Interfacial charge transfer in nanoscale polymer transistors

Jeffrey H. Worne\textsuperscript{1}, Rajiv Giridharagopal\textsuperscript{1}, Kevin F. Kelly\textsuperscript{1} and Douglas Natelson\textsuperscript{1,2}

\textsuperscript{1}Department of Electrical and Computer Engineering, Rice University, 6100 Main St., Houston, TX 77005 and
\textsuperscript{2}Department of Physics and Astronomy, Rice University

Abstract

Interfacial charge transfer plays an essential role in establishing the relative alignment of the metal Fermi level and the energy bands of organic semiconductors. While the details remain elusive in many systems, this charge transfer has been inferred in a number of photoemission experiments. We present electronic transport measurements in very short channel ($L < 100$ nm) transistors made from poly(3-hexylthiophene) (P3HT). As channel length is reduced, the evolution of the contact resistance and the zero-gate-voltage conductance are consistent with such charge transfer. Short channel conduction in devices with Pt contacts is greatly enhanced compared to analogous devices with Au contacts, consistent with charge transfer expectations. Alternating current scanning tunneling microscopy (ACSTM) provides further evidence that holes are transferred from Pt into P3HT, while much less charge transfer takes place at the Au/P3HT interface.

Keywords: organic semiconductors; band alignment; charge transfer; organic field-effect transistor; scanning tunneling microscopy
Understanding the band alignment between organic semiconductors (OSCs) and metal electrodes is of basic physical interest as well as significant technological importance\[1\]. Such energetic considerations are crucial for optimizing charge injection in organic light emitting diodes (OLEDs) and organic field-effect transistors (OFETs). Similarly, the relative band alignment at such interfaces also affects the open circuit photovoltage achievable in organic photovoltaic applications. Because interfacial dipoles can be used to modulate the effective work function of a metal surface, self-assembled monolayers (SAMs) of polar molecules have been used in both OLEDs and OFETs to engineer charge injection\[2, 3, 4\].

Conceptually the issue is straightforward, though very complicated in detail. In equilibrium the chemical potential throughout a metal/organic heterointerface must be constant. This condition is achieved through a combination of interfacial dipole formation, charge transfer, and the self-consistent solution of the electrostatics to equilibrate carrier drift and diffusion. Dipole formation and charge transfer lead to deviations from the Schottky-Mott limit, so that simple alignment of vacuum levels does not give an accurate picture of the true heterojunction energetics\[5\].

The relative alignment of levels is most readily measured via photoemission experiments. Recent investigations using electroluminescent polymers\[6\] as well as small molecule OSCs and conducting polymers\[7\] have shown Fermi level pinning at the metal-organic interface. For the case of holes, when the Fermi level of the metal is relatively far above the highest occupied molecular orbital (HOMO) of the molecules (the center of the disorder-broadened valence band, in the case of polymers), the system is in the Schottky-Mott limit. When the Fermi level approaches the HOMO, strong deviations from the Schottky-Mott limit are reported\[6\]. There are several suggested mechanisms for these deviations, including changes in the interfacial dipole (via chemical reactions or the “pillow effect”\[8\]), shifting of frontier orbitals and formation of induced states\[9\], and charge transfer involving polaron formation\[6, 10\]. Recent photoemission experiments examining metal/polymer interfaces show evidence of charge transfer into valence band tail states and accompanying band bending\[11\].

Other data consistent with charge transfer and band bending at similar interfaces has been seen in transport measurements\[4, 16\] of contact resistance in P3HT-based OFETs. In devices with high electrode work functions, hole injection remains Ohmic down to very low carrier densities, a natural result if interfacial charge transfer effectively dopes the contact interface. Similarly current-voltage characteristics in devices with (lower work function) electrode metals giving non-Ohmic injection are consistent with a nanoscale region (between 10 nm and 100 nm in extent)
of reduced mobility at the metal/organic interface\textsuperscript{12, 13, 14, 15}. This may be due to interfacial structural disorder in the organic semiconductor, but it would also be compatible with the formation of an effective depletion region near the contacts due to band bending. Conduction in P3HT takes place via hopping through a density of localized states in the valence band tail. A local change in the chemical potential farther into the band tail would simultaneously deplete holes and reduce the spatial density of available hopping sites, reducing the mobility as well.

