Semiconducting single-walled carbon nanotubes (SWNTs) with various \((m,n)\) indices, or chiralities, are quasi one-dimensional materials exhibiting characteristic optical absorption and photoluminescence (PL). The intrinsic photoluminescence of SWNTs is measured to be enhanced more than 10-fold. The dependence of fluorescence enhancement on metal–nanotube distance and on the surface plasmon resonance (SPR) of the gold substrate for various SWNT chiralities is measured to reveal the mechanism of enhancement. Surfactant-coated SWNTs in direct contact with metal exhibit strong MEF without quenching, suggesting a small quenching distance for SWNTs on the order of the van der Waals distance, beyond which the intrinsically fast nonradiative decay rate in nanotubes is little enhanced by metal. The metal enhanced fluorescence of SWNTs is attributed to radiative lifetime shortening through resonance coupling of SWNT emission to the reradiating dipolar plasmonic modes in the metal.

On the other hand, metal-enhanced fluorescence (MEF), a phenomenon reported for various organic fluorophores and quantum dots proximal to gold or silver substrates or other nanostructures, has not been observed with carbon nanotubes thus far. Here, we report the first case of MEF of surfactant-coated, water-soluble SWNTs on solution-grown Au films seeded by Au nanoparticles (called ‘AuAu films’). We find that surfactant coated SWNTs in contact with a metal substrate are immune to quenching, likely due to protection by the coating layer. To the contrary, we observed enhanced fluorescence of SWNTs by more than 10-fold. The enhancement factor monotonically decreases when the SWNTs are placed further away from the AuAu surface using ‘spacing’ layers (SEM, Figure 1b inset) showed small gold islands with small gaps in between the islands. The substrates have been shown recently to afford excellent surface-enhanced Raman scattering (SERS) properties for nanotubes and other molecules. Using 658 nm laser

Figure 1. Schematics showing SWNTs embedded in a PVP matrix on (a) quartz and (b) a AuAu film made on glass. The inset of (b) shows an SEM image of the nanostructured AuAu film. (c) NIR photoluminescence image of SWNTs on quartz and (d) on the AuAu film at the same emission intensity scale showing much enhanced fluorescence on AuAu. The inset in (c) shows the fluorescence image of (c) after amplifying the intensity by 4 times. (e and f) Emission spectra (at 658 nm excitation) for two individual nanotubes in (c) on quartz and in (d) on AuAu substrate respectively. Note that the two different chirality (10,3) and (7,6) tubes both have an absorption band near the 658 nm excitation.

SWNTs (average length \(\sim 400 \text{ nm}\)) coated by PEGylated phospholipid (DSPE-mPEG) in a stable water suspension were prepared by exchanging sodium cholate solubilized SWNTs into the DSPE-mPEG coating to afford relatively long SWNTs with an optimal quantum yield. The SWNT suspension was mixed with polyvinylpyrrolidone (PVP) and spin-coated onto both a quartz substrate (Figure 1a) and a AuAu substrate (Figure 1b) made by solution-phase seeding and growth of gold nanoparticles, as described in the Supporting Information (SI). The morphology of as-made AuAu substrates imaged by scanning electron microscopy (SEM, Figure 1b inset) showed small gold islands with small gaps in between the islands. The substrates have been shown recently to afford excellent surface-enhanced Raman scattering (SERS) properties for nanotubes and other molecules. Using 658 nm laser
excitation and an InGaAs 2D camera, we obtained fluorescence images of ensembles of SWNTs in the 0.9–1.7 µm emission range and detected significantly brighter fluorescence on the AuAu substrate (Figure 1d) than on bare quartz (Figure 1c) for the same density of SWNTs deposited on both substrates. By taking emission spectra for ensembles of SWNTs in the same field of view on both substrates (Figure S1), we found that this particular AuAu film enhanced the overall fluorescence intensity (integrated in the 0.9–1.7 µm range) of SWNTs by ∼8 times compared to SWNTs on bare quartz.

