The tuning of tuning: how adaptation influences single cell information transfer

Fleur Zeldenrust¹⁽²⁾*, Niccolò Calcini ²⁽²⁾, Xuan Yan³, Ate Bijlsma⁴, Tansu Celikel⁵

1 Donders Institute for Brain, Cognition, and Behaviour, Radboud University, Nijmegen - the Netherlands

2 Maastricht Centre for Systems Biology (MaCSBio), University of Maastricht, Maastricht, The Netherlands

3 Institute of Neuroscience, Chinese Academy of Sciences, Beijing, China

4 Department of Population Health Sciences / Department of Biology, Universiteit Utrecht, the Netherlands

5 School of Psychology, Georgia Institute of Technology, Atlanta - GA, U.S.A.

These authors contributed equally to this work.

* fleur.zeldenrust@donders.ru.nl

Abstract

Sensory neurons reconstruct the world from action potentials (spikes) impinging on them. To effectively transfer information about the stimulus to the next processing level, a neuron needs to be able to adapt its working range to the properties of the stimulus. Here, we focus on the intrinsic neural properties that influence information transfer in cortical neurons and how tightly their properties need to be tuned to the stimulus statistics for them to be effective. We start by measuring the intrinsic information encoding properties of putative excitatory and inhibitory neurons in $L^{2/3}$ of the mouse barrel cortex. Excitatory neurons show high thresholds and strong adaptation, making them fire sparsely and resulting in a strong compression of information, whereas inhibitory neurons that favour fast spiking transfer more information. Next, we turn to computational modelling and ask how two properties influence information transfer: 1) spike-frequency adaptation and 2) the shape of the IV-curve. We find that a subthreshold (but not threshold) adaptation, the 'h-current', and a properly tuned leak conductance can increase the information transfer of a neuron, whereas threshold adaptation can increase its working range. Finally, we verify the effect of the IV-curve slope in our experimental recordings and show that excitatory neurons form a more heterogeneous population than inhibitory neurons. These relationships between intrinsic neural features and neural coding that had not been quantified before will aid computational, theoretical and systems neuroscientists in understanding how neuronal populations can alter their coding properties, such as through the impact of neuromodulators. Why the variability of intrinsic properties of excitatory neurons is larger than that of inhibitory ones is an exciting question, for which future research is needed.

Author summary

Intracellular information transfer from synaptic input to output spike train is necessarily lossy. Here, we explicitly measure the mutual information between a neuron's input and spike output and show that information transfer is more lossy and heterogeneous for excitatory than for inhibitory neurons. By using computational modelling we show that the shape of the input-output curve as well as how fast a neuron adapts to its input collectively determine the rate of information loss. These insights will help both experimentalists and modellers in designing and simulating experiments that investigate how network coding properties can adapt to the environment, for instance through the effects of neuromodulators.

Introduction

Perception and other brain functions require information transmission and signal transformation at each processing step. Specifically, for perception, stimuli that impinge on sensory receptors are transferred via the brain stem and thalamus to cortical networks: each of these processing steps results in information transfer and compression, due to intracellular information transfer from synaptic input current to spike train. The spike train of a single neuron though, can contain only a limited amount of information about an incoming stimulus [1]. However, the working range of a neuron is typically limited, more limited than the range of inputs a neuron might receive. A neuron's ability to adapt its working range to the properties of the stimulus is crucial for its ability to transfer information about the stimulus to the next processing level [2-5]. For example, if the input amplitude is too low, a neuron that cannot adapt will not respond, whereas when the input amplitude is too large, a neuron that cannot adapt will enter depolarization block or its output firing rate will be saturated, both resulting in a neuron that does not respond adequately to changes in input and hence in a neuron that does not transfer information. Therefore, neurons need to continually adapt their working range (i.e. their excitability) in order to fit the dynamic range of the input. They can do this by reducing synaptic strength [6,7] or by shifting (gain shift) or widening (gain modulation) their intrinsic excitability [8–10]. This changing of the intrinsic input-output curves happens on different timescales: from fast (spike frequency adaptation [11]) to slow (homeostatic scaling, for reviews see [7, 12]). The dynamics of such adaptation mechanisms impact the effectiveness of the adaptation in relation to the stimulus dynamics: if the adaptation is too fast (relative to the input statistics), it has no practical effect, but if it is too slow, it is constantly saturated and has no dynamic effects. Here, we focus on the relatively fast adaptive changes in intrinsic excitability and ask how such mechanisms influence information transfer in cortical neurons and how tight their properties need to be tuned to the stimulus statistics for them to have an effect.

We start by measuring the intrinsic information encoding properties of putative excitatory (regular-spiking) and inhibitory (fast-spiking) neurons in L2/3 of the mouse barrel cortex. We measure the effects of several intrinsic neural characteristics on the information transfer from input current to output spike train, using a combination of ex-vivo experiments [13, 14] and computational modelling. We aim to unravel how both the threshold behaviour and the I-V curve shape of excitatory and inhibitory neurons affect information transfer, using a recently developed method to estimate the mutual information between input and output in an ex-vivo setup [15]. This method has several advantages: instead of the traditionally long (~ 1 hour) experiments that are needed to obtain a single mutual information estimate [16–20], this method needs only about 5 minutes of recording to obtain an information transfer estimate. Moreover, the properties of the input current can be adapted to fit different cell type properties, and it has an optimal observer model so that the measured information transfer can be compared with the 'optimal' Bayesian Neuron information transfer [21]. Using this method, we can simultaneously measure the information transfer from input current to

10

11

12

13

14

15

16

17

18

19

20

21

22

23

24

25

26

27

28

29

30

31

32

33

34

35

36

37

38

40

41

42

output spike train and assess intrinsic cell properties, thereby showing how intrinsic cell properties correlate with information transfer. In particular, putative excitatory neurons show high thresholds and strong adaptation, making them fire sparsely and resulting in a strong compression of information between input and output. Their intrinsic properties are quite heterogeneous, showing a large variability. Putative inhibitory neurons on the other hand have intrinsic properties that favour higher firing rates, corresponding to a higher information processing rate. Their response properties are more stereotypical than those of excitatory neurons.

The Bayesian neuron that is optimal for this task has two properties that distinguish it from a standard leaky integrate-and-fire model: 1) spike-frequency adaptation and 2) a non-linear I-V curve that results amongst others in the suppression of hyperpolarization. To untangle how these mechanisms influence information transfer, we turn to computational modelling. Firstly, we use an exponential integrate-and-fire (expIF) model with two types of adaptation: subthreshold adaptation [22,23] and threshold adaptation [24] and research the effects of these two types of adaptation on the information transfer in the aforementioned mutual information protocol. We find that subthreshold adaptation increases the information transfer if tuned well, whereas threshold adaptation increases the working range of the neuron over a broad range of parameters. So despite the fact that at first glance these to forms of adaptation appear to serve a similar purpose (i.e. reducing the firing rate of a neuron for strong stimuli), it turns out that their effects are quite different. Secondly, we assess the effects of changing the shape of the I-V curve (the right-hand-side of the membrane voltage equation). We model the effects of suppression of hyperpolarization by adding an instantaneous 'h-current' to the expIF neuron, the effects of an instantaneous subthreshold potassium current, and the effects of changing the leak conductance of the neuron. We find that a well-tuned subthreshold (but not threshold) adaptation, the 'h-current', and a properly tuned leak conductance can increase the information transfer of a neuron, whereas threshold adaptation can increase its working range.

Materials and methods

Experiments

All analyzed current clamp and simulation data and the code to analyze and simulate them can be found in this repository: https://doi.org/10.34973/4f3k-1s63. The voltage clamp data are part of the dataset of da Silva Lantyer et al. (2018) [13].

Ethics statement

Animals used were Pval-cre and SSt-cre mice from 9 to 45 weeks kept with unlimited access to water and food, housed in a 12-hour light/dark cycle. All experimental procedures were performed according to Dutch law and approved by the Ethical Committee for Animal Experimentation of Radboud University (RU DEC) as described before (for further details, see [25,26]). Each mouse was perfused with iced and oxygenated $(95\% O_2/5\% CO_2)$ Slicing Medium (composition in mM: 108ChCl, 3KCl, $26NaHCO_3$, $1.25NaH_2PO_4H_2O$, 25 Glucose. H_2O , $1CaCl_2.2H_2O$, $6MgSO_4.7H_2O$, 3 Na-Pyruvaat) under anaesthesia with 1,5ml Isoflurane.

Slice electrophysiology

The brain was covered in 2% agarose and submerged in a Slicing Medium after which it was sliced in 300 μM thickness using a VF-300 compresstome (Precisionary Instruments LLC) and then incubated for 30 min in 37°C artificial cerebrospinal fluid (ACSF,

44

45

46

47

48

49

50

51

52

53

54

55

56

57

58

59

60

61

62

63

64

65

66

67

68

69

70

71

72

73

75

76

77

78

79

80

81

82

83

84

85

composition in mM: 1200NaCL, 35KCL, $13MgSO_4.7H_2O$, $25CaCl_2.2H_2O$, 100Glucose. H_2O , $12.5NaH_2PO_4.H_2O$, $250NaHCO_3$), oxygenated ($95\%O_2/5\%CO_2$). The bath was then transferred to room temperature. Slices were allowed to accomodate to room temperature for 30 min and were kept in this bath until use. Slices were placed into the recording chamber under the microscope (Eclipse FN1, Nikon) and perfused continuously at a rate of 1 ml/min with the oxygenated ACSF at room temperature. Patch pipettes for whole-cell recordings were pulled from borosilicate glass capillaries, 1.0 mm outer diameter, 0.5mm inner diameter, on a pipette-puller (Sutter Instrument Co. Model P-2000), until an impedance of $8\pm 2 M\Omega$ for the tip was obtained. Pipettes were filled with a solution containing (in mM) $115CsMeSO_3$, 20CsCl, 10HEPES, $2.5MgCl_2$, $4Na_2ATP$, 0.4NaGTP, 10Na—phosphocreatine, 0.6EGTA, 5QX - 314(Sigma). The whole cell access was obtained after reaching the gigaohm seal and breaking the membrane. Upon entering the cell and the whole-cell mode, the membrane potential was kept fixed at -70mV, outside stimulation.

Input current generation

Data acquisition was performed with HEKA EPC9 amplifier controlled via HEKA's PatchMaster software (version 2.90x.2), and subsequent analysis with MatLab (Mathworks, v.2016b). Three types of experiments were performed: current clamp (CC) step-and-hold, current clamp (CC) frozen noise, and voltage clamp (VC).

The current clamp (CC) step-and-hold protocol was performed in every cell and used to distinguish between cell types, according to the firing rate and spike shape (Fig 1). The protocol consisted of clamping the neuron at a baseline current I_{baseline} , corresponding to the one required to keep its membrane at -70mV, and providing a 500ms long stimulus of fixed current value $I = I_{\text{baseline}} + (40pA * \text{step number})$, for a total of 10 steps, reaching a maximum current injected of $I_{\text{baseline}} + 400pA$. Between each current injection step, a 5.5s recovery window was allowed.