In this paper we present additional evidence that significant charge transfer takes place at the Pt/P3HT interface, leading to a population of mobile holes within a nanoscale distance of the interface. Measurements of Pt-based OFET device resistance as a function of channel length, $L$, show a decrease in resistance as $L$ is reduced below 100 nm. In the same devices this corresponds to nearly a factor of 10 increase in zero-gate-bias (ZGB) current over the same $L$ range. In contrast, identically prepared short channel Au-based devices show an increase in resistance, as well as ZGB currents orders of magnitude lower than in Pt devices. Devices with Au surfaces functionalized with SAMs known to increase the electrode effective work function show trends intermediate between the Au and Pt cases. Further measurements using alternating current scanning tunneling microscopy (ACSTM) show clear evidence for excess holes in thin P3HT films on Pt, in marked contrast to ACSTM measurements on identically prepared P3HT films on Au.

Devices are fabricated on $n^+$ silicon with 200 nm of thermally grown oxide as the gate dielectric. Electrodes are patterned using electron beam lithography, e-beam evaporation, and liftoff processing. Using two-step lithography, various channel lengths, $L$, are fabricated, from $\sim$50 nm to 5 $\mu$m, with widths, $w$ of 50 $\mu$m for short channel device studies. In addition, for comparison a structure of interdigitated electrodes with $L = 1 - 50$ $\mu$m and $w = 200$ $\mu$m is fabricated on the same chip in order to allow standard transmission line measurements of P3HT field effect mobility. Electrode films are 15 nm of either gold or platinum with a 1 nm titanium adhesion layer. Following lithography and liftoff, the devices are cleaned under ultraviolet exposure, 1 minute of oxygen plasma, and a 1 minute soak in piranha etch (3:1 $\text{H}_2\text{SO}_4$:$\text{H}_2\text{O}_2$ (30%)). For functionalized Au electrode devices, the piranha etch step is followed by self-assembly of fluorinated oligo(phenylene ethynylene) (F-OPE)\textsuperscript{4} by immersion for 24 h in a solution of F-OPE at 0.25 mg/mL concentration in 1:1 ethanol:chloroform under nitrogen gas, with standard thioacetate deprotection chemistry\textsuperscript{17}.

After the SiO\textsubscript{2} dielectric surface is treated with octadecyltrichlorosilane (OTS) (10 $\mu$L OTS in 8 mL hexadecane, assembled in the dark at room temperature for $\sim$12 hours), P3HT (0.1%
by weight in chloroform) is then spin-coated to a thickness of 5 nm (measured via AFM) over the entire substrate. Contact to the substrate is made, and the Si becomes the gate of the device. Figure 1b illustrates a typical short channel device. Devices are transferred to a variable temperature vacuum probe station immediately upon completion of spin coating. Devices remain in vacuum ($\sim 10^{-6}$ mTorr) for at least 1 hour before any measurements are performed. Individual OFETs are isolated from one another by using a probe tip to scratch away the P3HT film in a rectangle around each device to prevent stray current paths.

Using the interdigitated electrodes of fixed $w$ but varying $L$, device characteristics ($I_D$ vs. $V_{SD}$ at fixed $V_G$, where $I_D$ is drain current and $V_{SD}$ is source-drain voltage) are measured via a semiconductor parameter analyzer. In the shallow channel regime ($|V_{SD}| << |V_G|$) we find these characteristics to be linear, consistent with Ohmic injection. As in our previous work[4, 18] we use the transmission line approach to characterize the contact resistances and field-effect mobilities as a function of gate voltage, $V_G$. In the shallow channel regime at a particular $V_G$, the measured source-drain resistance $R = (\partial I_D/\partial V_{SD})^{-1}$ is plotted as a function of $L$ and found to be linear for long ($L > 1 \mu$m) devices. Channel resistance, $R_C$, is determined by the slope of this line and the effective contact resistance, $R_S$ is determined by extrapolating this line to zero channel length, as shown in Fig. 1a. This linearity observed in longer devices indicates that the overall film uniformity is good, even for the relatively thin P3HT layers used here.