We identified and characterized individual SWNTs on both quartz (Figure 1c) and AuAu (Figure 1f) substrates. Under laser excitation with rotating polarization, individual nanotubes exhibited fluorescence emission periodic to the polarization angle (periodicity with rotating polarization, individual nanotubes exhibited fluorescence emission on bare quartz. Due to the significantly enhanced fluorescence, we recorded emission spectra for individual SWNTs and assigned (m,n) indices (Figure 1e, 1f) based on established spectral assignments.

Figure S2), as shown previously with maximum emission occurring when the laser polarization was parallel to the nanotube axis. We recorded emission spectra for individual SWNTs and assigned (m,n) indices (Figure 1e, 1f) based on established spectral assignments. Due to the significantly enhanced fluorescence, individual SWNTs on the AuAu substrate were easily identified and characterized, with emission spectra showing much higher signal/noise ratios, smoother profiles, and less noisy baselines. The comparisons clearly suggested the advantages of using the AuAu substrate for the characterization of relatively weak fluorophores.

Figure 2. (a) A schematic drawing showing SWNTs on various substrates indicated. (b) A bar chart showing the decreasing PL intensity of SWNTs on different substrates, corresponding to those in the upper schematic.

To elucidate the mechanism of the observed fluorescence enhancement of SWNTs, we formed various spacer layers on AuAu substrates and measured the degree of SWNT fluorescence enhancement as a function of the thickness of the spacers including self-assembled monolayers (SAMs) of 1-propanethiol (thickness ∼0.5 nm) and 1-octadecanethiol (∼2 nm), and ∼5 nm Al2O3 and ∼10 nm Al2O3 made by ALD on cysteamine coated AuAu substrates (Figure 2a). For the same amount of SWNTs deposited onto these substrates, we observed a monotonic decrease of SWNT fluorescence (Figure S3) as the spacer layer thickness increased, with SWNTs on bare quartz exhibiting the weakest fluorescence (Figure 2b). This result suggested that the nanostructured Au film was responsible for the enhanced fluorescence of SWNTs. The enhancement effect reduced as SWNTs were spaced away from the underlying metal surface.

We attribute the enhanced fluorescence of SWNTs on our AuAu substrates to resonance coupling of nanotube emission with surface plasmons of the underlying AuAu film. Similar to the MEF of common fluorophores, coupling between nanotube fluorescence emission and the plasmon modes in metal structures shortens the radiative lifetime (1/Γr) of excited states, leading to a higher quantum yield η = Γr/(Γr + Γnr), where 1/Γnr = nonradiative lifetime). Such a coupling is well-known to reduce monotonically as fluorophores are placed away from the metal substrates by spacer layers, which reduces the enhancement effect as seen in our experiment. SWNTs are known to exhibit low intrinsic quantum efficiency on the order of ∼1–3% in the absence of any metal enhancement, corresponding to Γr ≪ Γnr and η ≈ Γr/Γnr. If the intrinsically high nonradiative decay rate Γnr is not further enhanced by metal, then any enhanced radiative decay Γr will translate into higher quantum efficiencies of SWNTs, by up to ∼8× as in Figures 1 and 2.

It is known that coupling between fluorophores and the nonradiating dark plasmon modes (nondipolar higher order modes) in metal could enhance the nonradiative decay rate and shorten the nonradiative lifetimes (1/Γnr) due to energy transfer, leading to reduced fluorescence emission or even quenching. In our experiments, no quenching was observed for the most proximal case of SWNTs on AuAu substrates without any intentionally placed spacers. However, even in this case there was an ∼3 nm distance between the SWNTs and gold corresponding to the radius of gyration of the DSPE-mPEG (molecular weight ∼5000 Da) polymer chains coated on SWNTs. This prevented the SWNTs from direct contact (within the van der Waals distance of ∼0.34 nm of graphitic materials) with gold and thus avoided quenching. We also coated SWNTs with a much smaller surfactant molecule cholate (molecular weight ∼400 Da) and also observed the MEF of these tubes on the AuAu substrate vs on quartz. These results suggest that the ‘quenching distance’ on metal surfaces for SWNTs seems to be shorter than that of most organic fluorophores (7–15 nm).