Information transfer was measured using the 'frozen noise' method introduced by Zeldenrust et al. [15]. To measure information transfer from input to spike train in short periods of time, instead of the long measurements needed for the traditional methods (see Introduction), the noisy input current injected into a neuron in the current clamp setting was generated as the output of an artificial neural network (ANN) that responded to a randomly appearing and disappearing preferred stimulus or 'hidden state' (Markov process) x: a binary variable that can take the values of 1 (preferred stimulus present, 'on-state') and 0 (preferred stimulus absent, 'off-state', see Fig 2A). This hidden state is switched on and off according to a Markov process with rates $r_{\rm on}$ and $r_{\rm off}$. The advantage of using such a binary hidden state stimulus is that there is no need to reconstruct the full input current (which is high-dimensional and therefore requires long recordings), but it is sufficient to reconstruct the binary stimulus, for which less data is needed. The N = 1000 neurons of the ANN fired Poisson spike trains, whose firing rates were modulated by the hidden state stimulus, so that each neuron fired with rate q_{on}^i when x = 1, and q_{off}^i when x = 0. These rates were drawn from a Gaussian distribution with mean μ_q (see Table 1) and standard deviation $\sigma_q = \sqrt{\frac{1}{8}\mu_q}$. Each spike was convolved with an exponential kernel with a unitary surface and a decay time of 5ms. The spike trains from different presynaptic neurons contribute to the output with weight $w_i = \log(\frac{q_{on}^i}{q_{off}^i})$. In order to choose the parameters of this input current two considerations needed to be made:

1. The Markov process had rates $r_{\rm on}$ and $r_{\rm off}$, which correspond to a switching time 136 constant $\tau_{\rm input} = \frac{1}{r_{\rm on}+r_{\rm off}}$: since excitatory neurons do not fire at rates higher 137 than a few Hz, but the inhibitory neurons show a much broader working range, up 138

90

91

92

93

94

95

96

97

98

99

100

101

102

103

104

105

106

107

108

109

110

111

112

113

114

115

116

117

118

119

120

121

122

123

124

125

126

127

128

129

130

131

132

133

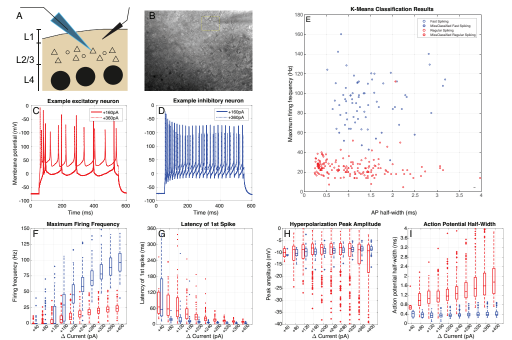


Fig 1. Cell Classification with the CC step-and-hold protocol. A, B: We selected neurons to record from the mouse somatosensory cortex (barrel cortex), in L2/3. Visually, the shape and size of soma were a good indicator of the cell type: smaller and roundish shapes would point towards fast-spiking neurons, while slightly larger and triangular shapes would point to regular spiking (putative excitatory) neurons. C: Example responses of an excitatory cell to a constant injected current. D: Example responses of an inhibitory cell to a constant injected current. E: Cell classification using agglomerative clustering based on the maximum firing frequency and spike width. Cells were classified as inhibitory (blue) when they had a small spike half-width combined with a high maximum firing rate and as excitatory (red) with a large spike half-width and low maximum firing rate. In pink the cell(s) where the agglomerative clustering and the initial classification disagreed (see Materials & Methods). F: Maximum firing frequency distribution for incremental current injection amplitudes for inhibitory (blue) and excitatory (red) neurons. G: Same as F, but for the latency of the first spike. H: After-hyperpolarization distribution. I: Spike half-width distribution. For threshold behaviour in the current-clamp step-and-hold protocol, see Supplementary Fig. S1.

> to 100 Hz (Fig 1), the information transfer could not be measured in the two 139 types of neurons with the same switching speed of the hidden state τ_{input} , but the 140 switching speed of the hidden state needed to be adapted to the working range of 141 the neuron type. Fortunately, the method allows for keeping the information 142 about the hidden state in the input constant while changing τ_{input} , by adjusting 143 the mean firing rate of the ANN μ_q (see Fig 2B). So the information transfer in 144 the two neuron types could still be compared by choosing a time constant τ_{input} of 145 50 or 250 ms for inhibitory/fast spiking or excitatory/regular spiking neurons 146 respectively, with matching values of μ_q of 0.5 Hz or 0.1 Hz respectively (see Table 147 1, so that the mutual information between the input current and the hidden state 148 (MI_I) was about 0.3 bit (Fig 2E). This target mutual information between the 149 input current and the hidden state was chosen so that the input current was 150 informative about the hidden state, but not too informative. 151

2. The input generated by the ANN responding to the Markov Process $(I_{\text{Markov}}(t))$ is dimensionless. Therefore, this dimensionless theoretical "input current" needed to be scaled to Ampère so that it could be injected into the neuron in a current clamp setup. Therefore, the injected current was defined as $I_{\text{injected}} = I_{\text{hold}} + I_{\text{scale}} I_{\text{Markov}}(t)$. Here, the neuron was clamped a baseline current I_{baseline} , corresponding to the current required to keep its membrane at -70mV, and I_{scale} was set at 2100 pA for excitatory and 700 pA for inhibitory cells (see Table 1).

The scripts for generating this current can be found in this GitHub repository: https://github.com/DepartmentofNeurophysiology/Analysis-tools-forelectrophysiological-somatosensory-cortex-databank.

Table 1. Input parameters for the ex-vivo experiments.

Parameter	Excitatory cells	Inhibitory cells
Number of artificial neurons N	1000	1000
Hidden state time constant τ_{input}	250 ms ($r_{\rm on} = 1.3$ Hz, $r_{\rm on} = 2.7$ Hz)	50 ms ($r_{\rm on} = 6.7$ Hz, $r_{\rm on} = 13.3$ Hz)
Average firing rate artificial neurons μ_q	0.1 Hz	0.5 Hz
Baseline input current I_{baseline}	(set so the cell was at -70 mV , see Fig 2)	(set so the cell was at -70 mV , see Fig 2)
Amplitude input current I_{scale}	2100 pA	700 pA
Analysis window size	100 s	20 s
Number of measured cells	144	72 (+ 9 control)
Number of trials	220	78 (+11 control)

Analysis

Cell classification

Cells were classified using the following procedure: before the frozen noise injection, for 165 each cell, the response to a current-clamp step (CC-step) protocol was recorded. From 166 these recordings, the maximum firing rate, the average spike-halfwidth, and the average 167 after hyperpolarization (AHP) amplitude were extracted (Fig 1). On-site, the cells were 168 classified by the experimenter based on the firing rate and the spike width. Based on 169 this initial classification the cell received the frozen noise input current with either 170 $\tau_{\text{input}} = 250 \text{ ms}$ (excitatory neurons) or $\tau_{\text{input}} = 50 \text{ ms}$ (inhibitory neurons). Offline, 171 the initial classification was verified using an agglomerative clustering protocol 172 (MATLAB 'clusterdata') to cluster the data into 2 groups (separated following Ward's 173 method(Ward, 1963)), according to the maximum firing rate and the average spike-half 174

6	/33
0	/ 33

163

152

153

154

155

156

157

158

159

160

161

162

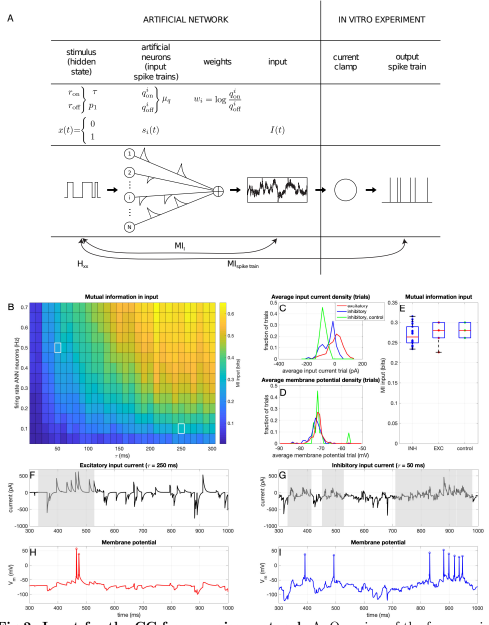


Fig 2. Input for the CC frozen noise protocol. A: Overview of the frozen noise method for input generation and measurement of mutual information (copied with permission from [15]) B: Mutual information between the input current and the hidden state, for different values of the switching speed of the hidden state (τ_{input}) and the average firing rate of neurons in the ANN (average over 10 trials). The white squares denote the used values for the input for the inhibitory (top left) and excitatory (bottom right) neurons. C: Average (over the trial) input current and D: membrane potential for all trials. Green data points/lines denote the control experiments where the inhibitory neurons received the input current that was otherwise given to the excitatory neurons. E: Mutual information between the hidden state and the input current, for all trials. Note that because frozen noise was used, every frozen noise trial was actually the same. Therefore, there are not many different realizations and hence not many different MI values. F: Example injected frozen noise current for an excitatory neuron. The grey shaded area corresponds to times when the hidden state was 1. G: Example injected frozen noise current for an inhibitory neuron. H: Example resulting membrane potential of an excitatory neuron. I: Example resulting membrane potential of an inhibitory neuron

width (normalized to zero mean and unit standard deviation) reached during the 175 CC-Step protocol (Fig 1E). There was a single cell where the initial classification and 176 the post-hoc classification were in disagreement (Fig 1E, pink star). We decided to keep 177 this cell in the original (inhibitory) group due to its position between the two clusters. 178

Calculation of mutual information

The mutual information between the hidden state and the input (MI_I) or a spike train $(MI_{\rm spike train})$ was estimated with the help of the hidden state x (see Input current generation and Fig 1A). The method was explained in detail in Zeldenrust et al. (2017) [15] and followed derivations from Denève (2008) [21] and Lochmann and Denève (2011) [27]). Code for how to calculate the mutual information can be found in the following repository: https://github.com/DepartmentofNeurophysiology/Analysis-tools-for-electrophysiological-somatosensory-cortex-databank as well as with the data.

In short, the mutual information was calculated using the following steps. The estimated log-odds ratio \hat{L} that the hidden state is 1, given the history of the input until now I(t) can be estimated by integrating the following differential equation (see [15,21] for the derivation):

$$\frac{d\hat{L}}{dt} = r_{\rm on}(1+e^{-\hat{L}}) - r_{\rm off}(1+e^{\hat{L}}) + I(t) - \theta, \tag{1}$$

where $\theta = \sum_{i=1}^{N} q_{\text{on}}^{i} - q_{\text{off}}^{i}$ is the constant offset of the input, which is chosen to equal to 0 by generating the input as explained before (drawing q_{on}^{i} and q_{off}^{i} from a Gaussian distribution). Using the estimate of the log-odds ratio from equation (1) over time, we can now estimate the conditional entropy by averaging over time:

$$\hat{H}_{xy} = \langle x \log_2 \left(\frac{1}{1 + e^{-\hat{L}}} \right) + (1 - x) \log_2 \left(1 - \frac{1}{1 + e^{-\hat{L}}} \right) \rangle_{\text{time}}.$$
 (2)

Because the hidden state follows a memory-less Markov process, its entropy at every moment in time equals

$$H_{xx} = P_1 \log_2(P_1) - (1 - P_1) \log_2(1 - P_1).$$
(3)

Here, $P_1 = \frac{r_{\text{on}}}{r_{\text{on}} + r_{\text{off}}}$ is the prior probability that the hidden state equals 1. With the canonical $MI = H_{xx} - H_{xy}$, the mutual information between the input and the hidden state can now be estimated. Similarly, the mutual information between a spike train and the hidden state can be estimated by integrating equation 1 where the input I(t) is now replaced by the spike train input given by 201

$$I_{\text{spike train}}(t) = w \cdot \rho(t), \tag{4}$$

where $\rho(t)$ is the spike train of the neuron and, and its weight w is given by

$$w = \log_2 \frac{\hat{q}_{\text{on}}}{\hat{q}_{\text{off}}} = \log_2 \left(\frac{\# \text{ spikes while } x = 1}{\# \text{ spikes while } x = 0} \cdot \frac{\text{total duration } x = 0}{\text{total duration } x = 1} \right).$$
(5)

Parameter $\theta = \sum_{i=1}^{N} q_{\text{on}} - q_{\text{off}}$ is calculated similarly based on the observed q_{on} and q_{off} in the spike train.

