Systems/Circuits

Serotonin Decreases the Gain of Visual Responses in Awake Macaque V1

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Serotonin, an important neuromodulator in the brain, is implicated in affective and cognitive functions. However, its role even for basic cortical processes is controversial. For example, in the mammalian primary visual cortex (V1), heterogenous serotonergic modulation has been observed in anesthetized animals. Here, we combined extracellular single-unit recordings with iontophoresis in awake animals. We examined the role of serotonin on well-defined tuning properties (orientation, spatial frequency, contrast, and size) in V1 of two male macaque monkeys. We find that in the awake macaque the modulatory effect of serotonin is surprisingly uniform: it causes a mainly multiplicative decrease of the visual responses and a slight increase in the stimulus-selective response latency. Moreover, serotonin neither systematically changes the selectivity or variability of the response, nor the interneuronal correlation unexplained by the stimulus ("noise-correlation"). The modulation by serotonin has qualitative similarities with that for a decrease in stimulus contrast, but differs quantitatively from decreasing contrast. It can be captured by a simple additive change to a threshold-linear spiking nonlinearity. Together, our results show that serotonin is well suited to control the response gain of neurons in V1 depending on the animal's behavioral or motivational context, complementing other known state-dependent gain-control mechanisms.

Key words: awake macaque; extracellular recordings; gain modulation; iontophoresis; serotonin; striate cortex

Significance Statement

Serotonin is an important neuromodulator in the brain and a major target for drugs used to treat psychiatric disorders. Nonetheless, surprisingly little is known about how it shapes information processing in sensory areas. Here we examined the serotonergic modulation of visual processing in the primary visual cortex of awake behaving macaque monkeys. We found that serotonin mainly decreased the gain of the visual responses, without systematically changing their selectivity, variability, or covariability. This identifies a simple computational function of serotonin for state-dependent sensory processing, depending on the animal's affective or motivational state.

Introduction

Perceptually guided behavior depends on context. Such context includes an animal's prior experience or knowledge of the environment and task, and its behavioral and motivational state

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(Harris and Thiele, 2011). The context dependence of perceptually driven behavior relies, in part, on the context-dependent neuromodulation of sensory processing (Hurley et al., 2004; Harris and Thiele, 2011). One mode of such neuromodulation involves subcortical nuclei that have widespread projections throughout the brain (Jacobs and Azmitia, 1992). These are ideally suited to modulate processing in extended networks according to changing behavioral–motivational conditions (Dayan, 2012).

One important neuromodulatory system is the serotonin system. Serotonin [5-hydroxytryptamine (5-HT)] in the brain has been implicated in a variety of affective, cognitive, and sensorimotor functions, but identifying an account of its computational role even for the intensely studied links to reward signaling has proved challenging (Ranade et al., 2014; Dayan and Huys, 2015). Similarly, the reported effects of serotonin on sensory processing have been heterogeneous (Waterhouse et al., 1990; Hurley et al., 2004; Petzold et al., 2009; Watakabe et al., 2009; Lottem et al.,

2016) and a simple computational account remains elusive. For example, in the primary visual cortex (V1) of anesthetized animals' bidirectional modulation of the responses (Watakabe et al., 2009) and variable effects on receptive field properties have been observed (Waterhouse et al., 1990; Hurley et al., 2004; Bachatene et al., 2013). However, despite this multitude of effects on sensory processing, the serotonergic effects on perceptually driven behavior have been surprisingly uniform and consistent with a decreased perceptual response (Davis et al., 1980; Dugué et al., 2014). Potential reasons for these apparent differences are that cortical neuronal modulation in some studies was examined using receptor-selective rather than the endogenous ligand (Watakabe et al., 2009), and these studies were all performed using anesthetized animals. Under anesthesia, substantial fluctuations in brain state that affect responses in the primary sensory cortex have been observed (Ecker et al., 2014). Moreover, anesthesia, such as isofluorane, can directly influence the activity of serotonergic neurons (Johansen et al., 2015). It may therefore be that in awake animals, where the serotonergic system may be in a more controlled state, a less diverse modulation would be seen.

We therefore set out to characterize the modulatory effect of serotonin on sensory processing in the awake animal. To this end, we focused on the macaque V1. The macaque V1 receives extensive projections from the dorsal and medial raphe nuclei, the major source of serotonin in the brain. Serotonergic input projections are most pronounced in the input layers of V1 (de Lima et al., 1988). This pattern is approximately mirrored by the laminar profile of the expression of serotonin receptors in V1 (Watakabe et al., 2009). From a functional perspective, such a bias toward the input layers would be expected, for example, for a mechanism involved in adjusting the gain of the sensory input, as previously suggested (Hurley et al., 2004; Disney et al., 2007). Here, we leveraged the extensive knowledge of the spatiotemporal tuning properties in V1 to quantify the serotonergic modulation on visual processing in awake monkeys along several stimulus dimensions.

Serotonergic neurons are thought to convey a variety of signals on different time scales, as found, for example, for their phasic versus tonic response components (Ranade and Mainen, 2009; Cohen et al., 2015; Fonseca et al., 2015; Hayashi et al., 2015; Correia et al., 2017), versus long-term (days) effects of phasic activation (Correia et al., 2017). In this study we focused on time scales that are consistent with the tonic component of the response by iontophoretically applying serotonin in the minute range time scale.

We find that across the population of neurons in V1 and across different stimulus dimensions, the serotonergic modulation is surprisingly simple: serotonin predominantly decreases the gain of the visual responses, with little change to the tuning properties. A simple additive change to a threshold-linear spiking nonlinearity can account for the observed modulation. Gain modulation is an important computation to change response levels without affecting tuning (Atallah et al., 2012). It has been implicated in the modulation by cognitive states, such as attention (McAdams and Maunsell, 1999), and is subject to cholinergic modulation at the cortical input (Disney et al., 2007). Our results show that serotonin is well suited to control the response gain of neurons in V1, potentially complementing these known gain control mechanisms.

Materials and Methods

Animals. Two adult male rhesus monkeys (Macaca mulatta; M, 8 kg, 11 years old; K, 12 kg, 7 years old; housed in pairs) participated in the experiments. Using aseptic techniques, the monkeys were implanted with a titanium head-post and titanium chambers over the operculum

of V1 under general anesthesia. All experimental procedures followed guidelines for animal experimentation and were approved by the local authorities, the Regierungspräsidium Tübingen, Germany.

Electrophysiological recordings and iontophoresis. We recorded extracellular single-unit activity in V1 while the animals performed a 2 s fixation task (fixation within 0.75° of a small 0.1° fixation dot on the center of the screen) for fluid rewards while we presented stimuli in the receptive field of the recorded unit. The positions of both eyes were recorded at 500 Hz using an infrared optical recording system (Eyelink 1000, SR Research). Experimental control and stimulus presentation was done using custom-written software in Matlab modified after Eastman and Huk (2012) using the psychophysics toolbox (Brainard, 1997; Pelli, 1997; Kleiner et al., 2007).

Recordings and iontophoresis were done using custom-made tungstenin-glass electrodes flanked by two pipettes as described previously (Thiele et al., 2006; Jacob et al., 2013). This electrode pipette was mounted inside a guide tube and inserted transdurally without a dura-piercing guide tube using a custom-made electric microdrive. Iontophoretic application was controlled by an MVCS iontophoresis system (NPI Electronic). Neuronal signals were amplified, digitized, and filtered (250 Hz to 5 kHz) with the Ripple Grapevine System (Ripple). Spike sorting was performed offline using the Plexon Offline Sorter. Spike clusters were computed based on a variety of features, including principal components, energy, peak, trough, and spike amplitude. Single-unit clusters were identified using the features that provided the best separation. Spike isolation was quantified by computing the isolation distance and L ratio (Schmitzer-Torbert et al., 2005). For 763 of 780 (98%) included experimental blocks, the isolation distance was >20 and the L ratio <0.1. For the remaining 17 (2%) experimental blocks that did not meet these criteria, unit isolation was verified by visual inspection. One barrel of the electrode pipette was filled with serotonin hydrochloride (Sigma-Aldrich; 10 mm in doubledistilled water; pH 3.5-3.8), the other with pH-matched saline (NaCl; 0.9%). The electrodes typically had impedances between 0.3 and 1.6 M Ω (measured at 1 kHz) and tip sizes of $10-15 \mu m$. The ejection current ranged between 2 and 50 nA (median, 10 nA) for serotonin and between 5 and 20 nA (median, 11 nA) for saline. To better quantify effects on the neuronal tuning properties, we aimed for relatively small modulatory effects. The values of the ejection currents used were therefore toward the lower end of the range of values previously used in the macaque cortex (Williams et al., 2002). In a subset of experiments, we used different ejection currents across blocks to examine the dose dependence of the effect (Fig. 1B). We typically used ejection currents less than or equal to the value for which the serotonergic modulation seemed to asymptote (Fig. 1B), which we observed at values between 20 and 30 nA, and this limited range of currents may have contributed to the homogeneity of our results. The retention current was -8 nA to prevent leakage from the drug barrels during the control conditions. The pipette resistance ranged from 10 to 150 M Ω , as was used previously (Ott et al., 2014). To minimize long-term effects of serotonin (Maya Vetencourt et al., 2008; Correia et al., 2017), we avoided recording from nearby locations in V1 in consecutive recording sessions.

