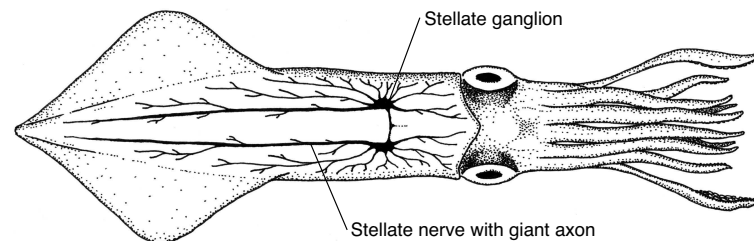


# Coherence effects in the response of neuronal models induced by correlations

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# Main motivations

- A neuron in the brain cortex is subject to a continuous synaptic bombardment of inputs, resembling a background noise  
(A. Destexhe, M. Rudolph, D. Paré - Nature Reviews - Neuroscience - 2003)
- Inputs are mainly originating from the cortex itself, the statistical properties of the neural response (input) can be (roughly) summarized as
  - Frequency range 0 – 200 Hz;
  - Distribution of interspike interval : approximately Poissonian;
  - Spike rate modulations (5 – 10 msec)(M.N. Shadlen & W.T. Newsome, J. Neuroscience - 1998)
- Neurons in the cortex, due to the high connectivity, can receive correlated inputs: correlations seem to be important for information processing. (level of attention )  
(E. Salinas & T.J. Sejnowski, J. Neuroscience - 2000)

How do noise and correlated inputs influence the response of single neurons ?

# Summary

- Brief introduction of the HH model
- Characterization of the stochastic stimulation protocol
- Analysis of the neuronal responses for different noise levels
- Looking for coherence in the neuronal response
- Influence of correlations on the coherent response
- Conclusions



# The Hodgkin-Huxley model



*Hodgkin*

The HH model reproduces the time evolution of the membrane potential and of the ionic currents measured experimentally for a giant squid axon.



*Huxley*

# The Hodgkin-Huxley model

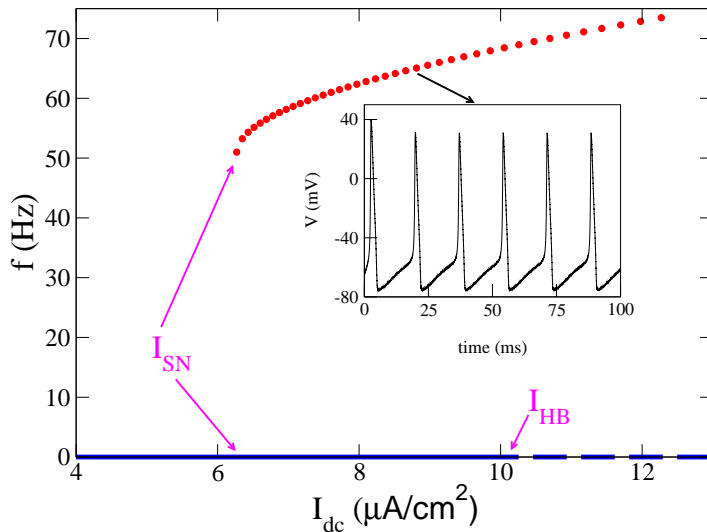


Hodgkin

The HH model reproduces the time evolution of the membrane potential and of the ionic currents measured experimentally for a giant squid axon.



Huxley



- $C = 1 \mu F/cm^2$  - Membrane capacitance
- $V$  - Membrane Potential ( $mV$ )
- $I_j$  - Ionic channel currents ( $\mu A/cm^2$ )
- $g_j$  - Maximal ionic conductances ( $mS/cm^2$ )
- $V_j$  - Ionic reversal potentials ( $mV$ )

$$C\dot{V} = \sum_j I_j + I_{syn} = -g_{Na}m^3h(V - V_{Na}) - g_Kn^4(V - V_K) - g_L(V - V_L) + I_{syn}$$

$$\dot{x} = \alpha_x - x(\alpha_x + \beta_x) \quad x = n, m, h \quad \text{gating variables}$$

$\alpha_x = \alpha_x(V)$  and  $\beta_x = \beta_x(V)$  are highly nonlinear functions.



# High-input regime

- Instead of a constant current  $I_{dc}$ , we consider  $N_E$  excitatory (EPSP) and  $N_I$  inhibitory postsynaptic inputs (IPSP), each corresponding to a voltage kick  $\Delta V = 0.5 \text{ mV}$ .
- These inputs originate from **uncorrelated** neurons emitting **Poissonian spike trains** with frequency  $\nu = 100 \text{ Hz}$ .
- This amounts to one excitatory (resp. inhibitory) Poissonian spike train with frequency  $\nu_E = N_e \times \nu \sim 10^4 - 10^5 \text{ Hz}$  (resp.  $\nu_I = N_I \times \nu$ ) for  $N_e \sim N_I \sim 100 - 1,000$ .
- Firstly **independent** inputs are considered, and then also the effect of **correlations** among the inputs is analyzed.

