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Phonon scattering limited performance of monolayer MoS$_2$ and WSe$_2$ n-MOSFET

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In this paper we show the effect of electron-phonon scattering on the performance of monolayer (1L) MoS$_2$ and WSe$_2$ channel based n-MOSFETs. Electronic properties of the channel materials are evaluated using the local density approximation (LDA) in density functional theory (DFT). For phonon dispersion we employ the small displacement / frozen phonon calculations in DFT. Thereafter using the non-equilibrium Green’s function (NEGF) formalism, we study the effect of electron-phonon scattering and the contribution of various phonon modes on the performance of such devices. It is found that the performance of the WSe$_2$ device is less impacted by phonon scattering, showing a ballisticity of 83% for 1L-WSe$_2$ FET for channel length of 10 nm. Though 1L-MoS$_2$ FET of similar dimension shows a lesser ballisticity of 75%. Also in the presence of scattering there exist a 21–36% increase in the intrinsic delay time ($\tau$) and a 10–18% reduction in peak transconductance ($g_m$) for WSe$_2$ and MoS$_2$ devices respectively.

I. INTRODUCTION

Two dimensional (2-D) materials such as the Transition Metal Dichalcogenides (MX$_2$: M=Mo, W; X=S, Se, Te) have emerged as a prospective channel material for the post-Si CMOS technology.$^{1,3–6}$ Among such MX$_2$ materials so far the MoS$_2$ and WSe$_2$ have been successfully realized into MOS devices experimentally.$^{1–3}$ Apart from monolayer flakes of such MX$_2$ materials, few layers thick MX$_2$ MOSFET have also been the focus of various device fabrication studies.$^{7–9}$

The role of electron-phonon scattering in such MX$_2$ layers do have a significant impact on the performance of such 2-D channel MOSFETs.$^{10–12}$ Recently Guo et al. have shown the effect of phonon scattering in monolayer MoS$_2$ MOSFETs.$^{12}$ According to recent simulations tungsten chalcogenide (e.g. WS$_2$) FETs are expected to outperform other MX$_2$ FETs in the ballistic limit.$^6$ However in the more realistic case, the devices often operate at the quasi-ballistic limit. In this regard it could be of interest to comparatively study the MoS$_2$ and WSe$_2$ MOSFET inclusive of the electron-phonon scattering. In this paper we study difference in performance of 1L-MX$_2$ (hereafter by MX$_2$ we refer to MoS$_2$ and WSe$_2$ only) channel 2-D planar n-MOSFET in the quasi-ballistic regime. The materials properties of the monolayer MX$_2$ channel are modeled by ab-initio methods. Thereafter using our in-house simulator based on the non-equilibrium Green’s function (NEGF) formalism, we calculate the performance parameters of the MX$_2$ n-MOSFETs. It is found that performance of WSe$_2$ devices are less impacted by phonon scattering compared to the MoS$_2$ counterparts.

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II. METHODOLOGY

Fig. 1 shows the schematic device structure of the planar 2-D MoS$_2$ FET considered for our studies. We consider a 1L MoS$_2$ or WSe$_2$ as the channel material. The 2-D channel is placed over an SiO$_2$/Si substrate. High-$\kappa$ HfO$_2$ of 2.5nm thickness is chosen as the gate dielectric. We consider highly doped (10$^{20}$ /cm$^3$) n$^{++}$ regions as the source/drain for the n-MOSFET. Such doping concentrations allow for good alignment of the source/drain fermi levels with the conduction band for the monolayer MX$_2$ FETs.$^5,6$

In the first step in our study we evaluate the different properties of the MX$_2$ channel material e. g. bandstructure, electron effective mass, phonon spectra and phonon density of states, by ab-initio simulations. For this purpose we employ density functional theory(DFT) in QuantumWise ATK.$^{13}$ We use a 16 $\times$ 16 $\times$ 1 Monkhorst-Pack k-grid$^{15}$ and employ the Local Density Approximation (LDA) exchange correlation function with the Perdew-Zunger(PZ) basis.$^{16}$ The DFT simulations are performed by relaxing the structures by optimizing the positions by a Broyden-Fletcher-Goldfarb-Shanno (BFGS)$^{17}$ Quasi-Newton optimization method in ATK with maximum force of 0.05 eV/$\text{Å}$. The energy cut-off value in our simulations is set at 75 Hartree.

The phonon dispersion and phonon DOS are calculated in ATK using a supercell based small displacement method also known as the frozen phonon calculations. For phonon calculations in ATK we use a 9 $\times$ 9 $\times$ 1 supercell. 72 small displacements, each of value 0.01 Å is applied to the supercell in the x and the y directions. From this the forces are calculated and subsequently the dynamical matrix of the system is found out, which is used in calculating the phonon properties.