For each unit we initially quantified the center of the receptive field from receptive-field profiles along a horizontal and vertical axis as described previously (Nienborg et al., 2004) by presenting an elongated rectangular grating (height, 3-5°; width, 0.2°) at different horizontal or vertical positions across the receptive field and its immediate surroundings. Subsequent stimuli were centered on the receptive field at a median eccentricity of 3.6° (range, 1.6 to 6.4°). We initially measured neuronal tuning curves without applying serotonin or saline to establish the baseline using the retention current. We then started to apply serotonin or saline using the ejection current in blocks and then remeasured a tuning curve during the application (note that our analyses of the time course of the response modulation across trials averaged over all experiments did not reveal a systematic difference between the baseline, saline control, and serotonin experiments, suggesting that the lead time of the serotonin application of several seconds was, on average, sufficient). Following the application of the drug, we remeasured the tuning in subsequent blocks using the retention current to obtain the recovery of the response if it was possible to maintain unit isolation. Thus, if it was possible to maintain unit

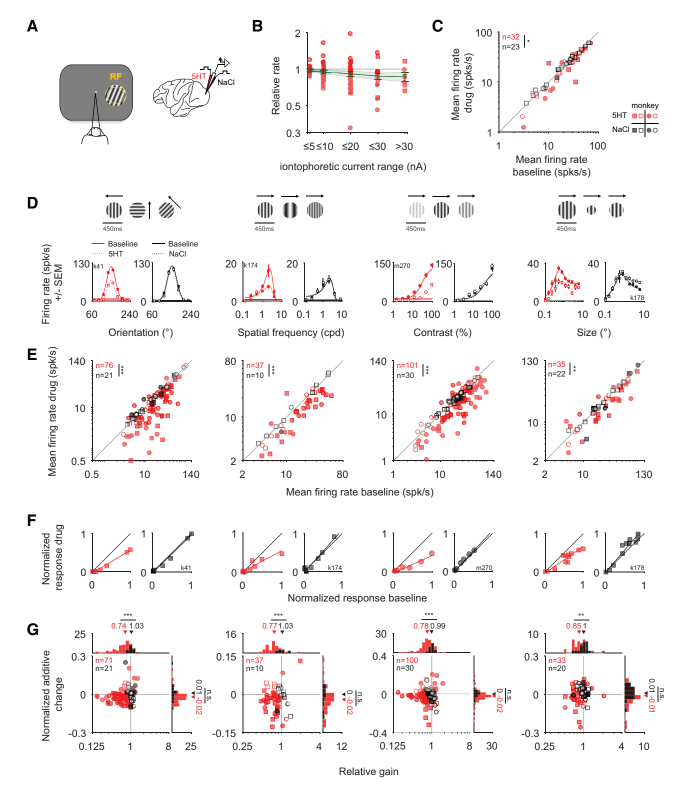


Figure 1. Iontophoretic application of serotonin leads mainly to a gain decrease. A, Schematic of the experimental setup. While the monkey performed a fixation task, grating stimuli were presented in a neuron's receptive field (RF) during blockwise iontophoretic application of serotonin (5HT; red) or pH-matched saline (NaCl; black). B, Dose dependence of the serotonergic modulation. The relative firing rate is plotted as a function of the applied iontophoretic current range of 90 serotonin blocks from n = 36 units. Each unit is shown only once per bin. The green line represents the geometric mean \pm SEM. C, Modulation for the subset of units for which the duration of unit isolation allowed for a full sequence of baseline, drug application, and recovery. Note that for four units for which tuning for > 1 stimulus dimensions was tested, only the first comparison is included. The median of the average firing rate decreased from 27 to 24 spikes/s for serotonin but not for saline application [20 spikes/s for the baseline, 21 spikes/s for saline application; p = 0.03, $p_{S-HT} = 32$, $p_{NACl} = 23$; 11 of 32 (34%) of units were significantly suppressed, 2 of 32 (6%) were significantly enhanced for serotonin application; 1 of 23 (4%) was significantly enhanced and suppressed for saline application]. p = 0.00, orientation, spatial frequency, contrast, and size were varied in blocks, and results are shown in the first, second, third, and last column, respectively. p = 0.00, p = 0.00

isolation and keep the monkey working, each block of drug application was followed by \geq 1 block of recovery before the next application of the drug. Full recovery was evaluated by observation during recording but verified statistically off-line: the responses to each stimulus parameter were z-scored to remove the stimulus-driven variability of the response, and the z scores for the baseline and recovery block were then required to be statistically indistinguishable (p > 0.05, Wilcoxon rank-sum test). The median time to achieve full recovery [possible for n = 90 serotonin experiments for which the inclusion criteria (see below) were met; this value includes multiple experiments per unit, e.g., for our dose-response measurements; Fig. 1B] after the serotonin application was 32 s (range, 15 to 1171 s). Note that these values reflect the upper bound of the times to full recovery since we occasionally inserted wait times of variable duration after the serotonin application experiment to ensure full recovery while keeping the animal motivated to perform additional trials afterward.

Stimuli. Visual stimuli were back-projected on a screen using DLP LED Propixx projector (1920×1080 pixels resolution; 30 cd/m^2 mean luminance; linearized gray values; run at 100 Hz/eye, combined with an active circular polarizer, DepthQ, run at 200 Hz) at a distance of 98 cm in front of the animals. The animals viewed the screen through passive circular polarizing filters monocularly or binocularly. Visual stimuli were generated in Matlab (Mathworks) using the Psychophysics toolbox (Brainard, 1997; Pelli, 1997; Kleiner et al., 2007).

Stimuli were circular drifting sinusoidal luminance gratings centered on and slightly exceeding a neuron's receptive field and presented for 450 ms (temporal frequency typically 7 Hz) binocularly or monocularly to the preferred eye. For each experimental block, either the direction (16 equally spaced values), the spatial frequency [eight logarithmically spaced values from 0.125 to 16 cycles per degree (cpd)], the contrast (typically seven logarithmically spaced values from 1.56 to 100%), or the size (typically 12 logarithmically spaced values from 0.3 to 8°) of the grating was pseudorandomly varied, randomly interleaved by blank stimuli, with all other parameters constant at approximately the preferred value for each unit. Within each experimental block each stimulus was typically presented 8–10 times.

For the subspace reverse correlation, we briefly flashed sinusoidal luminance gratings (flash duration typically 10 ms; 100% contrast; preferred spatial frequency; presented binocularly or monocularly to the preferred eye) that randomly varied in orientation (eight equally spaced values) and spatial phase (equally spaced values), randomly interleaved by blank stimuli (also 10 ms duration, presented with equal probability as each orientation).

Data analysis. All analyses were done in Matlab (Mathworks). For experiments using drifting gratings, we computed the mean firing rate during the 450 ms stimulus presentation. To obtain orientation tuning curves, we averaged the responses for directions 180° apart. Tuning curves were computed as the mean firing rate as a function of orientation, spatial frequency, contrast, and size, respectively, and fit with standard descriptive functions.

←

(Figure legend continued.) of saline. The size of the change in firing rate induced by serotonin or saline, respectively, differs statistically (for orientation: $p < 10^{-5}$; for spatial frequency: p <10⁻³; for contrast: $p < 10^{-3}$; for size: p = 0.001; all Wilcoxon ranked-sum tests). **F**, To quantify the size of the additive and multiplicative component of the change in the response (normalized to the peak response in the baseline condition), we performed linear (type II) regression on the tuning curves [baseline condition plotted on the abscissa against application of serotonin (red) or saline (black) on the ordinate]. The slope reflects the multiplicative change (relative gain) and the intercept the additive change. \boldsymbol{G} , For each cell, we compared the relative gain and the normalized additive change (normalized by the peak response in the baseline condition). Note that for all stimulus dimensions, the relative gain after applying serotonin is significantly smaller than in the control condition (for orientation: $p < 10^{-5}$, n = 71 for serotonin; n = 21 for saline; for spatial frequency: $p < 10^{-3}$; n = 37 for serotonin; n = 10 for saline; for contrast: $p < 10^{-4}$; n = 100 for serotonin; n = 30 for saline; for size: p = 0.003; n=33 for serotonin; n=20 for saline; all Wilcoxon ranked-sum tests). Circles and squares correspond to data from monkey M and K, respectively. Filled symbols represent units with significant mean response modulation (p < 0.05, Wilcoxon ranked-sum test). *: <0.05; **:<0.005; ***:<5*10 ⁻

Orientation tuning curves were fit with Gaussian functions for which all parameters were constrained to values >0. To avoid an overestimation of the amplitude resulting from placing the preferred orientation between two sparsely measured values, we further restricted the amplitude to be smaller than twice the peak value of the tuning curve.

Spatial frequency tuning curves were also fit with Gaussian functions, either in linear or logarithmic units, whichever resulted in better fits.

Contrast tuning curves were fit with the Naka–Rushton function (Albrecht and Hamilton, 1982): $R(c) = R_{\rm max} \, c^n / (c_{50}^n + c^n) + R_{\rm offset}$, where c is the stimulus contrast, c_{50} is the semisaturation contrast, and n is the exponent influencing the shape of the curve. All parameters were restricted to values >0, and additionally $R_{\rm max} + R_{\rm offset}$ could not exceed the peak response of the tuning curve to ensure that fits reached saturation within the contrast range of 0 to 100%. For the model comparison in Figure 6, we first fit the Naka–Rushton function to the baseline condition. In a second step, to account for the serotonin-induced modulation of the response, we allowed one parameter of this fit to the baseline to change: $R_{\rm max}$ for the response-gain model, or c_{50} for the contrast-gain model (Williford and Maunsell, 2006).