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For these frequencies the **net input spike count** within a temporal window  $\Delta T (\geq 1 \text{ msec})$  is essentially **Gaussian distributed** and it can be characterized by

$$\text{average } \mu = \nu(N_E - N_I)\Delta T \quad ; \quad \text{variance } V = \nu(N_E + N_I)\Delta T = \nu\sigma^2\Delta T$$

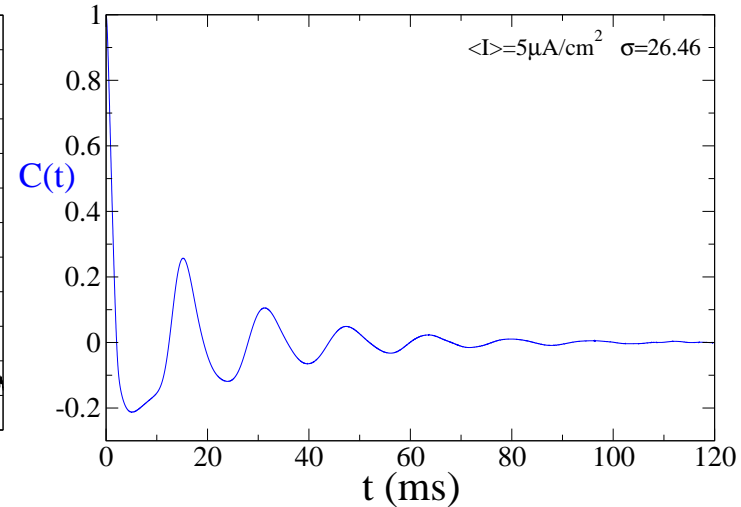
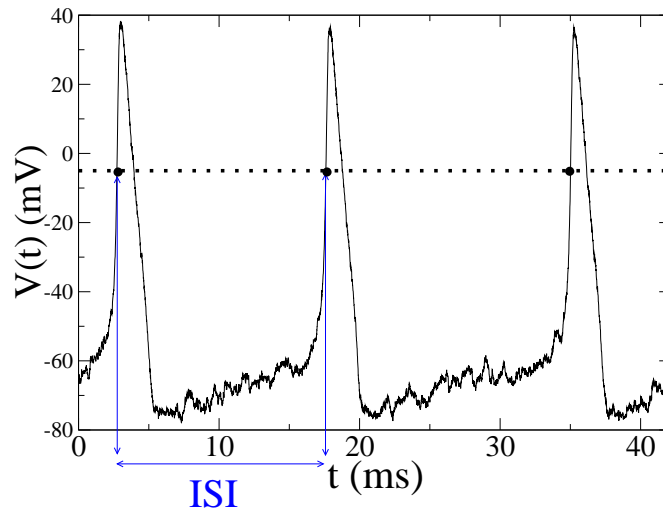
The response of the neuron is examined for **fixed** average input current

$$\bar{I} = C\Delta V\nu(N_E - N_I)$$

by varying only the **standard deviation of the noise**  $\sigma$



# Statistical and dynamical indicators



- ISI distribution  $\rightarrow P_{ISI}(t)$ ;
- $CV = \frac{Std(ISI)}{\langle ISI \rangle} \rightarrow$  coefficient of variation of the ISIs:  
 Poissonian distribution  $\rightarrow CV = 1$   
 regular sequence  $\rightarrow CV = 0$ ;
- $\tau_c = \int_0^\infty C^2(t) dt \rightarrow$  correlation time,  

