## Direct Determination of Energy Level Alignment and Charge Transport at Metal-Alq<sub>3</sub> Interfaces via Ballistic-Electron-Emission Spectroscopy

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Using ballistic-electron-emission spectroscopy (BEES), we directly determined the energy barrier for electron injection at clean interfaces of Alq<sub>3</sub> with Al and Fe to be 2.1 and 2.2 eV, respectively. We quantitatively modeled the sub-barrier BEES spectra with an accumulated space charge layer, and found that the transport of nonballistic electrons is consistent with random hopping over the injection barrier.

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Dramatic progress has been made in organic electronics, where organic-based devices are rivaling those based on conventional inorganic semiconductors [1]. Presently, efforts are underway to apply spintronic principles [2], which originated from inorganic materials, to carbon nanotubes, molecular monolayers, and bulk organic semiconductors for a new class of organic electronics that exploit the spin of charge carriers [3]. There is a controversy over whether the reported magnetoresistance in organic spin-valve devices indeed results from injection and coherent transport of spins on molecular levels [4]. These devices [5] consist of a nominally thick organic semiconductor layer (e.g., Alq<sub>3</sub>) sandwiched between two ferromagnetic metal (FM) electrodes; but their qualification as spin valves has been challenged, because similar devices indicate direct tunneling through structural nonuniformities rather than molecular transport, and because low hole mobility in truly nontunneling devices prevents efficient spin injection in the first place [6-8].

More importantly, the state of confusion over spin transport in organic media underscores a larger issue: gaps in understanding the properties of metal-organicsemiconductor interfaces have become impediments to properly describing the basic physics of charge transport in organic-based devices, and to employing organics as active materials in new frontiers of science. Despite Alq<sub>3</sub> being an archetypal organic electroluminescent material, interpretations of the charge transport characteristics in Alq<sub>3</sub> devices vary dramatically in the literature [9]. Differences in material and preparation conditions and interface characteristics can contribute to the discrepancies [9-11]. Wolf et al. [12] pointed out that the injection barrier height ( $\Delta$ ), i.e., the difference between the electrode Fermi level ( $\epsilon_F$ ) and the organic semiconductor's lowest unoccupied molecular orbital (LUMO) or highest occupied molecular orbital (HOMO), is a crucial parameter in determining the nature of charge transport in organic electronic devices. However, while the HOMO level and vacuum level offset can be reliably determined from ultraviolet photoemission spectroscopy (UPS) [13], it is generally understood that the conventional approach of estimating LUMO by adding the optical gap to the HOMO level neglects excitonic effects [11,14,15]. Unfortunately, different experimental techniques to directly measure LUMO have their own caveats and yield conflicting results. The Alq<sub>3</sub> HOMO-LUMO gap was determined to be 4.6 eV by inverse photoelectron spectroscopy (IPES) [14], but was given as 2.96 eV from scanning tunneling spectroscopy (STS) [15]. Zhan *et al.* [16] constructed an energy-level diagram that favored electron injection from Co and LSMO into Alq<sub>3</sub> by choosing the Alq<sub>3</sub> HOMO-LUMO gap determined from STS over that from IPES. They argued that IPES could have caused sample modifications with strong electron fluxes, although reports [17,18] exist that a scanning tunneling microscope (STM) tip might do just the same.

Ballistic-electron-emission spectroscopy (BEES) is an established technique for characterizing the band structure at metal-inorganic-semiconductor interfaces.[19] It was recently extended to measure  $\Delta$  at metal–organic semiconductor interfaces [17,18]. A schematic diagram of the BEES technique is shown in Fig. 1(a). Ballistic electrons tunnel from an emitter into a thin metal base in contact with an organic semiconductor; they can enter the LUMO if their energy exceeds  $\Delta$ . Since the energy of the electrons is the potential difference between the base and emitter  $(V_{\rm BE})$ ,  $\Delta$  is simply the threshold  $V_{\rm BE}$  at which the collector current  $(I_C)$  rises sharply. The emitter can be the tip of an STM, or an all-solid-state tunnel junction. The STM implementation offers spatial resolution, but there are concerns about the stability of the spectra and measurementinduced sample modifications [17,18]. STM-based BEES is also limited to interfaces where the base is on top of the organic semiconductor. Since metal deposition onto organic semiconductors tends to create interfacial gap states [11], care must be taken when associating the injection threshold in STM-based BEES with true molecular levels.