Size tuning curves were fit with a ratio-of-Gaussians function (Cavanaugh et al., 2002). All parameters had to be positive, and the width of the center Gaussian had to be less than or equal to that of the surround Gaussian function. The preferred size was defined as the smallest size that evoked 98% of the maximal response based on the fitted data (Nienborg et al., 2013). The suppression index was computed as the difference between the neuron's maximum response and the response to the maximum size, divided by the maximum response. Goodness of fit was quantified as variance explained and all fits had to explain ≥70% of the variance

For a subset of units we also measured temporal frequency tuning with stimulus presentations of 2 s each. We used these longer stimulus presentations to quantify the selectivity to spatial phase for each unit as the f1/f0 ratio (Skottun et al., 1991) measured at the preferred temporal frequency of that unit, where f1 corresponds to the amplitude of response modulation at the stimulus temporal frequency and f0 to the mean firing rate. For these analyses, eye movements within the fixation window were not factored out, which may have contributed to the modest phase selectivity across the population.

Orientation selectivity was quantified using the circular variance (Ringach et al., 2002). Direction selectivity was computed as a simple contrast metric $(r_{\theta}-r_{\theta-180})/(r_{\theta}+r_{\theta-180})$, where r_{θ} and $r_{\theta-180}$ correspond to the response at the neuron's preferred direction θ and that at 180° away from preferred, respectively.

Receptive field size was quantified as the mean of the equivalent widths (w; Bracewell, 1986) in both the vertical and horizontal directions: If A is the area under the horizontal or vertical receptive field profile, and h is the peak response of the receptive field profile, then w = A/h. The median equivalent width was 0.56° (range, 0.25 to 1.34°).

To evaluate significant response modulation during the serotonin or saline condition, we *z*-transformed the mean response for each stimulus condition and compared the *z* scores across all stimuli between the serotonin or saline condition and the baseline condition using Wilcoxon rank-sum test (two-sided, 5% significance threshold).

For the experiments with flashed gratings, we quantified the tuning curves using reverse correlation subspace analysis (Ringach et al., 1997). We smoothed the stimulus-triggered spike-density function (SDF) using a 4 ms boxcar. The SDFs were used to compute the mean number of spikes elicited by each frame to yield "orientation subspace maps," analogous to an approach described previously in the disparity domain (Nienborg and Cumming, 2009). SEs were estimated based on bootstrapping (1000 resamples). To estimate the dynamics of the orientationselective component of the response, we first computed the SD across the SDFs for each orientation. Response latencies were defined as the first point in time after frame onset for which the SD exceeded the half-height between the baseline variability and peak deviation (Lee et al., 2007). The baseline variability was computed as the average of the first 20 ms after frame onset. For inclusion of latency estimates, the orientation-selective response component had to exceed four times the baseline variability. Additionally, latencies were restricted to 25–100 ms after frame onset. To

ensure that changes in latency could not be explained by differences in the response variability between conditions, we estimated the dependence of the latency estimate on baseline variability by randomly subsampling the data in the baseline condition and computing latency for the subsampled data. The change in latency was then predicted from the change in baseline variability using linear regression, and the actually measured latency was corrected by the predicted change (see Fig. 5*D*). For comparison, we also quantified response latency using maximum likelihood estimation to identify the critical change point of the dynamics of the response (Friedman and Priebe, 1998). This approach assumes one Poisson process for the baseline response and one for the stimulusevoked component of the response, and we required significantly different Poisson processes for inclusion. This analysis yielded similar results.

Analysis of fixational eye movements. To examine potential effects on fixational eye movements, we explored the fixation precision as well as the amplitude and frequency of microsaccades. Microsaccades within the fixation window were identified as defined previously (Nienborg and Cumming, 2006, 2014; Clery et al., 2017). Fixation precision was defined by Cherici et al. (2012) as the area around the mean gaze position that encompassed the 75 th percentile of the gaze positions.

Noise correlations. Stimulus-independent covariability ["noise correlations," (Cohen and Kohn, 2011)] was calculated between the single-unit activity and the simultaneously measured multiunit activity recorded from the same electrode. Note that since this approach overestimates the absolute value of noise correlation (Ecker et al., 2010), we only explored the changes in noise correlation with serotonin, not the absolute value. To reduce the effect of slow fluctuations on nonstationarities (Ecker et al., 2014; Goris et al., 2014; Rabinowitz et al., 2015) resulting from the onset of the serotonin application, we removed the initial 20 stimulus presentations of each experimental block for this analysis. Responses for each stimulus condition were first z-scored, and noise correlations computed as the Pearson correlation coefficient of the z scores for single-unit and multiunit activity across all trials. Noise correlations have typically been found to depend on the neuronal spike rate (Cohen and Kohn, 2011). We estimated this dependence in our dataset for the baseline condition by linear regression and corrected the noise correlation by that predicted from the change in firing rate.

Fano factor. For each stimulus condition, we computed the Fano factor as the ratio between the variance and the mean response for each stimulus for each experimental block, excluding the initial 20 stimulus presentations to reduce variability from potential nonstationarities due to onset of serotonin. The average of the Fano factors for each stimulus was then calculated as the Fano factor for that unit.

Membrane potential-based model. The responses of the membrane potential-based model were explored for the stimulus used for orientation subspace mapping described earlier. The membrane potential was orientation selective and the selectivity described by a Gaussian function [amplitude, 20 mV; similar to empirically observed values (Priebe and Ferster, 2008)]. The width of the Gaussian function describing the orientation selectivity of the membrane potential (SD ranged from 10 to 40°) was chosen such that the bandwidth of the orientation subspace maps was within the range of those of the neuronal data in the baseline condition. The time-varying stimulus (a sequence of oriented gratings each flashed for 10 ms) induced fluctuations in membrane potential $V_m(t)$ that were convolved by a temporal kernel that consisted of a Gaussian temporal filter (chosen to be within the range of the neuronal data: SD, 6 ms; mean, 57 ms after stimulus onset to account for the lag of the response). Spike rates [k(t)] were derived from the following thresholdlinear function:

$$k(t) = \left\{ \begin{array}{cc} c \cdot (V_{\textit{m}}(t) - V_{\textit{thresh}}) & \textit{if } V_{\textit{m}}(t) > V_{\textit{thresh}} \\ 0 & \textit{otherwise} \end{array} \right.$$

where $V_m(t)$ denotes the membrane potential at time t, $V_{\rm thresh}$ is the spiking threshold, and c is a scalar value set to 15, approximating the value previously obtained for the cat striate cortex (Carandini, 2004). The spiking rate was converted to Poisson spike events, which were analyzed like the neuronal data.

The serotonergic modulation was imitated by a subtractive shift in the membrane potential (equivalent to changing $V_{\rm thresh}$). We explored shifts

over a range of 1 to 8 mV, similar to empirically observed changes in membrane potential in response to serotonin (Ko et al., 2016). We adjusted the number of stimulus repetitions to obtain a comparable level of baseline variability for the baseline and serotonin condition.

Inclusion criteria. For each unit, we required a minimum response to the neuron's preferred stimulus of 10 spikes/s, a minimum of four presentations per stimulus condition (except for the noise correlation and Fano factor analysis, where ≥ 8 presentations per stimulus condition were required), and that the neuron showed selectivity for the respective stimulus dimension (orientation, spatial frequency, contrast, size; evaluated by an ANOVA at a significance threshold of p < 0.01). To be included in the comparison of additive and multiplicative changes (Figs. 1, 2, 5), the type-II regression had to account for $\geq 70\%$ of the variance.

Results

Two macaque monkeys performed a standard fixation task while we recorded the activity of single units in their V1 during blockwise iontophoretic application of serotonin (Fig. 1A). We examined the effect of serotonin on the visual responses to drifting gratings that varied systematically in orientation, spatial frequency, contrast, or size, and to briefly flashed gratings of varying orientation (see Materials and Methods). We recorded a total of 265 single units in macaque V1 (118 from monkey M, and 147 from monkey K). To be included for further analysis, we required a minimum response to the neuron's preferred stimulus of 10 spikes/s, ≥4 presentations per stimulus condition, and that the neuron showed selectivity for the respective stimulus dimension as evaluated by an ANOVA at a significance threshold of p < 0.01. These criteria were passed by 229 units (108 for monkey M; 121 for monkey K). Of these, 206 (100 for monkey M; 106 for monkey K) were recorded with serotonin application and 65 (39 for monkey M; 26 for monkey K) with pH-matched saline (NaCl) application as control experiments (thus, in a subset of 42 units, we were able to measure responses for both serotonin and saline application in consecutive blocks; moreover, whenever possible, experiments along several stimulus dimensions were done on the same unit in consecutive blocks).

