Tailoring Drug Mobility by Photothermal Heating of Graphene Plasmons

Anh D. Phan,* Nguyen K. Ngan,* Do T. Nga, Nam B. Le, and Chu Viet Ha

A theoretical approach to quantitatively determine the photothermally driven enhancement of molecular mobility of graphene-indomethacin mixtures under infrared laser irradiation is proposed. Graphene plasmons absorb incident electromagnetic energy and dissipate them into heat. The absorbed energy depends on optical properties of graphene plasmons, which are sensitive to structural parameters, and concentration of plasmonic nanostructures. By using theoretical modelling, temperature gradients of the bulk drug with different concentrations of graphene plasmons are calculated. From these, the temperature dependence of structural molecular relaxation and diffusion of indomethacin are determined and how the heating process significantly enhances the drug mobility is found out.

1. Introduction

Photothermal effects have a wide range of applications such as solar steam generation,[1] thermo-optical data writing,[2] cancer treatment,[3,4] and destroying bacteria.[5] The incident electromagnetic field excites collective oscillations of free electrons, which are surface plasmons in metal-like materials. Plasmonic excitations allow confining light near the metal–dielectric interface and enhancing local fields. Thus, plasmonic materials can absorb light and turn it into heat. Investigating mechanisms underlying light-to-heat conversion not only provides insight into fundamental science, but also proposes novel practical applications. Very recently, photothermal plasmonic nanoparticles have been experimentally exploited to induce drug amorphization inside a table[6] upon laser irradiation. Compared with other hyperthermia methods (using thermal chambers, ultrasound, microwave, and magnetic nanoparticles associated with an alternating magnetic field),[7] the photothermal approach has a simple operation and an advantage to locally and selectively increase temperature instead of spreading the heat to larger zones. Compared to global heating, a localized heating effect controls on-demand release of drugs in abnormal cell regions at a desired time and location to improve pharmaceutical efficacy. In addition, global and uniform heating can damage to surrounding normal tissues and induce direct cell necrosis. Thus, it is a highly efficient technique and side effects are minimized. The laser heating increases temperature and molecular mobility of the drug and polymer in the tablet. Molecular mobility is characterized by molecule’s structural relaxation and diffusion, which can be measured using the broadband dielectric spectroscopy. The photothermal heating of drugs may be an answer to overcome poor aqueous drug solubility in oral drug delivery.[8,9] Recrystallization of pharmaceuticals during manufacturing or storage, which restricts the use of amorphous drugs, can be resolved by plasmonic heating. All challenging problems are of interest for the scientific and industrial community but have not been theoretically investigated.

Among many types of photothermal agents, graphene has received much attention since it has various intriguing behaviors.[10–13] Graphene exhibits strong light absorption with relatively low losses.[13] The number of free electrons in graphene is small and this leads to a significant reduction of the heat dissipation compared to noble metals.[13] Large inelastic losses reduce lifetimes of plasmon and optical confinement. It is easy to tune plasmonic properties of graphene by using doping, defects, external fields, and charge injection.[12]

Although graphene nanostructures are composed of carbon atoms which is more friendly with human body than conventional plasmonic materials, their toxicity still exists and strongly depends on shape, size, purity, concentration, and dose.[14–16] For example, wistar rats have no sign of toxicity when injected doses of 1 and 50 mg kg⁻¹ of 100-nm graphene nanodisks, however, a 100 mg kg⁻¹ dose killed two of eight rats after 2 weeks of
injection. Meanwhile, graphene nanostructures having size in the range of 500–1000 nm can cause cytotoxicity when studying the viability of RAW 264.7 macrophages injected by 20–100 μg mL⁻¹. Oxidation of graphene was found to be a main reason for toxicity in the lungs of mice. According to guidance for safety studies of Food and Drug Administration (FDA), a maximum safe starting dose is 100 μg per subject for clinical trials and the maximum dose 100 times larger than the clinical dose can be used for nonclinical studies. Previous in vivo experiments clearly confirmed the safety of the recommended clinical dose. Using higher doses can cause either several serious problems in some cases or minor side effects. Overall, graphene nanostructures having the 100-nm size possibly have less toxicity than their larger counterparts and health risks can be potentially found when investigated at the dose exceeding 100 μg per subject. The toxicity is governed by the complex interplay of physiochemical properties but it remains not well understood and further experimental investigations are necessary. This is why graphene-based nanomaterials have been received much attention to improve the biocompatibility and propose various new biological and pharmaceutical applications.