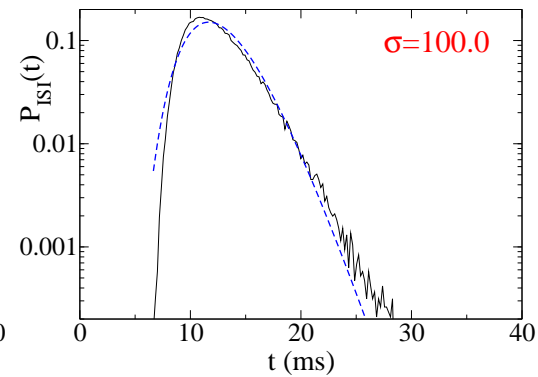
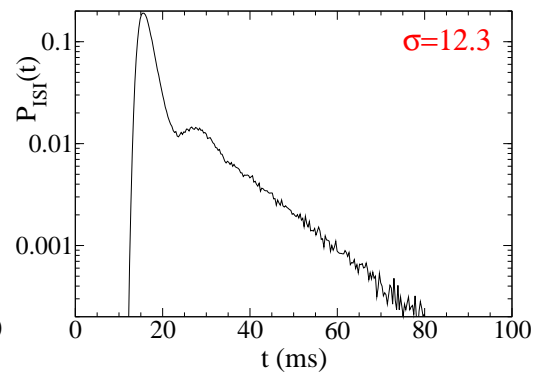
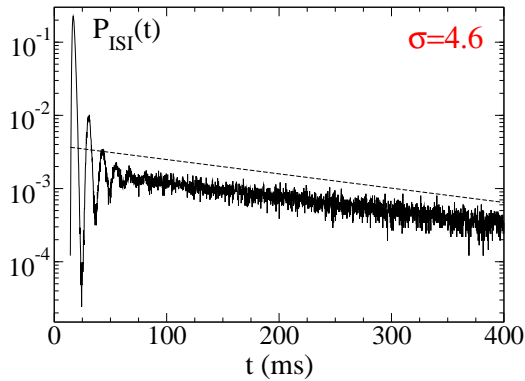
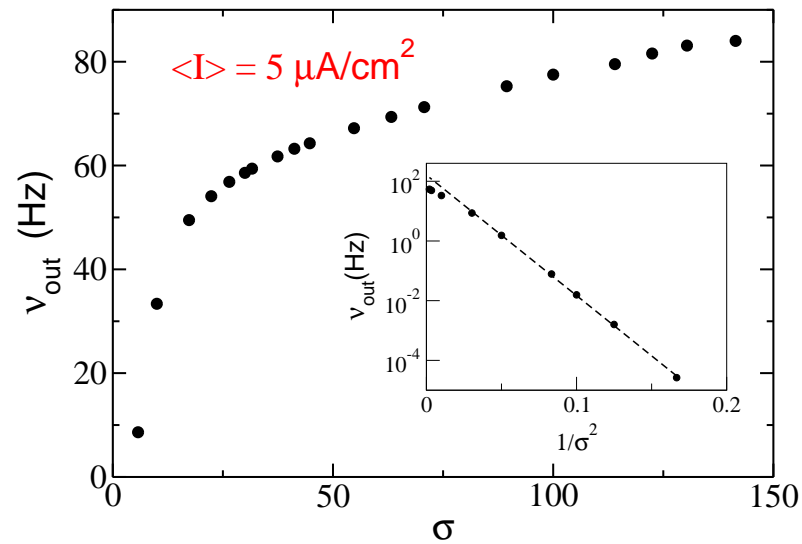
$$C(\tau) = \frac{\langle V(t+\tau)V(t) \rangle - \langle V \rangle^2}{\langle V^2 \rangle - \langle V \rangle^2}$$





# Response of the silent neuron

The HH neuron is in the silent state, i.e. the average input current  $\bar{I}$  is smaller than  $I_{SN}$ .



# Response of the silent neuron

## Firing activated by noise

Two mechanisms compete:

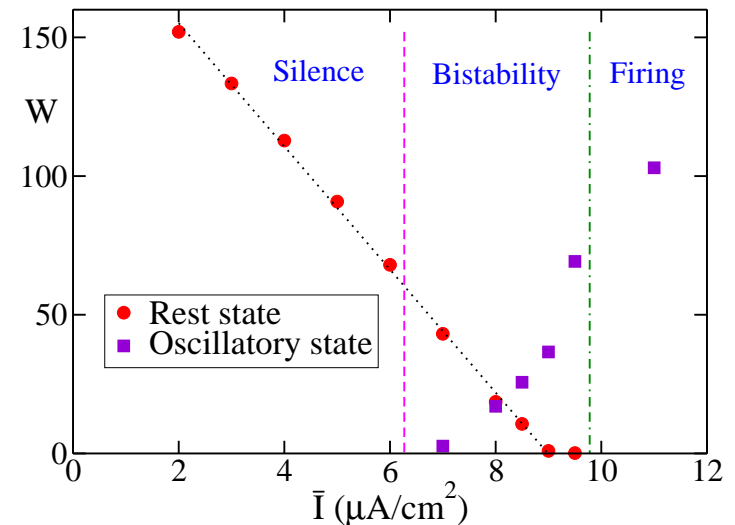
- the HH dynamics tends to relax towards the rest state;
- noise fluctuations lead the system towards an excitation threshold.

The dynamics of  $V(t)$  resembles the overdamped dynamics of a particle in a potential well under the influence of thermal fluctuations, and the firing times can be expressed in terms of the Kramers expression (for sufficiently small noise)

$$t_a \propto e^{W_S/\sigma^2}$$

the time distribution is Poissonian ( $CV = 1$ ).

- for  $\sigma < \sqrt{W_S}$  → Activation Process
- for  $\sigma > \sqrt{W_S}$  → Diffusive Dynamics



# Response of the silent neuron

## High noise limit

The effect of noise fluctuations on the neuron dynamics is twofold:

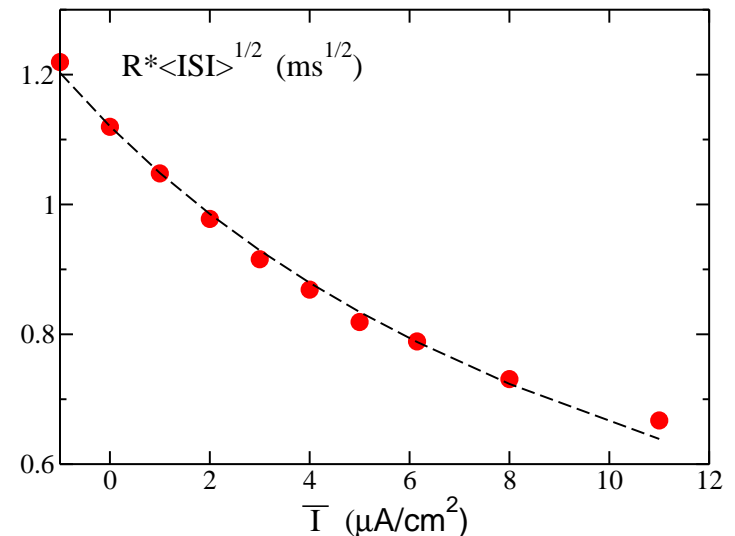
- a constant current  $\bar{I}$  driving the system;
- a stochastic term with zero average.