Serotonin predominantly decreases the responses in V1 by multiplicative changes of the tuning curves

The most salient consequence of the serotonin application was a substantial decrease in the visual responses. This effect was evident for the tuning curves for orientation, spatial frequency, contrast, and size, respectively, in four example neurons (Fig. 1D, red symbols). We found that these response changes could not be explained by the iontophoretic current application: control experiments with pH-matched saline (NaCl) did not result in such modulation of the neuronal responses (Fig. 1D, black symbols). Indeed, across the population, the mean firing rate in response to gratings of different orientations decreased for the serotonin condition (the median decrease was from 18 to 10 spikes/s) but not for the saline application (median values, 14 and 15 spikes/s, respectively), and the changes differed significantly between conditions [Fig. 1*E*; $p < 10^{-5}$, $n_{5\text{-HT}} = 76$, $n_{\text{NaCl}} = 21$; monkey M: $p < 10^{-3}$, $n_{5\text{-HT}} = 45$, $n_{\text{NaCl}} = 11$; monkey K: $p < 10^{-3}$, $n_{5\text{-HT}} = 10^{-3}$ 31, $n_{\text{NaCl}} = 10$; significant modulation for serotonin: 51 of 76 units (67%) were suppressed; 4 of 76 units (5%) were enhanced; significant modulation for saline: 4 of 21 units (19%) were suppressed; 3 of 21 units (14%) were enhanced]. These results were similar for gratings of varying spatial frequency, contrast, and size (Fig. 1E). For spatial frequency, the median average firing rate for serotonin decreased from 15 to 12 spikes/s and remained constant at 11 spikes/s for saline application [$p < 10^{-3}$, $n_{5-HT} = 37$, $n_{\text{NaCl}} = 10$; monkey M: p = 0.41, $n_{5\text{-HT}} = 11$, $n_{\text{NaCl}} = 2$; monkey

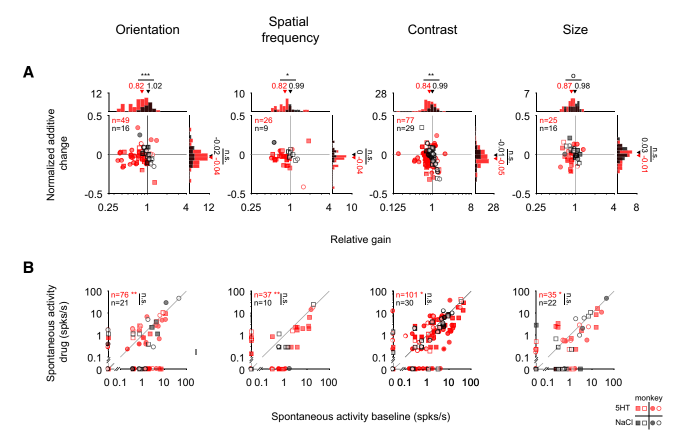


Figure 2. The gain decrease with serotonin does not merely reflect the iceberg effect. Symbols are the same as in Figure 1. Note that filled symbols reflect units with significant response modulation based on the full tuning curves as in Figure 1*E*. **A**, Comparing the gain and additive changes to the tuning curves as in Figure 1*G* but after restricting the datapoints to those whose responses were >0 spikes/s to explore the serotonergic modulation independent of the iceberg effect. In this reduced dataset, we also observe a significant gain decrease (relative gain for orientation: 0.82 for serotonin and 1.02 for saline, $p < 10^{-3}$, $n_{S-HT} = 49$, $n_{NaCl} = 16$; for spatial frequency: 0.82 for serotonin and 0.99 for saline, p = 0.016, $n_{S-HT} = 26$, $n_{NaCl} = 9$; for contrast: 0.84 for serotonin, 0.99 for saline, p = 0.003, $n_{S-HT} = 77$, $n_{NaCl} = 29$; for size: 0.87 for serotonin, 0.98 for saline, p = 0.075, $n_{S-HT} = 25$, $n_{NaCl} = 16$). **B**, The application of serotonin mainly resulted in a decrease of the blank response (blank interleaved in orientation ($p_{S-HT} = 0.005$, $n_{S-HT} = 76$; $p_{NaCl} = 0.35$, $n_{NaCl} = 21$), spatial frequency ($p_{S-HT} = 0.001$, $n_{S-HT} = 37$; $p_{NaCl} = 0.58$, $n_{NaCl} = 101$; $p_{NaCl} = 0.09$, $n_{NaCl} = 0.09$, $n_{NaCl} = 0.00$, $n_{S-HT} = 0.0$

K: p = 0.001, $n_{5-HT} = 26$, $n_{NaCl} = 8$; significant modulation for serotonin: 24 of 37 units (65%) were suppressed, 2 of 37 units (5%) were enhanced; significant modulation for saline: 1 of 10 units (10%) was suppressed]. For contrast tuning, the median firing rate decreased from 26 to 19 spikes/s for serotonin [the respective values for saline application were 23 and 22 spikes/s; $p < 10^{-3}$, $n_{5-HT} = 101$, $n_{NaCl} = 30$; monkey M: p = 0.04, $n_{5-HT} = 66$, $n_{NaCl} = 15$; monkey K: p = 0.004, $n_{5-HT} = 35$, $n_{NaCl} = 15$; significant modulation for serotonin: 58 of 101 units (57%) were suppressed, 8 of 101 (8%) units were enhanced; significant modulation for saline: 4 of 30 (13%) were suppressed]. Similarly, the mean firing rate decreased for the size tuning curves [29 to 25] spikes/s for serotonin; constant at 18 spikes/s for saline, p =0.001, $n_{5\text{-HT}} = 35$, $n_{\text{NaCl}} = 22$; monkey M: p = 0.65, $n_{5\text{-HT}} = 14$, $n_{\text{NaCl}} = 6$; monkey K: $p < 10^{-4}$, $n_{\text{5-HT}} = 21$, $n_{\text{NaCl}} = 16$; all Wilcoxon rank-sum tests; significant modulation for serotonin: 19 of 35 (54%) were suppressed, 5 of 35 (14%) were enhanced; significant modulation for saline: 2 of 22 (9%) were suppressed, 5 of 22 (23%) were enhanced].

In a subset of experiments, unit isolation was maintained and the animal worked for sufficiently long for a full sequence of baseline, drug application, and recovery. The results for this subset of experiments are shown in Figure 1C (same format as Fig. 1E), also supporting the significant decrease in firing when serotonin was applied. The proportion of units for which full, statistically defined (see Materials and Methods), recovery was

achieved was relatively small mainly because of sessions in which the monkeys were not motivated to continue to work or the unit isolation was lost. However, this relatively small proportion does not affect the interpretation of the main results of this paper, which relies on a comparison of the serotonergic modulation with that for pH-matched saline, and this comparison was robust when restricting the data to the subset of units with full recovery and saline control experiments (Fig. 1C). To additionally verify that a lack of recovery for some serotonin experiments did not impact the saline control experiments, we also performed two additional analyses. First, we compared the serotonergic modulation with that for saline for units for which only one substance (either serotonin or saline) per unit was applied. Second, we compared the serotonergic modulation for units with full recovery with the subset of the same dataset for which we also applied saline. For both analyses, we found a significant difference between the modulation for serotonin and that for saline (comparison across units: $p < 10^{-7}$, $n_{5-HT} = 53$, $n_{NaCl} = 23$ and comparison for units with full recovery: p = 0.04, $n_{5-HT} = 32$, $n_{NaCl} = 11$, respectively; note that for each unit only the first experiment per condition was included; Wilcoxon rank-sum tests). Moreover, our analysis of the responses to the interleaved blank stimuli revealed that the change in firing was not limited to the stimulus-driven response, but also observed in response to the blank stimuli (Fig. 2B).

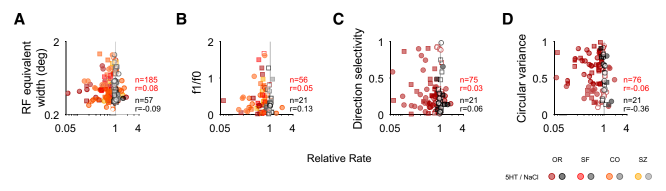


Figure 3. Probing the serotonergic modulation as a function of receptive field (RF) size and phase, direction, and orientation selectivity. **A–D**, Symbols are the same as in Figure 1. Shading represents the type of experiment for which the modulation by drug application [serotonin (5HT) or saline (NaCl)] was quantified. The modulation was quantified as the relative rate based on the average firing rates across the tuning curve (compare Fig. 1*E*) for the drug condition (compare Fig. 1*E*, ordinate), divided by that for the baseline condition (compare Fig. 1*E*, abscissa). Neither RF size, quantified as equivalent width (**A**); phase selectivity, quantified as f1/f0 (**B**); direction selectivity (**C**); nor orientation selectivity, quantified as circular variance (**D**), were correlated with the modulation by serotonin or saline. Note that low values for circular variance reflect strong orientation selectivity (all Spearman's rank correlation).

For the example tuning curves (Fig. 1D) serotonin seems to primarily scale down the tuning curve, indicative of a gain decrease (multiplicative effect). We then quantified across the population whether the effect of serotonin was primarily additive or multiplicative. To do so, we compared the tuning curves with and without serotonin application, fit by type-II regression (Fig. 1F). The slope and intercept of the regression line reflect the multiplicative (relative gain) and additive change. When comparing these values for each unit across the population, we found that the changes in the tuning curve were primarily accounted for by a gain decrease for each type of tuning curve. For orientation tuning curves, the median gain decreased to 74% after serotonin application, in contrast to saline (median relative gain for saline: 103%; $p < 10^{-5}$, $n_{5-HT} = 71$, $n_{NaCl} = 21$; monkey M: p < 0.01, $n_{5-HT} = 43$, $n_{NaCl} = 11$; monkey K: $p < 10^{-4}$, $n_{5-HT} = 28$, $n_{NaCl} = 11$ 10; Fig. 1G). Similarly, for spatial frequency, the gain reduced on average to 77% for serotonin (median relative gain for saline: 103%; $p < 10^{-3}$, $n_{5\text{-HT}} = 37$, $n_{\text{NaCl}} = 10$; monkey M: p = 0.23, $n_{5\text{-HT}} = 11$, $n_{\text{NaCl}} = 2$; monkey K: $p < 10^{-3}$, $n_{5\text{-HT}} = 26$, $n_{\text{NaCl}} = 26$ 8). For contrast tuning, the gain decreased on average to 78% for serotonin (for saline the median relative gain was 99%; $p < 10^{-4}$, $n_{5-HT} = 100, n_{NaCl} = 30$; monkey M: $p < 0.01, n_{5-HT} = 66, n_{NaCl} =$ 15; monkey K: $p < 10^{-4}$, $n_{5-HT} = 34$, $n_{NaCl} = 15$). As for the other stimulus dimensions, the median gain for size tuning was a decrease (to 85%) for serotonin while for saline the median relative gain was unchanged at 100% ($p=0.003, n_{5\text{-HT}}=33, n_{\mathrm{NaCl}}=20;$ monkey M: p = 0.78, $n_{5-HT} = 13$, $n_{NaCl} = 5$; monkey K: $p < 10^{-3}$, $n_{5\text{-HT}} = 20$, $n_{\text{NaCl}} = 15$; all comparisons Wilcoxon rank-sum tests). The normalized additive suppressive effect differed significantly from 0 for the serotonin but not for the saline conditions for orientation, spatial frequency, and contrast (for orientation: -0.02 for serotonin, $p < 10^{-3}$, n = 76; 0.01 for saline, p = 0.96, n = 21; for spatial frequency: median normalized additive change for serotonin was -0.02, p < 0.01, n = 37; change for saline was 0.00, p = 0.56, n = 10; for contrast the median normalized additive change for serotonin was -0.02, $p < 10^{-3}$, n = 100 and 0.00for saline, p = 0.45, n = 30; for size: the corresponding changes were -0.01 for serotonin, p = 0.62, n = 33, and 0.01 for saline, p = 0.26, n = 20; Wilcoxon signed-rank test) but given the variability in the control condition, the additive suppression for serotonin did not significantly exceed that for the control condition (for orientation: p = 0.12, $n_{5-HT} = 71$, $n_{NaCl} = 21$; for spatial frequency: p = 0.29, $n_{5-HT} = 37$, $n_{NaCl} = 10$; for contrast: p = $0.21, n_{5-HT} = 100, n_{NaCl} = 30;$ for size: $p = 0.20, n_{5-HT} = 33, n_{NaCl} = 0.20$

