Note that a linear trical activity, and (3) the initial increase in deoxyhemo-
relation corresponds to $b = 1$. However, the value $b =$ globin that precedes an increase in blood volume **1** falls far below the 95% confidence intervals for *b* counterbalanced by an equal initial decrease in oxyhesity shown above the plots indicating a significant nonline- moglobin.

can explain the disagreement between our results and previous studies that inferred a linear relationship. First, electrophysiological and hemodynamic signals were recorded simultaneously from the same cortical location (same barrel). Second, rapid and randomized eventrelated stimulus presentation methodology allowed a large number of stimulus trials (990 trials for each stimulus amplitude). This method is robust against slow drifts in baseline conditions. Data can therefore be acquired for long periods of time until the desired SNR level is reached. High reproducibility of the results allowed across-animal averaging. Likewise, high SNR and rich two-dimensional sampling of the hemodynamic data account for our ability to resolve the initial increase in Hb, which is accompanied by an initial decrease in HbO (Figure 2). This observation, which we interpret as reflecting a transition of HbO into Hb during the initial dip, Figure 3. Simultaneous Recordings of Neural and Hemodynamic is of particular importance for refining models of the
Signals (A) MUA (expressed in spikes per second) and LFP (expressed in (Buxton, 2001).
Units of SD compared

(B) Signal time courses of Hb, HbO, and HbT calculated from the trophysiological response. The largest amplitude satudata in Figure 1B. Note that the Hb scale is inverted. rated the electrophysiological response but was still not

cussed above. By transforming the measured electro- imposed.

delayed response components. The responses shown in Figure 3 nonlinearity, we obtain the following expression were integrated 0–25 ms (red), 25–300 ms (green), and 0–300 ms \int (blue) following the stimulus.

(E and F) Averaged, rectifyed MUA (E) and CSD (F) recorded from supragranular layers (depth of 0–400 m, red), granular layer (500– where *S¹* **is exactly the neurovascular coupling function ⁹⁰⁰ m, blue), and infragranular layers (900–2000 m, green). given in Equation 1.**

approximated by the following linear expression deflection amplitudes and velocities larger than those we used. It might be possible to reach hemodynamic (*ui* **plateau by adopting multiwhisker or electrical stimula-)***h***(***t***), (4) tion paradigms.**

It has been proposed that neural metabolism is more **closely linked to synaptic than spiking activity (Laurit- of delta functions, centered at the onset of each event; zen, 2001; Logothetis, 2002). We are unable to address see Dale, 1999),** *ui* **is the corresponding (time-integrated)** this question, since for the range of stimulus intensities electrophysiological response, and $h_i(t)$ is the event**used in the present study, LFP and MUA measures were related hemodynamic response function. Estimation of the** *hi* **highly correlated. However, our findings of continued (***t***) can be performed using standard linear estimahemodynamic increases beyond saturation of the elec- tion or deconvolution methods (Burock and Dale, 2000; trophysiological measures are difficult to reconcile with Friston et al., 1998).**

hemodynamic activity driven by metabolic demands related to synaptic currents. In this context, it is important to note that LFP and MUA represent derived measures of neuronal activity and may in fact be nonlinearly related to other parameters such as ion pumping, neurotransmitter release, and vasoactive processes. Excess metabolic demand might also be attributed to nonneural cortical elements, for example astrocytes, which have high oxygen metabolism and release vasoactive compounds in response to neural neurotransmitter release (Zonta et al., 2003). Coupling between neural activity and hemodynamic signals confined to a well-defined dynamic range has been observed previously using laser Doppler flowmetry (Nielsen and Lauritzen, 2001) and in glucose consumption experiments (Ackermann et al., 1984). In the latter, within a certain domain, increased metabolism was accompanied not only by unchanged but decreasing neural activity. This result was interpreted as reflecting the metabolic demands of activating inhibitory circuits.

Our findings would be consistent with a saturating nonlinear relationship between neurotransmitter release and synaptic currents (e.g., due to the depolarization of postsynaptic membranes toward the reversal potential), and a roughly linear relationship between neurotransmitter release and the hemodynamic response. More formally, we would propose the following model:

$$
y(t) = z(t) \otimes h(t), u(t) = S(z(t)), \qquad (2)
$$

where *y***(***t***) and** *u***(***t***) are the measured hemodynamic and electrophysiological signals, respectively, as a function of time,** *t***.** *h***(***t***) is a hemodynamic impulse response func-Figure 4. Saturation of Electrophysiological Measures tion, and z(***t***) represents the neuronal process underlying** Each point is an average of 990 stimulus trials. Average responses
were normalized to the maximal amplitude.
(A and B) MUA (A) and LFP (B) consistently saturate as a function
of stimulus intensity. Responses for each of th **(C and D) MUA and LFP saturate for both the transient and the physiological signals through the inverse of the static**