In this letter, we theoretically investigate effects of photothermal heating on molecular mobility in a graphene–drug composite under an infrared laser illumination (Figure 1a). Monolayer graphene plasmons are randomly dispersed in bulk indomethacin and considered as heat sources. Absorption and heat generation of this mixture depends on optical properties and density of graphene nanostructures. After determining the temperature dependence of structural relaxation and diffusion by the elastically collective nonlinear Langevin equation (ECNLE) theory, we calculate time-dependent temperature of the drug mobility upon laser irradiation.

2. Optical Properties and Photothermal Effects of Graphene Plasmons

When the wavelength of the incident electromagnetic field is much larger than the size of graphene plasmon, the quasi-static approximation can be exploited to analytically determine the absorption cross section of a single graphene plasmon by

\[ Q_{\text{abs}} = \frac{4\pi\omega}{\epsilon} \text{Im} \left[ \frac{\sqrt{\varepsilon_m} D^3 \zeta^2}{-i\omega D \frac{\sigma(\omega)}{\sigma_m} - 1} \right] \]

(1)

where \( \varepsilon_m \) is the dielectric constant of medium, \( \omega \) is the frequency of the incident field, \( D \) is the diameter of graphene nanodisks, \( \sigma(\omega) \) is the optical conductivity, and \( \zeta = 0.8508 \) and \( \eta = -0.0728 \) are morphological parameters. In mid-infrared regime, the monolayer graphene conductivity is

\[ \sigma(\omega) = \frac{e^2 |E_F|}{\pi \hbar^2 (\omega + i\tau)^{-1}} \]

(2)

where \( E_F \) is a chemical potential, \( e \) is the charge of electron, \( \hbar \) is the reduced Planck constant, and \( \tau \) is the carrier relaxation time. We choose \( E_F = 0.35 \text{ eV} \) and \( \hbar \tau^{-1} = 0.001 \text{ eV} \) to be consistent

![Figure 1.](image-url)
with ref. [27]. Typically, the full expression of \( \sigma(\omega) \) has two terms corresponding to interband and intraband transitions. However, the interband conductivity is extremely small in the mid-infrared regime. Thus, we ignore contributions of interband transitions in Equation (2).

Figure 1b shows influence of finite-size effects of graphene plasmon on the normalized absorption cross section calculated by Equation (1) and (2). Since there is no dielectric data for indomethacin the frequency range of 0.08–0.16 eV, we extrapolate dielectric data in ref. [29], which is measured in the frequency range of \( 10^{-2}–10^7 \) Hz. This extrapolation suggests that dielectric loss at higher frequencies is very small while the dielectric constant is \( \epsilon_\infty \approx 1.77 \). Thus, the absorption of the drug is extremely lowered and we assume to ignore it in calculating \( \sigma_{\text{abs}} \). The maximum values of the normalized absorption spectrum are \( \approx 5–7 \) although the size of graphene nanodisk increases from 60 to 160 nm. Graphene plasmons in this size range are safe for living subjects.[17,22] One can roughly estimate the maximum absorption, \( \sigma_{\text{abs}} \), of the plasmonic agents under irradiation of infrared laser light with ref. [27]. Typically, the full expression of \( \sigma(\omega) \) has two terms corresponding to interband and intraband transitions. However, the interband conductivity is extremely small in the mid-infrared regime. Thus, we ignore contributions of interband transitions in Equation (2).