20; Wilcoxon rank-sum test). Moreover, the absence of an additive suppressive effect seemed to at least partially result from the "iceberg effect" (Carandini and Ferster, 2000): as a consequence of the spiking nonlinearity, spike rates are restricted to values >0 such that subtractive changes are limited by the minimum response of the tuning curve. In support of this, we found a significant correlation between the size of the normalized additive change and the minimum response of the tuning curve for the baseline condition (r = -0.41, $p < 10^{-8}$, $n_{5-HT} = 198$; Spearman's rank correlation; for this comparison we combined data for different stimulus dimensions but included only one data point per unit; data not shown). To verify that the observed gain decrease was not merely a consequence of the iceberg effect, we repeated the regression analysis after removing all data for which the responses were 0. In the subset of units for which the regression fits to the reduced dataset met the inclusion criteria (see Materials and Methods), we also found a significant gain decrease (Fig. 2A). Together, the additive and multiplicative changes provided an excellent fit to the data. Indeed, this simple linear model accounted for 90% of the variance (averaged across all regression fits for serotonin in Fig. 1G; note that the quality of the fits was similar for the saline controls: 92%). These results suggest that a simple linear transformation, predominantly multiplicative, can account for the serotonergic modulation of visual responses along several visual dimensions.

Given that serotonin receptors are differentially expressed across layers and on different cell types in the macaque V1 (Watakabe et al., 2009) the overall homogeneity of the effect is surprising. We therefore wondered whether the serotonergic modulation was systematically related to parameters that have been reported to vary to a certain degree with layer, such as receptive field size, orientation, direction, and spatial-phase selectivity (Ringach et al., 2002; Gur et al., 2005). To test this, we compared each of these parameters with the strength of serotonergic modulation (Fig. 3). Our analyses revealed no correlation of the serotonergic modulation with these parameters (r = 0.08, p = 0.28, n = 185 for equivalent width; r = 0.05, p = 0.72, n = 56for f1/f0 ratio; r = 0.03, p = 0.77, n = 75 for direction tuning index; r = -0.06, p = 0.62, n = 76 for circular variance; Spearman's rank correlation with serotonergic modulation ratio for each). Moreover, we did not observe a systematic difference in the serotonergic modulation for narrow spiking compared with broad spiking (defined by a spike waveform width of \leq 200 or \geq 200 μ s, respectively) units (p = 0.49; n = 206; Wilcoxon rank-sum test). Together,

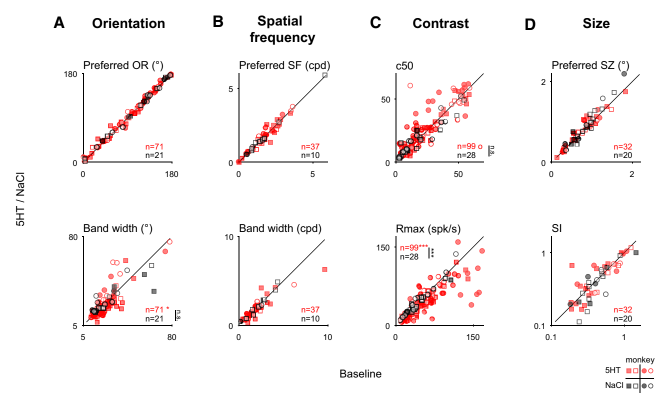


Figure 4. Serotonin does not systematically change the neuronal selectivity. Filled symbols reflect units with significant response modulation based on the full tuning curves as in Figure 1*E*. A–D, The neuronal tuning curves for orientation (A), spatial frequency (B), contrast (C), and size (D) were fit with descriptive functions (see Materials and Methods) to quantify changes in selectivity. Symbols are the same as in Figure 1. A, For orientation tuning, we compared the mean (preferred orientation, top) and the SD (orientation tuning width, bottom) of Gaussian fits. There was no significant change for either parameter that exceeded changes observed for saline (preferred orientation: p = 0.29, n = 71 for serotonin; p = 0.93, n = 21 for saline; p = 0.24 for the Wilcoxon ranked-sum test). B, For the spatial frequency tuning curves, neither the mean (preferred spatial frequency, top) nor the SD (bandwidth, bottom) of the fitted Gaussians differed significantly (mean: p = 0.50, n = 37 for serotonin; p = 0.92, n = 10 for saline; SD: p = 0.11, n = 37 for serotonin; p = 0.85, n = 10 for saline). C, For the contrast tuning, the contrast at half-maximum showed a nonsignificant trend toward higher values (top; C_{SO} ; p = 0.06, n = 99 for serotonin; p = 1.0, n = 28 for saline). The maximum response (R_{max} bottom) decreased significantly for serotonin, a = 28 for saline; Wilcoxon ranked-sum test). a = 0.30, a = 0.3

these analyses suggest that the observed serotonergic modulation was not restricted to a particular cell type or layer.

Serotonin leaves visual tuning properties largely unchanged

We next wondered whether the serotonin application additionally resulted in a systematic modulation of the visual encoding properties along one of the visual stimulus dimensions we explored. To this end we fit descriptive functions (see Materials and Methods) to the tuning curves and examined the effect of serotonin parametrized by these fits. For orientation tuning, the serotonergic modulation changed neither the preferred orientation (Fig. 4A, top; p = 0.29, $n_{5-HT} = 71$; monkey M: p = 0.34, $n_{5-HT} =$ 41; monkey K: p = 0.63, $n_{5-HT} = 30$) nor the orientation bandwidth (Fig. 4A, bottom; p = 0.05, $n_{5-HT} = 71$; monkey M: p =0.18, $n_{5-HT} = 41$; monkey K: p = 0.23, $n_{5-HT} = 30$; Wilcoxon paired sign-rank tests; note that the slight decrease in bandwidth did not significantly differ from the saline condition, p = 0.24, $n_{\text{NaCl}} = 21$). Similarly, serotonin did not alter the preferred spatial frequency (Fig. 4B, top; p = 0.50, $n_{5-HT} = 37$; monkey M: p =0.32, $n_{5-HT} = 11$; monkey K: p = 0.12, $n_{5-HT} = 26$) or spatial frequency bandwidth of the recorded neurons (Fig. 4B, bottom; p = 0.11, $n_{5-HT} = 37$; monkey M: p = 0.97, $n_{5-HT} = 11$; monkey K: p = 0.11, $n_{5-HT} = 26$; Wilcoxon paired sign-rank tests). However, the amplitude of the fits was significantly reduced for both orientation ($p < 10^{-4}$, $n_{5-HT} = 71$, $n_{NaCl} = 21$; monkey M: p <0.01, $n_{5\text{-HT}} = 41$, $n_{\text{NaCl}} = 11$; monkey K: p < 0.01, $n_{5\text{-HT}} = 30$, $n_{\text{NaCl}} = 10$) and spatial frequency ($p < 0.01, n_{5-\text{HT}} = 37, n_{\text{NaCl}} =$ 10; monkey M: p = 0.41, $n_{5-HT} = 11$, $n_{NaCl} = 2$; monkey K: p <0.01, $n_{5\text{-HT}} = 26$, $n_{\text{NaCl}} = 8$; all Wilcoxon sign-rank tests), as expected given the observed reduction in gain. Also as expected from the reduction in gain, we observed a significant reduction in R_{max} for contrast tuning (Fig. 4C, bottom; $p < 10^{-4}$, $n_{5\text{-HT}} = 99$, $n_{\text{NaCl}} = 28$; monkey M: p < 0.01, $n_{5\text{-HT}} = 66$, $n_{\text{NaCl}} = 14$; monkey K: $p < 10^{-3}$, $n_{5-HT} = 33$, $n_{NaCl} = 14$; Wilcoxon sign-rank test). Conversely, the sensitivity for changes in contrast (i.e., the steepness of the tuning curve parametrized by the exponent n in the Naka-Rushton function; see Materials and Methods), did not change (p = 0.15, n = 99; monkey M: p = 0.43, n = 66; monkey K: p = 0.19, n = 33; Wilcoxon paired sign-rank test; data not shown). Interestingly, we also found that the contrast that yielded half the maximum response (c_{50}) was slightly increased. We note that this trend did not reach statistical significance in the population and was only significant in one of the animals (p = 0.06, $n_{5-HT} = 99$; monkey M: p = 0.01, $n_{5-HT} = 66$; monkey K: p = 0.010.56, $n_{5-HT} = 33$). Nonetheless, this trend raises the question whether serotonin engages a similar mechanism as contrast gain control, which we will address in more detail below.

