## **RSC Advances**



PAPER View Article Online



Cite this: RSC Adv., 2017, 7, 46092

DOI: 10.1039/c7ra06323g

Accepted 4th September 2017

Received 6th June 2017

rsc.li/rsc-advances

# 1/f pink chaos in nanopores

Vishal V. R. Nandigana (10 \*\* and N. R. Alurub

Nanopores have been used for myriad applications ranging from water desalination, gas separation, fluidic circuits, DNA sequencing, and preconcentration of ions. In all of these applications noise is an important factor during signal measurement. Noisy signals disrupt the exact measuring signal in almost all of these applications. In this paper, we rationalize whether current oscillations should be classified only as noise or the physical disturbance in ionic charges has some other meaning. We infer that the physical disturbance in ionic charges and the current oscillations are not noise but can be chaos. Chaos is present in the system due to depletion of the ions, created by nonequilibrium anharmonic distribution in the electrostatic potential. In other words, multiple electric potential wells are observed in the nanoporous system. The multiple electric potential wells leads to bi-directional hopping of ions as the ions transport through the pore. The bi-directional hopping results in current oscillations. This paper suggests that chaos exists from a deterministic perspective and that there is no stochastic element leading to current oscillations. We prove this case by considering a simple oscillator model involving the electrostatic and dissipative forces in order to model ionic current. We observed current oscillations even in the absence of a stochastic noise force. Hence, we state that current oscillations in nanopores can be due to chaos as well and not necessarily due to noise. Furthermore, the color associated with the chaotic spectrum is not brown but pink, with 1/f type dynamics similar to the 1/f type pink noise presented by theorists and experimentalists. However, the 1/f type pink chaos exists due to deterministic current oscillations and not due to a stochastic fluctuating noise force.

## Introduction

Ionic logic-based circuits have witnessed a tremendous surge in recent years, ranging from single molecule sensing,2,3 DNA sequencing, 1,4-6 nanopower generators and electrolyte cells.8 The unsupported electrolyte is actively impelled across the ion membrane under an electric potential. One of the measuring signals is the current as ions are transported through the pore. Recently, it was found that the current signal has built-in oscillations as the ions traverse through the pore. These oscillations have been confirmed as noise by earlier theorists. 9-13 The noise is perceived as random statistical fluctuation in the electrical current signal. Variations in noise were explained earlier by the mechanistic motion of the charges due to surface defects, or else they were perceived as thermal fluctuations within the purview of the classical fluctuation-dissipation theorem.<sup>14</sup> Several instances of noise in membrane phenomenology occur due to the cooperative motion of ions due to random memory force. In other words, a noise force is used to model the mobility of the ions. Not only were the current oscillations perceived as noise but the resulting current spectra had a 1/f type pink noise in the low frequency regime. What causes the ionic membranes in electrolyte cells to exhibit pink noise is a subject for inspection. Several of these notions employ fluctuations in the inherent ionic mobility of the electrolyte.9 Another reason pertains to the stochastic variation in the protonation and deprotonation reactions near the charge site of the membrane surface. Finally, experimentalists have also argued that the membrane structural constituents also prompt the low frequency pink noise.10,12 Now, to what extent colored pink noise is the true state representation of the current dynamics under a dynamical driving/dissipative potential is a challenging question. Is it possible that physical disturbance in the ionic charges and current oscillations in the ion-selective membranes can be triggered by some phenomenon other than statistical fluctuation, namely noise? In this paper, we state that the physical disturbance in ionic charges is due to chaos, and current oscillations can be perceived as chaos as well and we need not restrict ourselves in claiming current oscillations as noise. We note that chaos occurs when the potential of mean force on a charged ion is anharmonic in the nonequilibrium state space. In other words, a multi-well potential acts on the charged ion when the ion is driven under an electric field. The anharmonic or multi-well potentials arise due to the depletion of ions owing to the charge conservation. The depleted ions

<sup>&</sup>quot;Department of Mechanical Engineering, Indian Institute of Technology, Madras, Chennai 600036, India. E-mail: nandiga@iitm.ac.in; Web: https://mech.iitm.ac.in/ meiitm/personnal/dr-vishal-v-r-nandigana/; Tel: +91-44-2257-4668

<sup>&</sup>lt;sup>b</sup>Department of Mechanical Science and Engineering, Beckman Institute for Advanced Science and Technology, University of Illinois at Urbana – Champaign, Urbana, Illinois 61801, USA

**RSC Advances** Paper

create an electric field barrier that prevents other ions from entering into the nanopore. This causes the ions to hop in a bidirectional manner before the ions enter into the nanopore. The bi-directional hopping of ions results in current oscillations leading to chaos. We argue that the oscillations observed in the current signal have a deterministic origin and do not arise due to stochastic fluctuations postulated by earlier theorists. In this study we also observe that the spectrum of the current oscillations is pink in color as it exhibits "1/f" type dynamics similar to pink noise.

In this work, we understand that the origin of chaos occurs when the nanomembrane is selective to ions. More precisely, the membrane is ideally selective to one particular type of ion which is opposite to the membrane charge. The disproportionate opposing charge has the space to confine a linear and anharmonic potential under an external electric field given by the mean field approximation. That means a multi-well potential exists in an ideal ion-selective nanoporous membrane when an electric field is applied on the ion. In the case of a non-ideal membrane, we cease to observe chaos. Now, the transport of charge over an electrostatic force in an ideal-membrane results in electrostatic potential instability due to the existence of multiple potential wells. We propose this mechanism as the 'Potential-charge momentum theorem'. We further demonstrate a simple anharmonic oscillator model, calling it the 'VN Oscillator model' which stands for Variational Non-equilibrium potential oscillator, to explain the chaos mechanism. The current oscillations in the ideally selective nanoporous membrane can be applied in the field of nanopore sensing for DNA sequencing and chemical reactions. Furthermore, the current oscillations can be understood to manipulate fluid flow in fluidic logic circuits in areas of combinatorial chemistry, biology and electronic applications.