Note that even though theoretically $MI \leq 0$, due to our approximation it can happen that our estimate of $H_{xy} > H_{xx}$, due to the integration method, and hence that we find a small negative value of the MI between a spike train and the hidden state. These are often cells that fire either at very low rates or have firing patterns that for 200

179

180

181

182

183

184

185

186

187

188

189

190

195

196

other reasons deviate from a Poisson-like response (i.e. cells that stop firing during the experiment). However, to maintain a complete overview of the data, we decided not to discard those that have a low firing rate. However, we did exclude files with negative values in the input current (where the wrong input current was saved) and files with a vanishing firing rate.

In this manuscript, we mostly report on the unitless 'Fraction of Information' (FI) in the output spike train:

$$FI = \frac{MI_{\rm spike train}}{MI_{\rm input}}.$$
(6)

214

215

219

220

221

222

223

224

225

226

227

The FI quantifies how much information about the hidden state is transferred from the input current to the spike train, and thus quantifies which fraction of the information is kept during the spike-generating process. 218

Threshold detection

The membrane potential threshold of each recorded spike in the Current Clamp (CC) experiments was determined from the experimentally recorded membrane potential using the method explained in [24]: in a window from 1 to 0.25 ms before each spike maximum, the earliest time in the window at which either the first derivative exceeded 18 mV/ms or the second derivative exceeded 140 mV/ms2 was designated as the threshold-time, and the threshold value was determined as the corresponding membrane potential of that time point.

ROC curves

A Receiver Operator Characteristic (ROC) curve shows how well a system can be 228 classified into two binary classes by comparing the number of correctly detected 229 positives or 'hits' to the number of false positives or 'false alarms' depending on a 230 threshold parameter. Here, we assumed every trial had its own threshold, and we 231 defined a 'hit' as a period during which the hidden state was 1, in which at least 1 232 action potential was fired, and a 'miss' as a period during which the hidden state was 1, 233 in which no action potentials were fired. Similarly, we defined a 'false alarm' as a period 234 during which the hidden state was 0, in which at least 1 action potential was fired, and 235 a 'correct reject' as a period during which the hidden state was 0, in which no action 236 potentials were fired. So each period in which the hidden state was 1, was either defined 237 as a 'hit' or a 'miss', and each period in which the hidden state was 0, was either 238 defined as a 'false alarm' or a 'correct reject'. The total number of hits was divided by 239 the total number of periods during which the hidden state was 1, which resulted in the 240 fraction of hits $0 \le f_h \le 1$. Similarly, the fraction of 'misses', 'false alarms', and 'correct 241 rejects' were defined as the fraction of periods during which the hidden state was 1 but 242 no spike was fired, the fraction of periods during which the hidden state was 0, but a 243 spike was fired and the fraction of periods during which the hidden state was 0, and no 244 spike was fired, respectively. We calculated the fractions of hits, misses, false alarms, 245 and correct rejects for each spike train, as well as for a corresponding Poisson spike 246 train of the same length and with the same number of spikes. Note that for these 247 Poisson spike trains, the hit fraction is actually below the hit fraction = false alarm 248 fraction line, due to the nature of the hidden state: because the hidden state is more 249 often 0 than 1 ($P_1 < 0.5$), a random spike will have a higher chance of occurring during 250 a period where the hidden state equals 0. Therefore, the false alarm fraction will be 251 higher than the hit fraction for Poisson spike trains. 252

Fitting of exponential functions

In the results, we fit saturating functions to how FI (eq (6)) depends on different input variables $x \in \{I_{\text{scale}}, r, r_n\}$:

$$FI(x) = FI_{\max}\left(\frac{2}{1 + e^{-\lambda_x(x - x_{\text{offset}})}} - 1\right),\tag{7}$$

where r is the firing rate of an output spike train, and $r_n = r \cdot \tau_{\text{input}}$ is the unitless firing rate normalized by the switching speed of the hidden state τ_{input} . We fit parameters FI_{max} and λ , and in the case of $x = I_{\text{scale}}$ also $I_{\text{scale, offset}}$ (in the case of $x \in \{r, r_n\}$ the offset value is set equal to 0). To fit these curves, we use Matlab's 'fit' function, which automatically calculates 95 % confidence intervals. When the data does not only saturate, but decreases again after the maximum, we include only data up to the maximum.

Calculation of membrane capacitance and conductance using dynamic IV curves

We used the derivation of Badel et al. (2008) [28] to calculate the membrane capacitance (C_m) and conductance (g_m) for each analysis window. In short, we calculated $\frac{dV_m}{dt}$ from the recorded traces, and calculated the variance of $\frac{I_{\text{inj}}}{C_m^*} - \frac{dV_m}{dt}$ between the values of $-76 \leq V_m \leq -74 \text{ mV}$ for different values of C_m^* . The membrane capacitance C_m was determined as the value of C_m^* for which the variance was minimized :

$$C_m = \arg\max_{C_m^*} \operatorname{Var}(\frac{I_{\text{inj}}}{C_m^*} - \frac{dV_m}{dt})$$

Next, we defined the membrane current as

$$I_m = I_{\rm inj} - C_m \frac{dV_m}{dt}$$

and binned the $I_m - V_m$ curve in bins of 5 mV and calculated the average for each bin. A linear fit was made for subthreshold voltage values ($-200 \le V_m \le -60$ mV), and the slope was defined as the membrane conductance g_m . We excluded files where this fit did not succeed (the value of g_m was found to be negative).

Spike-triggered average

The whitened and regularized spike-triggered average (STA) was calculated as

$$STA(t) = (X^T X \lambda \mu_{X^T X} I) \setminus (X^T \rho)$$
(8)

where X is a stimulus-lag matrix, where each row is the stimulus vector with a different $_{277}$ lag (see [29–32]), $X^T X$ is the correlation matrix and I is the identity matrix. Operator $_{1}^{278}$ (denotes multiplication with the inverse, and T denotes a transpose. Parameter λ is a regularization (i.e. smoothing) parameter which was set to 10 and $\mu_{(X}^T X)$ is the mean of the diagonal of the correlation matrix. Finally, ρ denotes the spike train. The resulting STA was normalized with the L_2 norm.

Following the derivation of Slee et al. (2005) [33], the inner product of all 283 spike-triggering stimuli with the STA was calculated for each trial (P(stimulus—spike), 284 the posterior distribution), as well as the inner product of the same number of 285 random-triggered stimuli (P(stimulus), the prior distribution. With the 286 random-triggered stimuli, the prior distribution of the input was calculated, and 287

270

253

256

257

258

259

260

261

262

263

264

275

> compared to the distribution of spike-triggering stimuli (the posterior distribution). With Bayes' law, the shape of the threshold function could be calculated:

$$P(\text{spike}|\text{stimulus}) \sim P(\text{stimulus}|\text{spike})/P(\text{stimulus})$$
 (9)

However, if the distributions are not smooth due to limited sampling, the threshold function cannot be calculated. Therefore, the difference in mean between the prior and posterior was calculated for each neuron, and the distribution of means over all neurons was shown.

To assess the variability between the STAs calculated for each cell, we calculated the inner product between all pairs of STAs of inhibitory cells and between all pairs of STAs of excitatory cells. Because excitatory cells fire less, the STAs are based on a lower number of spikes for excitatory cells than for inhibitory cells. This in itself could introduce a higher variability of the STAs. To control for this, we also calculated the STAs for the inhibitory cells based on a comparable number of spikes as for the excitatory cells: we matched each inhibitory trial to an excitatory trial and reduced the number of spikes by only including the first spikes until they had the same amount of spike, and discarding the rest. Subsequently, we calculated the STA based on this reduced number of spikes, normalized them, calculated the posterior and prior for these, and calculated the inner product between all pairs of these STAs.

Simulations

We performed two types of simulations: an optimal observer for this experiment, the optimal 'Bayesian neuron' [21], and a more biologically realistic exponential integrate-and-fire (expIF) neuron with subthreshold and/or threshold adaptation [22–24].

Optimal observer: Bayesian Neuron

Next to the possibility of information estimation in short time windows, the in vitro information transfer method [15] has another advantage: the availability of an optimal observer model. This 'Bayesian neuron' [21] is a spiking neuron model that optimally integrates evidence about the hidden state from the ANN described above. It is optimal given an efficient coding or redundancy reduction assumption: it only generates new spikes if those spikes transfer new information about the hidden state, that cannot be inferred from the past spikes in its spike train. In practice, the neuron performs a leaky integration of the input, in order to calculate the log-odds ratio L for the hidden state being 1 (NB Note the similarity with equation 1):

$$\frac{dL}{dt} = r_{\rm on}(1+e^{-L}) - r_{\rm off}(1+e^{L}) + I(t) - \theta,$$
(10)

where $r_{\rm on}$ and $r_{\rm off}$ are the switching speeds of the hidden state, and $\theta = \sum_{i} (i=1)^{N} q_{\rm on}^{i} - q_{\rm off}^{i}$ is the constant offset of the input generated by the ANN as before, which is chosen to be equal to 0 in this paper. The neuron compares this log-odds ratio from the input, L, with the log-odds ratio of the hidden state being 1 inferred from its own spike train, G:

$$\frac{dG}{dt} = r_{\rm on}(1+e^{-G}) - r_{\rm off}(1+e^{G}).$$
(11)

Each time the log-odds ratio based on the input (L) exceeds the log-odds ratio based on the output spike train (G) by an amount $\frac{\eta}{2}$, a spike is fired:

if
$$L > G + \frac{\eta}{2} : \begin{cases} a \text{ spike is fired} \\ G \to G + \eta \end{cases}$$
 (12)

For the optimal observer, the parameters of the Bayesian neuron $(r_{\rm on}, r_{\rm off}, \theta)$ are the 327 same as the ones chosen for the hidden state and the ANN for generating the input. As 328 the model is made as an optimal observer for this input, the input does not have to be 329 scaled ($I_{\text{baseline}} = 0$; $I_{\text{scale}} = 1$), making η the only free parameter of the Bayesian 330 neuron model. This precision parameter η describes the distance between the threshold 331 and the reset of the Bayesian neuron, or in other words, the precision with which G332 tracks L. This parameter is varied in order to obtain the different firing rates in the 333 Results section (from 0.25 to 6 in steps of 0.25). Note that this neuron model has a 334 form of threshold adaptation: if it did not spike for a long time, G decays to its prior 335 value $G_{\text{prior}} = \log \frac{r_{\text{on}}}{r_{\text{off}}}$. With each spike, G is increased by η , and more input (larger L) is needed to fire a spike, thereby reducing its firing rate. 336 337

The simulated neurons received the same frozen noise input as used in the experiments (see Input current generation), but unscaled ($I_{\text{baseline}} = 0$; $I_{\text{scale}} = 1$). The simulations were performed in Matlab, using a standard forward Euler with a time step of 0.05 ms.