Thereafter, we proceed to solve the Poisson and Schrödinger equations self-consistently for our MX$_2$ FET. The self-consistent solutions are carried out under the Non-Equilibrium Green’s Function (NEGF) formalism$^{12,18}$ with our in-house NEGF simulator.$^{19}$ The electron-phonon interaction in the quasi-ballistic regime is incorporated into our simulator by self-consistent Born approximation method within the framework of NEGF formalism.$^{12,18}$ In our solver, we construct the Green’s function using an effective mass Hamiltonian $H$ and the energy eigenvalue matrix $E$ of the system along with the self-energy matrix of the system $\Sigma$.$^{20,21}$

$$G(E) = [(E + i\varnothing^+)I - H - \Sigma(E)]^{-1}$$

In (1) $I$ is the identity matrix and $\varnothing^+$ is an infinitesimally positive quantity. The self-energy matrix $\Sigma$ consists of $\Sigma_C$ for the source and drain contacts and $\Sigma_{Sc}$ which is the scattering self-energy matrix.

$$\Sigma(E) = \Sigma_C(E) + \Sigma_{Sc}(E)$$

$$\Sigma_C(E) = \Sigma_1(E) + \Sigma_2(E)$$

The spectral density $A(E)$ is expressed as

$$A(E) = i[G(E) - G^\dagger(E)]$$
the Green’s correlation function for the n-type and p-type carriers exclusively can be expressed as $G^n$ and $G^p$, and spectral density can also be expressed in terms of these two as\textsuperscript{18}

$$A(E) = G^n(E) + G^p(E)$$

(5)

The broadening $\Gamma$ is obtained from the self-energy as

$$\Gamma(E) = i[\Sigma(E) - \Sigma^\dagger(E)]$$

(6)

in terms of the in-scattering and out-scattering self-energies of the system it takes the form

$$\Gamma(E) = \Sigma^{in}(E) + \Sigma^{out}(E)$$

(7)

For electron-phonon scattering we consider $G^n$, which is found as\textsuperscript{18}

$$G^n(E) = G(E)(\Sigma_C^{in}(E) + \Sigma_S^{in}(E))G^\dagger(E)$$

(8)

while $G^p$, which is found as

$$G^p(E) = G(E)(\Sigma_C^{out}(E) + \Sigma_S^{out}(E))G^\dagger(E)$$

(9)

In the self-consistent Born approximation approach the in and the out scattering self-energies are evaluated as

$$\Sigma^{in,out} = \Delta^{n,p} G^{n,p}$$

(10)

here $\Delta$ represents the spin relaxation tensor. The number density of the electrons in a 2-D system of cell volume $V_{xy}$, is

$$n(r) = g_s g_v \frac{1}{V_{xy}} \int_{-\infty}^{+\infty} \frac{G^n(r,E)dE}{2\pi}$$

(11)

similarly the number density for holes is given by

$$p(r) = g_s g_v \frac{1}{V_{xy}} \int_{-\infty}^{+\infty} \frac{G^p(r,E)dE}{2\pi}$$

(12)

g_s and g_v being the spin and the valley degeneracies, $r$ being the positional co-ordinate. This carrier (electron) density is evaluated self-consistently within the Poisson-Schrodinger solver.

The Green’s functions $G$, $G^{n,p}$ the in and out scattering self energies $\Sigma^{in,out}$ are all evaluated using a recursive algorithm with the self-consistent Born approximation approach, as described in detail by Nikonov \textit{et al.}\textsuperscript{18} The drain current is evaluated as\textsuperscript{8,19}

$$I_D = \frac{q}{h^2} \sqrt{\frac{m_l k_B T}{2\pi}} \int_{-\infty}^{+\infty} \left[ F_{-1/2}(\eta_1 - E_{k,x}) \right. \left. - F_{-1/2}(\eta_2 - E_{k,x}) \right] \Im(E_{k,x})dE$$

(13)

$m_l$ being the carrier effective mass in the transverse direction, $k_B$ is Boltzmann constant, $T$ is temperature, $E_{k,x}$ the energy of the conducting level, $F_{-1/2}$ is the Fermi integral of order $-1/2$, $\eta_1$ and $\eta_2$ are the chemical potentials of the source and drain respectively. $\Im(E)$ is the transmission matrix given as

$$\Im(E) = \text{trace}[A_1 \Gamma_2] = \text{trace}[A_2 \Gamma_1]$$

(14)

the subscripts 1 and 2 designating the source and the drain contacts. For calculating currents purely ballistic in nature, the scattering self-energy matrix $\Sigma_S$ in (2) is considered zero and the calculations for such a case is described in detail by Sengupta \textit{et al.}\textsuperscript{19} Among the measured parameters the ballisticity is expressed as a ratio between the drain currents including scattering $I_D$ and the purely
FIG. 2. The bandstructure of single layer MoS$_2$ and WSe$_2$.