In the present work, we applied BEES to directly determine the electron-injection barrier at metal-Alq<sub>3</sub> interfaces. We used large-area (compared to a STM-tip) Al<sub>2</sub>O<sub>3</sub> tunnel junctions for injecting ballistic electrons, and placed the emitter under the base so that Alq<sub>3</sub> was deposited on top of the metal in order to achieve clean and stable metal-Alq<sub>3</sub> interfaces. Whereas in conventional BEES the sub-barrier  $I_C$  is considered leakage and is ignored, we quantitatively modeled the sub-barrier BEES spectra with an accumulated space charge layer from ballistic injection. The presence of the space charge allowed us to distinguish between the usual organic charge injection mechanisms. We show that nonballistic charge injection at clean metal-Alq<sub>3</sub> interfaces is limited by random hopping of carriers from  $\epsilon_F$  over the injection barrier, and that there also exist uniformly-distributed gap states on which charge carriers can be transported.

We fabricated the Alq<sub>3</sub> BEES devices in a high-vacuum cluster deposition system [7] via thermal evaporation and shadow masking. The vacuum pressure during evaporations was  $< 1 \times 10^{-8}$  Torr; sample transfers between deposition chambers and mask changes were performed without breaking the vacuum. The emitter structure always consisted of an Al electrode and an Al<sub>2</sub>O<sub>3</sub> tunnel barrier. We used Al or Fe for the base, and Al or Au for the collector. In the following, we will denote a device by the materials in its base-collector structure, e.g.,



FIG. 1 (color). (Color figure) (a) Schematic energy-level diagram of BEES. (b) A photograph of an Al-Alq<sub>3</sub>-Al device. (c)  $I_E$ and (d)  $\Delta I_C$  at several values of  $V_{CB}$  are plotted against  $V_{BE}$ . The solid curves are fits described in the text. (e)  $I_C(0)$  vs  $V_{CB}$ . The solid curve is a fit to the Arkhipov model [22].

Fe-Alq<sub>3</sub>-Au for a device with an Fe base, an Alq<sub>3</sub> spacer and a Au collector. Shown in Fig. 1(b) is an optical micrograph of an Al-Alq<sub>3</sub>-Al device. The 12-nm thick Al emitter was first evaporated through a 20- $\mu$ m wide slit in a mask on the SiN substrate. It was oxidized by *in situ* plasmaassisted oxidation to form the Al<sub>2</sub>O<sub>3</sub> tunnel barrier. A 4-nm thick, 200- $\mu$ m wide base electrode was subsequently deposited at 45° with the emitter to complete the emitter-base tunnel junction. The 100-nm Alq<sub>3</sub> layer covered the entire sample. Finally, the 10-nm thick collector electrode was deposited through a 100- $\mu$ m wide mask, perpendicular to the base electrode. The device thus had an emission area ( $A_e$ ) of  $1.4 \times 10^3 \ \mu$ m<sup>2</sup>, and a collection area of  $2.0 \times 10^4 \ \mu$ m<sup>2</sup>.