#### Theory II.

#### Charge oscillator model

Before looking at the VN oscillator model, we first disclose a unique charge-transfer and potential-dissipation model to uncover the notion of chaos that is exhibited by ion-selective nanogating channels. To demonstrate this phenomenon, we discerned the idea of the Maxwell model,  $\nabla \cdot \vec{D} = \delta \rho_f$ , where  $\vec{D}$  is the electric displacement vector field accounting for the effects of free and bound charge within the open-ended state space and  $\delta \rho_{\rm f}$  is the free charge in space (ionic charge) confined within a given volume. The free charge is a derivative of the total charge,  $\delta \rho$ , which is the sum total of the free and bounded charges  $(\delta \rho_b)$ ;  $\delta \rho = \delta \rho_b + \delta \rho_f$ . The electric displacement field was demarcated using the constitutive relationship  $\vec{D} = \epsilon_0 \vec{E} + \vec{P}$ , where *P* is the polarization density whose divergence envisions the density of bound charges,  $\nabla \cdot \vec{P} = \delta \rho_b$ , and  $\epsilon_0$  is the dielectric permittivity of free space. The stretching and compressive nature of the electric field  $(\vec{E})$  triggered the total charge in the confined ion gating regime,  $\nabla \cdot \vec{E} = \frac{\delta \rho_{\rm f}}{\epsilon_0} - \frac{\nabla \cdot \vec{P}}{\epsilon_0}$ . The negative divergence in polarization density implied that the charge

transfer of the bound charge was directionally dissimilar to the free charge because of the dielectric polarization. In a linear, homogeneous and isotropic dielectric with the electric field instantaneously varying with time, the polarization density is constituted by  $\vec{P} = \epsilon_0 \chi \vec{E}$ , where  $\chi$  is the degree of polarization in conjunction with the electric field. The electric field caused the bound and free charges to polarize in space constituting Gauss's theorem,  $\nabla \cdot \epsilon_{\rm r} \vec{E} = \frac{\delta \rho_{\rm f}}{\epsilon_0}$ . The equation was arrived at by substituting the polarization density  $\vec{P}$  in the previous  $\nabla \cdot \vec{E}$ equation.  $\epsilon_r$  is the dielectric permittivity of the medium (water) and correlated with the degree of polarization,  $\chi$ , by  $\epsilon_{\rm r}=1+\chi$ . The notion of Maxwell's electric displacement field and electric flux conservation was conferred upon the ion gating membrane resulting in the governing equation  $\iiint_{\delta\varOmega} \nabla \cdot \epsilon_{\vec{r}} \vec{E} = \iiint_{\delta\varOmega} \frac{\delta \rho_{f}}{\epsilon_{0}}$ , where the electric field is related to the Coulomb potential  $(\phi)$ ,  $\vec{E} = -\nabla \phi$ , and  $\delta \Omega$  is the gating membrane. We radially averaged the governing equation to  $\mbox{give} \quad \frac{\partial}{\partial z} \bigg( \varDelta(z) \epsilon_{\rm r} \frac{\partial \hat{\phi}}{\partial z} \bigg) = - \frac{\varDelta(z)}{\epsilon_0} \bigg( \delta \hat{\rho}_{\rm f} + \frac{2\sigma_{\rm S}(z)}{R(z)} \bigg) \quad \mbox{after} \quad \mbox{invoking} \quad \label{eq:continuous}$ Gauss's law,  $\epsilon_r \nabla \phi \cdot \vec{n} = \frac{\sigma_s(z)}{\epsilon_0}$ , where  $\vec{n}$  is the unit normal to the

surface and z is the axial direction of the membrane system. The above area-averaged model was configured across the inhomogeneous system, namely along the nanopores and micropores.  $\Delta(z)$  is the cross-sectional area, accounting for the micro and nanogating membrane radius given by R(z), and  $\sigma_s(z)$ is the variational surface charge density of the nanogating membrane along with the microporous membrane. Now, we can inspect the transfer of charges with the mass transfer of ions. The dynamic mass transfer of ions is correlated by the diffusive ionic gradients along with the electric mobility polarization density formulated by  $c\Omega \vec{E}$ , where  $c\Omega$  is the degree of polarization of the ion required to translocate the ion in the presence of an electric field. We call this model the 'charge oscillator model'. The rate of change of the ionic charge that was radially averaged was given as

 $\varDelta(z)\frac{\partial \hat{c}_i}{\partial t} = \frac{\partial}{\partial z} \left( \varDelta(z) D_i \frac{\partial \hat{c}_i}{\partial z} \right) - \frac{\partial}{\partial z} (\varOmega_i \varDelta(z) \hat{c}_i z_i \hat{E}), \text{ where } \varOmega_i \text{ is the }$ mobility of the individual ions which is related to the diffusion

coefficient of each ion ( $D_i$ ) given by Einstein's relation,  $\Omega_i = \frac{D_i}{K_{\rm B}T}$ 

 $K_{\rm B}T$  is the thermal energy with  $K_{\rm B}$  being the Boltzmann constant and T being the absolute thermodynamic temperature. The concentration of ions within the control volume was radially averaged and defined as  $\hat{c}_i$ .  $z_i$  is the valence of each ion

and  $\hat{f} = \frac{1}{\Delta(z)} \int_0^r \int_0^{2\pi} (f(r,\theta)) r d\theta dr$  denotes the area-averaged mapping function. The current density in the system was given

as 
$$I_{\text{tot}} = F \sum_{i=1}^{n} z_i \Gamma_i$$
, where  $\Gamma_i = -D_i \frac{\partial \hat{c}_i}{\partial z} + \Omega_i z_i \hat{c}_i E$  is the total area-

averaged flux of each ion. The total current across the nanogating membrane was manifested from the current density by multiplication with the membrane cross-sectional area.