ballistic drain current $I_{D,Bal}$. The ON current $I_{ON}$ for the devices are calculated at $V_G = V_D = 0.4$ V. The intrinsic delay time $\tau$ is expressed as $\tau = (C_{gg} + C_f) \times V_{DD}/I_{ON}$, where $C_{gg}$ is the device gate capacitance expressed as series combination of geometrical oxide capacitance $C_{ox}$ and the channel quantum capacitance $C_Q$, $C_f$ is the fringing capacitance. $V_{DD}$ is the supply voltage (0.4 V).

III. RESULTS AND DISCUSSIONS

Fig. 2 shows the simulated bandstructure of the 1L MX$_2$ sheets. Our DFT simulations show a direct band gap at the K point of the hexagonal Brillouin zone for both the 1L-MoS$_2$ and 1L-WSe$_2$ sheets with band-gap values of 1.82 and 1.62 eV respectively. Such values are agreeable to ab-initio results in recent reports. The effective masses obtained in our studies (Table I) are also consistent with ab-initio simulation results reported elsewhere.

In Fig. 3, we show the calculated phonon bandstructures for the layered MX$_2$ sheets under consideration. The in-plane acoustic vibration modes of the MX$_2$, namely the longitudinal acoustic (LA) and the transverse acoustic (TA) modes have higher energy compared to the out-of-plane acoustic mode (ZA). These acoustic phonon modes in MX$_2$ are clearly separated from the optical phonon modes. The two lowest branches of the optical modes are non-polar (NP) in nature, followed by the transverse optical (TO) and the longitudinal optical (LO) modes. The mode(s) with very low dispersion at energies 50 meV in MoS$_2$ and 30 meV in WSe$_2$ are known as the homopolar (HP) modes. Fig. 4 shows the calculated phonon density of states of the monolayer MoS$_2$ and WSe$_2$.

Among the different modes we can consider the LA, TA and ZA modes to contribute to acoustic scattering, non-polar optical and TO and HP modes to contribute to the optical phonon

| material   | $m_l/m_0$ | $m_t/m_0$ |
|------------|-----------|-----------|
| 1L-MoS$_2$ | 0.4742    | 0.4738    |
| 1L-WSe$_2$ | 0.3376    | 0.3372    |
scattering, while the polar LO modes are responsible for the Fröhlich interaction mechanism of electron-phonon scattering. The phonon energies of the different modes at the CB minima of the electronic bandstructure are considered in order to evaluate the electron-phonon interaction matrix and subsequently the scattering self-energy matrix $\Sigma$ in the channel.\(^{18}\)

In Fig. 5, we show the effect of the electron-phonon scattering on the output characteristics of the device. A gate voltage $V_G = 0.4$ V is considered for these simulations. A channel length $L_{Ch} = 25$ nm is taken so as to show the phonon scattering contributions more clearly in the plot. We see for purely ballistic currents (shown in black) the drain current $I_D$ for 1L-WSe\(_2\) is higher than that for the 1L-MoS\(_2\) channel. The optical phonon scattering (comprising of non-polar, TO and HP modes) contributes the highest in the degradation of drain current. This is followed by that due to the Fröhlich interaction and the acoustic phonon contributions (due to LA, TA and ZA modes).

The phonon scattering seems to degrade carrier transport in MoS\(_2\) more than that in WSe\(_2\) devices. The reason behind this is the stronger electron–phonon coupling in MoS\(_2\), than the WSe\(_2\) monolayer. This is quantitatively represented by the higher values of the deformation potentials (DP) for the various phonon modes for MoS\(_2\) compared to WSe\(_2\) as obtained by Kaasbjerg et al.\(^{10}\) and Jin et al.\(^{22}\) For the $L_{Ch} = 25$ nm device, the net reduction in $I_D$ due to electron-phonon scattering effects are 18.6% for WSe\(_2\), and 26.8% for 1L-MoS\(_2\) devices. The variation of purely ballistic and the quasi-ballistic (inclusive of phonon scattering) ON currents with the device channel length is shown in Fig. 6. The ON currents are evaluated at $V_G = V_D = 0.4$ V. It is observed that the performance of the WSe\(_2\) device is less impacted by phonon scattering, showing a ballisticity of
FIG. 5. The output characteristics of MoS$_2$ and WSe$_2$ n-MOSFET with $L_{ch} = 25$ nm showing the contribution of different phonon modes to the scattering. Gate voltage $V_G = 0.4$ V.