ExpIF neuron with (sub)threshold adaptation and non-linear I-V curve

In order to obtain more biologically interpretable results, and to disentangle the subthreshold and threshold effects of adaptation, we used the expIF neuron with subthreshold [22, 23] and/or threshold [24] adaptation. The equations are given by

$$C_m \frac{dV_m}{dt} = (g_L(E_L - V_m) + g_L \Delta_T e^{\frac{(V_m - \theta)}{\Delta_T}} + I(t) - w + I_{\{h,K\}})$$
(13)

$$\tau_w \frac{dw}{dt} = a(V_m - E_L) - w \tag{14}$$

$$\tau_{\theta} \frac{d\theta}{dt} = \theta_{\infty} - \theta, \tag{15}$$

(16)

338

339

340

341

342

343

344

345

where w describes the subthreshold adaptation and the threshold θ decays to a steady state 346

$$\theta_{\infty} = p * (V_m - V_i) + V_t + K_a * \log(1 + e^{\frac{(V_m - V_i)}{k_i}}).$$
(17)

A spike is defined when V_m passes a cutoff value V_{cutoff} and is reset to a reset potential $V_r = V_t + 5\Delta_T$: ³⁴⁸

if
$$V_m > V_{\text{cutoff}}$$
:

$$\begin{cases}
\text{a spike is fired} \\
V_m \to V_r \\
w \to w + b
\end{cases}$$
(18)

Moreover, we added the following instantaneous currents to the right-hand side of the membrane potential equation (13) in order to simulate non-linearities in the I-V curve: ³⁵¹

$$I_h(V_m) = g_h k_{\infty,h}(V_m)(V_h - V_m)$$
 (19)

$$k_{\infty,h}(V_m) = \frac{1}{1 + e^{\frac{V_{\text{half}}^h - V_m}{k_h}}}$$
(20)

and

$$I_K = g_K k_{\infty,K} (V_m) (V_K - V_m) \tag{21}$$

$$k_{\infty,K}(V_m) = \frac{1}{1 + e^{\frac{V_{\text{half}}^K - V_m}{k_K}}}$$
(22)

> To assess the effect of adaptation, we simulated 4 parameter regimes: 1) no 353 adaptation, 2) subthreshold adaptation only, 3) threshold adaptation only, and 4) 354 combined adaptation (both subthreshold and threshold adaptation), with the 355 parameters given in Table 2. To assess the effect of a non-linear I-V curve, we simulated 356 3 parameter regimes: 1) we added an instantaneous hyperpolarization-activated 357 depolarizing current, similar to an h-type current: 'vary g_h ', 2) we added an 358 instantaneous depolarization-activated hyperpolarizing current, similar to a 359 subthreshold potassium current: 'vary g_K ', 3) and we varied the leak-conductance: 360 'vary g_L ', with the following parameters given in Table 3. Note that for large values of 361 $I_{\rm scale}$ the simulations diverge: the membrane potential diverges and no further spikes 362 are fired. These simulations are not included in the analyses. 363

$\mathbf{regime} \rightarrow$	no	subthreshold	threshold	combined
$\mathbf{parameter}\downarrow$	adaptation	adaptation	adaptation	adaptation
C_m	50 pF	50 pF	50 pF	50 pF
$E_L = V_r$	-70 mV	-70 mV	-70 mV	-70 mV
g_L	10 nS	10 nS	10 nS	10 nS
Δ_T	1 mV	1 mV	1 mV	1 mV
$ au_w$	n/a	varied	n/a	varied
a	0 nS	4 nS	0 nS	4 nS
b	0 nA	0.0805 nA	0 nA	0.0805 nA
$ au_{ heta}$	n/a	n/a	varied	varied
p	0	0	0	0
V_i	-67 mV	-67 mV	-67 mV	-67 mV
V_t	-63 mV	-63 mV	-63 mV	-63 mV
K_a	0 mV	0 mV	5 mV	5 mV
k_i	5 mV	5 mV	5 mV	5 mV
g_h	0 nS	0 nS	0 nS	0 nS
g_K	0 nS	0 nS	0 nS	0 nS
$I_{\rm baseline}$	0 nA	0 nA	0 nA	0 nA
$I_{\rm scale}$	varied	varied	varied	varied

Table 2. Parameters for the adaptive expIF model with (sub)threshold adaptation

Parameters for equations 13 - 18.

The simulated neurons received the same frozen noise input as used in the experiments (see Input current generation), but with a different scaling (see Tables 2 and 3). Simulations were performed in Brian 2 [34], using a standard forward Euler with a time step of 0.025 ms.

Results

The goal of this research was to explore the relationship between intrinsic excitability and information transfer. To that end, we first performed 'classical' step-and-hold current clamp experiments in the mouse barrel cortex. Next, we used the 'frozen noise' protocol [15] to measure information transfer together with adaptive properties in these two cell types. After that, we turn to computational modelling to entangle how different biophysical mechanisms influence information processing in model cells. Finally, we return to the experimental recordings to verify the results obtained from computational modelling.

376

364

365

366

367

368

369

370

371

$\mathbf{regime} \rightarrow$	vary g_h	vary g_K	vary g_L
$\mathbf{parameter} \downarrow$			
C_m	$50 \mathrm{pF}$	50 pF	50 pF
$E_L = V_r$	-70 mV	-70 mV	-70 mV
g_L	10 nS	10 nS	varied
Δ_T	1 mV	1 mV	1 mV
$ au_w$	n/a	n/a	n/a
$ au_{ heta}$	n/a	n/a	n/a
g_h	varied	0 nS	0 nS
$V_{\rm half}^h$	-82 mV	n/a	n/a nS
k_h	-9 mV	n/a	n/a
V_h	-30 mV	n/a	n/a
g_K	0 nS	varied	0 nS
$V_{ m half}^K$	-60 mV	n/a	n/a nS
k_K	9 mV	n/a	n/a
V_K	-70 mV	n/a	n/a
I_{baseline}	0 nA	0 nA	0 nA
$I_{\rm scale}$	varied	varied	varied

Table 3. Parameters for the expIF model with non-linear I-V curve

Parameters for equations 13 - 21.

Information transfer in inhibitory and excitatory neurons

Excitatory neurons fire at low rates

Whole-cell recordings were made from pyramidal cells and interneurons in layer 2/3(L2/3) of mouse barrel cortical slices [13]. Cells were classified as either 'excitatory' or 'inhibitory' based on their electrophysiological responses to a standard current-step protocol (Fig 1, see Materials & Methods). In response to depolarizing steps, excitatory neurons show strong spike-frequency adaptation, limiting their maximum firing rate (Fig 1 and Supplementary Table S1, see also [13]), whereas inhibitory neurons fire at much higher rates.

To measure the information transfer from input current to output spike train, 386 traditionally long (~ 1 hour) experiments were needed to obtain a single mutual 387 information estimate [16-20]. To estimate the information transfer in a shorter time 388 period we used a recently developed method [15] that uses the output of an artificial 389 neural network (ANN) to generate the frozen noise current input used in our ex-vivo 390 experiments (Fig 2A, see Materials & Methods). Such a frozen noise input constitutes 301 an optimum between giving naturalistic stimuli (as far as possible in an ex-vivo setup, 392 given that we do not have access to the spatiotemporal input distribution a cell would 393 normally receive), being able to assess information transfer, and being able to assess 394 membrane properties (which are typically only stably accessible in slice experiments). 395 The ANN responds to a randomly appearing and disappearing preferred stimulus or 396 'hidden state' x (Markov process). This hidden binary state can either be 'on' (i.e. 397 x = 1) or 'off' (i.e. x = 0), and switches randomly between these states with time 398 constant τ_{input} . The neurons in the ANN respond to this hidden state with 399 Poisson-generated spike trains, of which the firing rate depends on the hidden state (i.e. 400 each neuron *i* responds with a rate of q_{on}^i when x = 1, and a rate of q_{off}^i when x = 0). 401 The mutual information between the input current and the hidden state depends on three properties of the ANN: the number of neurons (N), the average firing rate of the 403 neurons (μ_q) , and the time constant of the hidden state (τ_{input}) . We can now compare 404 the mutual information between the input current and the hidden state with the mutual 405

377

378

379

380

381

382

383

384

385

information between the output spike train and the hidden state. This has the advantage that because the hidden state is low-dimensional (it has only two states), the mutual information can be estimated in a short time-window.

Because of the differences in maximum firing rate between the excitatory and 409 inhibitory cells, it was not possible to use the exact same frozen noise input current for 410 the two cell types: τ_{input} had to be large for the excitatory neurons (neurons firing at a 411 low rate cannot transfer information about a fast-switching stimulus, so the hidden 412 state had to switch slowly), but this is not the case for the inhibitory neurons (which 413 fire at high rates, so the hidden state can switch fast, i.e. a small value for τ_{input} should 414 be used). However, the information in the input could be kept constant by adapting the 415 firing rates of the neurons in the ANN μ_q (Fig 2, see also Materials & Methods). This 416 resulted in the parameters in Table 1 for the frozen noise experiments to generate the 417 input currents shown in Fig 2. 418

Inhibitory neurons show broadband information transfer; Excitatory neurons transfer less information and at low frequencies

By using the 'frozen noise protocol' as described before (see Materials & Methods 421 and [15]), the information transfer from the hidden state to the output spike train of a 422 single neuron can be estimated in a short time window. In order to obtain the 423 information transfer from the input current to the output spike train, we define the 424 unitless fraction of transferred information (FI) as the mutual information between the 425 spike train and the hidden state $(MI_{\text{spike train}} \text{ divided by the mutual information})$ 426 between the input current and the hidden state $(MI_{input}, see eq (6))$. The FI quantifies 427 how much information about the hidden state is transferred from the input current to 428 the output spike train, and thus quantifies which fraction of the information is kept 429 during the spike-generating process. In Fig 3, we show the FI as a function of the firing 430 rate r, for inhibitory (blue) and excitatory (red) neurons, and compare it to the FI431 obtained from the 'Bayesian Neuron' (BN) model [21] for which parameters (see 432 Materials & Methods) were optimized for the input generated for the excitatory neurons 433 (pink) or inhibitory neurons (turquoise). Excitatory neurons transfer more information 434 at low firing rates (< 8 Hz) compared to inhibitory neurons. This is due to our choice 435 of slower switching speed (i.e. large τ_{input}) of the hidden state for excitatory neurons: a 436 fast-switching hidden state cannot be properly tracked by neurons firing at a low firing 437 rate (see also [15]). To compare inhibitory and excitatory neurons, we normalized the 438 firing rate of each neuron relative to the switching speed of the hidden state: 439 $r_n = r \cdot \tau_{\text{input}}$ (unitless). The FI was plotted as a function of this normalized firing rate 440 in Fig 3B. The FI increases up to a maximal value at about $r_n = 1.5$, after which the 441 FI appears to decrease again. Apparently, at very high firing rates, the transferred 442 information goes down due to too many spikes during x = 0. We fitted a saturating 443 function (see Materials & Methods) to the measured values, where FI_{max} is the 444 saturation value and λ_{r_n} is the rate with which this saturation value is reached (both 445 unitless). We fitted the data up to $r_n = 1.5$, because we do not have a mathematical 446 description for the type of curve that saturates and then dips again (but note that all 447 panels and figures contain all data points, including those for $r_n > 1.5$). In Fig 3E and 448 F, the fit values and their 95% confidence intervals are shown. Inhibitory experimental 449 and BN values saturate around similar values ($FI_{max} = 0.65$ (0.64 - 0.66) and $FI_{max} =$ 450 0.64 (0.63 - 0.65) respectively), with experiments having a slightly lower rate ($\lambda_{r_n} = 5.8$ 451 (5.6 - 6.0) and 7.7 (7.3 - 8.0) respectively). Excitatory neurons saturate at lower 452 experimental values $(FI_{\text{max}} = 0.51 \ (0.48 - 0.54))$ and slightly lower BN values $(FI_{\text{max}} =$ 453 0.58 (0.54 - 0.63), and the saturation rates are also lower ($\lambda_{r_n} = 4.5$ (4.0 - 4.9) and λ_{r_n} 454 = 6.1 (5.0 - 7.2) respectively). This shows that in the case of the excitatory neurons, the 455 experimentally recorded spike trains transmit less information than the spike trains of 456

406

407

408

419

the BN, whereas in the inhibitory case, the model and experimental spike trains perform similarly. This means that inhibitory neurons perform close to optimal for representing the hidden state, whereas excitatory neurons do not. As a control, we presented the input for the excitatory neurons also to inhibitory neurons (Fig 3, green, $FI_{max} = 0.63$ $(0.52 - 0.75), \lambda_{r_n} = 2.6 (1.6 - 3.7)$); these inhibitory neurons fired at a higher normalized rate (Fig 3 C) and performed better than the excitatory neurons. In conclusion, putative interneurons transfer more information than putative excitatory neurons.