This extrapolation to infer $R_S$ assumes that the device properties are uniform down to arbitrarily short channel lengths. In practice this may not be true, for a number of reasons. For example, a change in polymer morphology near the metal contacts would lead to deviations from linearity in $R(L)$ at short distances as contact regions interact with each other[19]. Similarly, there is indirect evidence[12, 13, 14] that nanoscale regions of poor effective mobility can exist near contacts due to band bending and depletion effects. Note that the characteristic size of these regions is constrained experimentally[14] to less than $\sim 100$ nm, since larger regions would be detectable within the resolution of scanning potentiometry experiments. If two such depletion regions were to intersect for sufficiently short channel devices, one would expect the measured $R(L)$ to have an up-turn as $L \rightarrow 0$, as shown by the red line in Fig. 1a.

Figure 2 shows these $R$ vs. $L$ plots for three sets of Au, F-OPE/Au, and Pt devices at 300 K and $V_G = -70$ V for shorter channel devices. The resistance was measured via $V_{SD}$ sweeps from 0 to -500 mV, while gate voltages from 0 V to -70 V (in 10 V steps) were examined. The mobilities at $V_G = -70$ V, inferred from the length dependences, were $5.3 \times 10^{-2}$ cm$^2$/Vs.
(Au), $5.3 \times 10^{-2}$ cm$^2$/Vs (F-OPE/Au), and $8.7 \times 10^{-2}$ cm$^2$/Vs (Pt). Initial measured resistances from the 5 $\mu$m channel length devices span roughly an order of magnitude between the three different electrode materials, with the highest resistance coming from the gold electrodes and the lowest resistance from the platinum devices. As the channel length is reduced significantly below 1 $\mu$m, there are qualitative differences between the three types of devices. The Au-based devices have much-increased resistances, while the Pt-based devices have decreased resistances. The F-OPE/Au structures are intermediate in their small $L$ properties. We analyzed several ensembles of identically prepared devices, and all exhibited these trends. These trends continued to hold at other gate voltages and as $T$ was reduced. Channel lengths in the short-channel devices were measured via electron microscopy after electrical characterization, to ensure that electrode contamination during imaging did not affect the results.

At the smallest channel lengths there is significant device-to-device variability in $R$, much more so than in the longer channel devices. This appears to reflect that microscopic differences in the metal/organic interface can have significant local impact on the injection process and contact resistances, even when large-scale properties are uniform and well defined. However, even with this variability the overall trends and the systematic deviation between the different electrode types are clear.

One possibility is that the morphology of the P3HT may differ at the metal/P3HT interface for the three different electrode materials, leading to varying contact properties. To test this, we performed tapping-mode atomic force microscopy (AFM) scans of a 3 $\mu$m $\times$ 3 $\mu$m area that encompassed the electrode/P3HT interface for both platinum and gold electrodes. The results are shown in Figure 3. The topographic images show that the polymer film on the oxide adjacent to the electrode edge is smooth (rms roughness of 0.83 nm for the P3HT on oxide next to Au; 0.78 nm for the P3HT on oxide next to Pt) and largely featureless for both electrode materials. Using phase imaging, more detail is observable, with some indications of the fibril morphology sometimes observed in P3HT. However, there is little difference between the two images, at least down to the resolution of our microscope. This implies that gross morphology changes near the contacts are not responsible for the difference in contact resistance properties.

We suggest instead that the explanation for the difference lies in interfacial charge transfer between the electrodes and the P3HT due to the energetics of band alignment. Previous experiments have suggested that in the absence of unintentional doping from the environment the alignment between the Au Fermi level and the P3HT HOMO is such that a significant
hole injection barrier exists. In this case one may expect the tail of the P3HT valence band to be locally depleted in the vicinity of the of the interface. In contrast, measurements involving the interface between P3HT and higher work function systems such as Pt[4, 16] and F-OPE/Au[4] show much lower contact resistances and persistent Ohmic injection in field-effect structures even under treatment conditions where the bulk two-terminal conductivity of the P3HT is immeasurably small. Consistent with recent photoemission experiments[11], we suggest that the charge transfer responsible for pinning the Pt Fermi level above the P3HT HOMO populates the tail states with mobile holes, leading to comparatively enhanced conduction in the shortest devices.