The room-temperature BEES measurements were carried out with the devices sealed in darkness. Because of the large thickness and the relatively poor mobility of carriers in Alq<sub>3</sub>, it was necessary to apply a base-collector bias  $(V_{CB})$  to attain a measurable  $I_C$ . At each  $V_{CB}$  value,  $V_{BE}$ was ramped stepwise, and  $I_C$  was measured in the steady state. The rms noise level of our setup is ~100 fA. The devices were stable over repeated cycling of  $V_{CB}$  and  $V_{BE}$ ; they failed only when we unknowingly ramped  $V_{BE}$  past dielectric breakdown of the tunnel junctions. Since we limited the measurement polarity to electron-injection only, there was no issue of Alq<sub>3</sub> degradation from unbalanced hole injection [7,20].

Shown in Fig. 1(c) is the emitter current  $(I_E)$  as a function of  $V_{\rm BE}$  for the emitter tunnel junction in an Al-Alq<sub>3</sub>-Al device.  $I_E$  rises monotonically with  $V_{BE}$ . The collector current due to ballistic injection,  $\Delta I_C =$  $I_C(V_{BE}) - I_C(0)$ , is plotted in Fig. 1(d) for several values of  $V_{\rm CB}$ . At small  $V_{\rm CB}$  (< 4 V),  $\Delta I_C$  initially increases slowly with  $V_{\rm BE}$ , but rises much faster at higher  $V_{\rm BE}$ , as is expected when ballistic electrons have sufficient energy to overcome the injection barrier and enter the Alq<sub>3</sub> LUMO. However, when  $V_{CB} \ge 4$  V, a striking feature is seen:  $\Delta I_C$  actually decreases initially, before rising sharply at higher  $V_{\rm BE}$ ; it appears that the injection of ballistic electrons hampers the charge transport in the Alq<sub>3</sub> layer. To our knowledge, this effect has not been reported in any BEES study. From the position of the  $\Delta I_C$  minima, which remains constant for all values of  $V_{\rm CB}$ , we determine  $\Delta$  to be  $2.10 \pm 0.05$  V for electron injection at the Al-Alq<sub>3</sub> interface. A constant injection barrier height indicates that the image charge effect is absent. This could be because the relaxation time for high-energy (> 2 eV) ballistic electrons ( $\sim 20$  fs [21]) is far shorter than the minimum polaronic hopping time ( $\sim 100$  fs [10]); the image hole left behind by a ballistic electron is already filled before the electron hops onto the next molecular site.

Figure 1(e) shows the zero-emission collector current,  $I_C(0)$ , as a function of  $V_{CB}$ . This nonballistic electron current injected from the Al base via the application of  $V_{CB}$  is precisely the usual charge current in a Al-Alq<sub>3</sub>-Al

device. Comparing our results with those in the literature, we find that the current density in our device is about 2 orders of magnitude smaller than that in a similar device in Ref. [10], but is very similar to that in the Alq(2) device of Ref. [9] under reverse bias. The device in Ref. [10] had Al grown on Alq<sub>3</sub>, which likely had chemical reactions at the interface, making it appropriate to describe injection as a process of charge hopping out of the reacted interfacial sites. On the other hand, both ours and the device of Ref. [9] had Alq<sub>3</sub> grown on Al, which led to cleaner and more ideal interfaces, and thus a different charge injection mechanism.

Arkhipov *et al.* [22] modeled charge injection from metals into organic semiconductors as thermally assisted hopping of carriers from the metal Fermi level onto a Gaussian distribution of molecular levels, followed by either recombination or diffusive escape. The hopping injected current ( $I_{hop}$ ) is a function of  $\Delta$  and the applied electric field (*F*). Taking  $\Delta = 2.1$  eV, and the accepted literature values [10] for the bulk distribution width ( $\sigma = 0.13 \text{ eV}$ ) and intermolecular distance (a = 1 nm), we fitted the data points in Fig. 1(e) to the Arkhipov model and obtained  $\gamma = 1.3 \text{ nm}^{-1}$  for the inverse localization radius. Since the injected charge binds with the Alq<sub>3</sub> molecule to form a polaron, it is reasonable that  $\gamma$  is close to 1/a.