Inhibitory neurons perform well as classifiers

The setup with the hidden state makes it possible to show 'receiver-operator curves' 465 (ROCs): we define a 'hit' as a period during which the hidden state was 1 (up-state), in 466 which at least 1 action potential was fired, and a 'miss' as an up-state in which no 467 action potentials were fired. Similarly, we define a 'false alarm' as a period during which 468 the hidden state was 0 (down-state), in which at least 1 action potential was fired, and 469 a 'correct reject' as a down state in which no action potentials were fired. We then 470 define the 'hit fraction' as the number of hits divided by the total number of up-states, 471 and similarly the false alarm fraction for the number of false alarms divided by the total 472 number of down-states. In Fig 4A the results are shown, for the same five conditions as 473 discussed above. For each experiment, a control experiment was simulated by generating 474 a Poisson spike train with the same number of spikes as the original experiment. Note 475 that this 'control' is below the line hit fraction = false alarm fraction because the 476 hidden state is more often 0 than 1 $(P_1 = \frac{1}{3})$. Since the hidden state is longer in the '0' 477 state, the probability that a random spike occurs when the hidden state equals 0 is 478 higher, hence the probability of a false alarm is higher than the probability of a hit. 479

Inhibitory neurons perform comparably to the BN, as shown in Fig 4, whereas the 480 excitatory neurons perform less optimally than their model counterparts. We performed 481 control experiments where input currents generated for excitatory neurons were injected 482 into inhibitory neurons, (green triangles in Fig 3 and 4). The results suggest that 483 interneurons perform comparably to (on the same curve as) excitatory neurons, but 484 with a lower discrimination threshold (i.e. with a higher firing rate), which is in 485 agreement with our previous observation that inhibitory neurons responded with a 486 higher firing rate than excitatory neurons. Note that inhibitory neurons fire slightly less spikes during the up-states (Fig 4B) and the normalized firing rate in the up-state is 488 somewhat lower for the inhibitory neurons (Fig 4F). Since the excitatory neurons fire 489 more spikes during the down states (Fig 4C and 4G), this corresponds to a lower 490 efficiency for excitatory neurons and a worse performance on the binary classification 491 task (Fig 4A). Indeed, the number of spikes per down state (Fig 4C) and normalized 492 firing rate in the down state (Fig 4G) differs between inhibitory and excitatory neurons 493 (Supplementary Tables S2 and S3). Note that most 'incorrect' spikes are actually fired 494 shortly after a down switch (Fig 4H-K), so they might be 'correct' spikes that were a 495 few milliseconds too late. In conclusion, putative interneurons are better binary 496 classifiers than putative excitatory neurons. 497

Dynamic threshold of both neuron types

To assess how intrinsic properties of the putative interneurons and pyramidal cells correlate with their information transfer capabilities in this setup, we next assess the threshold adaptation of these neuron types. In Fig 5, we show the threshold behaviour of the inhibitory and excitatory neurons. The membrane potential threshold of each spike was determined based on the method of [24] (see Materials & Methods). We show the distribution of the membrane potential as a function of the inter-spike interval (ISI, Fig 5A and E). For both inhibitory and excitatory neurons, the membrane potential

498

499

500

501

502

503

504

505

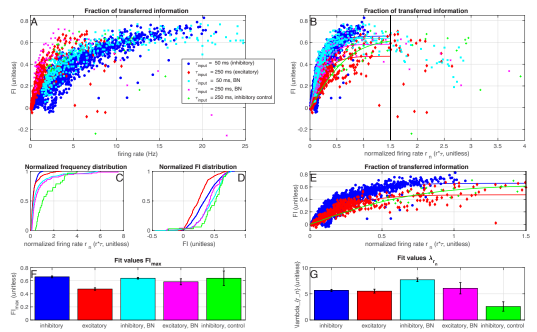


Fig 3. Inhibitory neurons transfer more information. A: Fraction of information kept during the spike generating process (FI, see eq. 6) as a function of the firing rate, for inhibitory neurons (blue) and excitatory neurons (red). In green, the control experiments where the inhibitory neurons received the input current that was normally given to the excitatory neurons ($\tau_{input} = 250 \text{ ms}$). In turquoise and pink, the simulations with the Bayesian Neuron (Materials & Methods, see Table 1 for parameter values). B: Fraction of information kept during the spike generating process, as a function of the normalized firing rate (normalized by the switching speed of the hidden state: $r_n = r \cdot \tau_{\text{input}}$, see Table 1). The solid lines denote fits of the data up to a normalized firing frequency of $r_n = 1.5$ (eq. (7), Materials and Methods). Colors/markers the same as in A. C: and D: Normalized firing frequency and FI distribution of the spike trains in all conditions. E: Zoom of B. F: and G: Fit values and their 95% confidence intervals (error bars) for parameters FI_{max} (F) and λ_{r_n} (G), see eq. (7). Data from 144 excitatory neurons (220 trials), 72 inhibitory neurons (78 trials) and 9 control inhibitory neurons (11 trials). NB Note that even though theoretically $MI \ge 0$, due to our approximation, our estimate of MI can take small negative values (see Materials & Methods).

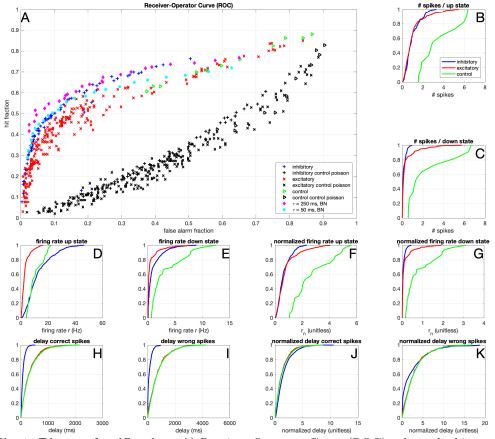


Fig 4. Binary classification. A) Receiver Operator Curve (ROC), where the hit rate was defined as the fraction of up-states, in which at least 1 action potential was fired. Similarly, the false alarm rate was defined as the fraction of down-states, in which at least 1 action potential was fired. In black the results for Poisson spike trains with firing rates matched to those of the experimental/simulation conditions are shown. B) Distribution of the number of spikes per period where the hidden state was 1 (up state), for inhibitory neurons (blue) and excitatory neurons (red). C) Same as B), but for periods where the hidden state was 0 (down state). D) Firing rate r distribution in the up-state. E) Firing rate r distribution in the down-state. F) Normalized firing rate r_n distribution in the up-state G) Normalized firing rate r_n distribution in the down-state. H) Delay (in ms) of each correct spike since the state switches from down to up. I) Delay (in ms) of each incorrect spike since the state switches from up to down. J) Normalized delay (delay/ τ , unitless) of each correct spike since the state switches from down to up. K) Normalized delay of each incorrect spike since the state switch from up to down. The results of hypotheses test for A-F are in Supplementary Table S2 and S3. Data from 144 excitatory neurons (220 trials), 72 inhibitory neurons (78 trials), and 9 control inhibitory neurons (11 trials).

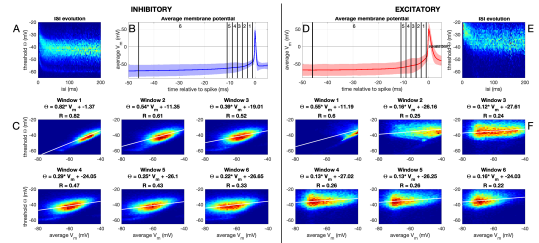


Fig 5. Dynamic threshold. A-C) Inhibitory neurons. D-F) Excitatory Neurons. A) Distribution of membrane potential threshold values (see Materials & Methods) for each inter-spike interval (ISI); normalized per ISI. B) Average spike shape (shaded region denotes standard deviation). Vertical lines denote the windows in C. C) Heatmap and regression for the relation between the threshold and the average membrane potential in the given window. D-F) Same as in A-C, but for excitatory neurons. This is all in the Frozen Noise protocol, for threshold behaviour in the current-clamp step-and-hold protocol, see Supplementary Fig. S1.

threshold goes up with short ISIs, as expected, and for long ISIs the threshold is low. 506 This effect has a long time scale (at least several tens of milliseconds), longer than 507 expected based on the relative refractory period alone (typically less than ten 508 milliseconds). The thresholds of excitatory neurons are almost 10 mV higher than those 509 of inhibitory neurons (Fig 5A,E, Supplementary Fig 1A-C). Next to the ISI, the 510 threshold also depends on the history of the membrane potential (Fig 5C,F): we 511 calculated the regression between the action potential threshold and the average 512 membrane potential in different windows preceding the spike. There is a strong 513 correlation between the threshold and the membrane potential immediately preceding 514 the spike for both neuron types, which reduces gradually with time before the spike. 515 However, for both neuron types, some relation between the membrane potential several 516 tens of milliseconds before the spike and the threshold is still visible. The current clamp 517 step protocol (Supplementary Fig 1) confirms the overall higher thresholds 518 (Supplementary Fig 1A,B) and strong spike-frequency adaptation (Supplementary Fig 519 1D) for excitatory neurons. The threshold adaptation rate however, shows significant 520 differences between fast spiking and regular spiking neurons at current injection 521 intensities ranging from +240 to +320 pA, while they do not show significant changes at 522 lower or higher intensities, possibly due to low firing rates or reaching a steady state 523 firing rate. (Supplementary Fig 1C, Table S1). 524

So in conclusion, both inhibitory and excitatory neurons show a dynamic threshold behaviour, with inhibitory neurons having much lower thresholds, so they can fire at high rates, whereas the dynamic threshold of excitatory neurons promotes low-frequency firing and shows stronger adaptation.

Information transfer in simulated neuron models

In the experimental data, we saw that both fast-spiking interneurons and regular spiking excitatory neurons transfer a significant amount of information about the

525

526

527

528

529

530

hidden state, not much less than the optimal Bayesian neuron, as they adapt their spike 532 threshold to the dynamics of the stimulus. The goal of this research was to explore the 533 relationship between intrinsic excitability and information transfer. The Bayesian 534 neuron that is optimal for this task has two properties that distinguish it from a 535 standard integrate-and-fire model: 1) spike-frequency adaptation and 2) a non-linear 536 I-V curve. To untangle how these mechanisms influence information transfer, we turn to 537 computational modelling. We use an exponential integrate-and-fire (expIF) model and 538 adapt both its adaptation and the shape of the IV-curve, to explore how these affect 539 information transfer. 540

Information transfer in neuron models with (sub)threshold adaptation

In the previous section, we saw that both inhibitory and excitatory neurons show a dynamic threshold behaviour, suggesting that both cell types have in theory the adaptation mechanisms that can influence information transfer, as is also present in the Bayesian Neuron. In biophysical models, spike-frequency adaptation can be implemented in different ways [11]. Particularly, in the expIF model, it has been implemented as either a subthreshold process [22,23] or as an adaptation of the spike threshold [24]. We research the effects of these two types of adaptation on the information transfer in the aforementioned mutual information protocol.