This interpretation is supported by other experimental signatures. First, we examine the low $V_{SD}$ transistor characteristics as a function of $V_G$ for short-channel devices with identical geometries. This is more revealing than $I_D - V_G$ transfer characteristics since the changing channel geometry as $L$ is reduced, high electric fields at large $V_{SD}$, and hysteresis in $V_G$ complicate the interpretation of inferred threshold voltages. A typical result is shown in Fig. 4, for Pt (top) and Au (bottom) devices with $L \approx 100$ nm at $T = 300 \text{K}$. Note that there is a large qualitative difference between these devices. The Pt device shows significant background conduction at $V_G = 0$, qualitatively similar to what is seen in the presence of doping. This conduction vanishes in adjacent, simultaneously fabricated Pt devices as channel lengths are increased into the micron range, showing that it is an effect of the contacts.

We investigate this further by examining the two-terminal source-drain conductance at $V_G = 0$ for the short channel devices ($L < 5 \mu m$) at $V_{SD} = -0.5 \text{V}$. Representative data are shown in Fig. 5. At the $5 \mu m$ limit the ZGB current is essentially the same for all devices. As the channel length decreases, however, the measured currents in devices with platinum contacts continue to increase by orders of magnitude, while the measured currents in devices with gold contacts do not follow this trend. Rather, the Au device $V_G = 0$ currents tend to be fairly constant over the measured channel length range. The current measured from the F-OPE/Au electrodes remains fairly constant and similar to that of gold over the longer channel lengths, but as the channel lengths decrease well below 200 nm, the zero-bias current displays a rapid increase. Since all of the devices of a given electrode composition are prepared simultaneously and measured in one run, these dependences of the $V_G = 0$ conduction on $L$ cannot arise from dopants from the environment.

To further examine charge transfer between the electrodes and the P3HT film, we used an extension of scanning tunneling microscopy (STM) known as alternating current scanning tunneling microscopy (ACSTM). STM and related spectroscopic techniques have long been used to char-
acterize the electronic properties of conducting polymer materials. More common spectroscopies provide high-resolution analysis of properties such as the density of states and work function, yet fail to provide capacitance versus voltage \((C - V)\) information, which is critical for understanding dopant effects. Unlike conventional STM, microwave frequency ACSTM can be used to obtain local \(C - V\) data.

In the ACSTM technique, microwave frequency radiation is applied to the tip-sample junction. By exploiting the nonlinear behavior of the junction using high frequencies, ACSTM can be used to acquire capacitance information. Specifically, the tunneling current response at microwave frequencies when imaging semiconductors reflects the \(\partial C/\partial V\) data\[22\]. In this way, ACSTM can provide high-resolution spatial capacitance information on semiconducting polymer films. The magnitude of the ACSTM signal ultimately reflects the carrier concentration in the substrate. For example, this microwave signal has been used previously in both STM and AFM mode to measure the dopant concentration in silicon\[23, 24, 25\].

For this ACSTM experiment a loop antenna geometry was used to apply microwave radiation to the junction similar to studies previously reported\[27\]. We used the difference frequency ACSTM technique\[23, 24, 25, 26\] to simplify detection of the high frequency modulation of the tunneling signal by allowing the use of conventional lock-in amplifier equipment. Care was taken to ensure that the ACSTM data on both samples were acquired under nearly identical scanning conditions\[28\]. The ACSTM signal was acquired at each point during scanning, thus producing \(\partial C/\partial V\) spectral images (magnitude and phase) and the topography image simultaneously. All scans were taken in ultrahigh vacuum at base pressures on the order of \(10^{-10}\) torr using a commercial RHK STM and mechanically cut Pt:Rh (80:20) tips. The STM was modified to include the antenna so that the loop antenna encircled the STM tip during scanning, similar to a method reported previously\[27\] and could be moved aside for tip and sample exchange. P3HT films were spin-coated onto Au- and Pt-coated \(\text{SiO}_2/\text{Si}\) substrates following the same procedure as in the fabrication of the OFET devices.