The dynamics of  $V(t)$  can therefore be described in terms of a Langevin process with a drift and the distribution of the first passage times is given by the inverse Gaussian distribution:

$$f(t) = \frac{\alpha}{\sqrt{2\pi\beta t^3}} e^{-\frac{(t-\alpha)^2}{2\beta t}}$$

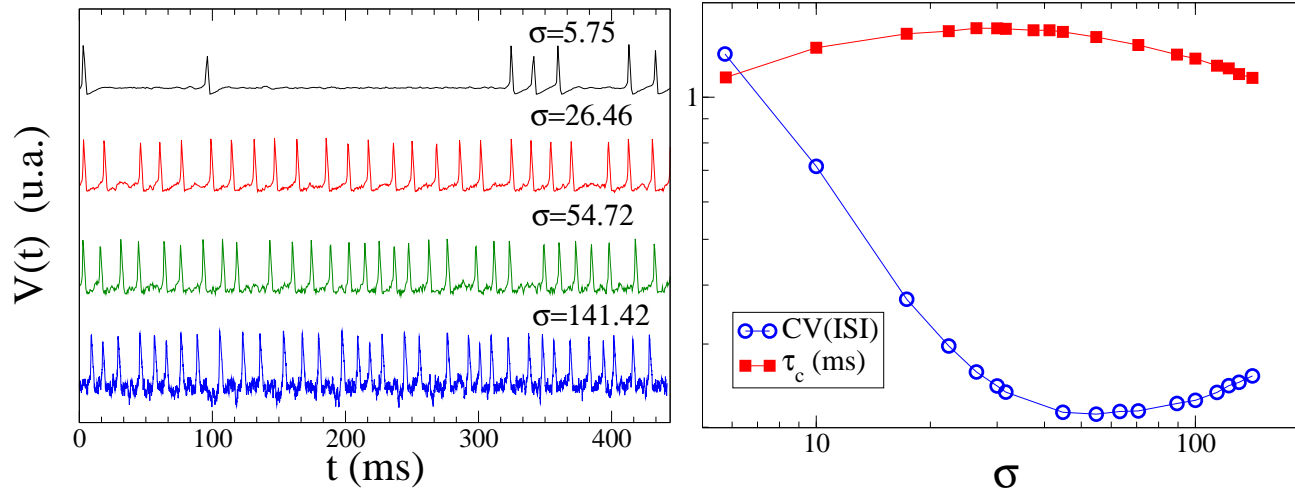
In this case the coefficient of variation should be given by

$$CV \propto \frac{\sigma}{(\bar{I} + I_0)\sqrt{\langle ISI \rangle}}$$



# Coherence resonance

## Coherence of the emitted spike trains



In the silent and bistable regime ( $I < 8\mu A/cm^2$ ):

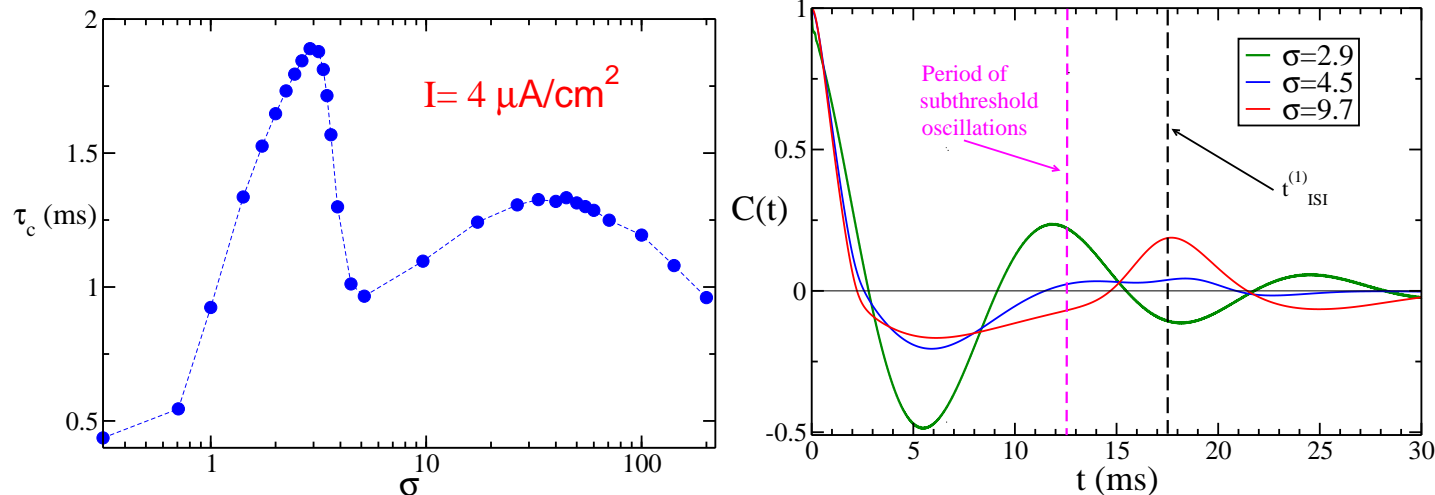
- by increasing  $\sigma$  the firing rate increases, the spike train becomes more regular (**Activation Process**);
- the maximal coherence is reached for an optimal  $\sigma$ -value;
- for higher noise amplitudes the noise influence even the duration of the single spike, the response becomes again more irregular (**Brownian motion + drift**).