In Fig 6C, we first note that the 'slow' input to the excitatory neurons is apparently more difficult to transfer than the 'fast' one: the exact same expIF model transfers less of the 'slow' input information (red) than of the 'fast' one (blue). Next, in Fig 6D-G, we show that adding threshold adaptation does not increase the amount of information that is transferred by the neuron. However, it does shift its working range towards higher values of the input amplitude I_{scale} , effectively increasing its working range. Contrasting, in Fig 6H-K, we show that adding subthreshold adaptation does increase the maximum information transfer when it is properly tuned, i.e. when the time constant of adaptation fits the input properties. However, too slow adaptation suppresses the firing rate too much (Fig 6J,K), resulting in a reduction of information transfer.

We ask whether the effects on information transfer are a result of a higher firing rate, or of a better detection. Therefore, we turn to the ROC curves discussed before. In Fig 7 we show that both forms of adaptation do not change the shape of the ROC curve. However, we do note that for the 'slow' input, the expIF neuron performs much worse than both the Bayesian neuron and the experimentally recorded neurons.

In conclusion, we see that subthreshold, but not threshold, adaptation can increase the maximum information transfer. Threshold adaptation, on the other hand, can increase the working range of the neuron. Moreover, an expIF neuron performs worse than both the Bayesian neuron and the experimentally recorded neurons. Since the Bayesian neuron differs from the expIF model in its IV curve, we next determine how information transfer is influenced by the shape of the IV curve.

The shape of the IV curve

We assess the effects of changing the shape of the I-V curve (the right-hand side of the 572 membrane voltage equation). The Bayesian neuron, tailor-made to transfer information 573 efficiently for this type of input, has two features that distinguish it from a classical 574 integrate-and-fire model: an adaptation mechanism, discussed in the previous 575 paragraph, and a non-linear IV-curve, as can be seen in Fig 8A: the amplitude of the 576 IV-curve increases exponentially when moving away from the steady-state value (dotted 577 vertical line). We add such non-linearities in the expIF neuron in a biologically realistic 578 way, to see how they would influence the classification (ROC curve) and the information 579 transfer. Firstly we model the effects of the suppression of hyperpolarization (i.e. 580

541

542

543

544

545

546

547

548

549

550

551

552

553

554

555

556

557

558

559

560

561

562

563

564

565

566

567

568

569

570

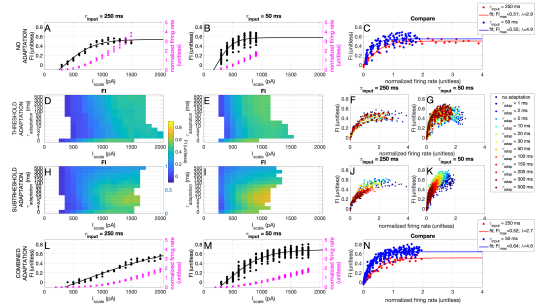


Fig 6. Effects of the dynamics of adaptation on information transfer. A-C) No adaptation. D-G) Threshold adaptation. H-K) Subthreshold adaptation. L-N) Combined adaptation. Left column: 'slow' input (time constant hidden state $\tau_{input} =$ 250 ms). Middle column: 'fast' input (time constant hidden state $\tau_{input} = 50$ ms). Right column: 'slow' and 'fast' input together. A) Fraction of information (FI, black)and normalized firing rate r_n (pink) as a function of the input amplitude I_{scale} for the expIF model without adaptation. B) same as A but for the 'fast' input. C) Fraction of information as a function of the normalized firing rate for the 'slow' (red) and 'fast' (blue) input for the expIF model without adaptation. D) Fraction of information (colorbar) as a function of the input amplitude I_{scale} for the expIF model with threshold adaptation with different adaptation time constants τ_{adap} (vertical axis) receiving the 'slow' input. E) Same as D, but for the 'fast' input. F) Fraction of information as a function of the normalized firing rate for the 'slow' input or the expIF model with threshold adaptation with different adaptation time constants τ_{adap} (colours). G) Same as F, but for the 'fast' input. H-K) Same as D-G, but for the expIF model with subtreshold adaptation. L-N) Same as A-C, but for the model with both threshold $(\tau_{\text{adap}} = 1 \text{ ms and subthreshold } (\tau_{\text{adap}} = 10 \text{ ms adaptation.})$

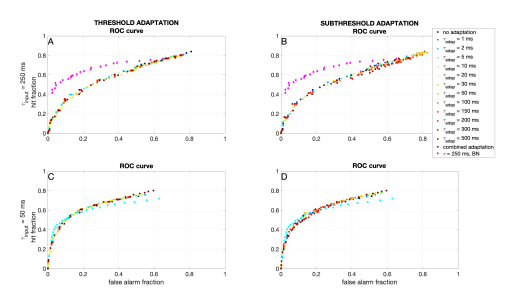


Fig 7. Effects of the dynamics of adaptation on binary classification. A) Receiver Operator Curve (ROC) (see also Fig 4) for the expIF neuron with threshold adaptation (colors denote time constant) receiving 'slow' input. Note that the adaptation does not change the shape of the ROC curve, and that the neuron performs much worse than the Bayesian neuron (pink). B) Same as A, but for subthreshold adaptation. C) Same as A, but for the neuron receiving 'fast' input and the response of the Bayesian neuron in turquoise. D) Same as C, but for the neuron with subthreshold adaptation.

increasing slope of the IV curve when hyperpolarizing the cell) by adding an instantaneous 'h-current' to the expIF neuron (see Materials & methods), as shown in Fig 8B. Next, we model the effects of the suppression of depolarization (i.e. increasing slope of the IV curve when depolarizing the cell) by adding an instantaneous subthreshold potassium current, as shown in Fig 8C. Finally, we also change the overall slope, but not the shape of the IV curve, by changing the leak conductance of the neuron, as shown in Fig 8D. In Fig 8F and J, we show that adding the 'h-current' (with conductance g_h) does not change the shape of the ROC curve. However, its effect is similar to lowering the detection threshold (i.e. the values shift over the curve towards higher hit and false alarm fractions). On the contrary, the addition of the potassium current (with conductance q_K) does not change the shape of the ROC curve, but its effect is similar to an increase in the detection threshold (i.e. the values shift over the curve towards lower hit and false alarm fractions, Fig 8G and K). Changing the overall slope of the IV curve (i.e. the 'leak conductance' g_L) does change the shape of the ROC curve (Fig 8H and L): for the slow input current ($\tau_{input} = 250 \text{ ms}$) it needs to be tuned to a lower value ($g_L \approx 1 \text{ nS}$) than for the faster input current ($\tau_{\text{input}} = 50 \text{ ms}$) for optimal information transfer and classification.

The effects seen in the ROC curves are confirmed by the information transfer measurements: in Fig 9A, D, G and J we show that adding an 'h-current' can strongly increase the information transfer of the expIF neuron, by increasing its firing rate. Adding a subthreshold instantaneous potassium current shows the opposite effect: it decreases both firing rates and information transfer (Fig 9B, E, H and K). Finally, the slope of the IV curve needs to be matched to the input statistics: the slow input needs a flatter IV-curve (lower g_L) than the fast input for information transfer (Fig 9C, F, I and L).

581

582

583

584

585

586

587

588

589

590

591

592

593

594

595

596

597

598

599

600

601

602

603

604

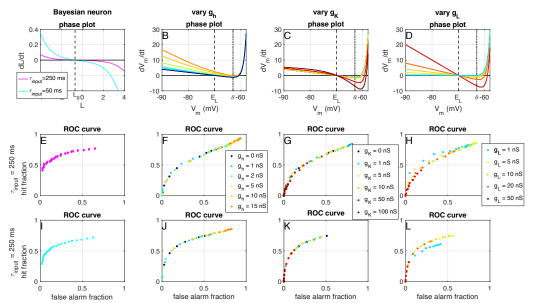


Fig 8. Effects of the IV-curve shape on binary classification. A-D) IV-curves (i.e. right-hand side of the membrane voltage equation) for the Bayesian Neuron (A) and for the expIF neuron (without adaptation) with added 'h-current' (B), 'subthreshold potassium current' (C) and while varying the membrane conductance (D), colours denote conductances (see legends below). E-H) Receiver Operator Curve (ROC) (see also Fig 4) for the expIF neuron with different IV curve shapes (colors denote time constant) receiving 'slow' input. Note that only g_L changes the shape of the ROC curve. I-L) Same as E-H, but for the neuron receiving 'fast' input.

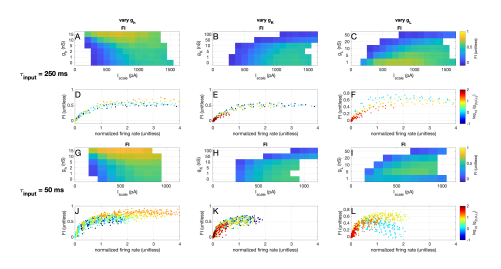


Fig 9. Effects of the IV-curve shape on information transfer. A) Fraction of transferred information FI as a function of the input amplitude I_{scale} for the expIF model with added instantaneous 'h-current' with different values of conductance g_h (vertical axis) receiving the 'slow' input. B) Same as A but for the subthreshold instantaneous 'potassium current' (g_K) . C) Same as A but for the leak conductance g_L . D-F) Fraction of information as a function of the normalized firing rate for the 'slow' input or the expIF model with different IV shapes (colours). G-L) Same as A-F but for the 'fast' input.

> In conclusion, we saw that the recorded excitatory neurons perform better for slow 606 input than the expIF model with or without adaptation; in fact, these neurons perform 607 similarly to the optimal simulated model (the Bayesian Neuron). Recorded inhibitory 608 neurons perform close to optimal for fast input, a result well captured with an expIF 609 model that includes an adaptation mechanism. Threshold adaptation increases the 610 working range of the expIF model, but does not increase, or even slightly reduces, the 611 amount of transferred information. Subthreshold adaptation, on the other hand, does 612 not increase the working range but does increase the maximum transferred information 613 if correctly tuned. Neither form of adaptation changes the shape of the ROC curve. The 614 slope of the IV curve does play an important role in the information transfer and needs 615 to be tuned to the statistics of the input. To check this conclusion, we will next assess 616 this statement in our experimental recordings. 617

Back to the recordings: dynamic IV curve unravels the relationship between membrane conductance and information transfer

We assess the relation between the slope of the IV-curve and the information transfer, 620 by determining the dynamic IV-curve [28] for each of our recordings (see Materials and 621 Methods). In figure 10, we show the fraction of transferred information (FI) as a 622 function of the membrane conductance g_m and membrane capacitance C_m (Fig 10 A 623 and B) and as a function of the membrane time constant τ_m (Fig 10 C and D). As in 624 the expIF model simulations, we can conclude that the fraction of transferred 625 information depends on the slope of the IV-curve: we see a clear inverse relation 626 between membrane conductance and transferred information. However, the recorded 627 neurons show quite a large variability of intrinsic properties, in particular the regular 628 spiking excitatory neurons. To assess how this large heterogeneity of excitatory neurons 629 influences their response properties, we calculate their spike-triggered averages. 630