$$
y(t) = S^{-1} \left(u(t) \right) \otimes h(t), \qquad (3)
$$

In order to relate this result to optical imaging or fMRI large enough to reach a plateau of the hemodynamic
response. In other sensory systems, such as the visual
and auditory systems, stimulus intensities cover many
orders of magnitude, and stimulus intensity is coded by
the ne

$$
y(t) = \sum_{i}^{N} x_{i}(t) \otimes h_{i}(t), \quad h_{i}(t) = S^{-1}(u_{i})h(t), \quad (4)
$$

where $x_i(t)$ is the event sequence for condition i (a sum

generalize to other brain areas, other stimulus para- racy and reliability of conclusions based on fMRI and digms, and the awake versus anesthetized state. Laurit- may greatly facilitate the integration of multiple imaging zen's group, using cerebellum as a model, has demon- modalities. strated an increase in the blood flow despite inhibition of Purkinje cells (Caesar et al., 2003). However, the large Experimental Procedures majority of the cells in the cerebellar cortex are granule
cells, and the increase in blood flow might correspond
to the increased activity in granule cell layer. Moreover,
approved by the Animal Care Committee. Five rats **blood supply in the cerebellum is much richer in the simultaneous recordings of intrinsic optical signals and electrophysgranule cell layer than in the molecular layer (Scremin, iological responses. Rats were anesthetized with urethane (1.2 g/kg, 1995), probably corresponding to great density of gran- i.p.), and an area of skull overlying the primary somatosensory cortex**

lus-response functions. The similarity of responses in for insertion of the recording electrode. In addition, two rats were plex, long-lasting, or multiwhisker, stimuli. Likewise, re- mals were initially anesthetized with ketamine hydrochloride (50 sponse properties of neurons are known to change as
a function of type and depth of anesthesia (Friedberg
et al., 1999; Martin et al., 2002). Further experiments in
the animal was supplemented by constant i.p. infusion of

as fMRI, positron emission tomography, near-infrared upper contacts showed stable recordings. spectroscopy, and electro/magnetoencephalography, offer great promise for noninvasive measurement of

brain function in space and time (Dale and Halgren,

2001). The findings presented in this paper provide a

2001). The findings presented in this paper provide a

method **hemodynamic measures through explicit mathematical rotated at 3 Hz, thus producing 18 independent illumination col**spatially localized stimuli and rich data sampling, we have
demonstrated a strong nonlinear relationship between
electrophysiological activity and the hemodynamic re-
sponse. Further elucidation of the neurovascular rela-
 tionship, including extension of our measurements to prior to averaging. Trials corresponding to each stimulus type were

Figure 5. Nonlinearity of Neurovascular Coupling

Each point is an average of 4950 stimulus trials from 5 animals. Before averaging animals, Σ MUA, Σ LFP, Hb, HbO, and HbT re**sponses to amplitude 9 was normalized to 1. The error bars represent intersubject standard error.**

 $(A \text{ and } B)$ $\sum MUA$ (A) and $\sum LFP$ (B) plotted as **a function of stimulus intensity.**

(C) ΣLFP as a function of ΣMUA.

(D) Hb (blue), HbO (red), and HbT (black) peak responses increase linearly with stimulus intensity.

(E) Σ **LFP** (right) and Σ MUA (left) as a function **of the peak percent change in HbO (red) and HbT** (black). A power-law function $f = ax^b$ **was fit to the data (solid lines). Note that the hemodynamic response continues to grow** beyond the plateau in Σ **LFP** and Σ MUA.

It has to be investigated whether our conclusions can higher primates, can be expected to improve the accu-

was thinned with a dental burr until transparent (\sim 100 μ m). A barrier
of petroleum ielly was built around the border of the thinned skull Using our stimulus paradigm, recordings of neural of performance of the dividend the border of the time skull
responses from different layers produce similar stimu-
skull over the center of a barrel, as determined by optic used for laminar electrophysiological recordings only. These ani-
mals were initially anesthetized with ketamine hydrochloride (50 **awake behaving animals will address these questions. primary somatosensory cortex and the dura matter were removed, The integration of modern imaging techniques, such and the electrode array was slowly inserted into the cortex until the**

principled basis for combining electrophysiological and lengths of 560, 570, 580, 590, 600, and 610 nm. The filterwheel modeling of the neurovascular coupling. Using brief and ors per second. Images were acquired by a cooled 12-bit CCD

components were projected out of the time course at every pixel

selectively averaged and blank (no stimulus) trial average was sub- References tracted. This is formally equivalent to temporal deconvolution, given randomization of stimuli and sufficient number of trials (Dale, 1999; Ackermann, R.F., Finch, D.M., Babb, T.L., and Engel, J., Jr. (1984). Dale and Buckner, 1997). Ratio images were calculated by dividing Increased glucose metabolism during long-duration recurrent inhibithe response (averaged 1.5–2.5 s following the stimulus) by the tion of hippocampal pyramidal cells. J. Neurosci. *4***, 251–264.**

baseline image (averaged from 7 s preceding the stimulus). Armstrong-James, M., Welker, E., and Callahan, C.A. (1993). The