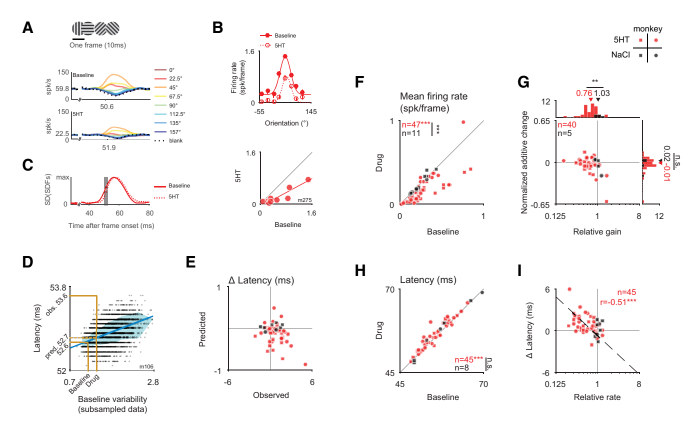


Figure 5. A slight modulation of the response dynamics by serotonin. Symbols are the same as in Figure 1. **A**, SDFs separated by orientation for one example neuron without (top) and with (bottom) application of serotonin obtained for random sequences of briefly (10 ms) flashed gratings of different orientations. **B**, The serotonergic application for this unit resulted in both a modest subtractive change and a divisive change of the tuning curve. **C**, To quantify the dynamics of the response, we computed the SD across the SDFs for each orientation. These SDs are superimposed for the baseline (solid) and serotonin condition. Latency was defined as the time to the half-maximum response. Note the slight increase in latency for the serotonin condition. **D**, In control analyses, we subsampled the data of the baseline condition to quantify changes in latency estimates due to changes in baseline variability. **E**, The predicted changes in latency from changes in baseline variability cannot account for the observed latency changes. Note the different scales of abscissa and ordinate. **F**, The mean firing rate for the orientation subspace stimulus decreases significantly with the application of serotonin (red, $p < 10^{-6}$, n = 47), but not for the saline (black, p = 0.70, n = 11). The change in mean firing with the iontophoretic application differs significantly between both conditions ($p < 10^{-4}$, n = 47 for serotonin, n = 11 for saline, Wilcoxon rank-sum test). **G**, The gain of the tuning curves decreases substantially more for the serotonin than for the saline condition (p = 0.006, n = 40 for serotonin, n = 5 for saline) while the subtractive change does not differ across conditions (p = 0.99, both Wilcoxon rank-sum test). **H**, The response latency increases slightly for serotonin ($p < 10^{-3}$, n = 45) but not for the saline condition (p = 0.55, n = 8). Note that due to the small sample for saline, this difference in modulation across conditions does not reach significance (p = 0.38).

Finally, we examined whether serotonin had a systematic effect on receptive field size, as suggested by previous work in anesthetized rats (Waterhouse et al., 1990). To address this question, we first compared the stimulus size for which the visual response was maximal ("preferred size") with and without application of serotonin. In contrast to the previous suggestion, we found no systematic change in the preferred size of the neurons (Fig. 4D, top; p = 0.70, $n_{5-HT} = 32$; monkey M: p = 0.41, $n_{5-HT} = 12$; monkey K: p = 0.30, $n_{5-HT} = 20$; Wilcoxon paired sign-rank tests). To explore the effect of center surround interactions, we also examined the degree to which the visual responses to large stimuli decreased the visual stimuli compared with stimuli of a neuron's preferred size ("suppression index"). Similar to the results for preferred size, we found no significant effect of serotonin on the neurons' surround suppression quantified by the suppression index (Fig. 4D, bottom; p = 0.54, $n_{5-HT} = 32$; monkey M: $p = 0.38, n_{5-HT} = 12$; monkey K: $p = 0.74, n_{5-HT} = 20$; Wilcoxon paired signed-rank tests).

Together these analyses corroborate our finding that the observed serotonergic modulation is dominated by a multiplicative change and modest additive change of the visual responses. Beyond that, serotonin leaves the receptive field properties largely unchanged.

Serotonin weakly increases the latency of the orientation selective response

We next wondered whether serotonin influenced the dynamics of the visual response, since previous work reported serotonergic influences on the dynamics of subcortical auditory processing in bats (Hurley and Pollak, 2005). To address this question, we used orientation subspace reverse correlation (Ringach et al., 1997), an approach that allows for detailed quantification of the dynamics of the orientation-selective response. The stimulus consisted of a random sequence of gratings of the same spatial frequency but different spatial phases and orientations, updated every 10 ms (see Materials and Methods). Figure 5A shows the average SDFs for each orientation for one example neuron for the baseline (top) and serotonin (bottom) blocks. We extracted tuning curves ("orientation subspace maps") from these SDFs (Nienborg and Cumming, 2009; Fig. 5B) and quantified the changes in tuning as additive and multiplicative changes using type-II regression. Similar to the results for drifting gratings, we found a significant decrease in the mean response (Fig. 5F; $p < 10^{-4}$, $n_{5-HT} = 47$, $n_{\text{NaCl}} = 11$; monkey M: p = 0.08, $n_{5\text{-HT}} = 34$, $n_{\text{NaCl}} = 3$; monkey K: $p < 10^{-3}$, $n_{5\text{-HT}} = 13$, $n_{\text{NaCl}} = 8$) that was largely explained by multiplicative changes in the tuning curves (Fig. 5G; relative gain: $p = 0.007, n_{5-HT} = 40, n_{NaCl} = 5$; monkey M: $p = 0.06, n_{5-HT} = 31$,

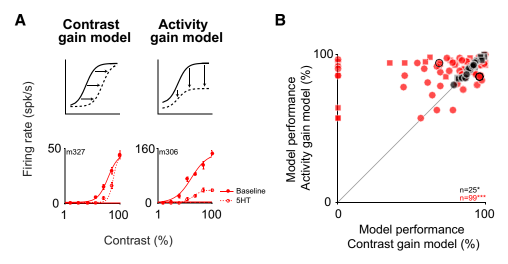


Figure 6. The serotonergic modulation cannot be accounted for by a contrast-gain model. $\textbf{\textit{A}}$, If the serotonergic modulation reflected a mechanism similar to contrast-gain control, the serotonergic modulation of the contrast tuning curve should result in a horizontal shift of the curve, without changing the response maximum (dashed line, top left), and an example neuron (bottom left) is compatible with such a change. Alternatively, under an activity-gain model (top right) serotonin application would result in a multiplicative downscaling of the tuning curve, as found for the example (bottom right). $\textbf{\textit{B}}$, Symbols are the same as in Figure 1. The activity-gain model in most cases explains a higher percentage of the variance than the contrast-gain model ($p < 10^{-6}$, n = 99; Wilcoxon paired test). The change in variance explained differs significantly (p = 0.02, n = 99 for serotonin, n = 25 for saline, Wilcoxon rank-sum test) between the serotonin (5HT; red) and the saline control condition (NaCl; black). Example units in $\textbf{\textit{A}}$ are marked by the black circles.

 $n_{\text{NaCl}} = 3$; monkey K: p = 0.15, $n_{5\text{-HT}} = 9$, $n_{\text{NaCl}} = 2$; additive chane: p = 0.99, $n_{5-HT} = 40$, $n_{NaCl} = 5$; monkey M: p = 0.86, $n_{5-HT} = 31$, $n_{NaCl} = 3$; monkey K: p = 0.44, $n_{5-HT} = 9$, $n_{NaCl} = 2$). To explore the dynamics of the orientation-selective response, we computed the SD across the SDFs over time (Fig. 5C). For the example cell, we find that the response latency for the orientationselective component of the response is slightly longer after serotonin application (Fig. 5A, C). Importantly, our control analysis (Fig. 5D, E; see Materials and Methods) reveals that this effect cannot be explained by the decreased signal-to-noise ratio owing to the reduced spike rate in the serotonin condition. Indeed, across the population, we observed a small but consistent and statistically significant increase in response latency (Fig. 5H; p < 10^{-3} , $n_{5-HT} = 45$; monkey M: p < 0.01, $n_{5-HT} = 33$; monkey K: p = 0.02, $n_{5-HT} = 12$), which was correlated with the size of the serotonergic modulation in firing rate (Fig. 5*I*; r = -0.51, p < 10^{-3} , $n_{5-HT} = 45$; monkey M: r = -0.56, $p < 10^{-3}$, $n_{5-HT} = 33$; monkey K: r = -0.20, p = 0.54, $n_{5-HT} = 12$).