Back to the recordings: Response heterogeneity of the Spike-Triggered Average

In Fig. 11, we show the normalized spike-triggered averages (STAs) for spikes of inhibitory (A and E) and excitatory neurons (C). The filter was whitehed and regularized (see Materials & Methods). Next, the projection values of spike-triggering and random currents were calculated (see Fig 11B for an example for 1 cell), and the distance between the means of the distributions for random and spike-triggering currents was calculated for each cell (Fig 11D). The average STAs for all inhibitory (Fig 11A, blue) and excitatory (Fig 11C, red) neurons were quite similar, but the traces for individual neurons (grey lines) were much more variable for excitatory neurons than inhibitory neurons. This indicates that the excitatory neurons have a higher variability in their feature selectivity of incoming current stimuli than inhibitory neurons, as was expected from the higher intrinsic variability discussed in the previous section. However, it is also possible that this is an effect of the lower number of spikes available for excitatory neurons. To control for this possibility, we calculated the STAs for spike trains of inhibitory neurons, where the number of spikes was reduced to match an 186 excitatory trial (Fig 11E, brown). For all three groups (inhibitory, excitatory, and inhibitory control spike trains) we calculated the inner product between all calculated STAs. Fig 11F shows the distributions of these inner products, and it is clear that both inhibitory full and control spike trains are much less variable (inner product closer to 1) than the excitatory spike trains (two-sample Kolmogorov-Smirnov test E-I p = 0, E-C p < 1e - 223, I-C p < 1e - 228). The distribution of all distances between the means is shown in Figs 11D. The distances between the distributions, measured in standard deviations of the prior (random triggered currents) distribution, are much higher for

618

619

631

632

633

634

635

636

637

638

639

640

641

642

643

644

645

646

647

648

649

650

651

652

653

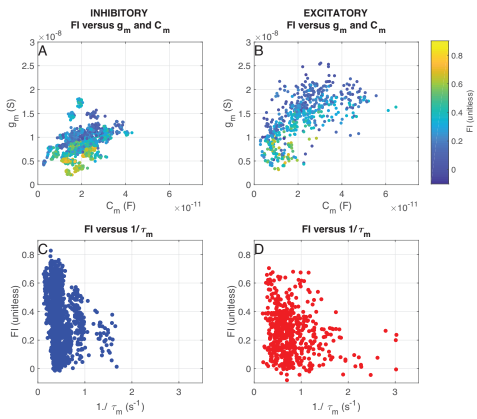


Fig 10. Effects of the dynamic IV-curve shape on information transfer of recorded neurons. A) Fraction of transferred information FI as a function of the membrane conductance g_m and capacitance C_m for inhibitory neurons. B) Same as A), but for excitatory neurons. C) Fraction of transferred information FI as a function of the inverse of the membrane time constant τ_m for inhibitory neurons. D) Same as C), but for excitatory neurons.

excitatory neurons than for inhibitory neurons, indicating that excitatory neurons are more selective (p-values two-sample t-test: E-I p < 1e - 28, E-C p < 1e - 24, I-C p = 0.14). In conclusion, excitatory cells fire less than inhibitory cells and are therefore more selective, but at the same time, there is more variability between excitatory neurons in what input features they respond to than between inhibitory cells.

Conclusion and discussion

In summary, we measured the differences in information transfer between (putative) 661 inhibitory interneurons and excitatory pyramidal cells in the cerebral cortex. We 662 utilised a technique in which the input current was generated by an Artificial Neural 663 Network (ANN), with each artificial cell firing Poisson spike trains whose firing rate was 664 modulated by the absence or presence of the stimulus [15]. We discovered that 665 excitatory cells are more selective due to their greater information compression. Inhibitory neurons exhibit a near-optimal response, transferring a great deal of 667 input-related information at relatively rapid rates. In a computational model, the 668 mechanisms that can explain such differences in information transfer were investigated. 669 We evaluated the effects of (sub)threshold adaptation and the IV curve's shape. We 670 discovered that adaptation increases information transfer (subthreshold adaptation) and 671 the working range (threshold adaptation). In addition, the shape of the IV-curve plays 672 a crucial role in determining the information transfer: the slope must correspond to the 673 input characteristics, and the suppression of hyperpolarization, such as by a 'h-current,' 674 can increase the information transfer. The effects are summarised in Table 4. Although 675 the current experimental data does not permit an explicit test of the effects of 676 (sub)threshold adaptation and/or 'h-current,' the relationship between information 677 transfer and the slope of the (dynamic) IV-curve (the membrane conductance) can be 678 evaluated. As predicted, we observe an inverse relationship between membrane 679 conductance and information transfer. Finally, we find that both the intrinsic 680 (membrane conductance) and response (STA, FI) properties of excitatory neurons are 681 more heterogeneous, compared to inhibitory neurons. 682

Mechanism	Max information transfer	Working range	ROC curve
threshold adaptation	unchanged/reduced	increased / shift to	unchanged
		higher amplitudes	
subthreshold adaptation	increased	unchanged	unchanged
	(if tuned properly)		
steepness	increased	depends on tuning	better detection
IV curve (g_L)	(if tuned properly)		if tuned properly
hyperpolarized part	increased at the cost of	shift to lower amplitudes	unchanged shape
IV curve (g_h)	higher firing rate		shift towards higher rates
depolarized part	decreased	shift to higher amplitudes	shape unchanged
IV curve (g_K)			shift towards lower rates

Table 4. Conclusions of the Exponential IF model simulations

It has been shown repeatedly, that the spiking behaviour of cortical neurons can be fitted relatively well with a simple threshold model with an extra feedback variable [22, 24, 35–45] and the heterogeneity in such cell properties has been investigated in excitatory (but not inhibitory) cells [46]. With this manuscript, we add a functional dimension to these basic properties of cortical spike initiation: We show how different mechanistic features of cortical cells can influence their information transfer and binary classification. Of course, we did not explore all mechanisms available to the

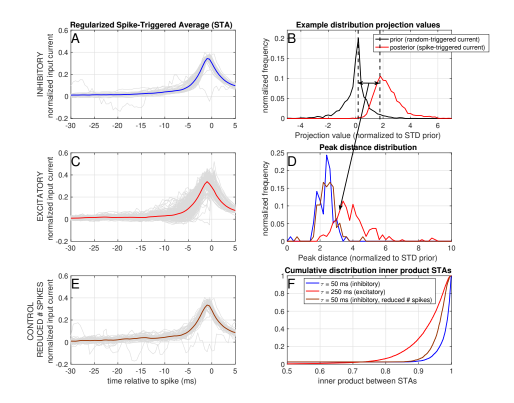


Fig 11. Linear filtering properties of recorded neurons. A) Whitened and regularized (see Materials and Methods) Spike-Triggered average (STA) for inhibitory neurons. The STAs for individual neurons are shown as thin grey lines, and the average over neurons is shown as a thick coloured line. B) Example of a prior (random triggered, black line) and posterior (spike-triggered, blue line) distribution of stimulus projection values for a single inhibitory neuron. C) Same as A), but for excitatory neurons. D) Distribution of the differences between the means (see arrow in B) between the prior and posterior distribution over all neurons.-E) Same as A), but for the reduced (i.e. fewer spikes) spike trains of the inhibitory neurons. F) Distribution of the inner products between the STAs for the three groups (note that because the STAs are normalized by the L2-norm, the maximal value of the inner product is limited to 1). Data from 144 excitatory neurons (220 trials) and 72 inhibitory neurons (78 trials).

cell. For instance, in this simplified cell model we could not assess the difference in 690 information transfer between 'type 1' (or integrator) and 'type 2' (or resonator) cells, as 691 this can only be modelled as a difference in the bifurcation from resting to spiking (a 692 saddle-node versus a Hopf bifurcation). Moreover, we used the expIF model as a proof-of-principle of the effects of different intrinsic cell properties on information 694 transfer and did not extensively fit the model to the experimental data. Indeed, the 695 recorded spike trains are better classifiers than even the best-performing expIF model 696 for the slow input current, suggesting that there are more relevant dynamic properties 697 that are not captured by such a simplified model. However, using such a simple setup 698 allows us to make several predictions that can be tested experimentally: we predicted 699 that 1) blocking 'h-currents' will decrease the amount of information that is transferred 700 2) blocking subthreshold potassium currents will not have such an effect, and 3) there is 701 an optimal range for the membrane conductance. 702

The heterogeneity of neuron properties has received much interest lately: for instance, it has been shown that heterogeneity in neural populations can increase coding robustness and efficiency [47], help optimize information transmission [48], increase network responsiveness [49], promote robust learning [50], help to control the dynamic repertoire of neural populations [51] and improve the performance on several tasks [52, 53]. Here, we show that in particular, the population of excitatory neurons of the barrel cortex shows a large variability in their intrinsic and response properties. Why the variability of the properties of excitatory neurons is larger than that of inhibitory ones is an exciting question, which is a subject for future experimental and computational evaluation. Moreover, the intrinsic properties of cortical neurons are under top-down influence by neuromodulators such as serotonin, acetylcholine and dopamine [54,55]. Using the protocol described herein, it will be possible to investigate how these neuromodulators affect the intrinsic neural properties and, consequently, their information transfer. This will help reveal how the specific actions of these neuromodulators on the intrinsic properties of specific cell classes affects information transfer in the cortex. By investigating the relationship between intrinsic neuron properties and information transfer, we can begin to predict the effect of top-down processes on cortical processing.

Supporting information

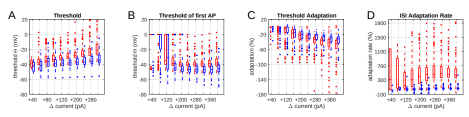


Fig 12. S1 Fig Threshold behaviour in the current clamp step-and-hold protocol. A) Thresholds of all spikes during the step protocol. B) Thresholds of the first spikes after the step current initiation. IC Threshold adaptation: difference in threshold between the first and the last spike of the response. D) Last ISI length relative to the first ISI of the response. Excitatory (red) and inhibitory (blue) neurons. NB Results for significance testing in table S1.

S1 Table. Supplementary Table S1: Statistical tests of the comparison between r22 excitatory and inhibitory neurons in the current clamp step protocol. r23

703

704

705

706

707

708

709

710

711

712

713

714

715

716

717

718

719

720

S2 Table. Supplementary Table S2: Statistical tests of the comparison between reactive rea

S3 Table. Supplementary Table S3: Statistical tests of the comparison between excitatory and inhibitory neurons receiving the control frozen noise stimulus (see main text Fig. 5). P-values were compared to a threshold 34 of 5% / 6 groups = 0.83 % (Bonferroni correction). 731

Acknowledgments

This work was supported by grants from the European Commission (Horizon2020, nr. 660328), European Regional Development Fund (MIND, nr. 122035) and the Netherlands Organisation for Scientific Research (NWO-ALW Open Competition, nr. 824.14.022) to TC and by the Netherlands Organisation for Scientific Research (NWO Veni Research Grant, nr. 863.150.25) to FZ. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript. 738

References

- Huang C, Englitz B, Reznik A, Zeldenrust F, Celikel T. Information Transfer and Recovery for the Sense of Touch. bioRxiv. 2020; p. 2020.12.08.415729. doi:10.1101/2020.12.08.415729.
- Brenner N, Bialek W, de Ruyter van Steveninck RR. Adaptive Rescaling Maximizes Information Transmission. Neuron. 2000;26(3):695–702.
- 3. Fairhall AL, Lewen GD, Bialek W, de Ruyter van Steveninck RR. Efficiency and Ambiguity in an Adaptive Neural Code. Nature. 2001;412:787–792.
- Laughlin S. A Simple Coding Procedure Enhances a Neuron's Information Capacity. Zeitschrift Fur Naturforschung Section C, Biosciences. 1981;36(9-10):910–912.
- Smirnakis SM, Berry MJ, Warland DK, Bialek W, Meister M. Adaptation of Retinal Processing to Image Contrast and Spatial Scale. Nature. 1997;386(6620):69–73. doi:10.1038/386069a0.
- Turrigiano GG, Leslie KR, Desai NS, Rutherford LC, Nelson SB. Activity-Dependent Scaling of Quantal Amplitude in Neocortical Neurons. Nature. 1998;391(6670):892–896. doi:10.1038/36103.
- Turrigiano GG, Nelson SB. Homeostatic Plasticity in the Developing Nervous System. Nature Reviews Neuroscience. 2004;5:97–107.
- Desai NS, Rutherford LC, Turrigiano GG. Plasticity in the Intrinsic Excitability of Cortical Pyramidal Neurons. Nature Neuroscience. 1999;2(6):515–520. doi:10.1038/9165.
- Remme MWH, Wadman WJ. Homeostatic Scaling of Excitability in Recurrent Neural Networks. PLoS Computational Biology. 2012;8(5). doi:10.1371/journal.pcbi.1002494.