The coupled electric displacement field and charge transfer governing equations were numerically solved using the finite

volume method in OpenFOAM<sup>15</sup> (Open Field Operation and Manipulation). The details regarding the solver implementation are discussed elsewhere. <sup>16-19</sup>

#### B. VN oscillator model

We now discuss another simple oscillator model proposed by us to understand the notion of chaos. We call this oscillator model the 'variational non-equilibrium potential oscillator' or VN oscillator. In this model, we assumed that the nanogating membrane was ideally selective to one type of ion, *i.e.*, the membrane allowed only ions of opposite polarity to that of the membrane charge. The model states that the mass transfer of each dissimilar ion  $(\vec{n_i})$  with respect to the charge of the ion  $(q_i)$  constituted the equation of motion of the

charged particle, manifesting current:  $\vec{I} = \sum_{i}^{N} \vec{n}_{i} q_{i}$ . The mass

transfer rate was mapped to the charge transfer rate  $\left(\dot{q}_{n}^{\ \ \delta V}=
ight.$ 

$$\frac{(q_n^{\delta V}(t+\Delta t)-q_n^{\delta V}(t))}{\Delta t}$$
 which was again mapped to the

Eulerian space inferred in the net ion motion  $\vec{I} = n_{\delta V} \delta V \times \left[ \frac{(q_n^{\delta V}(t+\Delta t) - q_n^{\delta V}(t))}{\Delta t} \right] = n_{\delta V} \delta V q_n^{\delta V}(t+\Delta t) \frac{1}{\delta \vec{l}} \lim_{\delta \vec{l} \to \vec{L}_{\rm nano}} \frac{\delta \vec{x}_n}{\delta t},$ 

where  $n_{\delta V}$  is the number of ionic charges in the control volume  $\delta V$ , varying across the nanopore and micropore.  $q_n^{\delta V}$  is the charge of the ionic particle,  $\Delta t$  is the time step and  $\delta l$  is the length of the gating membrane accounting for either the micro or nanomembrane. We propose a theorem called the 'potential charge-momentum theorem (PCM)' to rationalize the displacement of a charged particle,  $\vec{x}_n$ . The theorem solves Newton's law of motion for a coarse grained charged particle

given by  $m_{\delta V} \frac{\partial^2 \vec{x}_n}{\partial t^2} + \dot{\lambda} \frac{\partial \vec{x}_n}{\partial t} = F|_{t_n}$ , where  $m_{\delta V}$  is the mass of the ionic particle,  $\dot{\lambda} = 1/\Omega$  is the mass transfer rate of the confined momentized particle given as  $\left(\frac{K_B T}{D_b}\right)$ ,  $D_b$  is the bulk diffusion coefficient of the ionic particle,  $(K_B T)$  is the thermal energy of

coefficient of the ionic particle,  $(K_BT)$  is the thermal energy of the ionic particle, and  $F|_{t_n}$  is the mean force harnessed at time  $t_n$  by the ionic particle. The mean interaction potential force produced by electrostatic interaction is

$$\left.F\right|_{t_n} = rac{q_{
m s}q_n^{\delta V}}{4\pi\epsilon_0\epsilon_{
m r}|ec{x}_n-ec{x}_0|^2} + rac{q_n^{\delta V}V_0}{\left|ec{x}_{
m b2}-ec{x}_{
m b1}
ight|}, ext{ where } q_{
m s} = \sigma_{
m s}\delta S ext{ is the }$$

charge constituted by the surface of the nano and micro membrane and  $\delta S$  is the surface area of the membrane.  $\sigma_s$  is the surface charge density of the micro or nanogating membrane. An electrostatic potential,  $V_0$ , was applied across the micronanoporous architecture within a distance of  $|\vec{x}_{b2} - \vec{x}_{b1}|$ .  $\vec{x}_0$  is the position of the charged nano-membrane. The PCM model was marched in time for  $\Delta t$  using a well known velocity-verlet

algorithm:  $\vec{x}^n(t+\Delta t) = \vec{x}^n(t) + \dot{\vec{x}}(t)\Delta t + \frac{1}{2}\ddot{\vec{x}}(t)(\Delta t)^2$ . The ionic particle acceleration at  $(t+\Delta t)$  was configured using the interaction potential force at  $\vec{x}^n(t+\Delta t)$ . Finally, the velocity is

$$\dot{\vec{x}}(t+\Delta t) = \dot{\vec{x}}(t) + \frac{1}{2} \Big( \ddot{\vec{x}}(t) + \ddot{\vec{x}}(t+\Delta t) \Big) (\Delta t).$$
 Note, while

calculating the velocity  $\left(\vec{\vec{x}}(t)\right)$  and acceleration  $\left(\ddot{\vec{x}}(t)\right)$  we had to substitute the right length  $\delta l$  and surface charge  $\sigma_{\rm s}$  to account for the micro and nanopore membranes.