Now, graphene plasmons having the density, \( N \) (the number of nanostructures per volume), are randomly mixed with indomethacin drugs. In this work, we choose \( N = 5 \times 10^{13} – 3 \times 10^{14} \) nanodisks per m\(^3\). The averaged separation distance between two adjacent graphene plasmons is \( 1/N^{1/3} \approx 14.9 – 27 \mu \text{m} \gg D \). It means that nanostructures in the drug are fully isolated and no plasmonic coupling occurs. Since Equation (1) allows us to determine the in-plane optical absorption, we can effectively include effects of random orientation of graphene nanodisks on the graphene absorption by replacing \( \sigma_{\text{abs}} \) with \( 2\sigma_{\text{abs}}/3 \). From this, according to, an effective absorption coefficient of our binary mixture constructed using the Beer-Lambert law is \( \alpha(\omega) = 2N\sigma_{\text{abs}}/3 \). Photothermal effects of the plasmonic agents under irradiation of infrared laser light are exploited to control drug mobility. We consider the photo-thermal heating of bulk indomethacin of graphene plasmon with \( D = 120 \) nm when exposed by an infrared laser beam operating at \( \approx 0.1 \) eV (or 806.6 cm\(^{-1}\)) and the intensity \( I_0 = 1 \) W cm\(^{-2}\). Photothermal heating pass through these systems is absorbed by the plasmonic materials and converts from optical energy to thermal energy to increase the temperature of the surrounding medium. The light-to-heat conversion process is described using heat transfer equations without taking into account radiative and convective heat losses.

To calculate the spatial temperature profile in the bulk indomethacin of graphene plasmons under laser illumination, we use an analytic expression of thermal response of a semi-infinite substrate\(^{31,32} \) derived from a heat energy balance equation, which gives

\[
\Delta T(x, y, z, t) = \frac{I_0(1 - R)\alpha}{2\rho_\text{dr}c } \int_0^t \exp \left( -\beta^2 (x^2 + y^2) \right) \left[ \frac{e^{\beta z_\text{abs}}}{1 + 4\beta^2 \kappa^2} \left( e^{-\beta z_\text{abs}} \operatorname{erfc} \left( \frac{2\alpha \beta \kappa t - z}{2\sqrt{\kappa^2 t}} \right) \right) \right] \frac{1}{e^{\beta z_\text{abs}}} 
\]

where \( z \) is the depth direction parallel with the incident field, \( x \) and \( y \) are coordinates in the horizontal plane, \( R \) is the reflectivity of indomethacin, and \( \beta \) is the inverse of the laser spot radius. Suppose that the spot size is infinite, we have \( \beta = 0 \). The thermal diffusivity of the drug, \( \kappa = K_d/\rho_d c \), is calculated by the thermal conductivity, \( K_d = 0.18 \) W m\(^{-1}\)K\(^{-1}\)\(^{33} \) the mass density, \( \rho_d = 1213 \) kg m\(^{-3}\), and the specific heat capacity, \( c_d = 1200 \) J kg\(^{-1}\) K\(^{-1}\)\(^{33} \). Equation (3) has been used to quantitatively describe the photothermal heating of a solution of TiN nanoparticles and provide a good agreement with experiments\(^{31} \). We assume that the validation is still held in our calculations.

Based on refs. [34,35], the transmission of indomethacin at 806.6 cm\(^{-1}\) is about 97%. It suggests that \( R \) and the absorption of pure indomethacin are relatively small. Thus, we assume \( R \approx 0 \) in Equation (3) and the photothermal heating of graphene-drug composites are mainly caused by the absorbed energy in graphene plasmons. This is consistent with our assumption given to calculate the absorption cross section of graphene plasmons in Figure 1. The same performance (\( R \approx 0 \)) can be obtained when the frequency of laser and \( \omega_{\text{max}} \) lie in the range of 1800–2500 cm\(^{-1}\)\(^{34,35} \) or 0.22–0.31 eV. It means that the Fermi energy varies from 1.3 to 1.8 eV if the diameter of graphene plasmon is fixed at 120 nm. One can also reduce \( D \) to shift the plasmonic resonance to this desired range of frequency while \( E_F \) remains small.