The intrinsically high nonradiative decay rate Γnr of SWNT excited states appears to be only significantly enhanced when nanotubes are very close to metallic species, like in a bundle where metallic tubes intimately coexist. We conclude that SWNTs with noncovalent molecular functionalization are in fact relatively immune to quenching with a small quenching distance of ∼3 nm to metal surfaces.

We investigated the MEF of SWNTs on four different AuAu substrates (labeled as AuAu-I to -IV, Figure 3; see the SI for their synthesis) with increasing metal coverage or thickness on glass substrates, with higher optical absorbance and increasingly red-shifted surface plasmon resonance features (Figure 3f). Photo luminescence excitation (PLE) spectra of the same amount of SWNTs deposited on bare quartz (Figure 3a) and the four AuAu films (Figure 3b–e) revealed a monotonic increase in PL intensity for various chiral SWNTs (Figures 3 and 4), and the enhancement factor was up to 10–12× on the thinnest AuAu film (Figures 3e and 4d). Noteworthy was that a similar trend was observed for the MEF of indocyanine green (ICG) on silver substrates, where higher fluorescence was detected on thicker silver films with higher optical absorbance.

We analyzed the MEF of several groups of (m,n) SWNTs with similar excitation wavelengths but different emissions [e.g., (7,5), (7,6), and (10,3) SWNTs, Figure 3d] and correlated their MEF enhancement factors with the SPR absorbance at the emission wavelengths. Figure 4a–c show decreasing surface plasmons at
The (7,5), (7,6), and (10,3) SWNTs on the AuAu-III film showed similar enhancement factors but appeared to be slightly against the trend of the SPR absorbance/extinction curve (Figure 4c). This was attributed to the small variations in the extinction in the 1000–1400 nm range and that the intrinsic chirality dependence of MEF could be at play. We also analyzed SWNTs with similar emission wavelengths and different excitation wavelengths. The plotted EFs vs excitation wavelengths, alongside with SPR extinction curves for all AuAu substrates, show a much worse correlation (Figure S4) than that in Figure 4.

It is established that fluorescence enhancement of a molecule by metal depends on the size (and plasmonic frequency) of the metal structure. Stronger dipolar coupling between the molecule and plasmons in the metal prefers smaller metal sizes. On the other hand, larger metal structures, such as in the thicker AuAu films, are submicrometer structures composed of smaller nanocrystals, and they afford a stronger scattering or reradiating component of the optical extinction, leading to higher enhancement by reradiating the emission. These two competing factors often lead to an optimum metal size for the MEF of molecules. In our case with nanotubes of an average ~400 nm length as studied here, the monotonic increase in MEF as the AuAu film thickens suggests that stronger scattering and reradiating effects of surface plasmons are dominant and responsible for the enhanced SWNT fluorescence. We also studied the MEF of shorter nanotubes (~150 nm on average) with a much lower quantum yield (made by direct sonication in DSPE-mPEG surfactant) and observed even higher fluorescence enhancement factors (up to 40×) on the relatively thin AuAu-III film with many small gaps between the Au islands. On thicker and more continuous AuAu films, a decrease in enhancement factor was observed for these short tubes. These results suggested short SWNTs near the gaps of Au nanoislands may exhibit fluorescence enhancement due to electric field enhancement in addition to resonance plasmonic enhancement.

In summary, we observed the first metal enhanced fluorescence of SWNTs resulted from radiative lifetime shortening through resonance coupling of SWNT emission to plasmonic modes in the metal. The enhancement effect decreases monotonically as the separation between nanotubes and metal increases. Surfactant-coated SWNTs in direct contact with metal exhibit the strongest MEF without quenching, suggesting a small quenching distance on the order of the van der Waals distance, beyond which the intrinsically fast nonradiative decay rate in nanotubes is little changed. SWNT fluorescence enhancement monotonically increases with optical extinction of thicker gold films due to increased scattering and reradiating of stronger surface plasmons resonantly coupled to nanotube emission. We believe that metal enhanced fluorescence of SWNTs by >10 times will be welcoming news for these materials in fundamental and practical applications including sensing, detection, and imaging.

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Supporting Information Available: Experimental procedures and supportive data. This material is available free of charge via the Internet at http://pubs.acs.org.

References