The spectral data were converted to percent change maps for

Hb, HbO, and HbT using the modified B

microelectrodes (2–4 MΩ) placed in cortical layer IV or lower layer face, S.J. (2000). Linear coupling between functional magnetic reso-
Il/Ill in the center of a barrel. The position was identified by a selec-**heaps and** II/III in the center of a barrel. The position was identified by a selec-
tive response to the one whisker stimulated and by the cortical sensory cortex. Neuroscience 101, 803–806. **sensory cortex. Neuroscience** *101***, 803–806. tive response to the one whisker stimulated and by the cortical depth (400–500 m). The electrode was positioned only once for Barth, D.S., and Di, S. (1991). Laminar excitability cycles in neocoreach animal in order to preserve as much as possible of the thinned tex. J. Neurophysiol.** *65***, 891–898.** Scull necessary for good quality optical measurements. The signals

were amplified and filtered between 150 and 5000 Hz to record

MUA, and Hoehn-Berlage, M. (1999). Simultaneous recording of evoked poten-

MUA, and betwe **MUA and <code>SLFP</code>** Figure calculated as an integral of the area under the response curve
0–0.3 s after the stimulus onset. Averaged LFP curves were rectified
0–0.3 s after the stimulus onset. Averaged LFP curves were rectified
on the time a **formed using a linear array multielectrode with 24 contacts spaced Burock, M.A., Buckner, R.L., Woldorff, M.G., Rosen, B.R., and Dale, at 100 m (Ulbert et al., 2001a). Laminar signals were amplified and A.M. (1998). Randomized event-related experimental design allow** filtered between 500 and 5000 Hz to record MUA, and between 0.1 for extremely rapid presentation rates using functional MRI. Neuro**and 500 Hz to record LFP. CSD was full wave rectified and averaged report** *9***, 3735–3739. across the traces falling in the corresponding laminae (supragranu- Buxton, R.B. (2001). The elusive initial dip. Neuroimage** *13***, 953–958.** rat, granual, and Italya unit and Science of the total synaptor of consumer brane activity
experience and the synaptor of the total synaptor of the total synaptor and experiment activity-dependent increases in cerebral blo

Since 1 s ISI (see below) is sufficient for relaxation of the electro**physiological activity using a single deflection stimulus (Petersen, Dale, A.M., and Buckner, R.L. (1997). Selective averaging of rapidly**

Stimulation Paradigm

A single whisker was deflected by a computer-controlled piezoelec-

The stimulator, positioned 3 mm from the base of a

whisker, deflected a whisker upward and allowed a free return to

the resting po **of electrophysiological responses to different stimulus amplitudes. Eccles, J.C. (1951). Interpretation of action potentials evoked in the The angular velocity increased from 203/s (vertical displacement cerebral cortex. Electroencephalogr. Clin. Neurophysiol.** *3***, 449–464.** of 240 μ m, amplitude 1) to 969°/s (vertical displacement of 1200
 μ m, amplitude 9). Intervening stimulus amplitudes were spaced with

equal increments on a linear scale.

Example of a final scale.

We employed a fast, randomized a formulated stimulus presentation paradigm analogous to that used in event-related fMRI studies

(Burock et al., 1998; Wagner et al., 1998). We used a constant int **stimulus interval (ISI) of 1 s, including blank trials. The stimulus Fox, P.T., and Raichle, M.E. (1986). Focal physiological uncoupling** sequence was optimized for event-related response estimation effi**ciency using the approach described by Dale (1999). The inclusion sory stimulation in human subjects. Proc. Natl. Acad. Sci. USA** *83***,** of blank trials (or "null events") is equivalent to drawing the ISI **between nonblank events from a discrete random distribution (Bur- Friedberg, M.H., Lee, S.M., and Ebner, F.F. (1999). Modulation of**

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01A2, NIH R01 RR13609, NIH P41 RR14075, NIH R01 NS044623,

Trostig, R.D., Lieke, E.E., Ts'o

27–31. Electrophysiological Recordings

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(Ulbert et al., 2001b).

(Dale, A.M. (1999). Optimal experimental design for event-related

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 Since 1 s ISI (see below) is sufficient for relaxation o

2002), there was virtually no temporal overlap in the neural response. presented individual trails using fMRI. Hum. Brain Mapp. *5***, 329–340.**

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