Figure 2 shows numerical results for the temporal and spatial temperature of the graphene-drug mixtures calculated using Equation (3). Increasing the density of graphene plasmons not only leads to more absorption of optical energy in the mixture, but also shortens the penetration depth of electromagnetic fields. Thus, the absorbed energy is localized near the surface, and the light-to-heat conversion decays towards the drug center. This finding explains why the surface temperature monotonically grows with an increase of \( N \) from \( 5 \times 10^{13} \) to \( 3 \times 10^{14} \) disks m\(^{-3}\), while the temperature in the drug interior slightly decreases in comparison with that in the surface. Changing temperature changes molecular mobility in the drug.
3. Photothermal Induced Change of Molecular Dynamics of Drugs

To investigate the temperature dependence of structural relaxation and diffusion of amorphous materials, we use the ECNLE theory. In this theory, all glass-forming liquids are modeled as a hard-sphere fluid of the volume fraction \( \Phi \). A sphere of the fluid has the particle diameter, \( d \). Molecular mobility of a tagged particle is affected by two main factors: 1) the nearest-neighbor interactions and 2) cooperative motions beyond nearest neighbors. The former factor governs the local dynamics quantified by the dynamic free energy \( F_{\text{dyn}}(r) = F_{\text{ideal}}(r) + F_{\text{caging}}(r) \) with \( r \) being the displacement. \( F_{\text{ideal}}(r) \) and \( F_{\text{caging}}(r) \) correspond to the ideal fluid and localized state, respectively. The caging force acting on the tagged particle tightly depends on the density and structure of systems.

The inset of Figure 3a illustrates the function form of \( F_{\text{dyn}}(r) \) when the density is sufficiently large and the dynamical arrest occurs. The dynamical trapping of the tagged particle within a cage formed by its nearest neighbors is characterized by rising barrier in \( F_{\text{dyn}}(r) \). A particle cage radius is \( r_{\text{cage}} \approx 1.3 - 1.5 \) determined by the radial distribution function, \( g(r) \). Key physical quantities of the local dynamics are a localization length, \( r_L \), a barrier position, \( r_B \), a jump distance, \( \Delta r = r_B - r_L \), and a local barrier, \( F_B \). Then, the harmonic curvatures \( K_0 \) and \( K_B \) at \( r_L \) and \( r_B \), respectively, can be calculated.

Collective dynamic effects of particles beyond the cage on an escaping process of the tagged particle are distinct but strongly correlated to the local dynamics. Particles in the first shell need to reorganize to create extra space for hopping. This molecular reorganization triggers a displacement field, \( u(r) \), which is nucleated from the cage surface and radially propagates through the remaining medium. Thus, collective particles are vibrated as harmonic oscillators with a spring constant \( K_0 \). By using continuum mechanics analysis, the displacement field is given by \( u(r) = \Delta r_{\text{eff}} r_{\text{cage}}^2 / r^2 \) for \( r \geq r_{\text{cage}} \), where \( \Delta r_{\text{eff}} \) is an amplitude and its mathematical form is reported in Ref. [23,24]. From this, the elastic energy of an arbitrary oscillator is \( K_0 \Delta r_{\text{eff}}^2 r_{\text{cage}}^2 / d^3 \) and the total elastic energy of collective particles or the elastic barrier is

\[
F_e = \int_{r_{\text{cage}}}^{\infty} \rho g(r) K_0 u_r^2(r) \frac{2}{2} dr \\
\approx 12 \Phi K_0 \Delta r_{\text{eff}}^2 r_{\text{cage}}^2 / d^3
\]

where \( \rho = 6 \Phi / \pi d^3 \) is the number of particles per volume and \( g(r) \approx 1 \) for \( r \geq r_{\text{cage}} \).