A. Pikovsky & J. Kurths, PRL 78, 775 (1997) B. Lindner *et al.*, Phys Rep. 392 (2004) 321-424



# Coherence resonance

## Coherence of the subthreshold oscillations



A second coherence resonance is revealed by analyzing the correlations of the potential:

- for  $\sigma < 3$  almost no spikes are emitted, but the increase of noise leads to more and more regular subthreshold oscillations;
- for  $\sigma > 3$  the statistics of the emitted spikes is no more negligible and this decorrelates the signal;
- for  $\sigma > 10$  the dynamics is dominated by sequences of spikes and a second peak occurs related to the regularization of the spike trains.

S. Luccioli, T. Kreuz, A.T. Phys. Rev. E (2006)



# The FitzHugh-Nagumo Model

The FitzHugh-Nagumo (FHN) model is a two dimensional "simplification" of the HH model:

$$\begin{aligned}\dot{V} &= \phi\left(V - \frac{V^3}{3} - W\right) \quad ; \\ \dot{W} &= V + a_0 - I(t)\end{aligned}$$

where  $V$  is a voltage-like variable,  $W$  is a recovery variable and  $a$  is the bifurcation parameter.

For  $\phi = 100$  the **silent regime** is observed for  $a_0 > 1$ , while at  $a_0 < 1$  one has **periodic firing**.

$$I(t) = \Delta W_0 \left[ \sum_{k=1}^{N_e} \sum_l \delta(t - t_k^l) - \sum_{m=1}^{N_i} \sum_n \delta(t - t_m^n) \right]$$

We examine the FHN model subject to  $N_E$  (resp.  $N_I$ ) trains of excitatory (resp. inhibitory) post-synaptic potentials, in the **balanced** case (i.e. for  $N_E = N_I \equiv N$ ) where  $\langle I \rangle \equiv 0$  for  $a_0 = 1.05$  and  $\Delta W_0 = 0.0014$ .



# Correlations via shared inputs

- The **degree of correlation** among different synapses  $\rho$  is given by the average fraction of synapses delivering kicks at the same time;
- Correlations **ONLY** among either **excitatory** or **inhibitory** inputs are considered in the **balanced case**  $N_E = N_I \equiv N$ ;
- The superposition of  $N$  **correlated** ( $\rho$ ) Poissonian spike trains with rate  $\nu_0$  gives rise to a sequence of kicks of variable amplitude (**binomially distributed**) and with ISIs **Poissonian distributed** with rate  $\nu_x = \nu_0/\rho$ ;
- The uncorrelated **excitatory** (resp. **inhibitory**) inputs are small kicks delivered at **high rate**  $\nu_U = N\nu_0$  (this is an almost continuous background);
- The correlated **inhibitory** (resp. **excitatory**) inputs are large amplitude events delivered at a **much lower rate**  $\nu_x \ll \nu_U$ ;



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- The correlated **inhibitory** (resp. **excitatory**) inputs are large amplitude events delivered at a **much lower rate**  $\nu_x \ll \nu_U$ ;
- The effect of the **uncorrelated inputs** lead to a **renormalization** of the bifurcation parameter:  $\bar{a} = a_0 \pm (N\nu_0\Delta W_0)$ ;
- The influence of **correlated kicks** is embodied in **noise variance**  $\sigma^2 \simeq \rho_x \nu_0 N^2 \delta W_0^2$ ;

M.N. Shadlen & W.T. Newsome (1998) – E. Salinas & J. Sejnowski (2000)