## III. Simulation details

The simulated domain consisted of a nanopore of length  $L_{\rm n}=0.5~\mu{\rm m}$  and diameter  $d_{\rm n}=10~{\rm nm}$ . The diameter of the micropore was  $d_{\rm micro}=500~{\rm nm}$ , while the length of the micropore was  $L_{\rm micro}=4~\mu{\rm m}$ . Phosphate buffer of a constant concentration (0.39 mM NaH<sub>2</sub>PO<sub>4</sub> and 0.61 mM Na<sub>2</sub>HPO<sub>4</sub>) was used in all of the simulations. The temperature was  $T=300~{\rm K}$ . The diffusivities of Na<sup>+</sup>, H<sub>2</sub>PO<sub>4</sub><sup>-</sup> and HPO<sub>4</sub><sup>2-</sup> are  $1.33\times10^{-9}~{\rm m}^2~{\rm s}^{-1}$ , 0.879  $\times~10^{-9}~{\rm m}^2~{\rm s}^{-1}$ , and  $0.439\times10^{-9}~{\rm m}^2~{\rm s}^{-1}$ , respectively. We assumed the dielectric constant of the aqueous solution to be  $\epsilon_{\rm r}=80.^{21}$  We also assumed zero surface charge density on the walls of the micropore,  $\sigma_{\rm m}=0$ , as they were away from the nanopore membrane and could not have an influence on the transport. The charge on the walls of the nanopore was assumed to vary from  $\sigma_{\rm n}=0~{\rm mC}~{\rm m}^{-2}$  to  $\sigma_{\rm n}=-30~{\rm mC}~{\rm m}^{-2}$ . The voltage  $V_0$  was varied from  $V=0~{\rm V}$  to  $V=100~{\rm V}$ .

In the VN oscillator model, the mass  $m_{\delta V}=7\times 10^{-12}$  kg to match the charge oscillator model and the distance  $x_0=L_{\rm micro}, |\vec{x}_{\rm b2}-\vec{x}_{\rm b1}|=L_{\rm n}+2L_{\rm micro}.$  At time t=0, the initial charge was assumed to be equal to the surface charge,  $q_n^{\delta V}|_{t=0}=\sigma_{\rm n}\delta S$ , with  $\delta S=\pi d_{\rm n}L_{\rm n}.$  The time step used in the simulation was 1 ns. The total time of the simulation was 3 ms.

## IV. Results

Schematics of the dynamic charge oscillator model and VN oscillator model are shown in Fig. 1. On solving the charge oscillator model, we observed that the current oscillated with time. The oscillations were bi-directional, indicating that the ions hop back and forth as they enter into the nanopore. The results are shown in Fig. 2(a). Interestingly, the VN oscillator model also showed similar current oscillations where the ion hops bidirectionally similar to the charge oscillator model (see Fig. 2(a)). The magnitude of the oscillations was similar to that of the charge oscillator model, thus indicating that our notion of representing the charge dynamics as a simple potential-dissipative oscillator model is sufficient to capture the physics.

### A. Origin of chaos

We are now interested in answering the question 'why do these oscillations arise'? The oscillations are due to the depletion of ions near the micro-nano interface region, owing to the charge conservation. To elaborate, let us consider an ideal cation selective nanogating membrane as considered in this study. In an ideal cation selective nanopore, only cations are allowed to enter into the nanopore. When an electric field was applied from an anode to cathode, the anions were repelled from the cathode and from the cation-selective membrane (since the membrane allows only cations). Thus, the anions were accumulated or enriched near the cathode-nanogating membrane interface. To maintain neutrality, the cations were also enriched

Paper

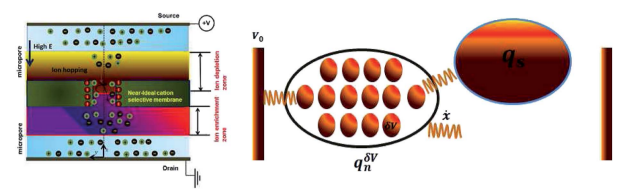


Fig. 1 A schematic physical view of the charge oscillator model (on the left) and the VN oscillator model (on the right). In the charge oscillator model, the following geometry for the micro-nanopore is considered. The nanogating membrane is a cylindrical nanopore of length  $L_n = 500$  nm, and diameter  $d_n = 10$  nm. The nanopore is leveraged with two microporous reservoirs of length  $L_{\text{micro}} = 4 \, \mu\text{m}$  and diameter  $d_{\text{micro}} = 500$  nm. A 1 mM phosphate buffer electrolyte cell is considered in the reservoir. The thermodynamic temperature is T = 300 K. In the VN oscillator model, the reduced mass  $m_{\delta V}$  is considered as  $7 \times 10^{-12}$  kg to match the charge oscillator model.