In Fig. 6, the topography and ACSTM image are both shown at various bias conditions for P3HT deposited on Au and Pt, respectively\[29\]. The ACSTM images are shown at the same relative color scale in units of mV, directly proportional to local \(\partial C/\partial V\) and therefore carrier concentration (as seen in doped Si devices\[23\]). Bear in mind that the carrier concentration seen in semiconductors in ACSTM depends to some degree on the bias conditions and gap geometry. The gap resistance was kept constant at \(~ 100 \text{ G}\Omega\) to eliminate the influence of tip height on the result-
ing data. A direct comparison of the ACSTM data on Au and Pt shows a marked contrast in carrier concentration. Averaged over the scan area, the carrier concentration is 25% larger for P3HT on Pt than on Au, but there is also a higher spatial variance on the Pt sample despite the comparable surface structure. This is consistent with the interpretation that there are more mobile holes in the P3HT layer on Pt. Measurements taken at multiple locations on each film confirm these results. In addition, the percentage difference between average ACSTM signal on Pt vs. Au increases to 69% as the tip bias relative to the sample is increased to 2 V. This shows that the carriers present on Au are easier to deplete relative to those on Pt. It is difficult to compare quantitatively the ACSTM data in these vertical layered structures with the situation in the transistor geometry. However, the ACSTM data presented here clearly indicate that Pt increases the carrier concentration in the P3HT film more than the Au layer.

Interestingly, both films show inhomogeneities in the magnitude and phase of their AC response that do not correlate in any immediately obvious way with sample topography, although Figs. 6A and 6C show that the surface features of the two films are similar. In particular the variation takes place on length scales that are not correlated with intrinsic structural properties of the underlying polycrystalline metal films, such as grain size.

The transport and ACSTM indications of charge transfer are consistent with simple calculations using a model for charge transfer developed by Paasch and Scheinert[30]. Their model was developed to explain charge transfer and band bending in thin layers of disordered organic semiconductor on top of metal films. This is precisely the situation of the ACSTM samples, and at zero source-drain bias should be reasonably applicable to the organic channel in the immediate vicinity of the source or drain electrodes. Assuming an exponential density of states for the organic semiconductor characterized by an energy scale \( k_B T_0 = 0.1 \) eV, the local potential in the organic semiconductor of layer thickness \( d \) as a function of distance \( x \) away from the metal interface is given by

\[
U(x) = U_d \pm k_B T_0 \ln \left[ 1 + \tan^2 \left( \frac{d - x}{2L_d} \exp \left( \frac{|U_d|}{2k_B T_0} \right) \right) \right],
\]

where \( U_d \) is the potential at \( x = d \), \( L_d \equiv \sqrt{\frac{\varepsilon \varepsilon_0 k_B T}{2e^2 n}} \) is the intrinsic screening length, and \( n = \sqrt{N_c N_v} \exp(-E_g/2k_B T_0) \) is the effective intrinsic carrier density[30]. Here \( \varepsilon \) is the relative dielectric constant of the organic semiconductor, \( E_g \) is the band gap of the organic semiconductor, \( N_c \) and \( N_v \) are the effective densities of states in the conduction and valence bands. The \( \pm \) sign is positive (negative) for hole(electron) accumulation. The Poisson equation can be used to find the
relationship between $d$ and $U_d$:

$$d = 2L_d \exp \left( -\frac{|U_d|}{2k_B T_0} \right) \arctan \left[ \exp \left( \frac{|U_s - U_d|}{k_B T_0} \right) - 1 \right]^{0.5}.$$  \hfill (2)

Here $U_s \equiv E_g/2 - \Phi_h$, where $\Phi_h$ is the hole injection barrier. Rough values used for our estimations are $N_c, N_v = 10^{21}$ cm$^{-3}$, $E_g = 2$ eV, $\epsilon = 3.24$, and $T = 300$ K.

Working with Eqs. (1, 2) and Poisson’s equation, we can determine the charge density in the organic semiconductor layer. Varying the metals involved affects the local potential $U(x)$ and carrier density by altering $\Phi_h$. Recalling previous photoemission results for P3HT on platinum and gold\[16\], for platinum with $\Phi_h = 0.6$ eV, the charge carrier density very near the interface is estimated to be $\sim 9.6 \times 10^{17}$ cm$^{-3}$; for gold with $\Phi_h = 1.2$ eV, we find $\sim -10^{18}$ cm$^{-3}$, with the negative sign indicating a depletion of hole density. These precise numbers should be viewed cautiously, since different surface preparations can alter the effective work functions of metals and the density of states values used are rough. The trend, however, from Au to Pt should be robust and these results are unlikely to change qualitatively upon inclusion of corrections for a Gaussian density of states for the polymer. For both the Pt, the $\Phi_h = 0.6$ eV value is consistent with prior photoemission experiments\[16\] as well as recent Fermi level pinning results for P3HT and estimates of polaron formation energy relevant to Pt/P3HT charge transfer\[6\]. The estimate of $\Phi_h$ for Au also comes from the same photoemission experiments\[16\]. The resulting calculation supports the claim that higher work function metals, with Fermi levels pinned relatively close to the valence band, locally dope the organic material very near the interface.