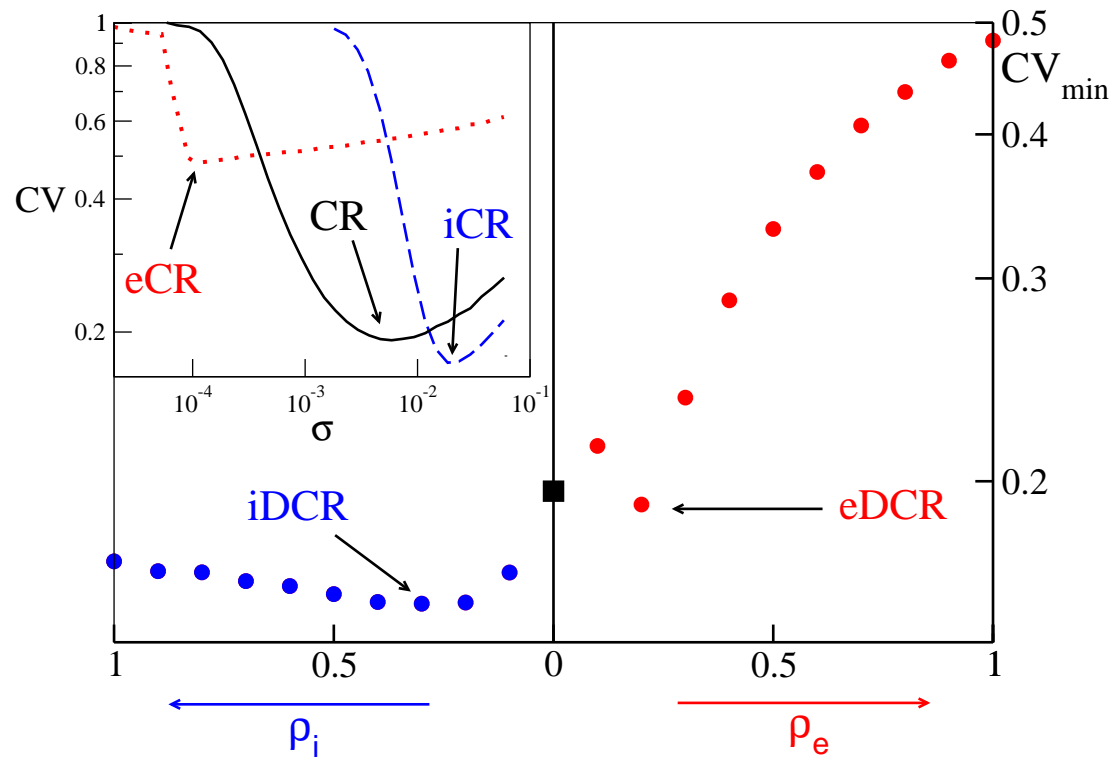




# Response to correlated inputs

We have studied the response of the (balanced  $N_E = N_I$ ) Fitz-Hugh Nagumo model in the silent regime for excitatory (resp. inhibitory) correlated inputs by varying **INDEPENDENTLY** correlation  $\rho$  and noise variance  $\sigma^2$ .

- Coherence Resonance (CR) is observed for any **excitatory** (resp. **inhibitory**) level of correlation at finite noise amplitude.
- Double Coherence Resonance (DCR) : an absolute CR with respect to noise and correlation can be identified for **excitatory** (resp. **inhibitory**) correlated inputs.

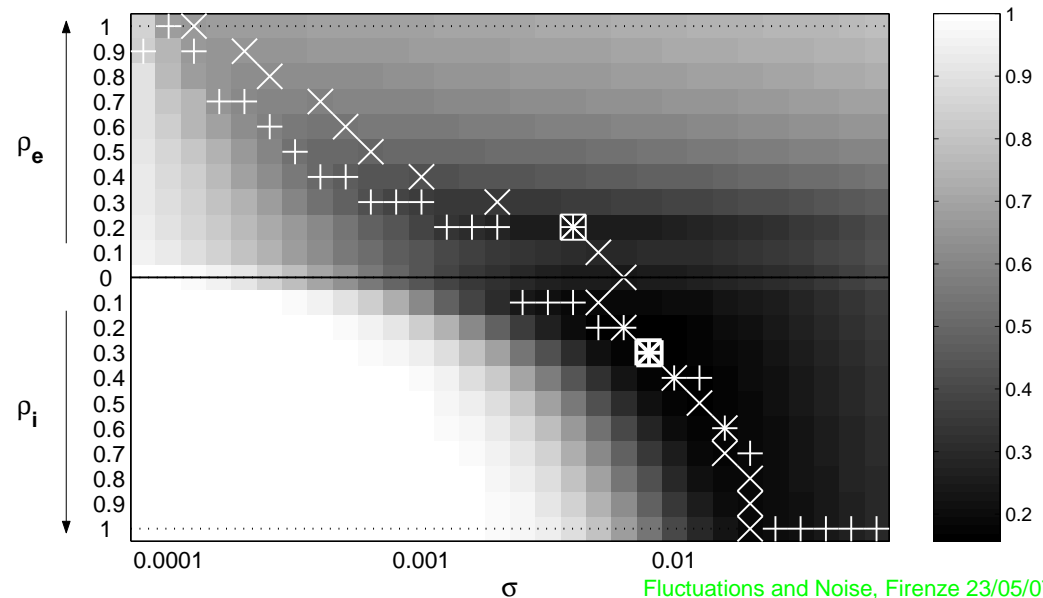


# Response to correlated inputs

- CR has been usually observed for **uncorrelated** inputs with respect to **noise intensity**  
Hu Gang et al., PRL 71, 807 (1993) – A. Pikovsky & J. Kurths, PRL 78, 775 (1997)
- A **second type of CR** has been also observed with respect to the **level of correlation** :
  - lasers – Buldú et al, PRE (2001)
  - digital circuits – Brugioni et al., PRE (2005)
  - chemical reactions – Beato et al. PRE (2005)
  - neuronal models – Casado PLA (1997)
- The origin of the **first CR** has been fully explained, while for the the **second CR** this remains unclear.