at this interface. At the anode-membrane interface the anions were repelled from the cation selective membrane due to electrostatic repulsion and attracted towards the anode due to electrostatic attraction, thus the anions were depleted at the anode-nanogating membrane interface. To maintain neutrality, the cations were also depleted at this interface. We observed that this depletion region was very strong for ideal ion-selective nanopores, resulting in non-equilibrium anharmonic potential and an electric barrier. The potential and the electric barrier creates an instability in the ionic charges near the interface. This instability results in bi-directional hopping of ions leading to chaos. From our understanding, this observation of bidirectional hopping of ions is made for the first time in the literature. A schematic of ion enrichment and ion depletion is shown in Fig. 2(b). We note that such oscillations were observed when the nanopore was ideally ion selective to one particular type of ion. Furthermore, we observed a stable current when the membrane was non-ideal, i.e., when the membrane allowed both cations and anions. We played with the ideality of the nanopore using the surface charge density. For nanoporous surface charge densities between  $\sigma_{\rm n}=0$  mC m $^{-2}$  and  $\sigma_{\rm n}=-3$ mC m<sup>-2</sup>, we did not observe current oscillations nor bidirectional hopping of the ions. Only for the nanoporous surface charge density of  $\sigma_n \ge -10 \text{ mC m}^{-2}$  did we observe an ideal ion-selective nature of the nanopore and current oscillations and chaos. Such large surface charge density nanopores can be assumed to be present for aluminum oxide nanopores.<sup>22</sup> We would like to stress the fact that neither the charge oscillator model nor the simple VN oscillator model takes into account any statistical fluctuating force to observe these oscillations. Even the earlier theorists' confirmation of necessitating the fluctuating ionic mobility or nano membrane surface charge fluctuating force to observe current oscillations is not considered in either of these models. The oscillator models manifest current oscillations and chaos in view of dynamic potential instability. Thus, in our study, we state that current oscillations can also be viewed from a chaos perspective and we need not

limit ourselves to analyzing current oscillations from a noise perspective.

When the physics of chaos occurs in nanogating membranes is the next interesting question one has to argue. The key to this lies in our proposed non-dimensional Nandigana number,  $N_v=\left(rac{\sigma_s\delta S}{4\pi\epsilon_0\epsilon_r x_\Delta V_0}
ight)$ . The Nandigana number is arrived at by non-dimensionalizing the VN oscillator model. We considered the following terms to non-dimensionalize the VN oscillator model: distance x is normalized by  $x_{\Delta} = 1$  nm; time t by  $t_{\Delta} = 1$ ns; diffusion coefficient D by  $D_{\Delta} = 1 \text{ nm}^2 \text{ ns}^{-1}$ ; charge q by  $q_{\Delta} =$  $\sigma_{\rm s}$ dS; and potential V by  $V_0$ . The derived Nandigana number correlates the influence of the surface charge density of the membrane with the applied electric potential. The scale of the Nandigana number should be of the order of 0.17 to visualize potential instability instigating chaos. At this Nandigana number, the surface charge density of the nanopore was -10mC  $m^{-2}$ , when the nanopore was ideally ion-selective. The relationship of the current with the Nandigana number is shown in Fig. 2(c) using the charge oscillator model. The figure shows that the ion dynamics were stable until a Nandigana number of 0.05294 and we see unstable ion dynamics for  $N_v >$ 0.05294. Under such a Nandigana number, the nanoporous surface charge density was greater than  $\neg 3$  mC m<sup>-2</sup>. To truly note that the current oscillations were indeed chaotic in nature, we calculated the positive Lyapunov exponent using the current oscillations. The notion of determining the Lyapunov exponent infers upon the dimensionality of the system. In this study, we focused our attention towards the calculation of the maximum Lyapunov exponent,  $\lambda^*$ , from the current-time map. We formed two state vectors,  $\vec{I}_{n1}$  and  $\vec{I}_{n2}$ , from the current signal at an initial time with a distance of  $||\vec{I}_{n1} - \vec{I}_{n2}|| = \delta_0 \ll 1$ . The ion trajectories from the initial distance travelled in time  $\Delta t$  were used to map the new state vector distance,  $\delta_{\Delta t} = ||\vec{I}_{n1+\Delta t}||$  $\vec{I}_{n2+\Delta t}$ . The maximum Lyapunov exponent ( $\lambda^*$ ) in this vector space was formed as  $\delta_{\Delta t} \approx \delta_0 e^{\lambda^* \Delta t}$ ;  $\delta_{\Delta t} \ll 1$  and  $\Delta t \gg 1$ . When  $\lambda^*$ was positive, we observed an exponential divergence of the current time series data, resulting in dynamic instability or

RSC Advances Paper

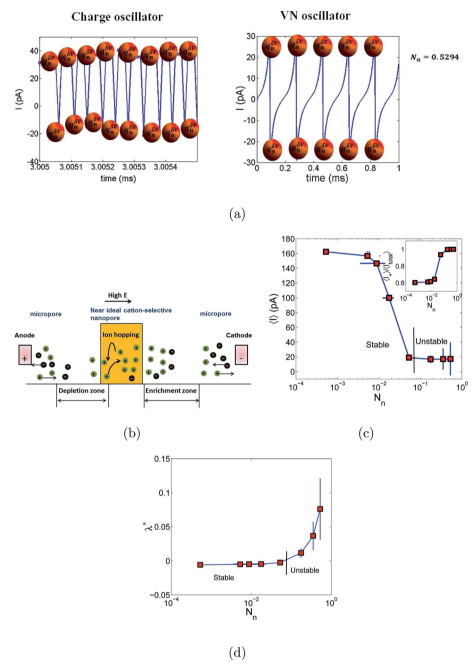


Fig. 2 (a) The dynamic similarity of the charge oscillator and VN oscillator at  $\mathbb{N}_n=0.5294$ . (b) Schematic of the ion enrichment and ion depletion zone in an ideal nanoporous membrane, and (c) variation of the computational current with the Nandigana number. The inset reveals the ideal cation-selective nature  $\left(L=\frac{\langle I_+\rangle}{\langle I_{\text{total}}\rangle}\right)$  of the nanomembrane with the Nandigana number. L=1 refers to the ideal cation-selective membrane. (d) Simulation results of the variation of the maximum Lyapunov exponent ( $\lambda^*$ ) with  $\mathbb{N}_n$ .

chaos. The maximum Lyapunov exponent was calculated from the current dynamics using TISEAN software. <sup>23</sup> The transition from a stable current signal to a chaotic current signal in terms of the maximum Lyapunov exponent is shown in Fig. 2(d). We observed chaos when the pore was ion selective. The transference number is defined as the current due to positive ions compared to the total current. We considered the positive ions since the pore was cation selective. We observed that when  $N_v$  >

0.05294, the majority of the current was contributed by the cation, as the transference number was almost equal to 1 (see Fig. 2(c)), thus indicating that the pore was ideally selective to one type of ion. At this transference number and Nandigana number, we observed the transition from stable current dynamics to unstable or chaotic current dynamics (see Fig. 2(c)).