83% for 1L-WSe$_2$ FET for channel length of 10 nm. Though 1L-MoS$_2$ FET of similar dimension shows a ballisticity of 75%. As the channel length increases, the electrons tend to encounter more number of scattering events and therefore the difference between the ballistic and quasi-ballistic currents tend to increase with the channel length. Here it must be mentioned that in our work we have included only the additional effect of electron-phonon scattering to the ballistic transport in MX$_2$ FET. Also we looked to focus our study on mostly short channel lengths as MX$_2$ as a channel material is essentially a candidate for post-Silicon sub-decananometer CMOS technology. In experimentally fabricated devices the channel dimensions vary from several hundred nanometer to even

FIG. 6. The ballistic current and the current including scattering effects for the MoS$_2$ and WSe$_2$ n-MOSFET of varying channel lengths.
a few microns with gate dielectric thickness of the order of few tens of nanometers. In case of such fabricated devices, a vast number of non-idealities like surface roughness, defects and impurity scattering, intrinsic ripples in 2-D membranes, interface strain, processing related defects etc. apart from electron-phonon scattering, come into play. Owing to these effects there is a large difference between experimental results and simulation studies that are being conducted by different groups. For theoretical studies conducted for nanoscale FET of MX2 material, the ON currents are in the range of 500 µA/µm for HfO2 top gated Schottky barrier MoS2 SBFET of LCh = 15 nm. For HfO2 top-gated MOSFET in the scaling limit of LCh = 5 nm the simulated ON current is reported as 238 µA/µm by Alam et al. In this context our simulated results for LCh = 10 nm, top gated HfO2 dielectric doped contact MoS2 MOFET shows ON current of 666 µA/µm which is consistent with other theoretical results based on similar structures and device geometry and gating.

The calculated values for the geometrical oxide capacitance $C_{ox}$ is 0.22 fF/µm, the channel quantum capacitance $C_Q$ at $V_G = 0.4V$ is 0.16 fF/µm and the fringing capacitance $C_f$ is 0.025 fF/µm. The capacitances are calculated by the methods described by Alam et al. For the 25 nm device, $\tau$ in the ballistic limit was calculated to be 0.67 ps for MoS2 and 0.53 ps for WSe2 FET. In case of MoS2 FETs scattering seems to affect delay time much more severely than the WSe2 FETs. As electron-phonon scattering is included the delay time increases significantly and the values increase to 0.91 ps and 0.64 ps for the MoS2 and the WSe2 FETs respectively.
The transconductance \((g_m)\) vs \(V_G\) characteristics in Fig. 7 of a 25nm device show the ballistic values of \(g_m\) for the MX\(_2\) are rather closely grouped together, while in the quasi-ballistic case the \(g_m-V_G\) curve are more spread out. This trend was earlier observed in the ON currents for ballistic and quasi-ballistic cases as well. The reason behind being the disparity in the amount of electron-phonon scattering among the two different MX\(_2\) materials. The values of peak \(g_m\) for the MX\(_2\) n-MOSFETs vary between 1226–1359\(\mu\)S/\(\mu\)m for the ballistic cases.

The energy and position resolved current spectra is shown in Fig. 8. It is clear that the WSe\(_2\) MOSFETs show a higher current density compared to the MoS\(_2\) ones. Also the extension of the region of highest current density within the devices show that, for the 1L-WSe\(_2\) this region (marked red in the colorbar) extends to a significant portion of the channel length. For 1L-MoS\(_2\) the length of this region diminishes gradually due to higher degree of electron phonon scattering in the channel for those structures. In all the cases there is no evidence of direct tunnelling between the source and drain, as is expected for channel lengths of \(L_{ch} = 25\) nm.

### IV. CONCLUSION

In this paper we study the effect of phonon scattering on the performance of monolayer (1L) MoS\(_2\) and WSe\(_2\) n-MOSFETs. Material properties of the channel are evaluated using the local density approximation (LDA) in density functional theory (DFT) using the Perdew–Zunger (PZ) exchange correlation. Thereafter using our in-house non-equilibrium Green’s function (NEGF) simulator, we study the effect of phonon scattering on the performance of such devices. Individual contributions of the different phonon modes to the scattering and the effect of varying channel length of the device on performance parameters such as ON current, ballisticty, intrinsic delay time and transconductance are studied in detail. It is found that performance of WSe\(_2\) devices are less impacted by phonon scattering compared to the MoS\(_2\) counterparts.

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