A **DCR** with respect to **correlation** and **noise** intensity is a new phenomenon.

- **X** → minima for fixed  $\rho$
- **+** → minima for fixed  $\sigma$



# Response to correlated inputs

## Strong excitatory correlation

The coherence phenomenon is now determined only by the **kick amplitude** and not by the properties of the asymptotic stochastic processes, since in the present case the output can be always described as a **Poissonian process with a refractory time**:

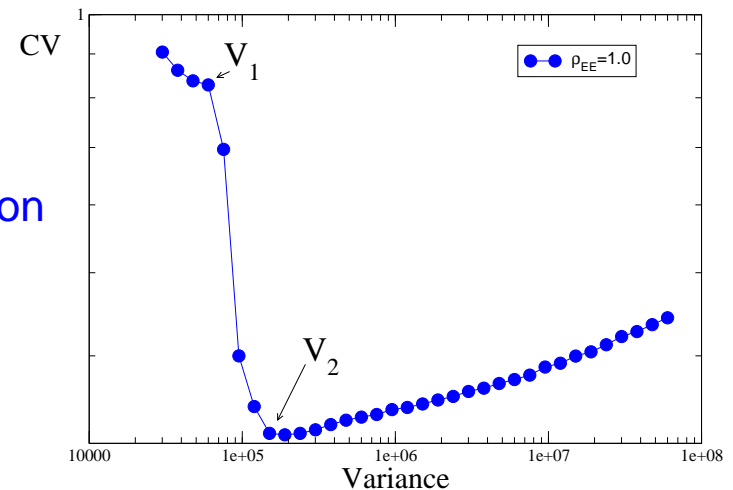
$$CV = 1 - \frac{T_{ref}}{\langle ISI \rangle}$$

For increasing variance ( $N$ ) the amplitude of the correlated kicks increases. and due to the uncorrelated (inhibitory) input the systems is more and more silent.

- for  $Variance < V_1 \rightarrow$  Activation Process
- for  $V_1 < Variance < V_2$   
1 kick may be sufficient to induce a spike emission
- for  $V_2 < Variance$   
One kick elicits always a spike -  $T_{ref}$  decreases

For  $Variance > V_2 \rightarrow \langle ISI \rangle = \rho_e / \nu_0$ ,

**1 : 1 synchronization** between input and output (apart from the refractory period).

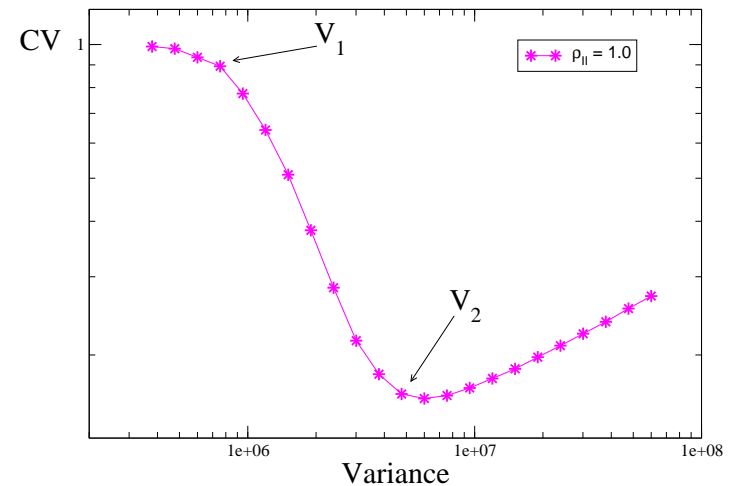


# Response to correlated inputs

## Strong inhibitory correlation

At large variances the dynamics is again ruled by the the **amplitude of the correlated kicks**, but at lower variances the inhibitory kicks are quite infrequent and their amplitude is not sufficient to influence the dynamics.

- for  $Variance < V_1$   
Silent regime - Activation Process -  $CV \simeq 1$
- for  $V_1 < Variance < V_2$   
Dynamics dominated by uncorrelated excitatory input leading the system in the repetitive firing regime
- for  $Variance > V_2$   
Each inhibitory kick induces a certain delay in the spike time of the neuron - A multimodal structure appears in the ISI distribution



Frequency of the correlated kicks ( $\nu_0$ )  $\ll$  Frequency of the uncorrelated kicks ( $N\nu_0$ )