- van Welie I, a van Hooft J, Wadman WJ. Homeostatic Scaling of Neuronal Excitability by Synaptic Modulation of Somatic Hyperpolarization-Activated Ih Channels. Proceedings of the National Academy of Sciences. 2004;101(14):5123–5128.
- 11. Gutkin BS, Zeldenrust F. Spike Frequency Adaptation. Scholarpedia. 2014;9(2):30643. doi:10.4249/scholarpedia.30643.
- Turrigiano G. Too Many Cooks? Intrinsic and Synaptic Homeostatic Mechanisms in Cortical Circuit Refinement. Annual Review of Neuroscience. 2011;34(1):89–103. doi:10.1146/annurev-neuro-060909-153238.
- da Silva Lantyer A, Calcini N, Bijlsma A, Kole K, Emmelkamp M, Peeters M, et al. A Databank for Intracellular Electrophysiological Mapping of the Adult Somatosensory Cortex. GigaScience. 2018;7(12):1–9. doi:10.1093/gigascience/giy147.
- Yan X, Calcini N, Safavi P, Ak A, Kole K, Zeldenrust F, et al. A Whole-Cell Recording Database of Neuromodulatory Action in the Adult Neocortex. bioRxiv. 2022;doi:10.1101/2022.01.12.476007.
- Zeldenrust F, de Knecht S, Wadman WJ, Denève S, Gutkin BS. Estimating the Information Extracted by a Single Spiking Neuron from a Continuous Input Time Series. Frontiers in Computational Neuroscience. 2017;11:49. doi:10.3389/fncom.2017.00049.
- Bialek W, Rieke F, de Ruyter van Steveninck RR, Warland D. Reading a Neural Code. Science. 1991;252(5014):1854–1857.
- 17. de Ruyter van Steveninck RR, Bialek W. Real-Time Performance of a Movement-Sensitive Neuron in the Blowfly Visual System: Coding and Information Transfer in Short Spike Sequences. Proceedings of the Royal Society of London Series B. 1988;234(1277):379–414.
- de Ruyter van Steveninck RR, Lewen GD, Strong SP, Koberle R, Bialek W. Reproducibility and Variability in Neural Spike Trains. Science. 1997;275:1805–1808. doi:10.1126/science.275.5307.1805.
- 19. Rieke F, Warland D, de Ruyter van Steveninck RR, Bialek W. Spikes: Exploring the Neural Code. Cambridge, Massachusets: MIT Press; 1997.
- 20. Strong SP, Koberle R, de Ruyter van Steveninck RR, Bialek W. Entropy and Information in Neural Spike Trains. Physical Review Letters. 1998;80(1):197–200.
- Denève S. Bayesian Spiking Neurons I: Inference. Neural Computation. 2008;20(1):91–117. doi:10.1162/neco.2008.20.1.91.
- Brette R, Gerstner W. Adaptive Exponential Integrate-and-Fire Model as an Effective Description of Neuronal Activity. Journal of neurophysiology. 2005;94(5):3637–42. doi:10.1152/jn.00686.2005.
- Gerstner W, Brette R. Adaptive Exponential Integrate-and-Fire Model. Scholarpedia. 2009;4(6):8427.
- Fontaine B, Peña JL, Brette R. Spike-Threshold Adaptation Predicted by Membrane Potential Dynamics In Vivo. PLoS Computational Biology. 2014;10(4):1–11. doi:10.1371/journal.pcbi.1003560.

- Kole K, Zhang Y, Jansen EJR, Brouns T, Bijlsma A, Calcini N, et al. Assessing the Utility of Magneto to Control Neuronal Excitability in the Somatosensory Cortex. Nature Neuroscience. 2020;23(9):1044–1046. doi:10.1038/s41593-019-0474-4.
- Kole K, Celikel T. Neocortical Microdissection at Columnar and Laminar Resolution for Molecular Interrogation. Current Protocols in Neuroscience. 2019;86(1):e55. doi:10.1002/cpns.55.
- Lochmann T, Denève S. Information Transmission with Spiking Bayesian Neurons. New Journal of Physics. 2008;10(5):055019. doi:10.1088/1367-2630/10/5/055019.
- Badel L, Lefort S, Brette R, Petersen CCH, Gerstner W, Richardson MJE. Dynamic I-V Curves Are Reliable Predictors of Naturalistic Pyramidal-Neuron Voltage Traces. Journal of Neurophysiology. 2008;99:656–666.
- Chichilnisky EJ. A Simple White Noise Analysis of Neuronal Light Responses. Network: Computation in Neural Systems. 2001;12:199–213.
- Paninski L. Convergence Properties of Some Spike-Triggered Analysis Techniques. In: Becker S, Thrun S, Obermayer K, editors. Advances in Neural Information Processing Systems 15. MIT Press; 2003. p. 189–196.
- Sharpee TO, Rust NC, Bialek W. Analyzing Neural Responses to Natural Signals: Maximally Informative Dimensions. Neural Computation. 2004;16(2):223–250. doi:10.1162/089976604322742010.
- Simoncelli EP, Paninski L, Pillow JW, Schwartz O. Characterization of Neural Responses with Stochastic Stimuli. In: Gazzaniga M, editor. The Cognitive Neurosciences. MIT Press; 2004. p. 1385.
- 33. Slee SJ, Higgs MH, Fairhall AL, Spain WJ. Two-Dimensional Time Coding in the Auditory Brainstem. The Journal of Neuroscience. 2005;25(43):9978–9988.
- Stimberg M, Brette R, Goodman DF. Brian 2, an Intuitive and Efficient Neural Simulator. eLife. 2019;8:1–41. doi:10.7554/elife.47314.
- Botella-Soler V, Deny S, Marre O, Tkačik G. Nonlinear Decoding of a Complex Movie from the Mammalian Retina. arXiv. 2016;q-bio(1605.03373v1):[q-bio.NC]. doi:10.1073/pnas.0709640104.
- Botella-Soler V, Deny S, Martius G, Marre O, Tkačik G. Nonlinear Decoding of a Complex Movie from the Mammalian Retina. PLOS Computational Biology. 2018;14(5):e1006057. doi:10.1371/journal.pcbi.1006057.
- Gerstner W, Naud R. How Good Are Neuron Models? Science. 2009;326(5951):379–80. doi:10.1126/science.1181936.
- Jolivet R, Lewis TJ, Gerstner W. Generalized Integrate-and-Fire Models of Neuronal Activity Approximate Spike Trains of a Detailed Model to a High Degree of Accuracy. Journal of neurophysiology. 2004;92(2):959–76. doi:10.1152/jn.00190.2004.
- Jones DL, Johnson EC, Ratnam R. A Stimulus-Dependent Spike Threshold Is an Optimal Neural Coder. Frontiers in Computational Neuroscience. 2015;9. doi:10.3389/fncom.2015.00061.

- Kobayashi R, Tsubo Y, Shinomoto S. Made-to-Order Spiking Neuron Model Equipped with a Multi-Timescale Adaptive Threshold. Frontiers in computational neuroscience. 2009;3(July):9. doi:10.3389/neuro.10.009.2009.
- Naud R, Gerstner W. Can We Predict Every Spike? In: Dilorenzo PM, Victor JD, editors. Spike Timing: Mechanisms and Function. November. CRC Press; 2013. p. 65–76.
- 42. Rauch A, La Camera G, Luscher HR, Senn W, Fusi S. Neocortical Pyramidal Cells Respond as Integrate-and-Fire Neurons to in Vivo-like Input Currents. Journal of neurophysiology. 2003;90(3):1598–612. doi:10.1152/jn.00293.2003.
- Rossant C, Goodman DFM, Platkiewicz J, Brette R. Automatic Fitting of Spiking Neuron Models to Electrophysiological Recordings. Frontiers in Neuroinformatics. 2010;4(2):1–10. doi:10.3389/neuro.11.002.2010.
- Rossant C, Goodman DFM, Fontaine B, Platkiewicz J, Magnusson AK, Brette R. Fitting Neuron Models to Spike Trains. Frontiers in neuroscience. 2011;5(February):9. doi:10.3389/fnins.2011.00009.
- Woo J, Kim SH, Han K, Choi M. Characterization of Dynamics and Information Processing of Integrate-and-Fire Neuron Models. Journal of Physics A: Mathematical and Theoretical. 2021;54(44):445601. doi:10.1088/1751-8121/ac2a54.
- Harrison PM, Badel L, Wall MJ, Richardson MJE. Experimentally Verified Parameter Sets for Modelling Heterogeneous Neocortical Pyramidal-Cell Populations. PLOS Computational Biology. 2015;11(8):e1004165. doi:10.1371/journal.pcbi.1004165.
- Zeldenrust F, Gutkin B, Denéve S. Efficient and Robust Coding in Heterogeneous Recurrent Networks. PLOS Computational Biology. 2021;17(4):e1008673. doi:10.1371/journal.pcbi.1008673.
- Haggard M, Chacron MJ. Coding of Object Location by Heterogeneous Neural Populations with Spatially Dependent Correlations in Weakly Electric Fish. PLOS Computational Biology. 2023;19(3):e1010938. doi:10.1371/journal.pcbi.1010938.
- di Volo M, Destexhe A. Optimal Responsiveness and Collective Oscillations Emerging from the Heterogeneity of Inhibitory Neurons. arxiv. 2020;.
- Perez-Nieves N, Leung VCH, Dragotti PL, Goodman DFM. Neural Heterogeneity Promotes Robust Learning. Nature Communications. 2021;12(1):5791. doi:10.1038/s41467-021-26022-3.
- Gast R, Solla SA, Kennedy A. Effects of Neural Heterogeneity on Spiking Neural Network Dynamics. arxiv. 2022;doi:10.48550/arXiv.2206.08813.
- 52. Shen G, Zhao D, Dong Y, Li Y, Zeng Y. Dive into the Power of Neuronal Heterogeneity. arxiv. 2023;.
- Doty B, Mihalas S, Arkhipov A, Piet A. Heterogeneous "Cell Types" Can Improve Performance of Deep Neural Networks. bioRxiv. 2021; p. 2021.06.21.449346. doi:10.1101/2021.06.21.449346.

- 54. Gittelman JX, Perkel DJ, Portfors CV. Dopamine Modulates Auditory Responses in the Inferior Colliculus in a Heterogeneous Manner. JARO - Journal of the Association for Research in Otolaryngology. 2013;14(5):719–729. doi:10.1007/s10162-013-0405-0.
- 55. Roach JP, Eniwaye B, Booth V, Sander LM, Zochowski MR. Acetylcholine Mediates Dynamic Switching Between Information Coding Schemes in Neuronal Networks. Frontiers in Systems Neuroscience. 2019;13:64. doi:10.3389/fnsys.2019.00064.