The serotonin-induced changes differ quantitatively from contrast gain, but are accounted for by a simple membrane potential-based model

The main serotonin-dependent changes of the tuning curves (i.e., a largely divisive change in the response and a slight increase in the response latency) are reminiscent of the divisive reduction and phase delay when lowering contrast (Carandini et al., 1997), which are accounted for by a model using divisive normalization (Heeger, 1992; Carandini et al., 1997). We therefore wondered whether the serotonin-induced changes mimic a reduction in contrast, suggesting it may engage a mechanism similar to contrast normalization. To test this hypothesis, we therefore compared the performance of two descriptive models, a contrast-gain model and an activity-gain model (Fig. 6A). For the contrast-gain model, the modulation by serotonin would only mimic a change in contrast, resulting in a horizontal shift of the tuning curve (Fig. 6A, left). Conversely, for the activity-gain model, the modulation by serotonin would result in a downscaling of the entire tuning curve (Fig. 6A, right). While the contrast-gain model provided a better fit to the data for a small number of cells in support of the hypothesis that the modulation by serotonin engages a similar mechanism as contrast, the activity-gain model performed substantially better in most cases (Fig. 6B; $p < 10^{-6}$, $n_{5\text{-HT}} = 99$; monkey M: p < 0.01, $n_{5\text{-HT}} = 66$; monkey K: $p < 10^{-4}$, $n_{5\text{-HT}} = 33$; Wilcoxon paired signed-rank test). This indicates that modulation by serotonin relies on a mechanism that differs from contrast-gain control.

Finally we wondered whether a simple membrane potentialbased model could account for the divisive change in the response and the slight increase in response latency. Specifically, we explored whether a linear change at the level of the membrane potential would suffice to account for the observed effects by serotonin (Fig. 7). This membrane potential-based model consisted of an orientation-selective response at the level of the membrane potential followed by a temporal low-pass filter to fit the orientation bandwidth and average latency of the neuronal response to the stimulus used for orientation subspace reverse correlation (see Materials and Methods; Fig. 7A). The time-varying response of the membrane potential was then passed through a threshold-linear spiking nonlinearity. To account for the effect of serotonin, a subtractive shift was applied to the membrane potential [Fig. 7A, $V_m(t)$], moving it further away from the spiking threshold. Note that, while biophysically different, in our model this shift is equivalent to increasing the spiking threshold (V_{thresh}) of the spiking nonlinearity. We found that this shift could account for the observed changes in gain, the additive changes (Fig. 7D), as well as the small increase in latency (Fig. 7 B, E). A simple subtractive change of a threshold-linear spiking nonlinearity can therefore capture the serotonin-induced modulation of the visual responses.

Serotonin has no systematic effect on response variability or covariability

Neuromodulators, such as acetylcholine, have been implicated in affecting not only the magnitude (Disney et al., 2007) but also the variability of sensory responses (Pinto et al., 2013), and it has been hypothesized that serotonin mediates an additional, acetyl-

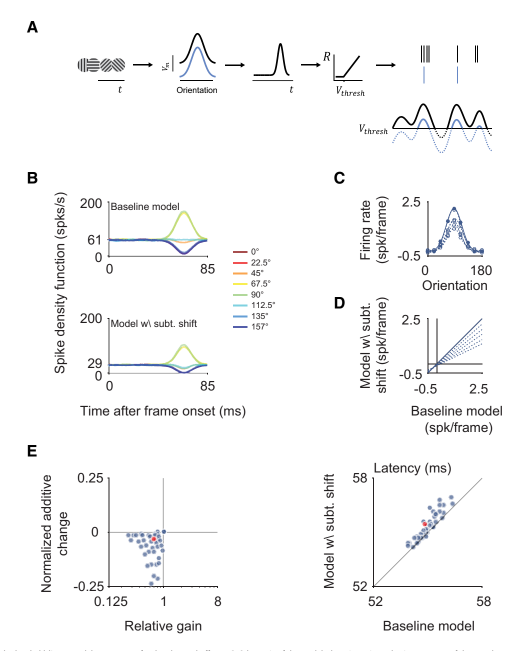


Figure 7. A simple threshold-linear model can account for the observed effects. **A**, Schematic of the model: the orientation-selective response of the membrane potential to the stimulus sequence is temporally low-pass filtered before being passed through a threshold-linear nonlinearity and a Poisson process to generate spikes. The only effect by serotonin is a subtractive shift to the membrane potential (blue lines) that is equivalent to a higher value of the threshold ($V_{\rm thresh}$). **B**, Example SDFs for the modeled baseline condition (top) and after a subtractive shift to the membrane potential aimed at mimicking the serotonin condition (bottom). **C**, Tuning curves for the baseline condition (solid) and for subtractive shifts of different sizes (dotted) are shown. **D**, Best fitting regression lines to quantify additive and subtractive changes are shown for different models. **E**, The subtractive change in the model leads to gain changes, additive changes (left), as well as latency shifts (right) in the range (blue data points) observed in our data (red circles).

choline-independent, mechanism of response desynchronization (Harris and Thiele, 2011). To test for this possibility, we measured the stimulus-independent response covariability ["noise-correlation" (Bair et al., 2001; Cohen and Kohn, 2011)] between the single-unit and the multiunit activity recorded on the same electrode. We note that measuring this correlation between single-unit and multiunit activity recorded on the same electrode has been shown to overestimate the absolute size of this correlation (Ecker et al., 2010). Nonetheless, this approach allows us to infer relative changes in noise correlation between the baseline and the serotonin condition. To reduce the effect of nonstationarities resulting from the onset of the serotonin application on noise correlations, we excluded the first 20 stimulus presenta-

tions of each experimental block for this analysis. Since fixational eye movements can affect the variability and covariability of visual neurons, we explored whether these differed systematically between the serotonin application and the saline controls. To this end, we compared the fixation precision (Cherici et al., 2012) as well as the frequency and amplitude of microsaccades within the fixation window (Fig. 8). For none of these metrics differed the modulation systematically between the serotonin and the saline application. Finally, we corrected noise correlation by that predicted from the change in firing rate (see Materials and Methods). In contrast to the hypothesis, we found no significant change in noise correlations for the serotonin condition (Fig. 9*B*; for orientation: p = 0.74, $n_{5-HT} = 63$ for serotonin; for spatial frequency:

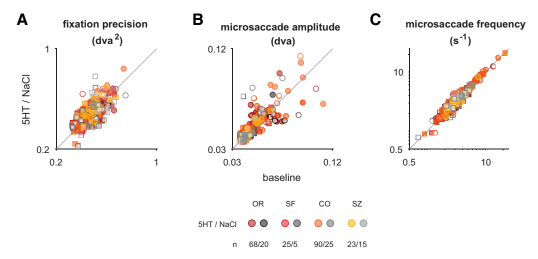


Figure 8. No systematic difference in eye movements between drug and baseline conditions. **A**, Fixation precision (see Materials and Methods) did not differ between conditions [median 0.49 degrees of visual angle squared (dva²) for both baseline and 5-HT application, respectively; median 0.49 dva² and 0.50 dva² for baseline and saline application, respectively; the change with drug application did not differ significantly between the saline and 5-HT experiments, p = 0.68]. **B**, Microsaccade amplitudes were similar between conditions (median 0.04 dva for baseline and serotonin application, respectively, $n_{S-HT} = 206$; median 0.04 dva for baseline and saline application, respectively, $n_{NACI} = 65$) and their changes did not differ significantly for serotonin versus saline application (p = 0.43). **C**, Similarly, the frequency of microsaccades did not change between conditions (median, $3.4 \, \text{s}^{-1}$ for baseline and serotonin application, respectively, $n_{S-HT} = 206$; median, $3.2 \, \text{and} \, 3.3 \, \text{s}^{-1}$ for baseline and saline application, respectively), and the changes did not differ between conditions (p = 0.96; $n_{S-HT} = 206$, Microsaccades and saline application, respectively), and the changes did not differ between conditions (p = 0.96; $n_{S-HT} = 206$, Microsaccades and saline application, respectively), and the changes did not differ between conditions (p = 0.96; $n_{S-HT} = 206$, Microsaccades and saline application, respectively, $n_{S-HT} = 206$; median, $n_{S-HT} = 206$; median, n

 $n_{5-HT} = 2$ for serotonin; for contrast: p = 0.07, $n_{5-HT} = 45$ for serotonin; for size: p = 0.59, $n_{5-HT} = 15$ for serotonin; all Wilcoxon paired signed-rank test). We also did not observe a systematic relationship between the size of effect on firing rate and noise correlation (Fig. 9D; for orientation: r = 0.07, p = 0.57, $n_{5-HT} =$ 63; for spatial frequency: $n_{5\text{-HT}} = 2$; for contrast: r = -0.02, p = 0.88, $n_{5\text{-HT}} = 45$; for size: r = -0.13, p = 0.66, $n_{5\text{-HT}} = 15$; all Spearman's rank correlation). Finally, we also explored the effect of serotonin on the variability of the sensory response, quantified as Fano factor, and observed no systematic change in the two conditions (Fig. 9A; for orientation: p = 0.77, $n_{5-HT} = 61$; for spatial frequency: $n_{5\text{-HT}} = 1$; for contrast: p = 0.81, $n_{5\text{-HT}} = 42$; for size: p = 0.17, $n_{5\text{-HT}} = 11$; all Wilcoxon paired signed-rank tests). Moreover, the change in mean firing rate was not associated with a systematic change in Fano factor (Fig. 9C; for orientation: r = -0.22, p = 0.08, $n_{5-HT} = 61$; for spatial frequency: $n_{5-HT} = 1$; for contrast: r = -0.14, p = 0.38, $n_{5-HT} = 42$; for size: r = 0.19, p = 0.58, $n_{5-HT} = 11$; all Spearman's rank correlation). Additionally, there was no systematic change in the difference in Fano factor in response to a neuron's preferred stimulus compared with that to a blank stimulus (Fig. 9E). Together, these analyses indicate that the application of serotonin leaves the response variability and covariability in macaque V1 largely unchanged.

Discussion

Here we combined extracellular recordings and iontophoresis in V1 of awake macaques to explore the modulatory effects of serotonin on visual processing. We found that across a variety of visual stimulus dimensions, the modulation by serotonin across the neuronal population was surprisingly uniform and (1) was dominated by a decrease in the neurons' response gain, (2) showed a slight slowing of the dynamics of the response, and (3) resulted in no systematic change of the neuronal variability, covariability, or stimulus selectivity. Our observed effects could be captured by a descriptive model in which serotonin caused a simple additive change at the level of the threshold-linear spiking nonlinearity.