We model the sub-barrier BEES spectra by considering what happens when ballistic electrons with energy less than  $\Delta$  are emitted into the base. Because of the energetic disorder in Alq<sub>3</sub>, some of these ballistic electrons can enter the molecular sites and accumulate near the base-Alq<sub>3</sub> interface. For a layer of space charge located at x = awith a density of *n*, along with the induced quasistatic image charge in the base and collector electrodes, the additional electrostatic potential created in the Alq<sub>3</sub> layer is

$$V_{\rm SC} = \begin{cases} \frac{ena}{\varepsilon\varepsilon_0} \left(1 - \frac{a}{L}\right) x \equiv F_1 x & x \le a, \\ \frac{ena}{\varepsilon\varepsilon_0} \left(\frac{a}{L}\right) (L - x) \equiv V_0 - F_2 x & x > a, \end{cases}$$
(1)

where x is the distance from the base electrode,  $\varepsilon$  is the dielectric constant,  $\varepsilon_0$  is the vacuum permitivity, and L is the Alq<sub>3</sub> thickness.

At  $x \le a$ , the space charge layer creates a retarding field  $F_1$  towards base electrode. In the steady state, due to charge conservation, the drifting of the accumulated electrons back to the base electrode under  $F_1$  mostly cancel the ballistic current impinging upon the base-Alq<sub>3</sub> interface. Therefore,

$$A_e n e \mu(F_1) F_1 = I_E \exp(-t/\lambda), \qquad (2)$$

where *t* is the base electrode thickness,  $\mu$  is the Poole-Frenkel field-dependent mobility as determined in Ref. [9], and the hot-electron attenuation length  $\lambda$  is ~10 nm [23] for Al. Solving Eqs. (1) and (2), we obtain *n* and plot it as a

function of  $V_{\text{BE}}$  in Fig. 2(a). Assuming single occupancy, 0.45% of the Alq<sub>3</sub> molecules in the first layer is charged at  $V_{\text{BE}} = 2$  V.

At x > a, the space charge layer raises the injection barrier by  $V_0$ , and the electric field in Alq<sub>3</sub> by  $F_2$ . The increased injection barrier impedes the hopping injection and reduces  $I_C$ . Since  $V_0$  is much smaller than  $\sigma$  ( $V_0 = 23$  mV at  $V_{BE} = 2$  V), variations in  $V_0$  due to disorder in the first Alq<sub>3</sub> layer do not affect  $I_C$  significantly.

On the other hand, some ballistically injected electrons also contribute to  $I_C$ . At the tail of the Gaussian LUMO distribution, the transport levels can be considered uniform in density. The ballistically injected electrons likely travel on all levels whose energies are less than  $V_{\rm BE}$ , the contribution to  $I_C$  being proportional to  $V_{\rm BE}$ . Consequently,  $\Delta I_C$ can be written as

$$\Delta I_C = [I_{\text{hop}}(\Delta + V_0, F + F_2) - I_{\text{hop}}(\Delta, F)] + CV_{\text{BE}}.$$
 (3)

The  $\Delta I_C$  curves in Fig. 1(d) are fitted to Eq. (3), with C as the sole fitting coefficient. In Fig. 2(b), we plot C as a function of  $V_{CB}$ . At low  $V_{CB}$ , C is nearly constant, suggesting that the transport is driven by diffusion, rather than by drift. At  $V_{CB} \ge 5$  V, C increases sharply. The increased transport of ballistically injected electrons overcomes the space charge-induced decrease in  $I_{hop}$  such that at  $V_{\rm CB} = 6$  V,  $\Delta I_C$  always increases with  $V_{\rm BE}$ . Comparing C at high  $V_{\rm CB}$  with the expression for charge current under a uniform distribution of traps  $J \propto$  $V_{\rm CB} \exp(2\varepsilon\varepsilon_0 V_{\rm CB}/N_n kTeL^2)$  [24], we estimated a trap density  $N_n$  of  $\sim 5 \times 10^{23}$  (m<sup>3</sup> eV)<sup>-1</sup>. It has been suggested that electrons in Alq<sub>3</sub> are self-trapping [25], i.e., the total trap density equals the molecular density, the estimated  $N_n$ value represents the trap density at  $\sim 4\sigma$  away from the center of the LUMO distribution.