Using the VN oscillator model, we could track the position of the charged ionic particle. Fig. 3(a) shows the ionic particle near Paper

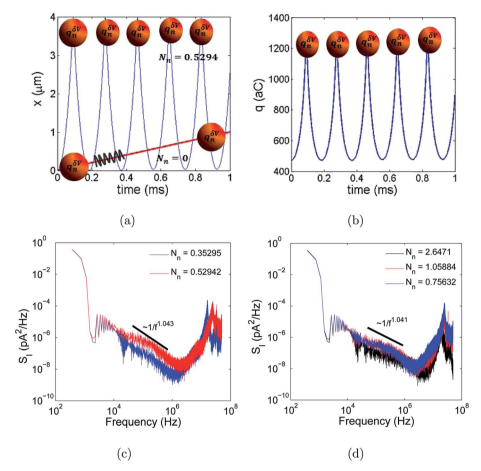


Fig. 3 Characterization of the position of a charged ionic particle. The figure illustrates (a) ionic oscillations due to electrostatic repulsion at  $\mathbb{N}_n =$ 0.5294 and the linear transport of the ion at  $\mathbb{N}_n = 0$ , and (b) the charge of the ionic particle at  $\mathbb{N}_n = 0.5294$ . The charge oscillates as the ion nears the nano membrane. The power spectral density demonstrating pink chaos with 1/f type dynamics for different Nandigana numbers (c and d).

the entrance of the micro-nanopore, i.e., when the particle reached  $x = 4 \mu m$ , the particle was repelled from the cation selective nanopores due to electrostatic interaction of the particle with the charged membrane and applied electric potential. There is electrostatic repulsion resulting in the ionic particle tracing back to its initial position, i.e., the particle moved back to x = 0. The charged particle tracing back to its initial position led to depletion of the ions near the micronanogating membrane junction, i.e., at  $x = 4 \mu m$  as shown in Fig. 3(a). This physics was observed when the Nandigana number = 0.5294. The action was repeated every time the charged particle moved near to the micro-nanogating membrane junction. Hence, the particle oscillated with time giving rise to instability and chaos. To get a better idea of the physics, we looked at the dynamics at a low Nandigana number = 0. At this Nandigana number, the charged particle did not interact with the membrane surface, since at this number the surface charge density of the nanomembrane was zero, allowing both cations and anions to enter into the pore. Thus, the electrostatic repulsion was absent at this Nandigana number, resulting in transport of the charged particle through the pore. We see a linear transport of the charged particle with time revealing that the particle was transported through the

micropore to the nanopore and to the other end of the reservoir. The dynamics are shown in Fig. 3(a). Hence, we could control the stability of the particle motion using our VN oscillator model. Using our VN oscillator model we also showed how the charge acted on the particle. Fig. 3(b) shows that as the particle neared the nanogating membrane, i.e. at  $x = 4 \mu m$ , the particle was repelled from the nanomembrane. Furthermore, the charge of the particle decreased due to electrostatic repulsion. Thus we see a decrease of the charge at  $x = 4 \mu m$  (see Fig. 3(a) and (b) together). When the particle reached the entrance, i.e., at x = 0, the charge increased on the particle due to electrostatic interaction of the particle with the voltage,  $V_0$ . Hence, as the particle oscillated between  $x = 4 \mu m$  and x = 0, the charge decreased and then increased periodically. The results are shown in Fig. 3(b). The charge oscillations resulted in current oscillations, leading to instability and chaos. The dynamics shown in Fig. 3(b) were at  $N_v = 0.5294$ . Thus, these two figures illustrate the depletion of ions and electrostatic repulsion experienced by the charged particle near the nano membrane.

### B. Power spectral density

Finally, we need to resolve the conundrum of whether chaos has color. To answer this query, we probed the current dynamics by **RSC Advances** Paper

reviewing the power spectra of the current dynamics. The temporal power spectral density  $(S_I)$  was analyzed in a frequency range of 0 to  $f_s = 1/\Delta t$  and was obtained using the Welch method.24 The current signal was divided into the longest sections to obtain close to, but not exceeding, 8 segments with 50% overlap. Each section was windowed with a Hamming window. The spectral estimate was evaluated by averaging the modified periodograms for a frequency range  $[0-f_s/2]$ . The power spectral density in Fig. 3(c) and (d) shows a power-law dependence  $(1/f^{\alpha})$  with the frequency. The exponent  $\alpha$  was close to 1 in this frequency range, similar to that of the pink noise phenomenon. However, we rationalized that the dynamics are pink chaos and not pink noise. This is because the pink chaos originates due to instability in the electrostatic potential which is a deterministic phenomenon in contradiction to needing a random memory force which was postulated by earlier theorists and experimentalists<sup>9-12</sup> when observing pink noise. Thus, the dynamics of the current are pink chaos according to our understanding.