We have presented a series of experiments examining the contact resistance and ZGB current in very short channel bottom-contact OFETs incorporating P3HT. The striking differences observed between devices with Pt and Au electrodes are consistent with expectations of metal/P3HT charge transfer inferred from photoemission experiments. In particular, it appears that a significant density of mobile holes are transferred from Pt into neighboring P3HT on the nanoscale. ACSTM measurements further support this conclusion, showing a significantly higher free carrier response in P3HT films on Pt compared to identically prepared films on Au. Further experiments, particularly those probing spatial scales comparable to small numbers of polymer chains, should be able to shed further light on the complex problem of charge transfer and Fermi level pinning, complementing spatially averaging techniques such as photoemission. Techniques like ACSTM point the way toward detailed quantitative assessment of local charge density, information that will be extremely useful in refining theoretical models of these important but complicated problems.
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FIG. 1: Top: Expected two-terminal resistance as a function of channel length, with deviations at very short channels expected if contact interface regions have properties that differ significantly from the bulk. Inset: Data on a series of devices with channel width 200 µm at $V_G = -50 \text{ V}$ and $T = 300 \text{ K}$, $V_{SD}$ from 0 to -1 V, showing the long-channel dependence. Of interest in our experiments is the very short channel limit.

Bottom: Electron micrograph of the center of a typical short channel device fabricated by electron beam lithography.
FIG. 2: Two-terminal resistance at low $V_{SD}$ at 300 K and $V_G = -70$ V for various series of OFETs incorporating different electrode materials. Note the greatly enhanced resistance in very short channel Au devices compared with Pt devices. Inset shows a more clear view of the short channel data.
FIG. 3: Atomic force microscopy images of P3HT films in contact with gold (A, C) and platinum (B, D) electrodes. There is no significant difference in P3HT morphology at the two interfaces, suggesting that the observed differences in electrical properties does not result from greatly varying amounts of disorder.
FIG. 4: Transistor characteristics at low $V_{SD}$ for identically prepared Pt (top) and Au (bottom) devices with channel lengths $\approx 144$ nm, width 50 $\mu$m at $T = 300$ K. These devices have essentially identical geometries, yet the Pt-based device shows significant background conduction even at $V_G = 0$. This qualitative difference vanishes as channel lengths exceed the few micron scale.
FIG. 5: Top: ZGB current (measured at $V_G = 0$, $V_{SD} = -0.5 \text{ V}$) at 300 K for the same series of devices (same symbols) in Fig. 2. Notice the great enhancement in ZGB current for extremely short channel Pt and FOPE-Au devices, much larger than simple geometric expectations would suggest. Bottom: Data acquired in a separately fabricated ensemble of Pt-based devices made using the nanogap method of Ref. [31].
FIG. 6: Topography (A, C) and ACSTM (B, D) magnitude images of P3HT films deposited on Au (A, B) and Pt (C, D), respectively. The simultaneously acquired ACSTM images are shown at the same relative color scale. All images were acquired at -1.0 V sample bias, 10 pA tunneling current, and are $473 \times 473$ nm$^2$. The scalebars on all images are 94.6 nm. The color bars are in units of nm for the topography images and mV for the ACSTM images. The greater magnitude in (D) indicates an excess of mobile holes for the P3HT/Pt case relative to the P3HT/Au case of (B).
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[28] Frequencies used were 300 and 300.004 MHz, approximately 3.7 dbm for the Pt sample and 3.0 dbm for the Au sample. The difference is due to the change in antenna position when switching between samples, and was adjusted by monitoring the input at the lock-in amplifier to ensure that the initial out-of-tunneling input signal strength at the difference frequency was the same for both samples. The same tip was used for both samples.

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