After obtaining the local and elastic barrier, one can calculate the structural relaxation time by Kramer’s theory\[^{[23-25]}\]

\[
\tau_{\alpha} = 1 + \frac{2 \pi}{\sqrt{K_0 K_B}} k_B T d^2 \exp\left( \frac{F_B + F_e}{k_B T} \right)
\]

where \( \tau_{\alpha} \) is a short relaxation time scale reported elsewhere. Note that Equation (5) provides \( \tau_{\alpha} \). Meanwhile, direct comparison with experiments requires the temperature dependence of \( \tau_{\alpha} \). Based on previous works, we use a density-to-temperature conversion or a thermal mapping: \( T = T_g + (\Phi_g - \Phi) / \beta_g \Phi_g \) with \( T_g \) being the experimental glass transition temperature (defined by \( \tau_{\alpha}(T = T_g) = 100 \) s in Figure 3a), \( \Phi_g = 0.6157 \), \( \beta_g = 12 \times 10^{-4} \) K\(^{-1} \), and \( \Phi_0 = 0.5 \). For indomethacin with low concentrations of graphene plasmons, \( T_g \approx 315 \) K\[^{[23]}\]. As shown in the mainframe of Figure 3a, theoretical ECNLE predictions for \( \tau_{\alpha}(T) \) quantitatively agree with experiment\[^{[38]}\] for a wide range of time. The relaxation is significantly sped up with heating and glassy dynamics in graphene-drug mixtures remains unchanged from that in the pure drug.

As shown in Figure 2, under laser illumination, photothermal effects of graphene plasmon increase temperature of binary mixtures and reduce the structural relaxation at room temperature. Figure 3b exhibits numerical results for \( \tau_{\alpha}(T = \Delta T(t)) \) near air–drug interface as a function of exposure time at several values of \( N \). At 1200 s of light exposure, the molecular relaxation is accelerated by approximately 3 and 9 orders of magnitude for \( N = 5 \times 10^{13} \) and \( 2 \times 10^{14} \) disks m\(^{-3} \), respectively. The structural relaxation in the drug interior can be easily calculated.
but interfacial dynamics is strongly related to molecular surface diffusion and drug solubility.

A relation between molecular diffusion constant and relaxation time is given by

\[ D(T) = \frac{\Delta r^2}{6 \tau_a(T)} \]  

By using data in Figure 3a and Equation (6), \( D(T) \) for the graphene-indomethacin mixtures is predicted. We contrast theoretical prediction and experimental data of several materials including indomethacin, \( \alpha \)-terphenyl, and trisnapthylbenzene in Figure 4a. Theory and experiment quantitatively agree with each other.

Laser-induced heating not only accelerates molecular dynamics, but also facilitates the diffusion. Numerical calculations for \( D(T + \Delta T(t)) \) with three densities of graphene plasmons are given in Figure 4b. Since \( D(T) \) is inversely proportional to \( \tau_a(T) \), these physical quantities vary in the same order of magnitude. One can estimate the variation of \( D(T) \) via \( \tau_a(T) \). In the future, the predictive diffusion constant can be used in Noyes-Whitney equation and its variants to calculate the solubility of materials.

4. Conclusions

In conclusion, we have investigated the relaxation time and diffusion of graphene-indomethacin mixtures under infrared laser irradiation. Dilute concentrations of graphene plasmons are randomly mixed in the drug. Based on the absorption cross section of individual graphene plasmons, the absorbed energy density and temperature rise are calculated when shining light. Increasing temperature makes pharmaceutical molecules move faster. Thus, the structural relaxation is sped up and the diffusion is enhanced when the photothermal heating occurs. We combine the ECNLE theory and laser-induced temperature rise prediction to quantitatively determine how plasmonic properties and density of graphene plasmons change the molecular mobility of drugs. This approach is a predictive model and could be applied to other plasmonic and amorphous materials.

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Conflict of Interest
The authors declare no conflict of interest.

Data Availability Statement
Research data are not shared.

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