Shown in Fig. 3(a) are  $I_C$  and  $dI_C/dV_{\rm BE}$  plotted against  $V_{\rm BE}$  for an Fe-Alq<sub>3</sub>-Au BEES device. From these curves, we obtain  $\Delta = 2.2 \pm 0.1$  eV for electron injection from Fe into Alq<sub>3</sub>. In Fig. 3(b), we show the schematic energy-level diagrams for Al-Alq<sub>3</sub> and Fe-Alq<sub>3</sub> interfaces. The difference between the Al  $\epsilon_F$  and Alq<sub>3</sub> HOMO level



FIG. 2. (a) Calculated *n* as a function of  $V_{\rm BE}$  for the Al-Alq<sub>3</sub>/Al device in Fig. 1. Inset: The position dependence of  $V_{\rm SC}$ . (b) The fitting coefficient *C* as a function of  $V_{\rm CB}$ .

was determined to be 2.7 eV via UPS [13]. Combining it with our BEES-determined  $\Delta = 2.1$  eV, we arrive at a value of 4.8 eV for the Alq<sub>3</sub> HOMO-LUMO gap, which is in close agreement with the peak-to-peak gap [14] obtained via IPES. Given that the HOMO- $\epsilon_F$  differences for Co-Alq<sub>3</sub> and LSMO-Alq<sub>3</sub> interfaces were measured via UPS [16] as 2.1 and 1.7 eV,, respectively, the electroninjection barriers are 2.7 and 3.1 eV at these interfaces. Such large barriers would make it highly unlikely that Co-Alq<sub>3</sub>-LSMO spin-valve devices can have electron transport. We note, however, that the electron-injection barriers determined via BEES are significantly higher than those defined from the bottom edges of the LUMO features in IPES [26]. In BEES, the upturn in  $I_C$  signifies ballistic electrons having sufficient energy to reach the collector electrode via the transport level; the lower-energy electrons, which enter the bottom edge of the LUMO distribution, are mostly reflected or become trapped, and are thus not transported through the thickness of the organic layer. Therefore, the different results from BEES and IPES highlight the need to properly define the injection barrier when examining charge transport in organic electronic devices.

The BEES technique has not been commonly employed to characterize molecular materials. By quantitatively modeling the sub-barrier BEES spectra, we have shown that it is a reliable method for directly determining the LUMO level. STM-based BEES has issues with spectra and sample stability; those concerns can however be alleviated in large-area tunnel junction-based BEES devices. More importantly, the basic scheme of BEES has been used to inject spin-polarized hot-electrons from tunnel junction



FIG. 3. (a)  $I_C$  and  $dI_C/dV_{\rm BE}$  vs  $V_{\rm BE}$  for an Fe-Alq<sub>3</sub>-Au device. (b) Schematic diagrams of energy-level alignment at Al-Alq<sub>3</sub> and Fe-Alq<sub>3</sub> interfaces.

emitters into inorganic semiconductors, exploiting the spin-filtering effect of the FM base [27]. A BEES device with FM base and collector electrodes would be an ideal structure for directly examining spin coherence in molecular materials.

In summary, we have determined the electron-injection barriers at clean Fe-Alq<sub>3</sub> and Al-Alq<sub>3</sub> interfaces using ballistic-electron-emission spectroscopy. By exploiting the interaction between the ballistic and nonballistic carriers, we were able to distinguish between charge injection mechanisms at the metal-Alq<sub>3</sub> interface. The transport of nonballistic electrons is consistent with random hopping over the interfacial energy barrier.

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