A surprisingly uniform effect despite receptor and cellular diversity

In this study, we focused on the serotonergic modulation in the awake macaque of functional tuning properties along four well characterized visual dimensions (orientation, spatial frequency, contrast, and size) that are encoded by neurons in V1. We focused on awake animals using the endogenous ligand while previous work examined the modulation to receptor-specific ligands in the striate cortex of anesthetized macaques and, in contrast with the predominantly suppressive effect we found, observed variable modulation (Watakabe et al., 2009). This apparent discrepancy may in part reflect a net effect between facilitation and suppression mediated by different receptor classes all activated by serotonin. Additionally, our recordings in the awake animal likely mimimized fluctuations in brain state found under anesthesia (Ecker et al., 2014) and avoided effects of anesthesia on serotonergic neurons (Johansen et al., 2015). Moreover, the focus on parametrized functional properties may have contributed to our ability to identify an effect—a gain decrease—that is surprisingly functionally uniform across the population and stimulus dimensions. This effect dominated despite variability on a neuron-byneuron basis, which was also observed previously in anesthetized animals (Waterhouse et al., 1990; Watakabe et al., 2009). In light of the known diversity of receptor expression on different cell types in the macaque V1 (Watakabe et al., 2009), this main observed effect is likely mediated by different cellular or network mechanisms. For example, in the input layers of V1, receptors 5-HT_{1B} and 5-HT_{2A} are expressed by the majority of excitatory neurons, but not by GABAergic neurons, and are typically coexpressed by the same neurons in layer IVC (Watakabe et al., 2009). The predominantly divisive effect of serotonin may therefore result from an interaction of the receptor activation in excitatory neurons. A previous study applied a 5-HT_{1A} agonist to layers IV and V in V1 of anesthetized macaques and also observed a decrease of multiunit activity (Rauch et al., 2008). Given the lack of 5-HT_{1A}-receptor expression in these layers in macaque V1

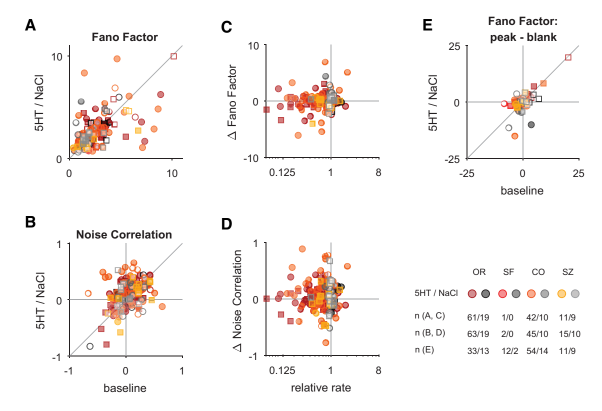


Figure 9. Serotonin application does not systematically change the Fano factor or noise correlation. Warm and gray colors depict experiments with serotonin (5HT) and saline (NaCl) application, respectively, as indicated. The number of units per experiment and condition included are specified. Filled symbols depict units with significant modulation (compare Fig. 1*E*). Circles and squares correspond to data from monkey M and K, respectively. *A*, The variability of the neuronal response quantified by the Fano factor does not change systematically for the application of serotonin orientation: p = 0.77, n = 61 for serotonin; p = 0.55, n = 19 for saline; spatial frequency: n = 1 for serotonin; n = 0 for saline; contrast: p = 0.81, n = 42 for serotonin; p = 0.13, n = 10 for saline; size: p = 0.17, n = 11 for serotonin; p = 0.57, n = 9 for saline; all Wilcoxon paired signed-rank test). Note the similar distribution for serotonin and saline application in A-D. B, The single-unit multiunit noise correlation (corrected for changes in firing rate; see Materials and Methods) is compared for the baseline condition (abscissa) and the application of serotonin or saline (ordinate). The serotonin application did not result in a significant change of noise correlation across the population (orientation: p = 0.74, n = 63 for serotonin; p = 0.15, n = 19 for saline; spatial frequency: n = 2 for serotonin; n = 0 for saline; contrast: p = 0.07, n = 45 for serotonin; p = 0.38, n = 10 for saline; size: p = 0.59, n = 15 for serotonin; p = 0.15, n = 19 for saline; spatial frequency: n = 2 for serotonin; n = 0 for saline; contrast: n = 0.07, n = 0.07,

(Watakabe et al., 2009), this decrease in activity may at least partially reflect a cross-activation of the densely expressed 5-HT_{1B} receptors. This modulation was thought to be mediated by a hyperpolarization of the membrane potential, analogous to the membrane potential-based model we present here. Serotoninmediated hyperpolarization of the membrane potential was also recently identified at the axon initial segment in auditory neurons in gerbils (Ko et al., 2016). Conversely, in layer II, where the receptors are expressed on a subset of both excitatory and inhibitory neurons, but typically not coexpressed (Watakabe et al., 2009), the decrease in gain may reflect network interactions between inhibition and excitation. Different circuit elements may also include vasoactive intestinal peptide-positive (VIP+) interneurons that express the 5-HT_{3A} receptor (Rudy et al., 2011), although in the macaque V1 5-HT_{3A}-receptor expression has only been observed for layers 5/6 (Watakabe et al., 2009), while VIP+ interneurons are most pronounced in layers 2/3 (Gabbott and Bacon, 1997). The same functional computation—gain control—may therefore be implemented by different components of the neuronal hardware within the same neuromodulator system serotonin, at least on the time scale we examined.

Indeed, since we applied serotonin in blocks in the minute time scale, the modulation likely mimics changes in the tonic level of discharge of serotonergic neurons, implicated in signaling the contextual valence on such slow time scales, and providing different signals from the phasic responses (Cohen et al., 2015). Interestingly, for short (<1 s) phasic serotonergic stimulation, a divisive modulation was previously observed only for the spontaneous response, not the sensory-driven response, in the olfactory bulb in anesthetized mice (Lottem et al., 2016), in contrast with our findings. Conversely, a decrease in gain of the sensory response in the mouse olfactory bulb was observed for sustained (>30 s) serotonergic activation (Petzold et al., 2009), similar to the modulation observed in this study.

Behaviorally, decreasing the gain of a sensory response can contribute to a delay or reduction in the response to sensory stimulation, as has been previously observed for a reduced startle response in rats (Davis et al., 1980; Sipes and Geyer, 1994), or reduced mechanosensory responsivity in mice (Dugué et al., 2014). Although these behavioral findings have typically been interpreted in the context of a serotonergic role for motor or emotional processing (Crockett et al., 2009; Cools et al., 2011; Correia et al., 2017), the observed decreased sensory gain suggests at least partially a sensory involvement. It may also reflect a sensory signature of how serotonin promotes waiting (Miyazaki et al., 2014;

Ranade et al., 2014; Fonseca et al., 2015) by lowering the salience of sensory input.

On this slow (minute) time scale, serotonin may be complementary to the action of cholinergic neuromodulation, which has been found to increase the gain of the visual input in macaque V1 (Disney et al., 2007), and has been linked to mediating spatial attention (Herrero et al., 2008). Notably, acetylcholine in macaque V1 was found to substantially modulate the gain but not the variability of the response (Herrero et al., 2008), a dissociation that is mirrored by our results here. It suggests that distinct mechanisms are available to modulate the variability and level of the sensory responses, although cognitive states, such as spatial attention, typically affect both (Cohen and Maunsell, 2009; Mitchell et al., 2009).

Potential implications for the serotonergic role in visual hallucinations

An influential perspective of perception is that it reflects an inference process (Helmholtz, 1867; Gregory, 1980; Lee and Mumford, 2003; Yuille and Kersten, 2006), in which internal beliefs about the world are combined with the incoming sensory evidence (Lee and Mumford, 2003; Yuille and Kersten, 2006). Mounting physiological evidence suggests that some of this combination of internal ("top-down") and external sensory signals occur already at the level of sensory neurons (Fiser et al., 2010; Nienborg and Roelfsema, 2015; Cumming and Nienborg, 2016). In this framework, psychiatric diseases such as schizophrenia (Friston, 2005) are associated with an imbalance between internally generated (top-down) and externally driven feed-forward signals. Hallucinations, for example, have been suggested to arise from an imbalance toward internally generated over externally driven sensory signals (Friston, 2005; Notredame et al., 2014; Jardri et al., 2016; Schmack et al., 2017). Such an imbalance may also explain, for example, visual hallucinations during visual impairment (Charles Bonnet syndrome; Gold and Rabins, 1989), when the feed-forward visual input is degraded. Similarly, decreasing the gain of the visual input could shift the balance toward the internally generated signals, and thus result in visual hallucinations. The decrease in gain by serotonin we found here may therefore shed light onto the mechanism by which the visual cortex is involved in hallucinations caused by serotonergic hallucinogens (Bressloff et al., 2002; de Araujo et al., 2012; Kometer et al., 2013; Carhart-Harris et al., 2016; Kometer and Vollenweider, 2016).

Conclusion

To our knowledge this is the first study to explore the role of serotonergic modulation of neuronal activity in the sensory cortex of awake animals. The modulatory effect we observed across the population was surprisingly homogeneous—a simple decrease in the response gain of the neural activity. Such gain modulation is an important component of the cortical computation (Salinas and Thier, 2000) that controls the responses without changing the receptive field properties. It is therefore well suited to modulate the responses according to the animal's internal state, e.g. influenced by the valence of the contextual environment (Cohen et al., 2015).

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