# Response to correlated inputs

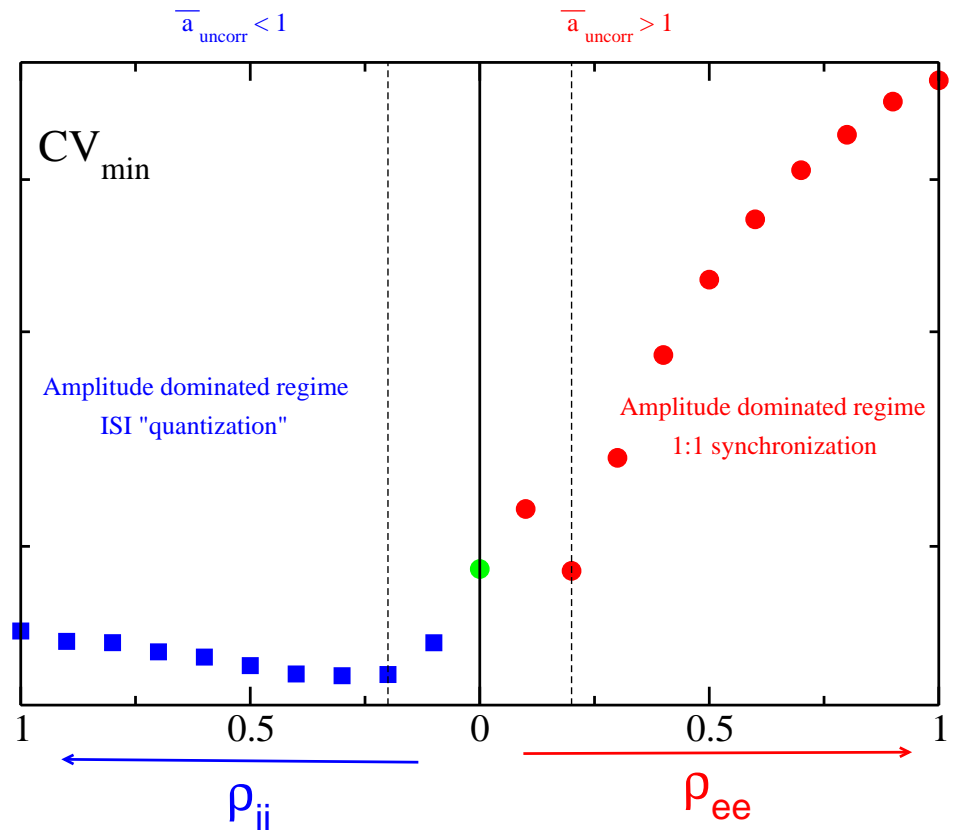
## Maximal Coherence

Absolute minima of CV corresponding to maximal coherence are observed at **finite noise and correlation** for both **inhibitory** and **excitatory** case.

The extrema of  $CV_{min}$  indicate the change in the mechanisms inducing CR, from amplitude dominated to **usual mechanisms** related to the crossover from activated to (biased) diffusive processes.

The  $CV_{min}$  associated to **inhibitory correlations** are **lower** since the system is driven in the repetitive firing regime by the uncorrelated **excitatory** inputs

$$\bar{a} = a_0 - (N\nu_0\Delta W_0)$$



T. Kreuz, S. Luccioli & A.T. PRL (2006)



# Conclusions

## ● Uncorrelated stochastic inputs

- The response of a **HH neuron** depends only on the **average** and the **variance** of the input, at least in the high input regime;
- at  $I < I_{SN}$  the neuronal firing, induced by the stochastic inputs, can be expressed as an **activation process** at low variances ( $\sigma^2$ ), while for large  $\sigma^2$  this process becomes essentially **diffusive**;
- **coherence resonance** can be observed in a large interval of currents in the silent and bistable regime whenever  $W_S > W_O$ ;
- a second **coherence resonance** (associated to **subthreshold oscillations**) coexists with the usual one;

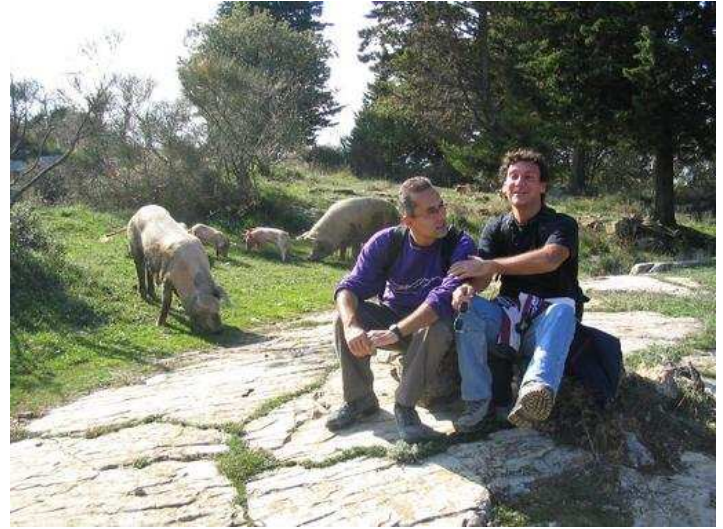
## ● Correlated stochastic inputs

- new mechanisms for the **coherence resonance** have been reported at high excitatory and inhibitory correlations;
- **maximal coherence** can be induced by an optimal combination of noise and correlation



# Credits

- **Stefano Luccioli** - Msc in Physics (2004-2005)
- Dynamics of realistic single neuronal models



- **Thomas Kreuz** - Marie Curie Fellow (2005-2006)
- Dynamical Entropies in Assemblies of Neurons



<http://www.fi.isc.cnr.it/users/alessandro.torcini/neurores.html>