### Current-charge phase maps

Similar to the VN oscillator model, the charge oscillator model also depicted interesting physics. The anharmonic distribution in the electrostatic potential leading to multiple potential wells is shown in Fig. 4(a). A section of the potential was sliced along the time plane, and we see the potential-time dynamics in Fig. 4(b). We observed oscillating potential dynamics due to depletion of ions near the micro-nanogating junction. The oscillations changed their sign, indicating that the particle changed its direction as it oscillated in the pore. At  $N_v$ 0.52942, the oscillations were not exactly bi-directional at every time interval, i.e., the oscillations were not periodical but still

we observed the particle change its direction, revealing the bidirectional hopping nature of the ionic particle. The phase map between the charged ionic particle and the potential (see Fig. 4(c)) shows a chaotic distribution of ions in space at a high  $N_{\rm v}$ . The current-charge phase map in Fig. 4(d) shows that the current oscillated between -20 pA to 100 pA when the charge was between 471.5 aC and 472 aC. The oscillations were chaotic and bi-directional due to the depletion of ions near the micronanogating membrane junction. The periodical bi-directional hopping of ions and the electric potential are shown in Fig. 4(e) and (f), respectively. These oscillations were observed at a relatively higher value of  $N_v = 1.058$ . The charge-potential phase map in Fig. 4(g) and the current-charge phase map in Fig. 4(h) show two circles indicating that the potential and current in these phase maps change their signs periodically resulting in bi-directional hopping of ions leading to instability and chaos.

We now think back to the VN oscillator model to derive analytical expressions for the charge transfer. We considered the VN oscillator model without taking into account the ion acceleration. The model reduced down to  $\dot{\lambda} \frac{\partial \vec{x}_n}{\partial t} = F|_{t_n}$ . Solving this equation analytically and substituting the velocity,  $\frac{\partial \vec{x}_n}{\partial t}$ , in the ionic current expression, we obtained the charge on the ionic particle using  $q_n^{\delta V}$  =

 $\frac{-\varDelta_{1}|x_{n}-x_{0}|^{2}|x_{b2}-x_{b1}|}{t(4\pi\epsilon_{0}\epsilon_{r}x_{0}|x_{n}-x_{0}|^{2}+\sigma_{s}\delta S|x_{b2}-x_{b1}|)+(\varDelta_{1}\varDelta_{2}|x_{n}-x_{0}|^{2}|x_{b2}-x_{b1}|)},$ where  $\Delta_1 = (4\pi\epsilon_0\epsilon_r\dot{\lambda}\delta l)$ ,  $\delta l$  varied along the ion particle displacement motion accounting for micropores and nanopores and  $\Delta_2$  was obtained by invoking the initial particle charge condition,  $q_n^{\delta V} = -Z\sigma_n \delta S$  at time t=0 and the position

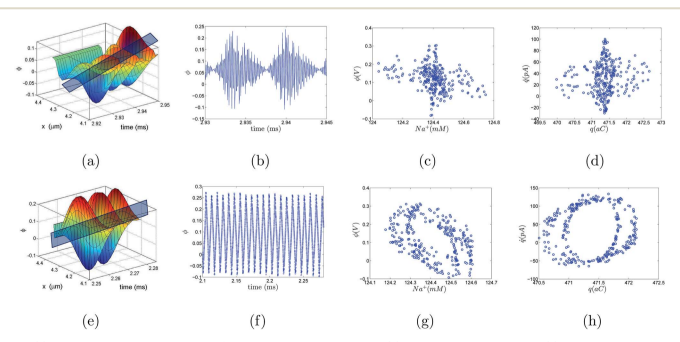


Fig. 4 (a) The space-time potential obtained from the charge oscillator model, (b) the potential-time dynamics, (c) the charge and potential phase map, and (d) the charge–current phase map at  $\mathbb{N}_n=0.5294$ . (e) to (h) The above plots at  $\mathbb{N}_n=1.0588$ 

of the charged particle at the origin,  $x_n = 0$  in the above analytical equation. Z was the valence of the ion transporting through the pore. Note, if the pore is selective for monovalent ions, Z is 1, and similarly for divalent ions, Z = 2. This results in

 $\Delta_2 = \frac{1}{Z\sigma_n dS}$ . This model also resulted in the bi-directional

hopping of charge and current when we assume the ions are displaced in a bi-directional manner. This illustrated that the acceleration of the ionic particle was not necessary to observe current oscillations and chaos. Only the electrostatic interaction of the ions with the membrane surface and the dissipative ionic particle force were necessary to observe the current oscillations.

## V. Conclusion

Paper

In conclusion, we show that for an ideally ion-selective nanopore, current oscillations can also exist due to chaos rather than noise. We observed chaos along with bi-directional hopping of ions due to depletion of ions at the micro-nanopore junction. The depletion of ions resulted in non-equilibrium instability in the electrostatic potential. Furthermore, the temporal power spectra map of the charged ion particle shows that the color of chaos is pink, with 1/f type dynamics similar to the 1/f type pink noise observed by fellow experimentalists. However, we note that pink chaos exists in nanopores and not pink noise as the oscillations observed are deterministic in nature. Thus, we think that, one can not argue that the current oscillations observed in experiments as mere noise, chaos can also be a new way of comprehending the current signals observed in experiments and other numerical simulations. We believe that the observation of bi-directional current hopping and current oscillations can lay pathways for nanopore sensing in areas of DNA sequencing and chemical reaction units along with paving new avenues to use current oscillations for fluidic logic circuits for flow manipulation in biology, combinatorial chemistry and electronic applications.

### Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

This work was supported by the National Science Foundation (NSF) under grants 1264282, 1420882 and 1506619 and the AFOSR under grant FA9550-12-1-0464. The authors gratefully acknowledge the use of the parallel computing resource provided by the University of Illinois. This work was done when V. N. was at the University of Illinois.

### References

Z. Chen, Y. Jiang, D. R. Dunphy, D. P. Adams, C. H. N. Liu,
 N. Zhang, G. Xomeritakis, X. Jin, N. R. Aluru, S. J. Gaik,
 H. W. Hillhouse and C. J. Brinker, DNA translocation through an array of kinked nanopores, *Nat. Mater.*, 2010, 9, 667.

- 2 S. Howorka and Z. Siwy, Nanopore analytics: sensing of single molecules, *Chem. Soc. Rev.*, 2009, **38**, 2360.
- 3 O. A. Saleh and L. L. Sohn, An artificial nanopore for molecular sensing, *Nano Lett.*, 2003, 3, 37.
- 4 J. J. Kasianowicz, E. Brandin, D. Branton and D. W. Deamer, Characterization of individual polynucleotide molecules using a membrane channel, *Proc. Natl. Acad. Sci. U. S. A.*, 1996, **93**, 13770.
- 5 H. Chang, F. Kosari, G. Andreadakis, M. A. Alam, G. Vasmatzis and R. Bashir, DNA-mediated fluctuations in ionic current through silicon oxide nanopore channels, *Nano Lett.*, 2004, 4, 1551.
- 6 J. B. Heng, C. Ho, T. Kim, R. Timp, A. Aksimentiev, Y. V. Grinkova, S. Sligar, K. Schulten and G. Timp, Sizing DNA using a nanometer-diameter pore, *Biophys. J.*, 2004, 87, 2905.
- 7 J. Feng, M. Graf, K. Liu, D. Ovchinnikov, D. Dumcenco, M. Heiranian, V. V. R. Nandigana, N. R. Aluru, A. Kis and A. Radenovic, Single-layer MoS<sub>2</sub> nanopores as nanopower generators, *Nature*, 2016, 536, 197–200.
- 8 E. V. Dydek, B. Zaltzman, I. Rubinstein, D. S. Deng, A. Mani and M. Z. Bazant, Overlimiting current in a microchannel, *Phys. Rev. Lett.*, 2011, **107**, 118301.
- 9 M. R. Powell, I. Vlassiouk, C. Martens and Z. S. Siwy, Nonequilibrium 1/f noise in rectifying nanopores, *Phys. Rev. Lett.*, 2009, **103**, 248104.
- 10 Z. Siwy and A. Fulinski, Origin of 1/f<sup>ect</sup> noise in membrane channel currents, *Phys. Rev. Lett.*, 2002, 89, 158101.
- 11 R. M. M. Smeets, U. F. Keyser, M. Wu, N. H. Dekker and C. Dekker, Nanobubbles in solid-state nanopores, *Phys. Rev. Lett.*, 2006, 97, 088101.
- 12 R. M. M. Smeets, N. H. Dekker and C. Dekker, Low-frequency noise in solid-state nanopores, *Nanotechnology*, 2009, 20, 095501.
- 13 R. M. M. Smeets, U. F. Keyser, N. H. Dekker and C. Dekker, Noise in solid-state nanopores, *Proc. Natl. Acad. Sci. U. S. A.*, 2008, 105, 417.
- 14 H. Nyquist, Thermal agitation of electric charge in conductors, *Phys. Rev.*, 1928, **32**, 110–113.
- 15 http://www.openfoam.com/, 2011.
- 16 V. V. R. Nandigana and N. R. Aluru, Understanding anomalous current-voltage characteristics in microchannel-nanochannel interconnect devices, *J. Colloid Interface Sci.*, 2012, 384, 162.
- 17 V. V. R. Nandigana and N. R. Aluru, Nonlinear electrokinetic transport under combined ac and dc fields in micro/ nanofluidic interface devices, *J. Fluids Eng.*, 2013, 135, 021201.
- 18 V. V. R. Nandigana and N. R. Aluru, Characterization of electrochemical properties of a micro-nanochannel integrated system using computational impedance spectroscopy (cis), *Electrochim. Acta*, 2013b, **105**, 514–523.
- 19 V. V. R. Nandigana and N. R. Aluru, Avalanche effects near nano-junctions, *Phys. Rev. E*, 2016, **94**, 012402.
- 20 L. Verlet, Computer experiments on classical fluids. i. thermodynamical properties of Lennard-Jones molecules, *Phys. Rev.*, 1967, **159**, 98–103.

- 21 B. Hille, *Ion Channels of Excitable Membranes*, Sinauer Associates Inc., Sunderland, MA, 3rd edn, 2001.
- 22 B. M. Venkatesan, D. Estrada, S. Banerjee, X. Jin, V. E. Dorgan, M. Bae, N. R. Aluru, E. Pop and R. Bashir, Stacked graphene- $Al_2O_3$  nanopore sensors for sensitive detection of DNA and DNA-protein complexes, *ACS Nano*, 2012, **6**, 441–450.
- 23 R. Hegger, H. Kantz and T. Schreiber, Practical implementation of nonlinear time series methods: the TISEAN package, *Chaos*, 1999, **9**, 413.
- 24 P. D. Welch, The use of fast Fourier transform for the estimation of power spectra: a method based on time-averaging over short, modified periodograms, *IEEE Trans. Audio Electroacoust.*, 1967, **15**, 70–73.